
From: Foss, William A.
Sent: Tuesday, January 28, 2020 4:39 PM
To: Watson, Susan
Cc: Mena, Kim
Subject: FW: June 2019 Draft Final LHAAP-03 RD/RAWP

TCEQ and EPA Approvals for the LHAAP-03 RD/RAWP below. EPA July 2, TCEQ July 10.

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Subject: RE: June 2019 Draft Final LHAAP-03 RD/RAWP

EXTERNAL SENDER

Good afternoon. TCEQ has also reviewed the June 2019 Draft Final LHAAP-03 RD/RAWP and the RTCs and accepts the document.

Thank you,

April

From: Mayer, Richard <mayer.richard@epa.gov>
Sent: Tuesday, July 2, 2019 1:25 PM
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Subject: RE: June 2019 Draft Final LHAAP-03 RD/RAWP

Good Afternoon Bill, EPA has reviewed the June 2019 Draft Final LHAAP-03 RD/RAWP and the RTCs and grants approval. Thank you.

From: Foss, William A. <William.Foss@aptim.com>
Sent: Friday, June 28, 2019 7:23 PM
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Subject: 06_2019 Draft Final LHAAP-03 RD/RAWP

Rich and April,

The *Draft Final Remedial Design/Remedial Action Work Plan, LHAAP-03, Former Waste Collection Pad, Building 722-P Paint Shop, Longhorn Army Ammunition Plant* has been uploaded to the portal under [Documents/Recent](#). Hard copies and CDs were sent today via UPS and should be delivered to you Monday. The document includes responses to the EPA comments dated June 3, 2019 and the TCEQ comments dated June 6, 2019 and revisions to reflect the responses. A PDF copy of the document is attached and it can also be accessed directly on the portal via the link below. Please let us know if you have any comments or questions. Thanks!

[06_19 DRAFT FINAL LHAAP-03 RD-RAWP](#)

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DEPARTMENT OF THE ARMY
LONGHORN ARMY AMMUNITION PLANT
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June 28, 2019

DAIM-ODB-LO

Mr. Rich Mayer
U.S. Environmental Protection Agency, Region 6
1201 Elm Street, Suite 500
Dallas, TX 75270-2102

**Re: Draft Final Remedial Design/Remedial Action Work Plan, LHAAP-03,
Former Waste Collection Pad, Building 722-P Paint Shop
Longhorn Army Ammunition Plant, Karnack, Texas, June 2019**

Dear Mr. Mayer,

One hard copy and one compact disc (CD) of the above-referenced document is being transmitted to you for your records. The document includes revisions based upon the Environmental Protection Agency's (EPA) comments on the Draft version received on June 3, 2019, and Texas Commission on Environmental Quality's (TCEQ) comments received on June 6, 2019. In accordance with Federal Facility Agreement, this Draft Final will be considered Final after 30 days without further comment. Response to comments on the Draft version of the document are included with this Draft Final.

The document was prepared by Bhate Environmental Associates, Inc., (Bhate) team, on behalf of the Army as part of Bhate's Performance Based Remediation contract for the facility. I ask that Kim Nemmers, Bhate's Project Manager, be copied on any communications related to the project.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in cursive script that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
Longhorn AAP Site Manager

Copies furnished:

- A. Palmie, TCEQ, Austin, TX (letter)
- P. Bruckwicki, Caddo Lake NWR, TX (1 hard copy and 1 CD)
- A. Williams, USACE, Tulsa District, OK (1 CD)
- R. Smith, USACE, Tulsa District, OK (Electronic only)

A. Sherman, USAEC, San Antonio, TX (1 CD)
K. Nemmers, Bhate, Lakewood, CO (1 hard copy and 1 CD)
P. Srivastav, APTIM, Houston, TX (letter)



DEPARTMENT OF THE ARMY
LONGHORN ARMY AMMUNITION PLANT
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June 28, 2019

DAIM-ODB-LO

Ms. April Palmie
Texas Commission on Environmental Quality
Superfund Section, MC-136
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Austin, TX 78753

**Re: Draft Final Remedial Design/Remedial Action Work Plan, LHAAP-03
Former Waste Collection Pad, Building 722-P Paint Shop
Longhorn Army Ammunition Plant, Karnack, Texas, June 2019**

Dear Ms. Palmie,

One hard copy and one compact disc (CD) of the above-referenced document is being transmitted to you for your records. The document includes revisions based upon the Environmental Protection Agency's (EPA) comments on the Draft version received on June 3, 2019, and Texas Commission on Environmental Quality's (TCEQ) comments received on June 6, 2019. In accordance with Federal Facility Agreement, this Draft Final will be considered Final after 30 days without further comment. Response to comments on the Draft version of the document are included with this Draft Final.

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The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in black ink that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
Longhorn AAP Site Manager

Copies furnished:

R. Mayer, USEPA Region 6, Dallas, TX (letter)
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R. Smith, USACE, Tulsa District, OK (Electronic only)
A. Sherman, USAEC, San Antonio, TX (1 CD)
K. Nemmers, Bhate, Lakewood, CO (1 hard copy and 1 CD)
P. Srivastav, APTIM, Houston, TX (letter)

**Response to Comments on
Draft Remedial Design / Remedial Action Work Plan
LHAAP-03 Former Waste Collection Pad, Building 722-P Paint Shop,
Longhorn Army Ammunition Plant, Karnack, Texas**

Document Date: 17 May 2019

Comment Date: 3 June 2019

Reviewer: Mr. Richard Mayer, USEPA

Respondent: Dr. Rose Zeiler

1. Respondent Concur (C), Does Not Concur (D), Takes Exception (E), or Delete (X)
2. Commenter Agrees (A) with response, or Does Not Agree (D) with response

Comment No.	Section, Page ref.	USEPA Comment	C, D, E, or X ¹	Response	A or D ²
1.	Appendix C	Appendix C titled <i>Well Completion Log for Monitoring Well 03WW01</i> appears to be a Well Completion Report. Please correct the title or provide the well completion log information.	C	The Appendix C fly sheet and the list of Appendices have been revised to “State of Texas Well Completion Report for Monitoring Well 03WW01.”	

**Response to Comments on
Draft Remedial Design / Remedial Action Work Plan
LHAAP-03 Former Waste Collection Pad, Building 722-P Paint Shop,
Longhorn Army Ammunition Plant, Karnack, Texas**

Document Date: 17 May 2019

Comment Date: 6 June 2019

Reviewer: Ms. April Palmie, TCEQ

Respondent: Dr. Rose Zeiler

1. Respondent Concur (C), Does Not Concur (D), Takes Exception (E), or Delete (X)
2. Commenter Agrees (A) with response, or Does Not Agree (D) with response

Comment No.	Section, Page ref.	TCEQ Comment	C, D, E, or X ¹	Response	A or D ²
1.	General	Please revise abbreviated dates. Example on page 1-1, Aug to August.	C	Dates have been revised as requested.	
2.	Pg. 1-2	First paragraph – add TCEQ before Risk Reduction Standard 3	C	Text has been revised as requested.	
3.	Table 1-1	Please make arsenic italic [or bold] and add note from ROD: Applicable RRS3 Industrial Soil MSC equals largest of calculated RRS3 MSC and background values. Italicized [or bold] text indicates value equals background.	C	Table 1-1 has been revised to show the arsenic value in italics and the note from the ROD has been added to the table as requested.	
4.	2.3, 2-2	Correct CLNWR acreage –7,100.	C	Text has been corrected to show the correct acreage.	

Comment No.	Section, Page ref.	TCEQ Comment	C, D, E, or X ¹	Response	A or D ²
5.	4.3.2, 6.2 and other relevant sections	Consider P&A of 03WW01 before excavation. The well could be replaced, if deemed necessary. After looking at year 4 results for LHAAP-58, it would also be acceptable to P&A 03WW01 and not replace the well.	D	Well 03WW01 is still needed as a performance monitoring well for the LHAAP-58 remedy. We will take appropriate measures to avoid damaging the well but will plug and abandon and replace it if it is damaged during the excavation.	
6.	Figure 4-3	Please revise the blue circle symbol description to “Existing SB results used as sidewall confirmation samples” [or similar] For notes 1 and 2 add reference to Table 4-1.	C	The description of the blue circle symbols has been revised to read “Existing soil boring sample to be used as a sidewall confirmation sample”. Reference to Table 4-1 has been added in parentheses in the 1 st and 2 nd notes.	
7.	Figure 4-4	See note regarding blue circle in previous comment, which also applies to yellow circles for floor samples. Add notes (like Figure 4-3) to describe confirmation sample process and reference to Table 4-1.	C	The descriptions for the blue and yellow symbols have been revised in accordance with the response to Comment #6. The callout to Table 4-1 and description of the new floor sample to be collected were added to the descriptions in parentheses.	



Draft Final
 Remedial Design and Remedial
 Action Work Plan, LHAAP-03 Former
 Waste Collection Pad, Building 722-P
 Paint Shop
 Longhorn Army Ammunition Plant
 Karnack, Texas



Prepared for U.S. Army Corps of Engineers, Tulsa District
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Contract No. W9128F-13-D-0012
 Task Order No. W9128BV17F0150
 Project No. 501032

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June 2019

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Appendix A	Analytical Data Reports from November 2018 Soil Samples
Appendix B	Sample Collection Logs for November 2018 Soil Samples
Appendix C	State of Texas Well Completion Report for Monitoring Well 03WW01

Acronyms and Abbreviations

AECOM	AECOM Technology Corporation
APTIM	Aptim Federal Services, LLC
ARAR	applicable or relevant and appropriate requirements
BERA	baseline ecological risk assessment
Bhate	Bhate Environmental, Inc.
BRAC	base realignment and closure
CERCLA	Comprehensive Environmental Response, Compensation, and Liability Act
COC	chemical of concern
CY	cubic yards
FFA	federal facility agreement
ft bgs	feet below ground surface
GPS	global positioning system
GWTP	groundwater treatment plant
IWWP	Installation-Wide Work Plan
Jacobs	Jacobs Engineering Group, Inc.
LHAAP	Longhorn Army Ammunition Plant
LUC	land use control
MATOC	Multiple Award Task Order Contract
MCLs	maximum contaminant levels
MEGA	Multiple Environmental Government Acquisition
mg/kg	milligrams per kilogram
MMRP	Military Munitions Response Program
MOA	memorandum of agreement
MSC	medium-specific concentration
NPL	National Priorities List
PPE	personal protective equipment
RAOs	remedial action objectives
RAWP	Remedial Action Work Plan
RD	remedial design
RI/FFS	Remedial Investigation/Focused Feasibility Study
ROD	record of decision
RRS2	Risk Reduction Standard 2

Acronyms and Abbreviations (continued)

RRS3	Risk Reduction Standard 3
SF	square feet
Shaw	Shaw Environmental, Inc.
SOP	standard operating procedure
SVOC	semivolatile organic compound
TAC	Texas Administrative Code
TCEQ	Texas Commission on Environmental Quality
U.S. Army	U.S. Department of the Army
UCL	upper confidence limit
USACE	U.S. Army Corps of Engineers
USC	United States Code
USEPA	U.S. Environmental Protection Agency
USFWS	U.S. Fish and Wildlife Service
VOC	volatile organic compound

1.0 INTRODUCTION

The U.S. Army Corps of Engineers (USACE), Tulsa District, contracted Bhate Environmental, Inc. (Bhate), under the Omaha Multiple Environmental Government Acquisition (MEGA) National Small Business Multiple Award Task Order Contract (MATOC) Environmental Remediation Services with Military Munitions Response Program (MMRP), Task Order No. W9128BV17F0150 to conduct environmental restoration of LHAAP-03 at Longhorn Army Ammunition Plant (LHAAP). The Bhate Team is comprised of Bhate and Aptim Federal Services, LLC (APTIM). This Remedial Design (RD) and Remedial Action Work Plan (RAWP) was prepared to describe the design elements selected to implement the remedy for LHAAP-03 described in the Final Record of Decision (ROD) (Bhate 2018a) and the actions necessary to implement them.

1.1 Site Description

LHAAP is approximately 14 miles northeast of Marshall, Texas and approximately 40 miles west of Shreveport, Louisiana (**Figure 1-1**). The installation occupies approximately 1,300 of its former 8,416 acres between State Highway 43 at Karnack, Texas and the western shore of Caddo Lake. The facility can be accessed via State Highways 43 and 134.

LHAAP was placed on the Superfund National Priorities List (NPL) on August 9, 1990. Activities to remediate contamination began in 1990. After its listing on the NPL, the U.S. Department of the Army (U.S. Army), the U.S. Environmental Protection Agency (USEPA), and the Texas Water Commission (now the Texas Commission on Environmental Quality [TCEQ]) entered into a Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA) §120 Federal Facility Agreement (FFA) for remedial activities at LHAAP. The FFA became effective December 30, 1991. LHAAP operated until 1997 when it was placed on inactive status and classified by the U.S. Army Armament, Munitions, and Chemical Command as excess property. LHAAP has been under the administrative control of the Base Realignment and Closure (BRAC) Division of the Army since 2003 and is Defense Environmental Restoration Account funded. The majority of LHAAP has been transferred by the U.S. Army to the U.S. Fish and Wildlife Service (USFWS) for management as the Caddo Lake National Wildlife Refuge (Bhate 2018a).

LHAAP-03 is located approximately 50 feet to the west of former Building 722-P (**Figure 1-2**). Building 722-P was used for paint spraying and polyurethane spray coating of various items. LHAAP-03 was a waste collection site (originally identified as a 16-foot by 15-foot-area) outside of the paint shop at Building 722-P, which was at the Maintenance Shop Area located within the boundary of LHAAP-35A (58) (see **Figures 1-2**).

The ROD identified two chemicals of concern (COCs) for human health in soil, as shown in **Table 1-1**. No chemicals of potential ecological concern were identified. The remedy selected in the ROD included excavation and off-site disposal of soil. The human health cleanup levels were set at concentrations equal to the largest of the calculated TCEQ Risk Reduction Standard 3 (RRS3) industrial soil medium-specific concentrations (MSCs) based on the site-specific soil standard for groundwater protection, and background soil concentration (AECOM 2013a). These cleanup levels are shown in **Table 1-1**.

Table 1-1
Chemicals of Concern and Cleanup Levels for LHAAP-03

Media	Chemicals of Concern (Human Health)	Cleanup Levels ^a
Soil	Arsenic Lead	5.9 mg/kg 180 mg/kg

Notes:

^a Applicable RRS3 Industrial Soil MSC equals largest of the calculated RRS3 MSC and background values. *Italicized text indicates the cleanup level equals background*

mg/kg – milligrams per kilogram

1.2 Selected Remedy

The final remedy is identified in the Proposed Plan (AECOM 2013b) that has been reviewed and approved by the regulatory agencies and placed in the Administrative Record file for LHAAP. The Final LHAAP-03 ROD (Bhate 2018a) signed by the Army and USEPA selected excavation and offsite disposal as the remedy for arsenic and lead contaminated soil at LHAAP-03 as summarized in Section 1.4 of the Final ROD. Following remedial action, COC concentrations will be reduced to achieve chemical-specific applicable or relevant and appropriate requirements (ARARs) (based on the commercial/industrial land use scenario). The remaining arsenic and lead concentrations in soil will not pose an unacceptable direct contact risk to humans or ecological receptors at this site. The selected remedy for LHAAP-03 protects groundwater by preventing migration of COCs into groundwater at concentrations that could possibly result in local exceedances of the maximum contaminant levels (MCLs) for arsenic and lead.

The target soil remediation area for LHAAP-03 is contained entirely within the boundaries of a larger site, LHAAP-35A (58). Because LHAAP-03 is small and entirely contained within LHAAP-35A (58) and its land use control (LUC) boundary, the management strategy is to address the LHAAP-03 groundwater remedy and LUCs as being indistinguishable from and included with the LHAAP-35A (58) remedy. This was approved in the Explanation of Significant Differences for the LHAAP-35A (58) ROD (Bhate 2018b).

The final remedy addresses arsenic- and lead-contaminated soil that, based upon testing and modeling, have the potential to result in local groundwater arsenic and lead concentrations exceeding their respective MCLs. The soil cleanup levels selected are protective of the groundwater for arsenic and lead. Surface water runoff to surface water streams is not shown as a pathway for the site, and there is no indication of concentrations at the surface that present an unacceptable risk to human or ecological receptors that could create a risk via that pathway (Bhate 2018a). To address ecological risk, LHAAP-03 was grouped with several other sites as part of the Industrial Sub-Area. The Baseline Ecological Risk Assessment (BERA) (Shaw 2007b) concluded that no unacceptable risk was present in the Industrial Sub-Area; therefore, no further action is needed at LHAAP-03 for the protection of ecological receptors (Bhate 2018a).

Because LHAAP-03 lies wholly within the LUC boundary of LHAAP-35A (58), all LUCs, as well as the non-residential land use notification, applicable to LHAAP-35A (58) will be applicable to LHAAP-03 in the same way and to the same extent (Shaw 2010 & Bhate 2018b). Therefore, the remedy selected for LHAAP-03 does not include any specific provisions for LUCs (Bhate 2018a).

CERCLA Five-Year Reviews specific to LHAAP-03 will be implemented following completion of the soil remedy to evaluate whether the remedy remains protective of human health and the environment. CERCLA Five-Year Reviews for LHAAP-03 will be addressed as part of the remedial action for LHAAP-35A (58). All monitoring and reporting requirements associated with CERCLA five year reviews, will be met under LHAAP-35A (58) (Bhate 2018b).

The ROD identified three areas to be excavated based on previous soil sample exceedances from 2006 through 2008 as the conceptual target remediation areas. The ROD also notes that the excavation extents would be refined as part of the pre-excavation sampling. These areas are described below:

- **Area A.** Approximately 25-feet-wide by 25- to 35-feet-long (approximately 560 square feet [SF]) by 2-feet-deep
- **Area B.** Approximately 5-feet by 5-feet-around 03SB15 (within Area A footprint) by at least 7-feet-deep (bottom excavation depth is not defined)
- **Area C.** Approximately 5-feet by 5-feet-around 03SB11 (within Area A footprint) by 7-feet-deep

1.3 Remedial Action Objectives

The remedial action objective (RAO) developed for LHAAP-03 and outlined in the Final ROD (Bhate 2018a) is to protect human health and the environment by preventing lead and arsenic contaminated soils from potentially leaching into the underlying groundwater. Per the ROD's RAOs, and consistent with the National Contingency Plan, COCs and cleanup levels must be set. These are shown in **Table 1-1**.

1.4 Document Organization

This document is composed of the following sections:

- **Section 1.0:** “Introduction” summarizes the site background, proposed remedy, and RAOs.
- **Section 2.0:** “Site Characteristics” summarizes the geology and hydrogeology of the site, as well as a summary of the nature and extent of contamination.
- **Section 3.0:** “Soil Sampling through November 2018” summarizes the results of the prior samples as well as the samples collected in November 2018 as part of the pre-excavation sampling as described in the ROD that form the basis for the design elements described in subsequent sections.
- **Section 4.0:** “Soil Excavation Remedial Design” describes the design basis for the planned excavation of contaminated soil.
- **Section 5.0:** “Remedial Action Work Plan” describes the tasks to implement the design for soil excavation.
- **Section 6.0:** “Schedule” provides a list of activities and anticipated durations for the work plan tasks.
- **Section 7.0:** “References” provides citations for the documents used as references.

This work plan also includes the following appendices supporting the main text:

- **Appendix A** includes the Analytical Data Reports for the November 2018 soil samples collected.
- **Appendix B** includes the Sample Collection Logs from the November 2018 soil sample collection event.
- **Appendix C** includes the State of Texas Well Completion Report for monitoring well 03WW01.

1.5 Deviations from the Installation-Wide Work Plan

There is a planned deviation from the Final Installation-Wide Work Plan (IWWP) (Bhate 2018c), Section 3.8.3, Verification of Excavation Limits. The IWWP indicates that verifications samples shall be collected from every 1,000 SF of excavation floor and from each wall. The largest planned excavation for LHAAP-03 is 300 SF, and the smallest is 36 SF. Because of the small excavation areas and shallow depths, the resulting exposed surface area is expected to be less than 400 SF. Therefore, it is proposed to verify the excavation limits both laterally and vertically with one composite confirmation sample. Each wall and floor will not be individually tested. See **Section 4.4** of this document for more details of the planned confirmation sampling.

Contract No. W9128F-13-D-0012, Task Order No. W9128BV17F0150 • Draft Final • Rev 0 • June 2019

REMEDIAL DESIGN AND REMEDIAL ACTION WORK PLAN, LHAAP-03 FORMER WASTE COLLECTION PAD, BUILDING 722-P PAINT SHOP



Contract No. W9128F-13-D-0012, Task Order No. W9128BV17F0150 • Draft Final • Rev 0 • June 2019

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2.0 SITE CHARACTERISTICS

2.1 Geology and Hydrogeology

The subsurface geology at LHAAP consists primarily of a thin veneer of Quaternary alluvium overlying Tertiary age formations of the Wilcox and Midway Groups. The Wilcox Group underlies most of the LHAAP installation (Jacobs 2001). The thickness of the Wilcox Group varies from 350 feet in the northwest corner of the installation to 130 feet along the eastern side near Caddo Lake. This formation consists of interbedded fine-to medium-grained sand, silt, and clay. The shallow soils at LHAAP-03 include interbedded layers of silty clays and sands (Shaw 2007a).

2.2 Nature and Extent of Contamination

From 2006 through 2008, soil samples were collected at LHAAP-03 soil operable unit at depths ranging from surface (0 to 0.5 feet below ground surface [ft bgs]) to 15 ft bgs. The collected samples were analyzed for metals, volatile organic compounds (VOCs), and semivolatile organic compounds (SVOCs). The analytical results were compared to TCEQ Risk Reduction Standard 2 (RRS2) and RRS3 MSCs (Title 30 Texas Administrative Code [TAC] Chapter 335) corresponding to commercial/industrial land-use. The RRS2 and RRS3 MSCs are risk-based values developed to protect human health and groundwater resources that are protective at a target risk level of 1×10^{-6} for carcinogens and a target hazard quotient of 1 for non-carcinogens, as described in the *Final Remedial Investigation – Focused Feasibility Study* (AECOM 2013a).

Arsenic and lead were detected in one or more samples from borings 03SB01 through 03SB15, and 03SB17 at concentrations exceeding their respective RRS2 MSCs. These data indicated that there is a potential for the metals-contaminated soil to contaminate the groundwater (Shaw 2009). RRS3 MSCs were developed using the soil attenuation model, according to the TCEQ Risk Reduction Rules (30 TAC §335 and updates), included in the Remedial Investigation/Focused Feasibility Study (RI/FFS), Appendix B (AECOM 2013a).

The calculated 95 percent upper confidence limits (UCLs) of the mean concentrations in soil for arsenic and lead were compared to their respective calculated RRS3 MSCs. This comparison indicated that arsenic and lead exceeded their respective applicable RRS3 industrial soil MSCs and they were identified as COCs in soil. **Section 3.0** discusses the sample results and location in more detail.

2.3 Current and Future Land Use

LHAAP is located near the unincorporated community of Karnack, Texas. Karnack is a rural community with a population of approximately 775 people. The incorporated community of Uncertain, Texas, approximate population 205, is located to the northeast of LHAAP on the edge of Caddo Lake and is a resort area and an access point to Caddo Lake. The industries in the surrounding area consist of agriculture, timber, oil and natural gas production, and recreation.

LHAAP has been an industrial facility since 1942. Production activities and associated waste management activities continued until the facility was determined to be in excess of the U.S. Army's needs in 1997. The plant area has been relatively dormant since that time. LHAAP is surrounded by a fence (except on the border with Caddo Lake) with an access gate that is locked after daylight hours, which restricts public access. The fence now represents the National Wildlife Refuge boundary. The public can access most of the facility during the day, with additional fencing and signage restricting access from some environmental sites.

The reasonably anticipated future use of LHAAP-03 is part of a national wildlife refuge. This anticipated future use is based on a Memorandum of Agreement (MOA) (U.S. Army 2004) between the USFWS and the U.S. Army. That MOA documents the transfer process of the LHAAP acreage to USFWS to become the Caddo Lake National Wildlife Refuge and will be used to facilitate a future transfer of LHAAP-03. Presently the Caddo Lake National Wildlife Refuge occupies approximately 7,100 acres of the 8,416-acre former installation. In accordance with the National Wildlife Refuge System Administration Act of 1966 and its amendments (16 USC 668dd), the land will remain as a national wildlife refuge unless there is a change brought about by an act of Congress, or the land is part of an exchange authorized by the Secretary of the Interior. Neither of these events terminating LHAAP-03's use as a wildlife refuge are reasonably foreseeable (Bhate 2018a).

2.4 Current and Future Surface Water Use

There is no surface water body present within LHAAP-03. Land at LHAAP-03 is relatively flat; water from heavy rains would eventually drain to the southern branch of Goose Prairie Creek, approximately 800 feet to the southwest. Goose Prairie Creek flows into Caddo Lake. Caddo Lake is a large recreational area that covers 51 square miles and has a mean depth of 6 feet. The watershed of the lake encompasses approximately 2,700 square miles. It is used extensively for fishing and boating and provides drinking water supply for multiple cities/towns. The anticipated future uses of surface water are the same as current uses.

2.5 Current and Future Groundwater Use

Groundwater in the drinking water aquifer (250 to 430 ft bgs) near LHAAP is currently used as a drinking water source. The drinking water aquifer should not be confused with the “deep zone” groundwater, which extends only to a depth of approximately 151 ft bgs. The deep zone groundwater and the drinking water aquifer are distinct from each other, and there is no connectivity between the contaminated zone and the drinking water aquifer. There are six active water supply wells near LHAAP that are completed in the drinking water aquifer (**Figure 1-2**).

Groundwater under LHAAP-03 is indistinguishable from and is included with the site wide LHAAP-35A (58) groundwater (Bhate 2018a).



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3.0 PRE-EXCAVATION SOIL SAMPLING

Pre-excavation sampling was conducted prior to preparation of the RD/RAWP to better define the excavation limits and to pre-characterize the waste for offsite disposal of the soil. The intent was to define the limits of the excavation based on the samples with concentrations below the cleanup levels and to use the results as confirmation samples. The sampling approach was outlined in the *Technical Memorandum – Pre-Excavation Sampling at LHAAP-03 Former Waste Collection Pad Building, 722-P Paint Shop, Longhorn Army Ammunition Plant, Karnack, Texas* (Bhate 2018d) and was approved by the regulators in October 2018. Soil samples were collected on November 29, 2018 at pre-determined locations and intervals as described in the Pre-Excavation Sampling Technical Memorandum (Bhate 2018d). Analytical results from the November 2018 pre-excavation sampling were all below the RRS3 MSCs. Because of the low concentrations, a pre-excavation waste characterization sample analysis was not performed. The analytical data from the November 2018 sampling event is provided in **Table 3-1** and shown on **Figure 3-1**. Analytical data reports are included as **Appendix A**, and Sample Collection Logs are provided as **Appendix B**.

The data from November 2018 was used along with the prior data from 2006 through 2008 to re-evaluate the excavation areas shown in the ROD. The 2006 through 2008 and the ROD excavation areas are shown on **Figure 3-1**. Results that exceeded the RRS3 MSC are yellow highlighted on **Figure 3-1**. The 2006 through 2008 sampling had been conducted at seventeen locations. The area sampled from 2006 through 2008 covered approximately 2,100 SF (or less than 1/10th of an acre) and identified an area of approximately 560 SF for remediation (Bhate 2018a). Samples were collected from 0 to 0.5 ft bgs, 3 to 4 ft bgs, and 5 to 7 ft bgs at each location except at 03SB11, which also had deeper intervals collected at 9 to 10 ft bgs and 14 to 15 ft bgs. At several of the locations, arsenic and lead concentrations were below the RRS3 MSCs in all sampled intervals. The maximum detected concentrations of arsenic and lead in soil were 32.7 and 6,760 milligrams per kilogram (mg/kg), respectively, in boring 03SB11 at the 6 to 7 ft bgs interval during the 2006 sampling event. The deeper interval samples were below the RRS3 MSC.

A groundwater monitoring well (03WW01) was installed at location 03SB11, where deeper intervals were sampled. Concentrations at depths below the 0 to 0.5 ft bgs interval exceeded the RRS3 MSC at only one other location, 03SB15, where just arsenic was detected above the RRS3 MSC. At 03SB15, the arsenic concentration at 6 to 7 ft bgs was 7.62 mg/kg, just above the RRS3 MSC of 5.9 mg/kg. Data from the November 2018 sampling did not indicate any other locations with deeper contamination. **Figure 3-1** shows the November 2018 and the prior 2006 through 2008 sample results.



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4.0 SOIL EXCAVATION REMEDIAL DESIGN

The design elements of the excavation are provided in the following sections. The details of the implementation of the design are included in **Section 7.0**.

4.1 Design Criteria

The soil design criteria define the goals to be achieved and are based on the RAOs of protection of human and ecological receptors from contaminated soil and the protection of human health by preventing further potential degradation of groundwater. The ROD identified COCs for the soil to groundwater pathway as shown in **Table 1-1**. The design for soil includes excavation of the contaminated soil until the cleanup levels are attained or groundwater is encountered. If the contamination is present at or below groundwater level, excavation will be terminated at the depth where saturated conditions are encountered.

4.2 Performance Objective

The performance objective is to remove the contaminated soil above the RRS3 MSCs. For areas that are not already defined by existing samples, confirmation samples will be collected and analyzed for the COCs after excavation to verify the contaminated soils were removed and cleanup levels were attained. Soil concentrations in few surface soil samples and a couple of subsurface samples exceeded the RRS3 MSC. The surface area locations (0 to 2 ft bgs) are shown on **Figure 4-1** and the subsurface locations (6 to 7 ft bgs) are shown on **Figure 4-2**.

4.3 Excavation Areas

The proposed excavation areas to address the soil to groundwater pathway are shown on **Figures 4-1** and **4-2**. The excavation depths anticipated to remove the contaminated soil at various locations within the footprint of LHAAP-03 are 1 or 8 ft bgs. The groundwater elevation typically is approximately 25 ft bgs and is not expected to be encountered during excavation.

4.3.1 Excavation Area Sample Summary

Excavation Area A (Areas A1 and A2)

Review of all results indicate that the majority of the contamination is in the 0 to 0.5 foot bgs interval with concentrations of lead and arsenic above the RRS3. The individual sample locations with lead and arsenic concentrations exceeding their respective RRS3 MSC values are shown with yellow highlighting on **Figure 3-1**. Additional samples were collected in November 2018 along the perimeter and within the target remedial Area A described in the ROD (**Figure 4-1**). The results from the 2018 and historic samples were used to define the lateral extent of the excavations.

For Area A1, there were 9 sample locations within 4 feet of the Area A1 excavation perimeter that were below the RRS3 MSC and are proposed to be used to indicate that the 0 to 1 foot bgs excavation side walls are clean. The excavation floor of Area A1 will have a confirmation sample collected.

For Area A2, the excavation is small in both width and depth (~140 cubic feet). Because the excavation area is so small, only one composite will be collected for the entire excavation. A composite sample consisting of a grab from each sidewall and the floor will be used for the confirmation sample for this 1-foot excavation area. This is a deviation from the IWWP sampling protocol as described in **Section 1.5**.

Excavation Areas B and C

Two soil borings, 03SB15 and 03SB11, with samples collected at deeper intervals from 2006 through 2008 had prior results above the RRS3 MSC. The ROD identified remediation to be conducted around these two areas with refinements based on the pre-excavation sampling (Bhate 2018a). The November 2018 samples collected from 0 through 9 ft bgs around these two areas confirm that these were isolated occurrences.

For Area B, a sample collected from 03SB15 in 2007 had arsenic slightly above the RRS3 MSC at only the 6 to 7 ft bgs interval. Three samples were collected in November 2018 at distances from 4 to 7 feet away from the 03SB15 location at intervals from 0 to 2 ft bgs, 3 to 4 ft bgs, 6 to 7 ft bgs, and one from 8 to 9 ft bgs. **Figure 4-2** shows the proximity of the results that make the reconfigured triangular Area B. None of the arsenic or lead results from the November 2018 samples had results above the RRS3 MSC. The triangular area shown on **Figure 4-2** will be excavated to a depth of 8 feet, and the three sample results defining the corners of the triangle will be used to define the horizontal limits of the excavation and a confirmation sample will be collected from the floor of the excavation.

For Area C, 03SB11 collected in 2007 had the highest concentrations of arsenic and lead detected at LHAAP-03 at the 6- to 7-foot-interval. In 2008, during the installation of 03WW01 adjacent to 03SB11, additional soil samples were collected at 8 to 9 ft bgs and 14 to 15 ft bgs and did not have any RRS3 MSC exceedances. In November 2018, samples were collected approximately 4 feet to the northwest and southeast of 03SB11 at 0 to 2, 3 to 4, and 6 to 7 ft bgs intervals (**Figure 4-2**). None of the 2018 sample results were above the RRS3 MSCs. Because the two samples were so close to 03SB11, the excavation at Area C will be approximately 8-feet by 8-feet-horizontally. The results from the 2008 sample from 8 to 9 ft bgs are being used to define the vertical limit of the excavation to 8 ft bgs and will be used as the floor confirmation sample. The two 2018 sample results from 6 to 7 ft bgs will also be used as the confirmation samples for the side walls.

Because the excavation is so small, an effort will be made to keep the monitoring well, 03WW01, in place. The screened interval of the well is below 18 ft bgs and the annular space was sealed with cement from 0 to 16 ft bgs. The 03WW01 State of Texas Well Completion Report is included in **Appendix C**.

4.3.2 Excavation Depth and Extent

Approximately 40 in-place cubic yards (CY) of soil will be excavated from Areas A, B, and C. The excavation areas proposed after the pre-excavation sampling are as follows:

- **Area A.** This area was split into two subareas from the ROD Area A—Area A1 and Area A2 as shown on **Figure 4-1**.
 - For Area A1, the excavation area was developed by surrounding the locations that were above the RRS3 MSC at 0 to 0.5 ft bgs and extending the horizontal limit to within 5 feet of sample locations that had results below the RRS3 MSCs at 0 to 0.5 ft bgs or 0 to 2 ft bgs. An initial excavation depth of 1 foot is planned since none of the 0 to 2 ft bgs November 2018 samples were above the RRS3 MSC.
 - For Area A2, the contamination appears to be isolated at 0 to 0.5 ft bgs, and a small excavation is planned around it. The estimated horizontal area is depicted on **Figure 4-1**, and the initial vertical excavation depth is 1 foot bgs.
- **Area B.** This area was defined in the ROD as a 5-foot-wide by 5-foot-deep by 7-foot-deep excavation. The area has been redefined by the sample results to a triangular area, as shown on **Figure 4-2**. The area will be excavated to a depth of 8 feet.
- **Area C.** This area has remained unchanged from the ROD and will be excavated as an 8-foot by 8-foot-square excavation approximately 8-foot-deep centered around monitoring well 03WW01. Because the excavation is small, and the vertical limit is known, an effort will be made to save monitoring well 03WW01 and not abandon it initially. The grout seal around the well extends down to 16 feet bgs and should remain intact and protected with careful excavation around the well. However, if it is damaged during excavation, the well will be plugged and abandoned and reinstalled by a Texas licensed well driller.

4.4 Confirmation Sampling

Confirmation samples will be collected to define the final excavation limits. If contamination is present in a confirmation sample, the excavation limits will be extended in increments of 1 foot vertically and horizontally until confirmation samples are below the RRS3 MSC.

The IWWP (Bhate 2018c) indicates that each confirmation sample will be a 5-point composite soil sample collected from every 1,000 SF of the excavation floor area and of each sidewall. However, a deviation from the 5-point composite methodology is proposed to allow the use of existing soil samples to define the excavation limits. Because of the small excavation areas and the abundance of samples with results below the RRS3 MSC in the small area at LHAAP-03, the use of existing samples and a deviation to the IWWP has been proposed for confirmation sampling. For excavations where the limits were set by existing samples below the RRS3 MSC, no additional confirmation samples will be collected. For small excavations, a single composite sample representing the walls and floor will be collected. **Figure 4-3** shows the existing sample locations and intervals to be used for confirmation samples for the 1-foot excavation areas, and **Figure 4-4** shows the existing locations to be used for the Areas B and C excavations. **Table 4-1** summarizes the samples (existing and proposed) to be used for excavation confirmation.

The following proposed sampling procedure for small excavation areas (less than 200 SF of floor area and sidewall height of less than 10 feet) is a deviation from the IWWP. It is proposed to collect a single 5-point composite sample from such excavations by collecting a grab sample from each of the four sidewalls and the floor. This alternate sampling will be conducted at Area A2. If the composite sample result is above the cleanup levels, then additional grab samples may be collected from each wall to determine which direction to overexcavate. If no additional sampling is conducted prior to over excavation, each wall and the floor will be overexcavated by 1 foot. The Area A1 excavation floor sampling will be performed in accordance with the IWWP.

4.5 Waste Characterization and Disposal

The excavated soil will be disposed at an off-site landfill. Waste characterization samples will be collected at a rate of one sample for every 1,000 CY of excavated soil to characterize the waste. It is anticipated that soil will be classified as non-hazardous. Samples will be analyzed for the analytes required by the selected waste disposal facility.

4.6 Backfill and Site Restoration

Following the receipt of clean confirmation samples for a given excavation area, clean fill dirt will be placed in the excavation and compacted with the backhoe/excavator bucket to prevent settling. The fill dirt will be suitable for future vegetation growth. Borrow source material will be considered clean if VOCs and metals are below the TCEQ RRS2 MSCs for industrial use. It is estimated that less than 100 CY of fill dirt will be brought on site. Borrow source samples will be collected at a rate of one per 1,000 CY of borrow soil. The site will be reseeded with a native grass and wildflower mix with guidance from USFWS.

4.7 Other Design Considerations

Several of the bridges at LHAAP have weight limits. Routes for trucks filled with waste soil as well as for trucks bringing clean soil from borrow sources will be selected with coordination from USFWS to avoid the bridges if possible. Additional gates may need to be unlocked to allow access to and from LHAAP from another gate besides the main gate in Karnack to avoid bridges where weight limits may be exceeded with loaded trucks.



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5.0 LAND USE CONTROL PLAN

LHAAP-03 lies wholly within the LUC boundary of LHAAP-35A (58) (see **Figure 1-2**). All LUCs, as well as the non-residential notification, applicable to LHAAP-35A (58) will be applicable to LHAAP-03 in the same way and to the same extent. Therefore, the remedy selected in the ROD does not include specific provisions for LUCs (Bhate 2018a).



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6.0 REMEDIAL ACTION WORK PLAN

6.1 Field Activities

This section describes the remedial action field activities planned at the LHAAP-03. Prior to initiation of the field activities performed as part of this RD/RAWP, the regulators will be notified at least 10 days in advance. Site-specific activities are described in associated subsections. The field activities to be conducted under this Work Plan are outlined below:

- Mobilization and Site Setup
- Surveying
- Soil Excavation and Disposal
- Confirmation Soil Sampling
- Backfilling and Site Restoration
- Waste Management
- Decontamination
- Site Restoration

In general, the field activities will be conducted in accordance with the IWWP for LHAAP (Bhate 2018c).

6.1.1 Mobilization and Site Setup

Prior to the mobilization of subcontractors to LHAAP sites, work locations for overhead and ground level accessibility will be evaluated. In areas that have excessive vegetation and/or tree growth, a backhoe or other appropriate earthmoving equipment will be used to clear the areas to allow equipment access. After coordinating with underground utility locators for utility clearances, excavation locations and areas that require surface soil removal will be located and staked. Utility location and clearance for intrusive activities will be conducted in accordance with Section 3.1 of the IWWP (Bhate 2018c).

Appropriate personnel, subcontractors, and equipment necessary to perform specific task(s) will be mobilized to the site. A permanent decontamination station is located at the on-site LHAAP-18/24 groundwater treatment plant (GWTP) and can accommodate large equipment. Temporary decontamination pads will be constructed as needed at approved on-site locations to decontaminate equipment and prevent cross-contamination between sites. Wash water will be contained and transported to the GWTP for disposal when necessary.

Improvement of the site access road using gravel or other materials to reduce the slopes may be performed to allow trucks and heavy equipment to enter and exit the site safely. USFWS will be consulted regarding any improvements to minimize impact to the surrounding terrain to the degree possible.

6.1.2 Surveying

During the excavation, the locations of soil confirmation samples and limits of excavation will be surveyed using global positioning system (GPS) equipment. Use of GPS equipment will be coordinated with USACE in accordance with Section 3.3 of the IWWP (Bhate 2018c). A professional land surveyor licensed in the State of Texas will survey the final horizontal excavation limits. After backfilling is complete, the new vertical elevation of the top of casing for 03WW01 (or 03WW01R if replaced) will be surveyed based on the North American Vertical Datum of 1988 to the nearest 0.01 foot.

6.1.3 Soil Excavation and Disposal

The limits and plans for excavation are described in **Section 4.0** of this Plan. Excavation of the soil generally consists of preparing the site, excavating the soil, transporting and disposing the soil, collecting confirmation samples, surveying the excavation limits, backfilling, and restoring the site. Excavations performed under this remedial action are expected to be less than one acre in size and are not subject to the storm water pollution prevention requirements of Chapter 26 of the Texas Water Code and TCEQ General Permit TXR15000. However, best practices (silt fencing, berms, etc.) will be used where appropriate to prevent excess runoff of sediment from the excavation and stockpile areas.

6.1.3.1 Site Preparation

The areas to initially be excavated will be established prior to mobilization of the excavation personnel. A GPS will be used to delineate and mark the excavation areas shown on **Figures 4-1** and **4-2**. The potential limits of excavation will then be physically marked with survey stakes, pin flags, paint, or other appropriate marking. Clearing of the vegetation in the excavation area will largely be conducted using conventional equipment. A temporary decontamination station may be constructed on site as needed.

An area between the site and the existing roads to the north and east will be cleared for construction of a temporary soil staging/stock pile area. This will allow the transport vehicles to stay on the road during loading activities. A temporary staging area consisting of berm with polyethylene sheeting will be constructed for stockpiling soil prior to collecting waste characterization samples and off-site transport and disposal.

6.1.3.2 Excavation and Soil Handling

After the initial excavation limits are established, excavation will begin. Vertical excavation will stop if groundwater is encountered. Areas B and C with an excavation depth of more than 4 feet will not be benched and no personnel will enter these excavations. All activities will be conducted from the surface. See **Section 6.2** below for monitoring well excavation or abandonment and reinstallation for the well located within excavation Area C.

The excavated soil will be stockpiled in the designated area. Composite samples will be collected from the stockpiled material at a rate of 1 sample per 1,000 CY and analyzed for the necessary analytes to obtain acceptance at the selected disposal facility. Each composite sample will consist of equal parts of five samples collected at evenly spaced locations within the stockpile.

The fully-characterized excavation stockpile soil will be placed into transport trailers or dump trucks for immediate transport from the site to the disposal facility. Licensed transporters will be used to haul the excavated soil to the pre-approved landfill for non-hazardous disposal. The excavated soil may be staged on plastic sheeting adjacent to the excavation while awaiting loading. The excavated soil stockpile will be protected from rainfall runoff and erosion by covering it with plastic sheeting. As an alternative to stockpiles, roll-off boxes may be supplied by the landfill and used in place of the stockpiles.

Loading of trucks will be coordinated with USFWS to ensure that load ratings for bridges along the haul route out of the refuge (if any) are not exceeded.

6.1.3.3 Confirmation Soil Sampling

After the initial excavation, confirmation samples will be collected from the sidewalls and floor of the excavation in the areas where existing samples are not available for use as confirmation samples. The confirmation samples will be tested for the contaminants. Excavation will continue until concentrations in the soil are less than the site-specific cleanup levels.

As described in **Section 4.4**, in small areas a composite wall/floor sample may deviate from the IWWP and will be collected by combining discrete samples collected from each of the four walls and the floor. GPS coordinates of each discrete sample location that comprises the composite confirmation sample will be collected. Vertical wall height will be manually measured and recorded. Each sample location will be numbered sequentially in order of collection, labelled on a map, and identified using the following nomenclature:

03WFXXX-ZZ-MMDDYY or 03FLXXX-ZZ-MMDDYY

The number 03 represents the site (LHAAP-03); WF indicates that it is small excavation with composite of both the wall and floor, WL indicates a sidewall sample, while FL indicates a floor sample; XXX represents the unique sample number; ZZ indicates excavation sidewall height or the average depth below ground surface of the excavation floor; and MMDDYY is the date of sample collection.

If contaminants are detected above their cleanup levels during the confirmation sampling, the area will be over-excavated. In the case of the small excavation where a wall and floor composite sample is collected, the excavation will be overexcavated approximately one foot deeper and sideways on all walls. This will continue until confirmation samples demonstrate the contaminants remaining in the soil are below their cleanup level or until groundwater is encountered.

Even though not expected, in the event that groundwater is encountered, and a floor sample cannot be collected, a linear 5-point composite sample will be collected from each excavation sidewall just above the groundwater interface to represent the floor area above the groundwater. If the linear 5-point composite sidewall sample is above the cleanup level, then additional excavation of the sidewall will be conducted to the groundwater interface depth, and over-excavation step outs and confirmation sampling would continue until the confirmation sample results are below the cleanup levels. The confirmation samples will confirm that the vadose zone soil identified as exceeding the cleanup levels would be removed.

6.1.3.4 Backfilling and Site Restoration

The backfill operations will proceed after excavation confirmation samples are clean for a definable area. For Area C, the excavations will be immediately backfilled to maintain the structural integrity of monitoring well 03WW01. The excavation areas will be backfilled with fill material that is certified to meet the site-specific cleanup levels. The backfill will be placed in 1-foot lifts to allow proper compaction with a backhoe/excavator bucket. After backfilling is complete, the area will be graded, with a mound approximately one foot above finished grade to promote positive drainage and allow for some soil settling without creating a depression. The top six inches will have a soil that will be suitable for vegetative growth. The well pad for 03WW01 (or 03WW01R if replaced), the protective casing with lock, and pipe bollards will be installed. The well identification will be clearly marked. The surface will be reseeded with a native grass and wildflower seed mixture selected in coordination with USFWS.

6.2 Monitoring Well Excavation or Abandonment and Installation

Monitoring well 03WW01 is in the 8-foot depth excavation area. The well pad will be broken up and removed along with the pipe bollards prior to initiating excavation. The protective well

casing will also be removed. The grout plug above the bentonite seal will remain in place. The excavation will extend down to 8 feet taking care to not impact the cement around the pipe. Supports may be placed around the well stickup during excavation if needed for support. A portion of the area will be excavated to 8 feet and will be immediately backfilled. Hand tools may be used to scrape any dirt from the cement near the 6 to 7 ft bgs area. A physical barrier (such as polyethylene sheeting) will be placed against the cut wall of the excavation prior to backfilling. The next portion of the excavation will remove the soil up to the barrier and will remove the physical barrier ensuring that contaminated soil was removed from Area C. Hand compaction may be conducted around the well taking care to not damage the well.

If the structural integrity of the well is damaged, the well will be abandoned by pulling out the well casing (or overdrilling if the casing cannot be removed) prior to grouting the borehole. If this fails, the well casing will be grouted in place and the portion extending into the excavation will be cut at least 2 ft bgs. Once the excavation is backfilled, a new well, 03WW01R, will be installed adjacent to the location and screened with the same interval to replace 03WW01. If needed, the well abandonment will be conducted in accordance with Section 3.9 of the IWWP, and the shallow well installation with stick up completion and its development will be completed in accordance with Section 3.2 of the IWWP (Bhate 2018c). The State of Texas Well Completion Report for 03WW01 is included in **Appendix C**.

6.3 Waste Management

Wastes generated during the project are anticipated to include waste water, soil, personal protective equipment (PPE), sampling equipment, and miscellaneous trash. These wastes will be managed in accordance with Section 3.7 of the IWWP (Bhate 2018c).

6.4 Decontamination

Decontamination of equipment will be performed in accordance with Standard Operating Procedure (SOP) A1 in Appendix A of the IWWP (Bhate 2018c).



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7.0 POST-REMEDIAL MONITORING AND REPORTING

7.1 Remedial Action Completion Report

A Remedial Action Completion Report will be submitted upon completion of the excavation, confirmation sampling, backfilling, and waste disposal to document the activities performed to complete the remedial action.

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8.0 SCHEDULE

Table 8-1 shows the estimated duration for each major site activity and timeline. Weather and unknown site conditions could affect this schedule.

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Contract No. W9128F-13-D-0012, Task Order No. W9128BV17F0150 • Draft Final • Rev 0 • June 2019

REMEDIAL DESIGN AND REMEDIAL ACTION WORK PLAN, LHAAP-03 FORMER WASTE COLLECTION PAD, BUILDING 722-P PAINT SHOP

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Tables

Table 3-1
November 2018 Soil Sample Analytical Results, LHAAP-03

			03SB18		03SB19				03SB20							
			03SB18-0.0-2.0		03SB19-0.0-2.0		03SB19-3.0-4.0		03SB19-6.0-7.0		03SB20-0.0-2.0		03SB20-0.0-2.0-FD		03SB20-3.0-4.0	
			11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018	
			0 - 2		0 - 2		3 - 4		6 - 7		0 - 2		0 - 2		3 - 4	
			REG		REG		REG		REG		REG		FD		REG	
Parameter	Units	RRS3 MSCs	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
METALS																
Arsenic	mg/kg	5.9	1.56		2.63		1.14		2.99		1.54		4.05		3.9	
Lead	mg/kg	180	6.15		9.52		7.89		5.71		8.26		11.3		16.3	

Notes:

FD - field duplicate

ft bgs - feet below ground surface

ID - identification

J - estimated value

mg/kg - milligrams per kilogram

RRS3 MSCs - Risk Reduction Standard 3 Medium-Specific Concentrations

REG - regular sample

Val Qual - validation qualifier

Table 3-1
November 2018 Soil Sample Analytical Results, LHAAP-03

Location Code			03SB21				03SB22						03SB23					
Sample ID			03SB21-0.0-2.0		03SB21-3.0-4.0		03SB22-0.0-2.0		03SB22-3.0-4.0		03SB22-3.0-4.0-FD		03SB22-6.0-7.0		03SB23-0.0-2.0		03SB23-3.0-4.0	
Sample Date			11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018	
Sample Interval Depth (ft bgs)			0 - 2		3 - 4		0 - 2		3 - 4		3 - 4		6 - 7		0 - 2		3 - 4	
Sample Purpose			REG		REG		REG		REG		FD		REG		REG		REG	
Parameter	Units	RRS3 MSCs	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
METALS																		
Arsenic	mg/kg	5.9	1.74	J	0.974		1.79		4.61		0.709		2.63		1.93		4.16	
Lead	mg/kg	180	10.2		4.97		9.07		8.66		6.26		7.62		12.3		7.12	

Notes:

FD - field duplicate

ft bgs - feet below ground surface

ID - identification

J - estimated value

mg/kg - milligrams per kilogram

RRS3 MSCs - Risk Reduction Standard 3 Medium-Specific Concentrations

REG - regular sample

Val Qual - validation qualifier

Table 3-1
November 2018 Soil Sample Analytical Results, LHAAP-03

			03SB24		03SB25		03SB26							
			03SB24-0.0-2.0		03SB25-0.0-2.0		03SB26-0.0-2.0		03SB26-0.0-2.0-FD		03SB26-3.0-4.0		03SB26-6.0-7.0	
			11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018	
			0 - 2		0 - 2		0 - 2		0 - 2		3 - 4		6 - 7	
			REG		REG		REG		FD		REG		REG	
Parameter	Units	RRS3 MSCs	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
METALS														
Arsenic	mg/kg	5.9	1.66		1.42		5.83		1.92		0.74		2.94	
Lead	mg/kg	180	14.3		8.02		8.6		8.38		4.14		6.96	

Notes:

FD - field duplicate

ft bgs - feet below ground surface

ID - identification

J - estimated value

mg/kg - milligrams per kilogram

RRS3 MSCs - Risk Reduction Standard 3 Medium-Specific Concentrations

REG - regular sample

Val Qual - validation qualifier

Table 3-1
November 2018 Soil Sample Analytical Results, LHAAP-03

			03SB27								03SB28					
			03SB27-0.0-2.0		03SB27-3.0-4.0		03SB27-6.0-7.0		03SB27-8.0-9.0		03SB28-0.0-2.0		03SB28-3.0-4.0		03SB28-6.0-7.0	
Location Code																
Sample ID			03SB27-0.0-2.0		03SB27-3.0-4.0		03SB27-6.0-7.0		03SB27-8.0-9.0		03SB28-0.0-2.0		03SB28-3.0-4.0		03SB28-6.0-7.0	
Sample Date			11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018		11/29/2018	
Sample Interval Depth (ft bgs)			0 - 2		3 - 4		6 - 7		8 - 9		0 - 2		3 - 4		6 - 7	
Sample Purpose			REG		REG		REG		REG		REG		REG		REG	
Parameter	Units	RRS3 MSCs	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
METALS																
Arsenic	mg/kg	5.9	1.41		2.84		5.02		2.33		1.59		0.565		1.6	
Lead	mg/kg	180	8.66		7.36		7.29		4.67		8.27		5.29		5.96	

Notes:

FD - field duplicate

ft bgs - feet below ground surface

ID - identification

J - estimated value

mg/kg - milligrams per kilogram

RRS3 MSCs - Risk Reduction Standard 3 Medium-Specific Concentrations

REG - regular sample

Val Qual - validation qualifier

Table 4-1
Proposed Excavation Sampling Summary

Excavation Area	Excavation Depth (feet)	Estimated Total Volume (CY)	Estimated Floor Area (square feet)	Existing Sidewall Samples	Existing Floor Samples	Estimated Additional Samples Required
A1	1	12	300	03SB18, 03SB12, 03SB25, 03SB24, 03SB23, 03SB28, 03SB27, 03SB15, 03SB26	None	1 floor
A2	1	1	25	None	None	1 composite of four sidewalls and floor
B	8	8	25	03SB15, 03SB19, 03SB26	None	1 floor
C	8	19	64	03SB22 and 03SB28	03SB11 sample intervals 9-10 ft bgs and 14-15 feet bgs	None
	Estimated Total Volume (CY)	40			Total # Samples to be Collected	2 floor and 1 composite for small excavation

Notes:

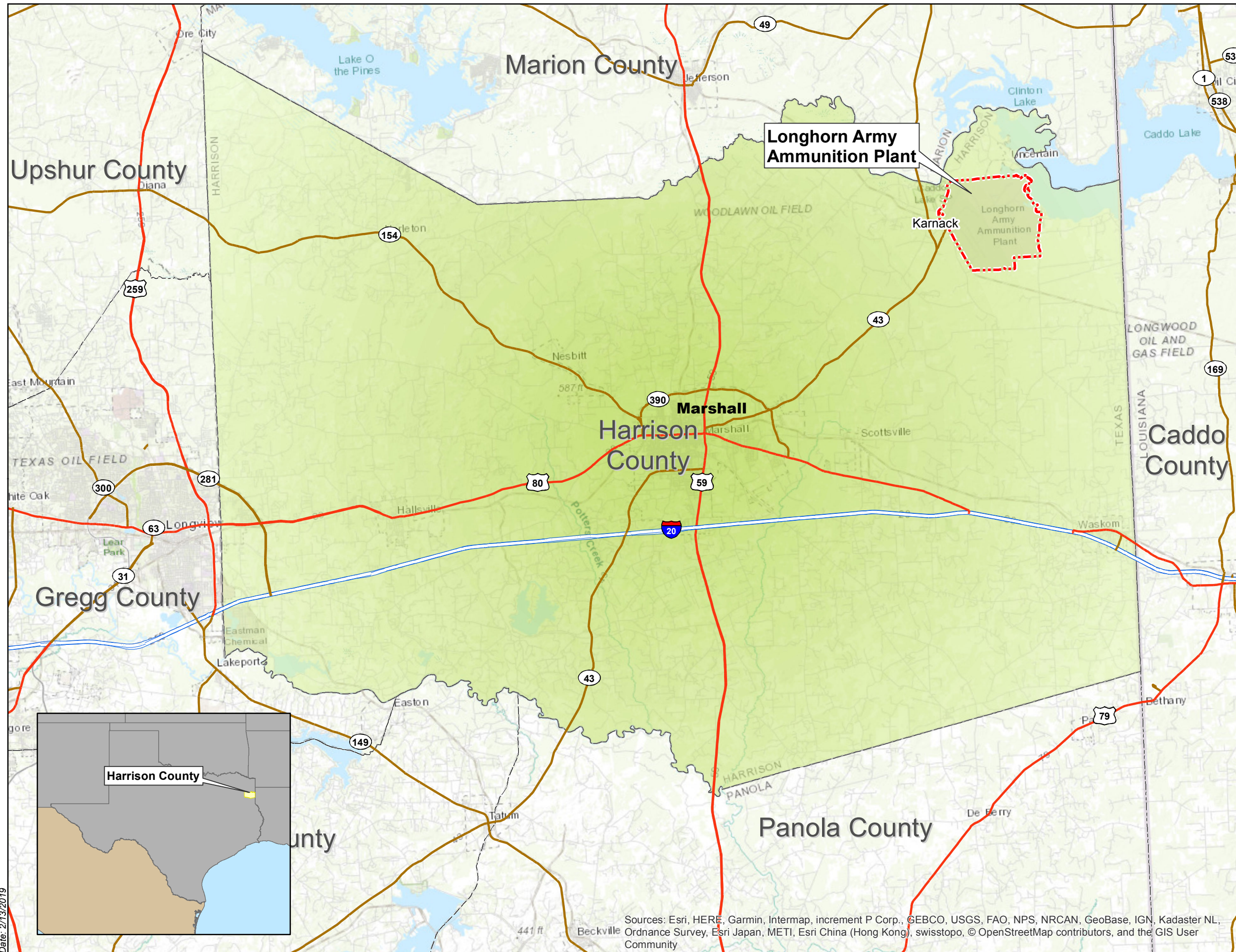
bgs - below ground surface

CY - cubic yards

Table 8-1
Schedule for Major Site Activities

Activities	Duration
Site Preparation and Mobilization	5
Utility Clearance	1
Excavation	3
Waste Characterization and Confirmation Sampling (includes analysis)	5
Loading Stockpiled Soil, T&D, Over-excavation, Confirmation Sampling	5
Site Backfill and Restoration	2
If needed, well replacement	1
Well pad construction and bollard replacement	1
Surveying (Excavation limits and well pad/top of casing)	1
Demobilization	2
Total number of days:	26

Figures



Longhorn Army Ammunition Plant

Karnack

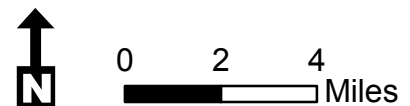
Marshall

Harrison County

Caddo County

Gregg County

Panola County



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TULSA DISTRICT
TULSA, OKLAHOMA

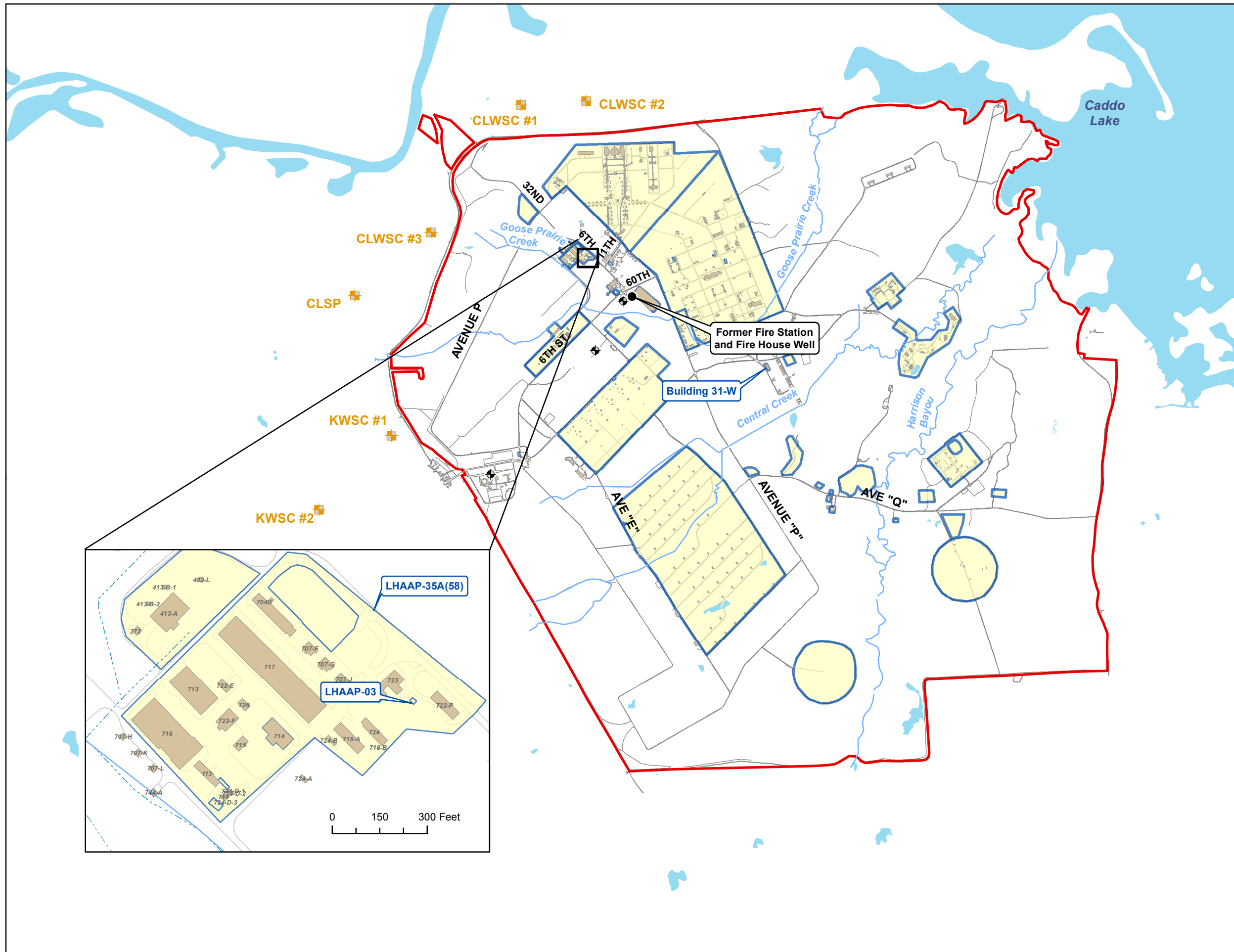


Figure 1-1

LHAAP Location Map
LHAAP-03
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

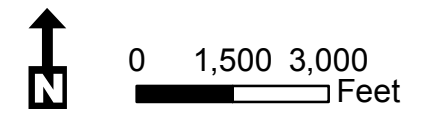
Sources: Esri, HERE, Garmin, Intermap, increment P Corp., GEBCO, USGS, FAO, NPS, NRCAN, GeoBase, IGN, Kadaster NL, Ordnance Survey, Esri Japan, METI, Esri China (Hong Kong), swisstopo, © OpenStreetMap contributors, and the GIS User Community

Date: 2/13/2019



- Fire House Well
- ⊕ Water Supply Well Locations
- ⊕ Public Water Supply Well Locations
- Streams
- Buildings
- ▭ LHAAP Boundary
- Lake/Pond
- ▭ LHAAP-03 Site Boundary
- Roads

KWSC – Karnack Water Supply Corporation
 CLWSC – Caddo Lake Water Supply Corporation
 CLSP - Caddo Lake State Park



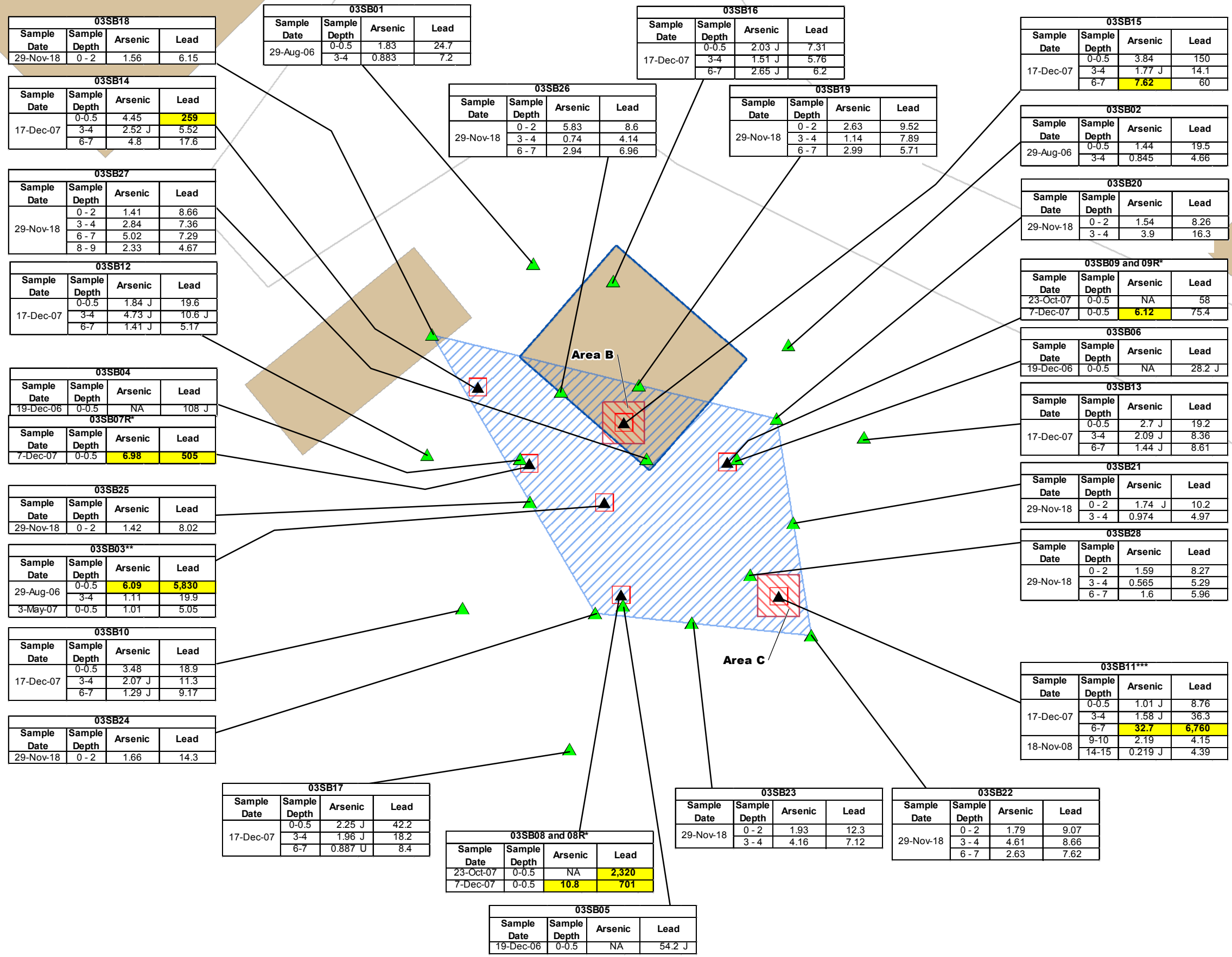
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 TULSA, OKLAHOMA



Figure 1-2

Site Location Map
 LHAAP-03

LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS



- ▲ Soil Boring
- Above Cleanup Level
- ▲ Soil Boring All Sampled Intervals below Cleanup Level
- Target Remediation Area B and Area C from ROD (Excavation Depth to 7 feet bgs)
- Target Remediation Area A from ROD (Excavation Depth to 2 feet bgs)
- Roads
- Fomer Buildings/Structures
- Site Boundary

Note:

1. Excavation depths and areas indicated are from the ROD Record of Decision.
2. All units in milligrams per kilogram (mg/kg)
3. J - Estimated concentration
U - Below detection limit
NA - Parameter not analyzed
4. bgs - below ground surface
5. RRS3 MSCs - Risk Reduction Standard 3 Medium-specific Concentrations
6. Yellow highlighting indicates results exceeding RRS3 MSCs
7. * Resample of the same location at a later date. Note that the original result from 03SB07 was rejected.
8. ** An additional sample called 03SB03-01-Total was collected in May 2007 from the original sample 03SB03-01 location and subjected to both total and the Synthetic Precipitation Leaching Procedure analyses
9. *** In Nov 2008, a soil boring was offset from the original boring location, additional samples were collected at deeper intervals, and well 03WW01 was installed in Nov 2008.

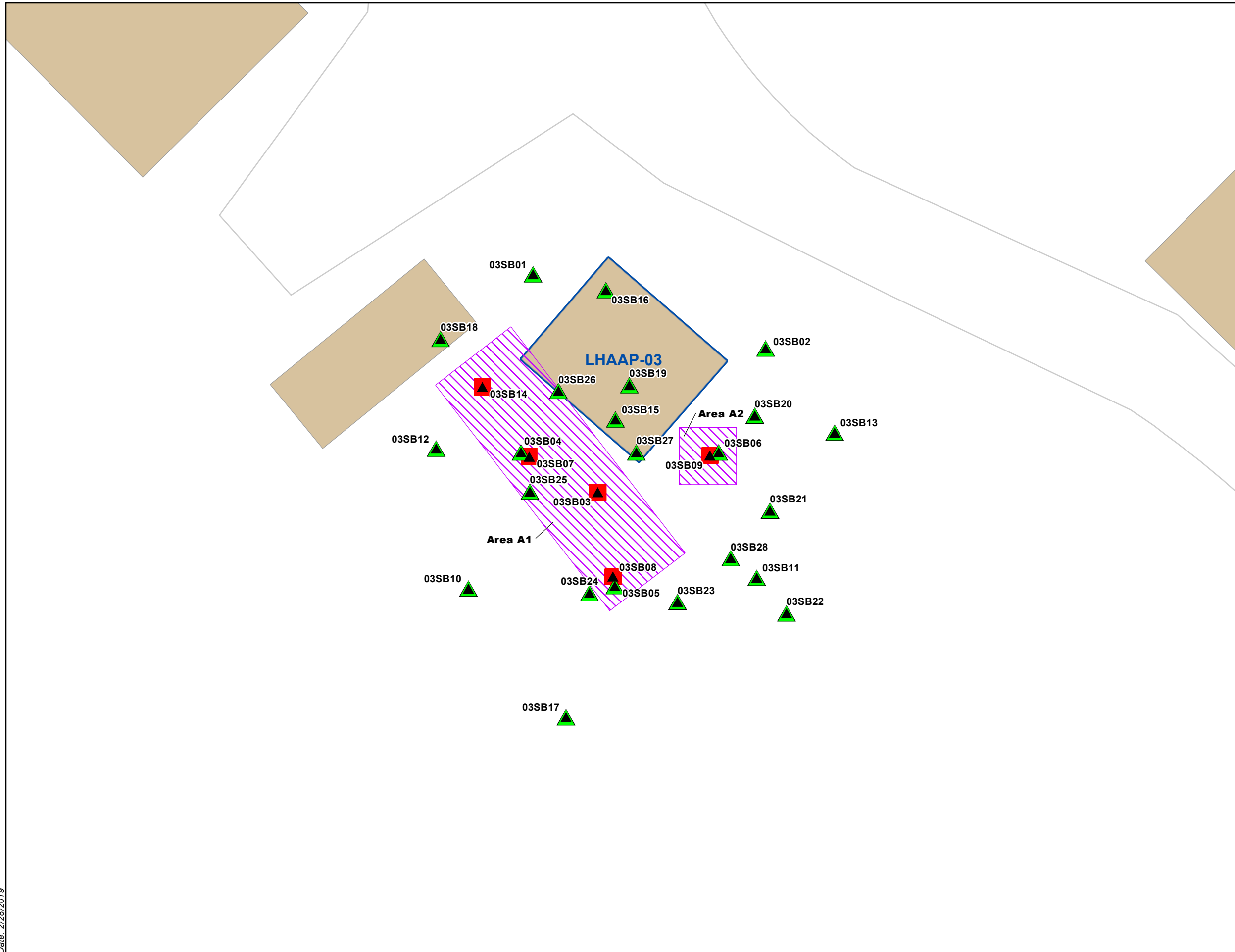
Parameter	Units	RRS3 MSCs
Arsenic	mg/kg	5.9
Lead	mg/kg	180



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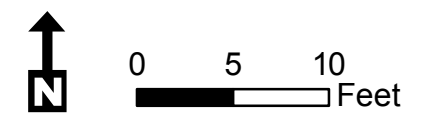


Figure 3-1
Soil Boring Locations and Results
LHAAP-03
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS



- ▲ Soil Boring
- ▲ 0-0.5 or 0-2 foot bgs Sample below RRS3 MSC
- 0-0.5 foot deep soil contamination above RRS3 MSC to be removed
- ▨ 1 ft Excavation Depth
- Roads
- Buildings
- Site Boundary

Note:
 1. bgs - below ground surface
 2. RRS3 MSCs - Risk Reduction Standard 3 Medium-specific Concentrations



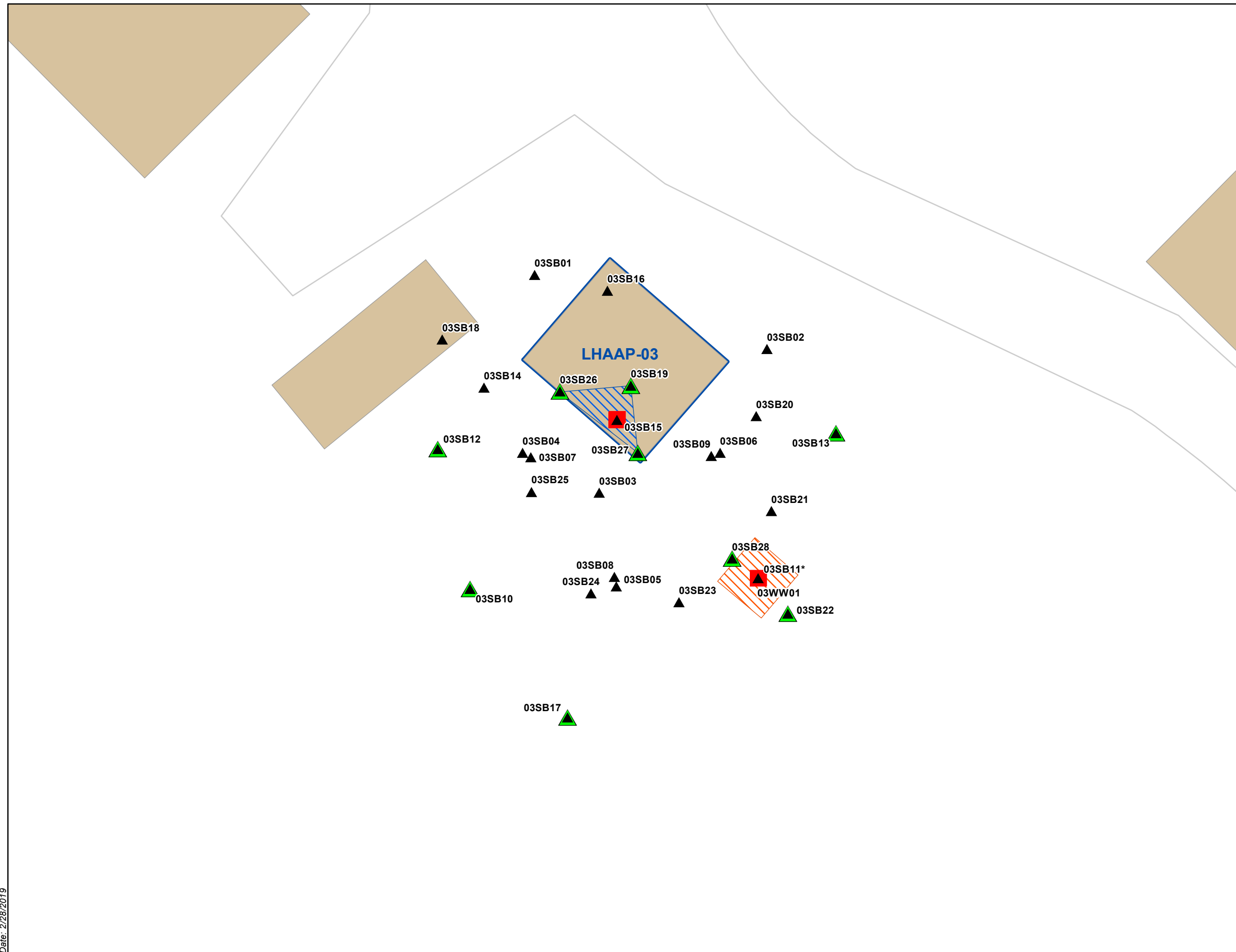
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Figure 4-1

Proposed 1-Foot Excavation Areas
 LHAAP-03
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS

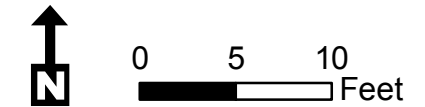
Date: 2/28/2019



- 6-7 foot deep soil contamination above RRS3 MSC to be removed
- ▲ Below Cleanup Level at 6-7 ft bgs
- ▲ Soil Boring not Sampled at 6-7 ft bgs
- Area B (Excavation 8 feet bgs)
- Area C (Excavation 8 feet bgs)
- ⊕ Shallow Monitoring Well
- Roads
- Former Buildings/Structures
- Site Boundary

Note:
 1. bgs - below ground surface
 2. RRS3 MSCs - Risk Reduction Standard 3 Medium-specific Concentrations

* 03SB11 sample intervals from 9-10 feet bgs and 14-15 feet bgs were below RRS3 MSC



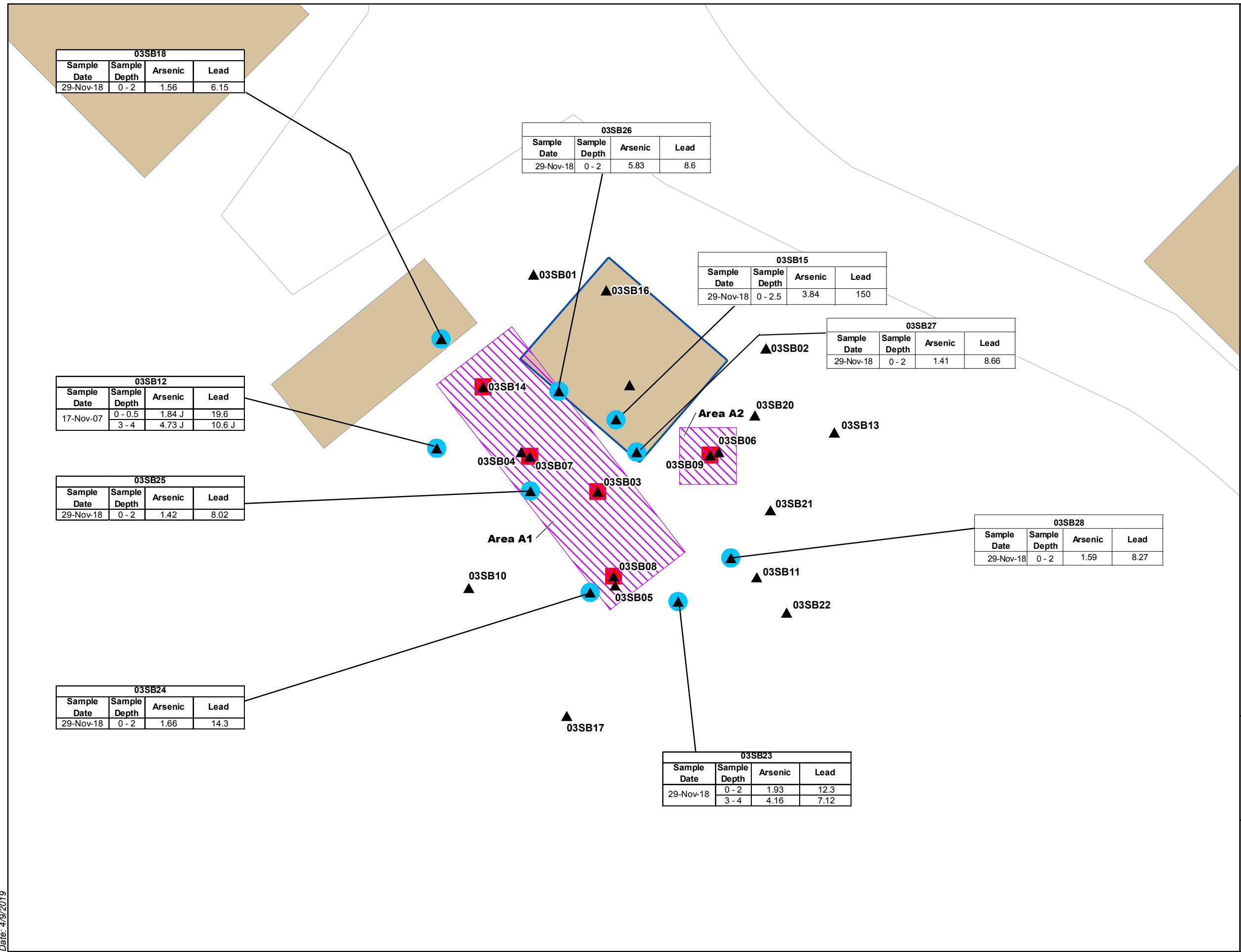
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Figure 4-2

Proposed 8-Foot Excavation Area
 LHAAP-03
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS

Date: 2/28/2019



03SB18			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.56	6.15

03SB26			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	5.83	8.6

03SB15			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2.5	3.84	150

03SB27			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.41	8.66

03SB12			
Sample Date	Sample Depth	Arsenic	Lead
17-Nov-07	0 - 0.5	1.84 J	19.6
	3 - 4	4.73 J	10.6 J

03SB25			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.42	8.02

03SB28			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.59	8.27

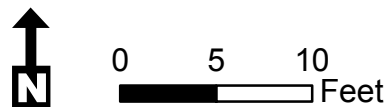
03SB24			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.66	14.3

03SB23			
Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.93	12.3
	3 - 4	4.16	7.12

- ▲ Soil Boring
- Existing Soil Boring Sample to be Used as a Sidewall Confirmation Sample
- Above Cleanup Level
- ▨ Initial Excavation Depth 1-Foot
- Roads
- Former Buildings/Structures
- Site Boundary

Note:
 1. A composite floor confirmation sample will be collected for Area A1 (See Table 4-1).
 2. A single composite confirmation sample will be collected for Area A2 with a grab from each sidewall and floor (See Table 4-1).
 3. All units in milligrams per kilogram (mg/kg)
 4. J - Estimated concentration
 U - Below detection limit
 NA - Parameter not analyzed
 5. bgs - below ground surface

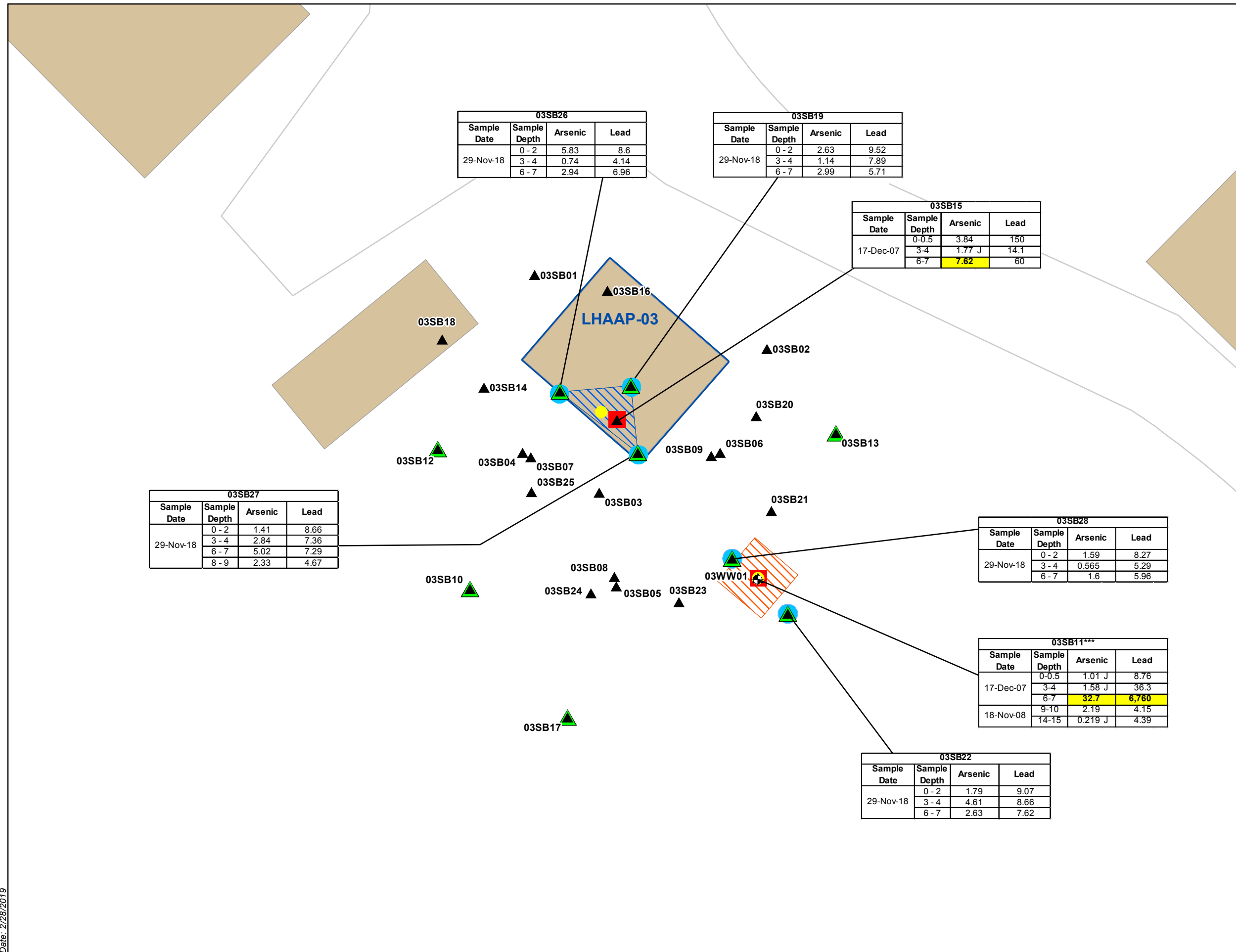
Parameter	Units	RRS3 MSCs
Arsenic	mg/kg	5.9
Lead	mg/kg	180



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Figure 4-3
 Proposed Confirmation Samples
 1-Foot Excavation Areas
 LHAAP-03
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS



- ◆ Shallow Monitoring Well
- ▲ Soil Boring
- ▲ Below Cleanup Level at 6-7 ft bgs
- 6-7 foot deep soil contamination above RRS3 MSC
- Existing Soil Boring Sample to be Used as a Sidewall Confirmation Sample (See Table 4-1)
- Floor Confirmation Sample (Existing soil boring sample for Area C and new location to be collected for Area B) (See Table 4-1)

- 8 ft Excavation Limit**
- ▨ Area B (8-foot excavation)
 - ▨ Area C (8-foot excavation)
 - Roads
 - Buildings
 - ▭ Site Boundary

Note:

1. Excavation depths indicated are from the Record of Decision.
2. All units in milligrams per kilogram (mg/kg)
3. J - Estimated concentration
- U - Below detection limit
- NA - Parameter not analyzed
4. bgs - below ground surface
5. RRS3 MSCs - Risk Reduction Standard 3 Medium-specific Concentrations
6. Yellow highlighting indicates results exceeding RRS3 MSCs
7. *** In Nov 2008, a soil boring was offset from the original boring location, additional samples were collected at deeper intervals, and well 03WW01 was installed in Nov 2008.

Parameter	Units	RRS3 MSCs
Arsenic	mg/kg	5.9
Lead	mg/kg	180



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TULSA, OKLAHOMA

Figure 4-4
Proposed Confirmation Samples
8-Foot Excavation Areas B and C
LHAAP-03
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.41	8.66
	3 - 4	2.84	7.36
	6 - 7	5.02	7.29
	8 - 9	2.33	4.67

Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	5.83	8.6
	3 - 4	0.74	4.14
	6 - 7	2.94	6.96

Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	2.63	9.52
	3 - 4	1.14	7.89
	6 - 7	2.99	5.71

Sample Date	Sample Depth	Arsenic	Lead
17-Dec-07	0-0.5	3.84	150
	3-4	1.77 J	14.1
	6-7	7.62	60

Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.59	8.27
	3 - 4	0.565	5.29
	6 - 7	1.6	5.96

Sample Date	Sample Depth	Arsenic	Lead
17-Dec-07	0-0.5	1.01 J	8.76
	3-4	1.58 J	36.3
	6-7	32.7	6,760
18-Nov-08	9-10	2.19	4.15
	14-15	0.219 J	4.39

Sample Date	Sample Depth	Arsenic	Lead
29-Nov-18	0 - 2	1.79	9.07
	3 - 4	4.61	8.66
	6 - 7	2.63	7.62

Appendix A

Analytical Data Reports from November 2018 Soil Samples



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

December 18, 2018

Susan Huang
Aptim Environmental & Infrastructure, Inc.
2500 City West Blvd., Suite 1700
Houston, TX 77042

Work Order: **HS18120004**

Laboratory Results for: **Longhorn Army Ammunition Plant LHAAP-03**

Dear Susan,

ALS Environmental received 36 sample(s) on Nov 30, 2018 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read "Raj. P. Modashia", enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
Work Order: HS18120004

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS18120004-01	03SB18-0.0-2.0	Soil		29-Nov-2018 10:00	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-02	03SB19-0.0-2.0	Soil		29-Nov-2018 10:18	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-03	03SB19-3.0-4.0	Soil		29-Nov-2018 10:20	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-04	03SB19-6.0-7.0	Soil		29-Nov-2018 10:24	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-05	03SB19-8.0-9.0	Soil		29-Nov-2018 10:26	30-Nov-2018 09:40	<input checked="" type="checkbox"/>
HS18120004-06	03SB20-0.0-2.0	Soil		29-Nov-2018 10:30	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-07	03SB20-0.0-2.0-FD	Soil		29-Nov-2018 10:30	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-08	03SB20-3.0-4.0	Soil		29-Nov-2018 10:35	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-09	03SB20-6.0-7.0	Soil		29-Nov-2018 10:38	30-Nov-2018 09:40	<input checked="" type="checkbox"/>
HS18120004-10	03SB21-0.0-2.0	Soil		29-Nov-2018 10:45	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-11	03SB21-3.0-4.0	Soil		29-Nov-2018 10:50	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-12	03SB21-6.0-7.0	Soil		29-Nov-2018 10:55	30-Nov-2018 09:40	<input checked="" type="checkbox"/>
HS18120004-13	03SB22-0.0-2.0	Soil		29-Nov-2018 11:05	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-14	03SB22-3.0-4.0	Soil		29-Nov-2018 11:10	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-15	03SB22-3.0-4.0-FD	Soil		29-Nov-2018 11:10	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-16	03SB22-6.0-7.0	Soil		29-Nov-2018 11:13	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-17	03SB22-8.0-9.0	Soil		29-Nov-2018 11:15	30-Nov-2018 09:40	<input checked="" type="checkbox"/>
HS18120004-18	03SB23-0.0-2.0	Soil		29-Nov-2018 11:30	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-19	03SB23-3.0-4.0	Soil		29-Nov-2018 11:33	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-20	03SB23-6.0-7.0	Soil		29-Nov-2018 11:35	30-Nov-2018 09:40	<input checked="" type="checkbox"/>
HS18120004-21	03SB24-0.0-2.0	Soil		29-Nov-2018 11:45	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-22	03SB25-0.0-2.0	Soil		29-Nov-2018 11:40	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-23	03SB26-0.0-2.0	Soil		29-Nov-2018 10:05	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-24	03SB26-0.0-2.0-FD	Soil		29-Nov-2018 10:05	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-25	03SB26-3.0-4.0	Soil		29-Nov-2018 10:10	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-26	03SB26-6.0-7.0	Soil		29-Nov-2018 10:13	30-Nov-2018 09:40	<input type="checkbox"/>

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
Work Order: HS18120004

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS18120004-27	03SB26-8.0-9.0	Soil		29-Nov-2018 10:15	30-Nov-2018 09:40	<input checked="" type="checkbox"/>
HS18120004-28	03SB27-0.0-2.0	Soil		29-Nov-2018 11:55	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-29	03SB27-3.0-4.0	Soil		29-Nov-2018 12:00	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-30	03SB27-6.0-7.0	Soil		29-Nov-2018 12:05	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-31	03SB27-8.0-9.0	Soil		29-Nov-2018 12:08	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-32	03SB28-0.0-2.0	Soil		29-Nov-2018 12:15	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-33	03SB28-3.0-4.0	Soil		29-Nov-2018 12:18	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-34	03SB28-6.0-7.0	Soil		29-Nov-2018 12:20	30-Nov-2018 09:40	<input type="checkbox"/>
HS18120004-35	03SB28-8.0-9.0	Soil		29-Nov-2018 12:24	30-Nov-2018 09:40	<input checked="" type="checkbox"/>
HS18120004-36	Trip Blank	Water	ALS- 111418-61	29-Nov-2018 00:00	30-Nov-2018 09:40	<input checked="" type="checkbox"/>

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
Work Order: HS18120004

CASE NARRATIVE**Metals by Method SW6020****Batch ID: 135452**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

Batch ID: 135451**Sample ID: 03SB27-3.0-4.0 (HS18120004-29MS)**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB18-0.0-2.0
 Collection Date: 29-Nov-2018 10:00

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-01
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.56		0.0661	0.0944	0.472	mg/Kg	1	14-Dec-2018 19:53
Lead	6.15		0.0123	0.0944	0.472	mg/Kg	1	14-Dec-2018 19:53

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB19-0.0-2.0
 Collection Date: 29-Nov-2018 10:18

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-02
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	2.63		0.0645	0.0921	0.460	mg/Kg	1	14-Dec-2018 19:55
Lead	9.52		0.0120	0.0921	0.460	mg/Kg	1	14-Dec-2018 19:55

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB19-3.0-4.0
 Collection Date: 29-Nov-2018 10:20

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-03
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.14		0.0639	0.0913	0.456	mg/Kg	1	14-Dec-2018 19:58
Lead	7.89		0.0119	0.0913	0.456	mg/Kg	1	14-Dec-2018 19:58

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB19-6.0-7.0
 Collection Date: 29-Nov-2018 10:24

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-04
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	2.99		0.0682	0.0975	0.487	mg/Kg	1	14-Dec-2018 20:00
Lead	5.71		0.0127	0.0975	0.487	mg/Kg	1	14-Dec-2018 20:00

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB20-0.0-2.0
 Collection Date: 29-Nov-2018 10:30

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-06
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.54		0.0684	0.0977	0.489	mg/Kg	1	14-Dec-2018 20:02
Lead	8.26		0.0127	0.0977	0.489	mg/Kg	1	14-Dec-2018 20:02

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB20-0.0-2.0-FD
 Collection Date: 29-Nov-2018 10:30

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-07
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	4.05		0.0669	0.0955	0.478	mg/Kg	1	14-Dec-2018 20:04
Lead	11.3		0.0124	0.0955	0.478	mg/Kg	1	14-Dec-2018 20:04

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB20-3.0-4.0
 Collection Date: 29-Nov-2018 10:35

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-08
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018	Analyst: JCJ	
Arsenic	3.90		0.0681	0.0973	0.486	mg/Kg	1	14-Dec-2018 20:07
Lead	16.3		0.0126	0.0973	0.486	mg/Kg	1	14-Dec-2018 20:07

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB21-0.0-2.0
 Collection Date: 29-Nov-2018 10:45

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-10
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.74		0.0642	0.0917	0.458	mg/Kg	1	14-Dec-2018 20:16
Lead	10.2		0.0119	0.0917	0.458	mg/Kg	1	14-Dec-2018 20:16

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB21-3.0-4.0
 Collection Date: 29-Nov-2018 10:50

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-11
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	0.974		0.0640	0.0915	0.457	mg/Kg	1	14-Dec-2018 20:27
Lead	4.97		0.0119	0.0915	0.457	mg/Kg	1	14-Dec-2018 20:27

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB22-0.0-2.0
 Collection Date: 29-Nov-2018 11:05

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-13
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.79		0.0660	0.0943	0.472	mg/Kg	1	14-Dec-2018 20:29
Lead	9.07		0.0123	0.0943	0.472	mg/Kg	1	14-Dec-2018 20:29

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB22-3.0-4.0
 Collection Date: 29-Nov-2018 11:10

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-14
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018	Analyst: JCJ	
Arsenic	4.61		0.0665	0.0950	0.475	mg/Kg	1	14-Dec-2018 20:31
Lead	8.66		0.0124	0.0950	0.475	mg/Kg	1	14-Dec-2018 20:31

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB22-3.0-4.0-FD
 Collection Date: 29-Nov-2018 11:10

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-15
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018	Analyst: JCJ	
Arsenic	0.709		0.0660	0.0943	0.471	mg/Kg	1	14-Dec-2018 20:33
Lead	6.26		0.0123	0.0943	0.471	mg/Kg	1	14-Dec-2018 20:33

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB22-6.0-7.0
 Collection Date: 29-Nov-2018 11:13

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-16
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018	Analyst: JCJ	
Arsenic	2.63		0.0672	0.0959	0.480	mg/Kg	1	14-Dec-2018 22:30
Lead	7.62		0.0125	0.0959	0.480	mg/Kg	1	14-Dec-2018 22:30

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB23-0.0-2.0
 Collection Date: 29-Nov-2018 11:30

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-18
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018	Analyst: JCJ	
Arsenic	1.93		0.0648	0.0926	0.463	mg/Kg	1	14-Dec-2018 22:32
Lead	12.3		0.0120	0.0926	0.463	mg/Kg	1	14-Dec-2018 22:32

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB23-3.0-4.0
 Collection Date: 29-Nov-2018 11:33

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-19
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	4.16		0.0645	0.0922	0.461	mg/Kg	1	14-Dec-2018 22:35
Lead	7.12		0.0120	0.0922	0.461	mg/Kg	1	14-Dec-2018 22:35

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB24-0.0-2.0
 Collection Date: 29-Nov-2018 11:45

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-21
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.66		0.0661	0.0944	0.472	mg/Kg	1	14-Dec-2018 22:37
Lead	14.3		0.0123	0.0944	0.472	mg/Kg	1	14-Dec-2018 22:37

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB25-0.0-2.0
 Collection Date: 29-Nov-2018 11:40

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-22
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.42		0.0681	0.0972	0.486	mg/Kg	1	14-Dec-2018 22:39
Lead	8.02		0.0126	0.0972	0.486	mg/Kg	1	14-Dec-2018 22:39

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB26-0.0-2.0
 Collection Date: 29-Nov-2018 10:05

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-23
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	5.83		0.0684	0.0977	0.489	mg/Kg	1	14-Dec-2018 22:41
Lead	8.60		0.0127	0.0977	0.489	mg/Kg	1	14-Dec-2018 22:41

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB26-0.0-2.0-FD
 Collection Date: 29-Nov-2018 10:05

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-24
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.92		0.0674	0.0963	0.481	mg/Kg	1	14-Dec-2018 22:44
Lead	8.38		0.0125	0.0963	0.481	mg/Kg	1	14-Dec-2018 22:44

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB26-3.0-4.0
 Collection Date: 29-Nov-2018 10:10

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-25
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	0.740		0.0646	0.0923	0.461	mg/Kg	1	14-Dec-2018 22:46
Lead	4.14		0.0120	0.0923	0.461	mg/Kg	1	14-Dec-2018 22:46

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB26-6.0-7.0
 Collection Date: 29-Nov-2018 10:13

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-26
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	2.94		0.0680	0.0971	0.486	mg/Kg	1	12-Dec-2018 20:26
Lead	6.96		0.0126	0.0971	0.486	mg/Kg	1	12-Dec-2018 20:26

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB27-0.0-2.0
 Collection Date: 29-Nov-2018 11:55

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-28
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.41		0.0680	0.0971	0.486	mg/Kg	1	12-Dec-2018 20:29
Lead	8.66		0.0126	0.0971	0.486	mg/Kg	1	12-Dec-2018 20:29

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB27-3.0-4.0
 Collection Date: 29-Nov-2018 12:00

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-29
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	2.84		0.0642	0.0917	0.459	mg/Kg	1	12-Dec-2018 20:31
Lead	7.36		0.0119	0.0917	0.459	mg/Kg	1	12-Dec-2018 20:31

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB27-6.0-7.0
 Collection Date: 29-Nov-2018 12:05

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-30
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	5.02		0.0640	0.0915	0.457	mg/Kg	1	12-Dec-2018 20:42
Lead	7.29		0.0119	0.0915	0.457	mg/Kg	1	12-Dec-2018 20:42

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB27-8.0-9.0
 Collection Date: 29-Nov-2018 12:08

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-31
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	2.33		0.0660	0.0943	0.471	mg/Kg	1	12-Dec-2018 20:53
Lead	4.67		0.0123	0.0943	0.471	mg/Kg	1	12-Dec-2018 20:53

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB28-0.0-2.0
 Collection Date: 29-Nov-2018 12:15

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-32
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	1.59		0.0655	0.0936	0.468	mg/Kg	1	12-Dec-2018 20:55
Lead	8.27		0.0122	0.0936	0.468	mg/Kg	1	12-Dec-2018 20:55

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB28-3.0-4.0
 Collection Date: 29-Nov-2018 12:18

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-33
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018		Analyst: JCJ
Arsenic	0.565		0.0642	0.0917	0.458	mg/Kg	1	12-Dec-2018 20:58
Lead	5.29		0.0119	0.0917	0.458	mg/Kg	1	12-Dec-2018 20:58

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: Longhorn Army Ammunition Plant LHAAP-03
 Sample ID: 03SB28-6.0-7.0
 Collection Date: 29-Nov-2018 12:20

ANALYTICAL REPORT

WorkOrder:HS18120004
 Lab ID:HS18120004-34
 Matrix:Soil

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
METALS BY SW6020A	Method:SW6020					Prep:SW3050A / 11-Dec-2018	Analyst: JCJ	
Arsenic	1.60		0.0675	0.0965	0.482	mg/Kg	1	12-Dec-2018 21:00
Lead	5.96		0.0125	0.0965	0.482	mg/Kg	1	12-Dec-2018 21:00

Note: See Qualifiers Page for a list of qualifiers and their explanation.

WEIGHT LOG

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
WorkOrder: HS18120004

Batch ID: 135451 **Method:** METALS BY SW6020A **Prep:** 3050_I_LOW

SampID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS18120004-26	1	0.5149	50 (mL)	97.11
HS18120004-28	1	0.5147	50 (mL)	97.14
HS18120004-29	1	0.545	50 (mL)	91.74
HS18120004-30	1	0.5466	50 (mL)	91.47
HS18120004-31	1	0.5304	50 (mL)	94.27
HS18120004-32	1	0.5343	50 (mL)	93.58
HS18120004-33	1	0.5454	50 (mL)	91.68
HS18120004-34	1	0.5183	50 (mL)	96.47

Batch ID: 135452 **Method:** METALS BY SW6020A **Prep:** 3050_I_LOW

SampID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS18120004-01	1	0.5298	50 (mL)	94.38
HS18120004-02	1	0.543	50 (mL)	92.08
HS18120004-03	1	0.5479	50 (mL)	91.26
HS18120004-04	1	0.513	50 (mL)	97.47
HS18120004-06	1	0.5116	50 (mL)	97.73
HS18120004-07	1	0.5233	50 (mL)	95.55
HS18120004-08	1	0.514	50 (mL)	97.28
HS18120004-10	1	0.5455	50 (mL)	91.66
HS18120004-11	1	0.5465	50 (mL)	91.49
HS18120004-13	1	0.53	50 (mL)	94.34
HS18120004-14	1	0.5262	50 (mL)	95.02
HS18120004-15	1	0.5305	50 (mL)	94.25
HS18120004-16	1	0.5212	50 (mL)	95.93
HS18120004-18	1	0.54	50 (mL)	92.59
HS18120004-19	1	0.5425	50 (mL)	92.17
HS18120004-21	1	0.5296	50 (mL)	94.41
HS18120004-22	1	0.5143	50 (mL)	97.22
HS18120004-23	1	0.5117	50 (mL)	97.71
HS18120004-24	1	0.5193	50 (mL)	96.28
HS18120004-25	1	0.5419	50 (mL)	92.27

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
WorkOrder: HS18120004

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID 135451	Test Name : METALS BY SW6020A			Matrix: Soil		
HS18120004-26	03SB26-6.0-7.0	29 Nov 2018 10:13		11 Dec 2018 13:27	12 Dec 2018 20:26	1
HS18120004-28	03SB27-0.0-2.0	29 Nov 2018 11:55		11 Dec 2018 13:27	12 Dec 2018 20:29	1
HS18120004-29	03SB27-3.0-4.0	29 Nov 2018 12:00		11 Dec 2018 13:27	12 Dec 2018 20:31	1
HS18120004-30	03SB27-6.0-7.0	29 Nov 2018 12:05		11 Dec 2018 13:27	12 Dec 2018 20:42	1
HS18120004-31	03SB27-8.0-9.0	29 Nov 2018 12:08		11 Dec 2018 13:27	12 Dec 2018 20:53	1
HS18120004-32	03SB28-0.0-2.0	29 Nov 2018 12:15		11 Dec 2018 13:27	12 Dec 2018 20:55	1
HS18120004-33	03SB28-3.0-4.0	29 Nov 2018 12:18		11 Dec 2018 13:27	12 Dec 2018 20:58	1
HS18120004-34	03SB28-6.0-7.0	29 Nov 2018 12:20		11 Dec 2018 13:27	12 Dec 2018 21:00	1
Batch ID 135452	Test Name : METALS BY SW6020A			Matrix: Soil		
HS18120004-01	03SB18-0.0-2.0	29 Nov 2018 10:00		11 Dec 2018 13:33	14 Dec 2018 19:53	1
HS18120004-02	03SB19-0.0-2.0	29 Nov 2018 10:18		11 Dec 2018 13:33	14 Dec 2018 19:55	1
HS18120004-03	03SB19-3.0-4.0	29 Nov 2018 10:20		11 Dec 2018 13:33	14 Dec 2018 19:58	1
HS18120004-04	03SB19-6.0-7.0	29 Nov 2018 10:24		11 Dec 2018 13:33	14 Dec 2018 20:00	1
HS18120004-06	03SB20-0.0-2.0	29 Nov 2018 10:30		11 Dec 2018 13:33	14 Dec 2018 20:02	1
HS18120004-07	03SB20-0.0-2.0-FD	29 Nov 2018 10:30		11 Dec 2018 13:33	14 Dec 2018 20:04	1
HS18120004-08	03SB20-3.0-4.0	29 Nov 2018 10:35		11 Dec 2018 13:33	14 Dec 2018 20:07	1
HS18120004-10	03SB21-0.0-2.0	29 Nov 2018 10:45		11 Dec 2018 13:33	14 Dec 2018 20:16	1
HS18120004-11	03SB21-3.0-4.0	29 Nov 2018 10:50		11 Dec 2018 13:33	14 Dec 2018 20:27	1
HS18120004-13	03SB22-0.0-2.0	29 Nov 2018 11:05		11 Dec 2018 13:33	14 Dec 2018 20:29	1
HS18120004-14	03SB22-3.0-4.0	29 Nov 2018 11:10		11 Dec 2018 13:33	14 Dec 2018 20:31	1
HS18120004-15	03SB22-3.0-4.0-FD	29 Nov 2018 11:10		11 Dec 2018 13:33	14 Dec 2018 20:33	1
HS18120004-16	03SB22-6.0-7.0	29 Nov 2018 11:13		11 Dec 2018 13:33	14 Dec 2018 22:30	1
HS18120004-18	03SB23-0.0-2.0	29 Nov 2018 11:30		11 Dec 2018 13:33	14 Dec 2018 22:32	1
HS18120004-19	03SB23-3.0-4.0	29 Nov 2018 11:33		11 Dec 2018 13:33	14 Dec 2018 22:35	1
HS18120004-21	03SB24-0.0-2.0	29 Nov 2018 11:45		11 Dec 2018 13:33	14 Dec 2018 22:37	1
HS18120004-22	03SB25-0.0-2.0	29 Nov 2018 11:40		11 Dec 2018 13:33	14 Dec 2018 22:39	1
HS18120004-23	03SB26-0.0-2.0	29 Nov 2018 10:05		11 Dec 2018 13:33	14 Dec 2018 22:41	1
HS18120004-24	03SB26-0.0-2.0-FD	29 Nov 2018 10:05		11 Dec 2018 13:33	14 Dec 2018 22:44	1
HS18120004-25	03SB26-3.0-4.0	29 Nov 2018 10:10		11 Dec 2018 13:33	14 Dec 2018 22:46	1

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
WorkOrder: HS18120004

QC BATCH REPORT

Batch ID: 135451		Instrument: ICPMS04		Method: SW6020						
MBLK	Sample ID: MBLK-135451	Units: mg/Kg		Analysis Date: 12-Dec-2018 19:57						
Client ID:	Run ID: ICPMS04_329054	SeqNo: 4861941	PrepDate: 11-Dec-2018	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	0.100	0.500								U
Lead	0.100	0.500								U
LCS	Sample ID: LCS-135451	Units: mg/Kg		Analysis Date: 12-Dec-2018 19:59						
Client ID:	Run ID: ICPMS04_329054	SeqNo: 4861942	PrepDate: 11-Dec-2018	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	9.119	0.500	10	0	91.2	80 - 120				
Lead	9.458	0.500	10	0	94.6	80 - 120				
MS	Sample ID: HS18120004-29MS	Units: mg/Kg		Analysis Date: 12-Dec-2018 20:35						
Client ID: 03SB27-3.0-4.0	Run ID: ICPMS04_329054	SeqNo: 4861959	PrepDate: 11-Dec-2018	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	11.66	0.462	9.244	2.842	95.4	75 - 125				
Lead	16.52	0.462	9.244	7.362	99.1	75 - 125				
MSD	Sample ID: HS18120004-29MSD	Units: mg/Kg		Analysis Date: 12-Dec-2018 20:38						
Client ID: 03SB27-3.0-4.0	Run ID: ICPMS04_329054	SeqNo: 4861960	PrepDate: 11-Dec-2018	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	11.67	0.464	9.28	2.842	95.1	75 - 125	11.66	0.0452	20	
Lead	16.98	0.464	9.28	7.362	104	75 - 125	16.52	2.74	20	
PDS	Sample ID: HS18120004-29PDS	Units: mg/Kg		Analysis Date: 12-Dec-2018 20:40						
Client ID: 03SB27-3.0-4.0	Run ID: ICPMS04_329054	SeqNo: 4861961	PrepDate: 11-Dec-2018	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	11.46	0.459	9.174	2.842	93.9	75 - 125				
Lead	16.07	0.459	9.174	7.362	94.9	75 - 125				

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
WorkOrder: HS18120004

QC BATCH REPORT

Batch ID: 135451		Instrument: ICPMS04		Method: SW6020						
SD	Sample ID: HS18120004-29SD	Units: mg/Kg		Analysis Date: 12-Dec-2018 20:33						
Client ID: 03SB27-3.0-4.0	Run ID: ICPMS04_329054	SeqNo: 4861958	PrepDate: 11-Dec-2018	DF: 5						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	Limit Qual	
Arsenic	3.005	2.29					2.842	5.71	10	
Lead	7.315	2.29					7.362	0.645	10	

The following samples were analyzed in this batch:

HS18120004-26	HS18120004-28	HS18120004-29	HS18120004-30
HS18120004-31	HS18120004-32	HS18120004-33	HS18120004-34

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
WorkOrder: HS18120004

QC BATCH REPORT

Batch ID: 135452		Instrument: ICPMS04			Method: SW6020					
MBLK	Sample ID: MBLK-135452	Units: mg/Kg			Analysis Date: 14-Dec-2018 19:49					
Client ID:		Run ID: ICPMS04_329261	SeqNo: 4866984	PrepDate: 11-Dec-2018	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	0.100	0.500								U
Lead	0.100	0.500								U
LCS	Sample ID: LCS-135452	Units: mg/Kg			Analysis Date: 14-Dec-2018 19:51					
Client ID:		Run ID: ICPMS04_329261	SeqNo: 4866985	PrepDate: 11-Dec-2018	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	9.338	0.500	10	0	93.4	80 - 120				
Lead	9.804	0.500	10	0	98.0	80 - 120				
MS	Sample ID: HS18120004-10MS	Units: mg/Kg			Analysis Date: 14-Dec-2018 20:20					
Client ID: 03SB21-0.0-2.0		Run ID: ICPMS04_329261	SeqNo: 4866998	PrepDate: 11-Dec-2018	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	10.17	0.459	9.184	1.743	91.8	75 - 125				
Lead	19.21	0.459	9.184	10.17	98.5	75 - 125				
MSD	Sample ID: HS18120004-10MSD	Units: mg/Kg			Analysis Date: 14-Dec-2018 20:22					
Client ID: 03SB21-0.0-2.0		Run ID: ICPMS04_329261	SeqNo: 4866999	PrepDate: 11-Dec-2018	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	11.17	0.464	9.287	1.743	102	75 - 125	10.17	9.37	20	
Lead	21.34	0.464	9.287	10.17	120	75 - 125	19.21	10.5	20	
PDS	Sample ID: HS18120004-10PDS	Units: mg/Kg			Analysis Date: 14-Dec-2018 20:25					
Client ID: 03SB21-0.0-2.0		Run ID: ICPMS04_329261	SeqNo: 4867000	PrepDate: 11-Dec-2018	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Arsenic	10.9	0.458	9.166	1.743	99.9	75 - 125				
Lead	20	0.458	9.166	10.17	107	75 - 125				

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
WorkOrder: HS18120004

QC BATCH REPORT

Batch ID: 135452		Instrument: ICPMS04		Method: SW6020						
SD	Sample ID: HS18120004-10SD	Units: mg/Kg			Analysis Date: 14-Dec-2018 20:18					
Client ID: 03SB21-0.0-2.0	Run ID: ICPMS04_329261	SeqNo: 4866997	PrepDate: 11-Dec-2018	DF: 5						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	Limit	Qual
Arsenic	1.71	2.29					1.743	0	10	J
Lead	10.41	2.29					10.17	2.42	10	

The following samples were analyzed in this batch:

HS18120004-01	HS18120004-02	HS18120004-03	HS18120004-04
HS18120004-06	HS18120004-07	HS18120004-08	HS18120004-10
HS18120004-11	HS18120004-13	HS18120004-14	HS18120004-15
HS18120004-16	HS18120004-18	HS18120004-19	HS18120004-21
HS18120004-22	HS18120004-23	HS18120004-24	HS18120004-25

ALS Houston, US

Date: 18-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: Longhorn Army Ammunition Plant LHAAP-03
WorkOrder: **HS18120004**

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
North Carolina	624-2018	31-Dec-2018
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	22-Dec-2018
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019

Sample Receipt Checklist

Client Name: CBI-Houston
 Work Order: HS18120004

Date/Time Received: **30-Nov-2018 09:40**
 Received by: **PJM**

Checklist completed by: Pablo Martinez 1-Dec-2018 Reviewed by: RJ Modashia 3-Dec-2018
 eSignature Date eSignature Date

Matrices: **SOIL, WATER** Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- Chain of custody present? Yes No
- Chain of custody signed when relinquished and received? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- TX1005 solids received in hermetically sealed vials? Yes No N/A
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 2.1C/2.5C UC/C IR # 11
 Cooler(s)/Kit(s): 44426
 Date/Time sample(s) sent to storage: 12/1/18 9:20

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

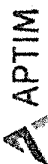
Login Notes: Trip Blank not listed on CoC, logged in on hold

Client Contacted: _____ Date Contacted: _____ Person Contacted: _____

Contacted By: _____ Regarding: _____

Comments:

Corrective Action:



COC ID: LHAAP03-NOV2018-ALSHT-1811-29

TURNAROUND TIME: normal

RUSH: Page 1 of 3

PROJECT/CLIENT INFO		LABORATORY		OTHER INFO	
Facility Name Longhorn AAP	Lab Name ALS Laboratories	Email Invoice To FedInvoices@aptim.com			
Project Number 501032	Lab Contact RJ Modashia	Email Report To Susan.Huang@aptim.com			
Address LHAAP-03 1203-B East Grand Avenue PMB 202	Email RJ.Modashia@alsglobal.com	Mail Reports To Susan Huang			
City/Postal Code Marshall 75670	City/Postal Code Houston 77099	Address 4005 Port Chicago Highway, Suit 200	City Concord	State CA	Country USA
Phone Number 713.243.7264	Phone Number 281.575.2279 or 281.530.5656	Postal Code 94520			
Project Manager Praveen Srivastava		Shipping Company Fedex			

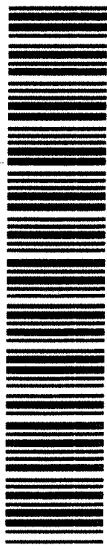
SAMPLE DETAILS

ANALYSIS REQUESTED

Sample ID	Location	Start Depth	End Depth	Depth Unit	Field Matrix	Date	Time (24hr)	# Of Cont.	ANALYSIS	Metals (As & Pb) by 6020
035B18-0.0-2.0	035B18	0	2	FT	S ₀₁	11/28/18	1000	1		X
035B19-0.0-2.0	035B19	0	2	FT			1018	1		X
035B19-3.0-4.0	035B19	3	4				1020	1		X
035B19-6.0-7.0	035B19	6	7				1024	1		X
035B19-8.0-9.0	035B19	8	9				1026	1		X
035B20-0.0-2.0	035B20	0	2				1030	1		X
035B20-3.0-4.0	035B20	3	4				1035	1		X
035B20-6.0-7.0	035B20	6	7				1038	1		X
035B21-0.0-2.0	035B21	0	2				1045	1		X
035B21-0.0-2.0-M1	035B21	0	2				1045	1		X
035B21-0.0-2.0-M10	035B21	0	2				1045	1		X
035B21-3.0-4.0	035B21	3	4				1050	1		X
035B21-6.0-7.0	035B21	6	7				1055	1		X
035B22-0.0-2.0	035B22	0	2				1105	1		X
035B22-3.0-4.0	035B22	3	4				1110	1		X

HS18120004

Aptim Environmental & Infrastructure, Inc.
Longhorn Army Ammunition Plant LHAAP-29



HOLD SAMPLE

HOLD SAMPLE

RELINQUISHED BY/AFFILIATION	DATE/TIME	ACCEPTED BY/AFFILIATION	DATE/TIME
<i>[Signature]</i>	11/29/18 1400		11/30/18 09:40

ADDITIONAL COMMENTS/SPECIAL INSTRUCTIONS

Cooler 44426
Temp 2.1
1/2/11
CFO.4



COC ID: LHAAP03-NOV2018-ALSHT-1811-24
PROJECT/CLIENT INFO
 Facility Name Longhorn AAP
 Project Number 501032
 LHAAP-03
 Address 1203-B East Grand Avenue
 PMB 202
 City Marshall
 Postal Code 75670
 Phone Number 713.243.7264
 Project Manager Praveen Srivastava

LABORATORY
 Lab Name ALS Laboratories
 Lab Contact RJ Modashia
 Email RJ.Modashia@alsglobal.com
 Address 10450 Stanciliff Rd., Suite 210
 City Houston
 State TX
 Country USA
 Postal Code 77099
 Phone Number 281.575.2279 or 281.530.5656

RUSH: Page 2 of 3
OTHER INFO
 Email Invoice To FedInvoices@aptim.com
 Email Report To Susan.Huang@aptim.com
 Mail Reports To Susan Huang
 Address 4005 Port Chicago Highway, Suit 200
 City Concord
 State CA
 Country USA
 Shipping Company **Fedex**

SAMPLE DETAILS

Sample ID	Location	Start Depth	End Depth	Depth Unit	Field Matrix	Date	Time (24hr)	# Of Cont.	Metals (As & Pb) by 6020
035B22-3.0-4.0-FD	035B22	3	4	FT	Soil	11/29/18	1110	1	X
035B22-6.0-7.0	035B22	6	7				1113	1	X
035B22-8.0-9.0	035B22	8	9				1115	1	X - HOLD SAMPLE
035B23-0.0-2.0	035B23	0	2				1130	1	X
035B23-3.0-4.0	035B23	3	4				1133	1	X
035B23-6.0-7.0	035B23	6	7				1135	1	X - HOLD SAMPLE
035B24-0.0-2.0	035B24	0	2				1145	1	X
035B25-0.0-2.0	035B25	0	2				1140	1	X
035B26-0.0-2.0	035B26	0	2				1095	1	X
035B26-0.0-2.0-FD	035B26	0	2				1005	1	X
035B26-3.0-4.0	035B26	3	4				1010	1	X
035B26-6.0-7.0	035B26	6	7				1013	1	X
035B26-8.0-9.0	035B26	8	9				1015	1	X - HOLD SAMPLE
035B27-0.0-2.0	035B27	0	2				1155	1	X
035B27-3.0-4.0	035B27	3	4				1200	1	X
035B27-3.0-4.0-MS	035B27	3	4				1200	1	X

ANALYSIS REQUESTED

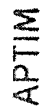
RELINQUISHED BY/AFFILIATION	DATE/TIME	ACCEPTED BY/AFFILIATION	DATE/TIME
<i>AMW</i>	11/29/18 1900	PM / MS	11/30/18 09:40

HS18120004
 Aptim Environmental & Infrastructure, Inc.
 Longhorn Army Ammunition Plant LHAAP-29



ADDITIONAL COMMENTS/SPECIAL INSTRUCTIONS

*Center 44426
 Temp 2.1
 12.11
 Cfo.4*



COC ID: LHAAP03-NOV2018-ALSHT-1811-29
PROJECT/CLIENT INFO
 Facility Name: Longhorn AAP
 Project Number: 501032
 Address: 1203-B East Grand Avenue
 PMB 202
 City: Marshall
 State: TX
 Country: USA
 Postal Code: 75670
 Phone Number: 713.243.7264
 Project Manager: Praveen Srivastav

LABORATORY
 Lab Name: ALS Laboratories
 Lab Contact: RJ Modashia
 Email: RJ.Modashia@alsglobal.com
 Address: 10450 Stanciliff Rd., Suite 210
 City: Houston
 State: TX
 Country: USA
 Postal Code: 77099
 Phone Number: 281.575.2279 or 281.530.5656

RUSH: Page 3 of 3
OTHER INFO
 Email Invoice To: Fedinvoices@aptim.com
 Email Report To: Susan.Huang@aptim.com
 Mail Reports To: Susan Huang
 Address: 4005 Port Chicago Highway, Suit 200
 City: Concord
 State: CA
 Country: USA
 Postal Code: 94520
 Shipping Company: **Fedex**

SAMPLE DETAILS										ANALYSIS REQUESTED									
Sample ID	Location	Start Depth	End Depth	Depth Unit	Field Matrix	Date	Time (24hr)	# Of Cont.	Meats (As & Pb) by 6020	ANALYSIS	RELINQUISHED BY/AFFILIATION	DATE/TIME	ACCEPTED BY/AFFILIATION	DATE/TIME					
035B27-3.0-4.0-MSD	035B27	3	4	FT	S&I	11/29/18	1200	1	X		mm	11/29/18 1900	OM / ALS	11/29/18 09:40					
035B27-6.0-7.0	035B27	6	7				1205	1	X										
035B27-8.0-9.0	035B27	8	9				1208	1	X										
035B28-0.0-2.0	035B28	0	2				1215	1	X										
035B28-3.0-4.0	035B28	3	4				1218	1	X										
035B28-6.0-7.0	035B28	6	7				1220	1	X										
035B28-8.0-9.0	035B28	8	9	✓	✓		1224	1	X	→ HOLD SAMPLE									

HS18120004

Aptim Environmental & Infrastructure, Inc.
 Longhorn Army Ammunition Plant LHAAP-29

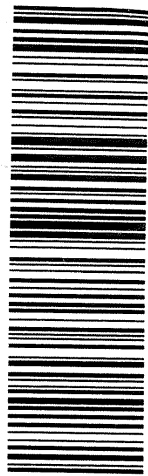


ADDITIONAL COMMENTS/SPECIAL INSTRUCTIONS	RELINQUISHED BY/AFFILIATION	DATE/TIME	ACCEPTED BY/AFFILIATION	DATE/TIME
	mm	11/29/18 1900	OM / ALS	11/29/18 09:40

*Colector 44426
 Temp 2-1
 11/21
 CF-0.4*

 <p>ALS 10450 Stanciliff Rd., Suite 2 Houston, Te 77099 Tel. +1 281 5656 Fax. +1 281 5887</p>	<p>Date: 11/20/16 Name: [Handwritten] Company: [Handwritten]</p>	<p>Shipped By: [Handwritten] Date: 11/20/16</p>
---	--	--

FedEx
 TRK# 0221 4380 9534 7170
AB SGRA 44426 TX-US IAH 77099
 FRI - 30 NOV 10:30A
 PRIORITY OVERNIGHT



4475872 11/29 552J2/E6P/DC65

Appendix B

Sample Collection Logs for November 2018 Soil Samples



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB18**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB18-0.0-2.0

DATE/TIME: 11/29/2018 / 10:00

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB19**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03S819-3.0-4.0

DATE/TIME: 11/29/2018 / 10:20

Sample Interval:
3 - 4 Ft


Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB19**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03S819-6.0-7.0

DATE/TIME: 11/29/2018 / 10:24

Sample Interval:
6 - 7 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB19**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03S819-8.0-9.0

DATE/TIME: 11/29/2018 / 10:26

Sample Interval:
8 - 9 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB19**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB19-0.0-2.0

DATE/TIME: 11/29/2018 / 10:19

Sample Interval:
0 - 2 Ft


Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB19**Project No: **501032**

Sampler(s):

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB19-3.0-4.0

DATE/TIME: 11/29/2018 / 10:20

Sample Interval:
3 - 4 FT


Sampling Method:

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
No COC Specified	None		

Sampler:		
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB19**Project No: **501032**

Sampler(s):

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB19-6.0-7.0

DATE/TIME: 11/29/2018 / 10:24

Sample Interval:
6 - 7 FT


Sampling Method:

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
No COC Specified	None		

Sampler:		
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB20**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB20-0.0-2.0

DATE/TIME: 11/29/2018 / 10:30

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes: 03SB20-0.0-2.0-FD

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB20**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB20-0.0-2.0-FD

DATE/TIME: 11/29/2018 / 10:30

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: FD

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB20**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB20-3.0-4.0

DATE/TIME: 11/29/2018 / 10:35

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB20**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB20-6.0-7.0

DATE/TIME: 11/29/2018 / 10:38

Sample Interval:
6 - 7 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB21**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB21-0.0-2.0

DATE/TIME: 11/29/2018 / 10:45

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes: 03SB21-0.0-2.0-MS and MSD

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB21**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB21-3.0-4.0

DATE/TIME: 11/29/2018 / 10:50

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB21**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB21-6.0-7.0

DATE/TIME: 11/29/2018 / 10:55

Sample Interval:
6 - 7 Ft


Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB22**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB22-0.0-2.0

DATE/TIME: 11/29/2018 / 11:05

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB22**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB22-3.0-4.0

DATE/TIME: 11/29/2018 / 11:10

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes: 03SB22-3.0-4.0-FD

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB22**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB22-3.0-4.0-FD

DATE/TIME: 11/29/2018 / 11:10

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: FD

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB22**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB22-6.0-7.0

DATE/TIME: 11/29/2018 / 11:13

Sample Interval:
6 - 7 Ft


Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB22**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB22-8.0-9.0

DATE/TIME: 11/29/2018 / 11:15

Sample Interval:
8 - 9 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB23**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB23-0.0-2.0

DATE/TIME: 11/29/2018 / 11:30

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB23**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB23-3.0-4.0

DATE/TIME: 11/29/2018 / 11:33

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB23**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB23-6.0-7.0

DATE/TIME: 11/29/2018 / 11:35

Sample Interval:
6 - 7 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB24**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB24-0.0-2.0

DATE/TIME: 11/29/2018 / 11:45

Sample Interval:
0 - 2 Ft


Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB25**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB25-0.0-2.0

DATE/TIME: 11/29/2018 / 11:40

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB26**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB26-0.0-2.0

DATE/TIME: 11/29/2018 / 10:05

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes: 03SB26-0.0-2.0-FD

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB26**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB26-0.0-2.0-FD

DATE/TIME: 11/29/2018 / 10:05

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: FD

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB26**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB26-3.0-4.0

DATE/TIME: 11/29/2018 / 10:10

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB26**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB26-6.0-7.0

DATE/TIME: 11/29/2018 / 10:13

Sample Interval:
6 - 7 Ft


Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB26**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB26-8.0-9.0

DATE/TIME: 11/29/2018 / 10:15

Sample Interval:
8 - 9 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB27**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB27-0.0-2.0

DATE/TIME: 11/29/2018 / 11:55

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB27**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB27-3.0-4.0

DATE/TIME: 11/29/2018 / 12:00

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes: 03SB27-3.0-4.0-MS and MSD

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB27**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB27-6.0-7.0

DATE/TIME: 11/29/2018 / 12:05

Sample Interval:
6 - 7 Ft


Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB27**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB27-8.0-9.0

DATE/TIME: 11/29/2018 / 12:08

Sample Interval:
8 - 9 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
-----------------	---	--------------



Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB28**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB28-0.0-2.0

DATE/TIME: 11/29/2018 / 12:15

Sample Interval:
0 - 2 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB28**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB28-3.0-4.0

DATE/TIME: 11/29/2018 / 12:18

Sample Interval:
3 - 4 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB28**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB28-6.0-7.0

DATE/TIME: 11/29/2018 / 12:20

Sample Interval:
6 - 7 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Sample Collection Log

1 of 1

Project Name: **Longhorn AAP**Location ID: **03SB28**Project No: **501032**Sampler(s): **William Foss**

FIELD CONDITIONS

SAMPLING INFORMATION

Sample No: 03SB28-8.0-9.0

DATE/TIME: 11/29/2018 / 12:24

Sample Interval:
8 - 9 Ft

Sampling Method: DP

Sample Purpose: REG

Sample Matrix: SO

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
LHAAP03-NOV2018-ALSHT-181129	None	METALS	SW6020A

Sampler:		William Foss
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Appendix C

State of Texas Well Completion Report for Monitoring Well 03WW01

STATE OF TEXAS WELL REPORT for Tracking #162475

Owner:	Longhorn Army Ammunition Plant	Owner Well #:	03WW01
Address:	Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Grid #:	35-23-6
Well Location:	Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Latitude:	32° 41' 10" N
Well County:	Harrison	Longitude:	094° 09' 15" W
Elevation:	No Data	GPS Brand Used:	Garmin e-trex
Type of Work: New Well		Proposed Use: Monitor	

Drilling Date: Started: **11/18/2008**
Completed: **11/18/2008**

Diameter of Hole: Diameter: **8.25 in From Surface To 30 ft**

Drilling Method: **Hollow Stem Auger**

Borehole Completion: Gravel Packed From: **18 ft to 30 ft**
Gravel Pack Size: **20/40**

Annular Seal Data: 1st Interval: **From 16 ft to 18 ft with 1 Bentonite (#sacks and material)**
2nd Interval: **From 0 ft to 16 ft with 5 Cement (#sacks and material)**
3rd Interval: **No Data**
Method Used: **Tremmie Pipe**
Cemented By: **Driller**
Distance to Septic Field or other Concentrated Contamination: **No Data**
Distance to Property Line: **No Data**
Method of Verification: **No Data**
Approved by Variance: **No Data**

Surface Completion: **Surface Sleeve Installed**

Water Level: Static level: **No Data**
Artesian flow: **No Data**

Packers: **No Data**

Plugging Info: Casing or Cement/Bentonite left in well: **No Data**

Type Of Pump: **No Data**

Well Tests: **No Data**

Water Quality: Type of Water: **No Data**
Depth of Strata: **No Data**
Chemical Analysis Made: **No Data**
Did the driller knowingly penetrate any strata which contained undesirable constituents: **No**

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the log(s) being returned for completion and resubmittal.

Company Information: **ETTL Engineers & Consultants Inc.**
1717 E. Erwin

Tyler , TX 75702

Driller License Number: **2126**

Licensed Well Driller Signature: **H. Douglas Hinds**

Registered Driller Apprentice Signature: **No Data**

Apprentice Registration Number: **No Data**

Comments: **No Data**

IMPORTANT NOTICE FOR PERSONS HAVING WELLS DRILLED CONCERNING CONFIDENTIALITY

TEX. OCC. CODE Title 12, Chapter 1901.251, authorizes the owner (owner or the person for whom the well was drilled) to keep information in Well Reports confidential. The Department shall hold the contents of the well log confidential and not a matter of public record if it receives, by certified mail, a written request to do so from the owner.

Please include the report's Tracking number (Tracking #162475) on your written request.

**Texas Department of Licensing & Regulation
P.O. Box 12157
Austin, TX 78711
(512) 463-7880**

DESC. & COLOR OF FORMATION MATERIAL

From (ft) To (ft) Description
0-9.5 Sandy lean clay - brown
9.5-15.5 Silty sand - brown-gray
15.5-23.5 Clayey sand - brown
23.5-30 Sandy clay - gray

CASING, BLANK PIPE & WELL SCREEN DATA

Dia.	New/Used	Type	Setting From/To
2	New	PVC Sch. 40	0 - 20
2	New	PVC Sch. 40 - slotted	20 - 30 0.010"



DEPARTMENT OF THE ARMY
LONGHORN ARMY AMMUNITION PLANT
POST OFFICE BOX 220
RATCLIFF, AR 72951

July 11, 2019

DAIM-ODB-LO

Mr. Rich Mayer
U.S. Environmental Protection Agency
Federal Facilities Section R6
1445 Ross Avenue
Dallas, TX 75202-2733

**Re: Draft Final 2018 Remedial Action Operation Report, Landfill 12 (LHAAP-12),
Longhorn Army Ammunition Plant, Karnack, Texas, July 2019**

Dear Mr. Mayer,

One hard copy replacement cover page and one compact disc (CD) of the above-referenced document are being transmitted to you for your review. There were no EPA or TCEQ comments on the Draft document. In accordance with Federal Facility Agreement, this Draft Final will be considered Final after 30 days without further comment.

The document was prepared by Bhate Environmental Associates, Inc., (Bhate) team, on behalf of the Army as part of Bhate's Performance Based Remediation contract for the facility. I ask that Kim Nemmers, Bhate's Project Manager, be copied on any communications related to the project.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in cursive script that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
Longhorn AAP Site Manager

Copies furnished:

- A. Palmie, TCEQ, Austin, TX (letter)
- P. Bruckwicki, Caddo Lake NWR, TX (1 hard copy and 1 CD)
- A. Williams, USACE, Tulsa District, OK (1 CD)
- R. Smith, USACE, Tulsa District, OK (electronic only)
- A. Sherman, USAEC, San Antonio, TX (1 CD)
- K. Nemmers, Bhate, Lakewood, CO (1 CD)
- P. Srivastav, APTIM, Houston, TX (letter)



DEPARTMENT OF THE ARMY
 LONGHORN ARMY AMMUNITION PLANT
 POST OFFICE BOX 220
 RATCLIFF, AR 72951

July 11, 2019

DAIM-ODB-LO

Ms. April Palmie
 Texas Commission on Environmental Quality
 Superfund Section, MC-136
 12100 Park 35 Circle, Bldg D
 Austin, TX 78753

**Re: Draft Final 2018 Remedial Action Operation Report, Landfill 12 (LHAAP-12),
 Longhorn Army Ammunition Plant, Karnack, Texas, May 2019**

Dear Ms. Palmie,

One hard copy replacement cover page and one compact disc (CD) of the above-referenced document are being transmitted to you for your review. There were no EPA or TCEQ comments on the Draft document. In accordance with Federal Facility Agreement, this Draft Final will be considered Final after 30 days without further comment.

The document was prepared by Bhate Environmental Associates, Inc., (Bhate) team, on behalf of the Army as part of Bhate's Performance Based Remediation contract for the facility. I ask that Kim Nemmers, Bhate's Project Manager, be copied on any communications related to the project.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

Rose M. Zeiler, Ph.D.
 Longhorn AAP Site Manager

Copies furnished:

R. Mayer, USEPA Region 6, Dallas, TX (letter)
 P. Bruckwicki, Caddo Lake NWR, TX (1 hard copy and 1 CD)
 A. Williams, USACE, Tulsa District, OK (1 CD)
 R. Smith, USACE, Tulsa District, OK (Electronic only)
 A. Sherman, USAEC, San Antonio, TX (1 CD)
 K. Nemmers, Bhate, Lakewood, CO (1 CD)
 P. Srivastav, APTIM, Houston, TX (letter)



Draft Final
**2018 Remedial Action Operation
 Report, Landfill 12 (LHAAP-12)**
 Longhorn Army Ammunition Plant
 Karnack, Texas



Prepared for:

U.S. Army Corps of Engineers, Tulsa District
 Contracting Division
 2488 East 81st Street
 Tulsa, Oklahoma 74137-4290

Prepared by:



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 Birmingham, Alabama 35205
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Aptim Federal Services, LLC
 2500 CityWest, Suite 1700
 Houston, Texas 77042

Contract No. W9128F-13-D-0012
 Task Order No. W9128BV17F0150
 Project No. 501032
 Rev 0
 July 2019

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2018 REMEDIAL ACTION OPERATION REPORT, LANDFILL 12 (LHAAP-12)

Contract No. W9128F-13-D-0012, Task Order No. W9128BV17F0150 • Draft Final • Rev 0 • July 2019

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Acronyms and Abbreviations

µg/L	micrograms per liter
AECOM	AECOM Technical Services, Inc.
APTIM	Aptim Federal Services, LLC
bgs	below ground surface
Bhate	Bhate Environmental Associates, Inc.
BRAC	Base Realignment and Closure
DCE	dichloroethene
ERS	Environmental Remediation Services
IRA	interim remedial action
LHAAP	Longhorn Army Ammunition Plant
LUC	land use control
MATOC	Multiple Award Task Order Contract
MCL	maximum contaminant level
MEGA	Multiple Environmental Government Acquisition
MMRP	Military Munitions Response Program
MNA	monitored natural attenuation
No.	number
RA(O)	Remedial Action Operation
RAOs	remedial action objectives
RD	remedial design
ROD	record of decision
Shaw	Shaw Environmental, Inc.
TCE	trichloroethene
U.S.	United States
USACE	U.S. Army Corps of Engineers
VC	vinyl chloride
VOC	volatile organic compound

Contract No. W9128F-13-D-0012, Task Order No. W9128BV17F0150 • Draft Final • Rev 0 • July 2019

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1.0 INTRODUCTION

The U.S. Army Corps of Engineers (USACE), Tulsa District, on behalf of the U.S. Army Base Realignment and Closure (BRAC) Division contracted Bhate Environmental Associates, Inc. (Bhate) under the Omaha Multiple Environmental Government Acquisition (MEGA) National Small Business Multiple Award Task Order Contract (MATOC), Environmental Remediation Services (ERS) with Military Munitions Response Program (MMRP), Task Order No. W9128BV17F0150, to conduct environmental restoration at multiple sites at the former Longhorn Army Ammunition Plant (LHAAP). The Bhate Team is comprised of Bhate and Aptim Federal Services, LLC (APTIM). APTIM is conducting the groundwater monitoring and site inspection activities as described in the *Final Remedial Design Addendum Landfill 12 (LHAAP-12)* (Shaw 2007). LHAAP-12 was transferred to the U.S. Fish and Wildlife Service as part of the Caddo Lake National Wildlife Refuge in 2014. This report, summarizing the results of the December 2018 groundwater sampling, the July 2018 landfill integrity inspections and repairs, and the January 2019 well integrity inspections, is the 11th Remedial Action Operation (RA[O]) Report since the Remedial Design (RD) Addendum was finalized in 2007 (Shaw 2007).

1.1 Remedial Action Objectives

Work completed in 2018 as part of the remedial action objectives (RAOs) at LHAAP-12 consisted of the following components: physical inspections and repairs; groundwater monitoring; and concentration trend analysis. Physical inspections are conducted annually, at a minimum, to confirm compliance with land use control (LUC) objectives, which consist of ensuring the integrity of the existing landfill cover, and ensuring no exposure to trichloroethene (TCE)-contaminated groundwater.

The objectives for groundwater monitoring at LHAAP-12 are to:

- Determine the effectiveness of natural attenuation in reducing contaminant concentrations over time
- Evaluate plume migration to ensure that TCE-contaminated groundwater does not impact nearby surface water
- Perform trend analysis of groundwater contaminants of concern: TCE, cis-1,2-dichloroethene (DCE), and vinyl chloride (VC)
- Perform a comparison of the contaminants of concern concentration trends relative to the maximum contaminant levels (MCLs)

The remedy at LHAAP-12 is functioning as intended and all human risks are currently under control. This report summarizes the results of groundwater sampling and the landfill integrity inspections and repairs.

1.2 Site Description

LHAAP-12 was established in 1963, covers approximately seven acres, and is 500 feet southeast of Central Creek, which eventually drains into Caddo Lake. LHAAP-12 was used for the disposal of industrial solid wastes, possibly containing small quantities of hazardous constituents generated at LHAAP. From 1978 until its closure in April 1994, the landfill was used continuously for the disposal of non-hazardous industrial solid waste, including cafeteria waste, chemical waste, petroleum-contaminated soil, and asbestos. The construction of a landfill cap over the site was completed in 1998 as part of an interim remedial action (IRA). The IRA is consistent with U.S. Environmental Protection Agency presumptive remedy guidance. The final remedy for LHAAP-12 is protective of human health and the environment and consists of LUCs in conjunction with monitored natural attenuation (MNA). LUCs include those set in place by the *Final Record of Decision, Landfill 12 (LHAAP-12)* (Shaw 2006) for landfill cap maintenance and restriction of groundwater use.

2.0 OPERATIONS AND MAINTENANCE ACTIVITIES AT LHAAP-12

Operation, maintenance and physical inspection activities at LHAAP-12 consist of:

- Maintenance of the integrity of the landfill cap and repairs to desiccation cracks, erosion, or gulying upon observance.
- Maintenance of the vegetative cover on the landfill cap, including periodic mowing.
- Maintenance of the signage around the landfill cap.
- Prohibition of any activities that would affect the integrity of the cap.
- Prohibition of any activities that would cause exposure to contaminated groundwater.

Inspections of the well field are conducted annually, and the information is recorded on well inspection forms (**Appendix A**). Maintenance or repairs are then conducted, as needed, based on observed conditions.

Mowing the sampling area for brush and weeds is conducted on a routine basis, and as needed, to access the well field. Mowing was conducted on 3 May 2018.

The RD Addendum (Shaw 2007) stipulated physical inspections to be conducted annually and annual groundwater sampling to monitor the effectiveness of natural attenuation in reducing contaminant concentrations over time. Inspections are actually conducted more frequently with periodic drive-by oversight to ensure compliance with the above requirements. The effectiveness of MNA was initially evaluated quarterly in the *Draft Annual Remedial Action Operation Report, Years 1 and 2, Landfill 12 (LHAAP-12)* (Shaw 2011) and has been evaluated annually beginning in the 3rd year of RAO. Groundwater monitoring was included as a component of the remedy for LHAAP-12 identified in the Record of Decision (ROD) (Shaw 2006).

2.1 Physical Inspection Summary

During the July 2018 inspection, the landfill appeared to be in good condition. Four holes were observed in the vegetative cover; otherwise, the vegetative cover was observed in good condition and well maintained during routine mowing with no excessive cracking or desiccation cracks. No change in land or groundwater use was observed at the site, and the use of the site is consistent with the LUCs mandated by the ROD (Shaw 2006). The holes in the

vegetative cover were repaired on 1 Feb 2019 by filling them with soil and the areas were allowed to vegetate naturally. Photos of the signs and the repaired holes in the cover are provided in **Appendix A**. Signs were all visible around the landfill. A follow-up well inspection was conducted in January 2019 to document the condition of the wells. No damage to bollards, pads, or protective casings was observed, and minimal encroachment of weeds or brush on the well pads was observed. Well head locks continue to be in good condition. The January 2019 well inspection forms are provided in **Appendix A**.

The official 2018 physical inspection of the LHAAP-12 cap was completed on 26 Jul 2018 and the LUC Inspection and Maintenance Log is included in **Appendix B**.

2.2 Groundwater Monitoring and Analytical Results

The purpose of sampling was to develop data to support the effectiveness of natural attenuation in reducing contaminant concentrations over time, to evaluate plume migration, and to ensure that TCE-contaminated groundwater does not impact nearby surface water. As described by the Groundwater Sampling Plan included in the RD Addendum (Shaw 2007), annual groundwater sampling of three on-site monitoring wells (12WW20, 12WW21, and 12WW24) and two downgradient compliance monitoring wells (12WW22 and 12WW23) was to be conducted annually beginning in the third year of the RA(O) (i.e., 2010). As recommended in the *Draft Final 2014 Remedial Action Operation Report* (AECOM 2015), the sampling frequency of monitoring well 12WW22 was changed to once every five years and sampling was discontinued at well 12WW23. The locations of these groundwater monitoring wells are shown in **Figure 2-1**. The sampling event in December 2018 included the three on-site wells 12WW20, 12WW21, and 12WW24.

All monitoring wells were in good condition and no evidence of turbidity or silt accumulation was identified during the sampling and gauging event. The groundwater elevations from the three wells sampled in December 2018 are included in **Table 2-1**. The remaining wells were inadvertently not gauged during the December 2018 sampling event. Therefore, in January 2019, an additional gauging event was conducted during the physical inspection of the wells which included eight wells at LHAAP-12 (12WW01, 12WW02, 12WW05, 12WW20, 12WW21, 12WW22, 12WW23, and 12WW24). The groundwater elevation measurements collected in January 2019 indicate an easterly groundwater flow as shown on **Figure 2-2**. The groundwater elevation in January 2019 was slightly higher than in December 2018. The hydraulic gradient based on January 2019 data was 0.006 feet vertical per foot horizontal. The well siltation measurements from the January 2019 gauging event have been included in **Appendix A**. Based on these observations, the wells at LHAAP-12 do not need additional repair.

Groundwater samples were collected in December 2018 using the low-flow method in accordance with Bhate's *Standard Operating Procedures, Groundwater Sampling Procedures* (Bhate 2018). The objective of using low-flow sampling methods is to collect representative samples of the groundwater in the formation adjacent to the well screen, eliminating the mixing of stagnant water above and below the well screen. After the water quality parameters stabilized and were within acceptable ranges, samples were collected at the same low flow rate. Groundwater monitoring well sampling forms were completed for each monitoring well sampled and are included in **Appendix C**. Laboratory analytical results are included in **Appendix D**.

Annual groundwater samples were collected on 4 Dec 2018 from 12WW20, 12WW21, and 12WW24, and were analyzed for volatile organic compounds (VOCs) according to SW-846 Method 8260. No VOCs were detected in monitoring wells 12WW20 and 12WW21. In monitoring well 12WW24, TCE was the only VOC that exceeded its respective MCL of 5 micrograms per liter ($\mu\text{g/L}$). TCE concentration at monitoring well 12WW24 reduced from 83 $\mu\text{g/L}$ in December 2017 to 48 $\mu\text{g/L}$ in December 2018. Analytical results for the samples are shown in **Table 2-2**. Historical TCE concentrations for LHAAP-12 wells are shown in **Table 2-3**. Historic contaminants of concern concentrations for monitoring well 12WW24 are shown on **Table 2-4**.

The MNA evaluation completed as part of the *Final 2013 Five-Year Review Report* (AECOM 2014a), concluded that TCE degradation was occurring via anaerobic reductive dechlorination. Although an increasing trend in TCE concentration was observed in well 12WW24 in December 2013 and January 2015, concentrations have decreased over the past four sampling events, December 2015, December 2016, December 2017, and December 2018 (see **Figure 2-3** for 12WW24). TCE concentrations appear to inversely correlate with fluctuating water levels because samples collected when the water table is high tend to have lower contaminant concentrations as shown on **Figure 2-3**. The presence of cis-1,2-DCE and VC indicates that degradation by anaerobic reductive dechlorination is occurring (see **Table 2-4** and **Figure 2-3**). However, the concentration trends for these degradation products mimic those of TCE, suggesting that dilution and dispersion is the dominant natural attenuation pathway. Groundwater TCE concentrations for all wells can be found in **Table 2-3**. The December 2018 concentrations are shown on **Figure 2-4**.

Statistical analysis of the concentration profile for TCE in 12WW24 using Mann-Kendall trend analysis was completed for the data collected from 2006 to 2018. The results are shown in **Appendix E**. The Mann-Kendall trend test indicates a statistically significant decreasing trend at the 99% confidence level for TCE concentrations in monitoring well 12WW24. A drought persisted between 2011 and 2012. This resulted in decreased groundwater levels and well 12WW24 being dry and not being sampled in 2012. After the groundwater level in 12WW24

had recovered, TCE concentrations increased during the December 2013 and January 2015 sampling events, before declining during the December sampling events in 2015, 2016, 2017 and 2018. Additional annual sampling is necessary for the next five year review when the trends will continue to be evaluated for statistical significance and a decrease of sampling frequency. The decreasing contaminant trend provide evidence that the plume is stable and natural attenuation is occurring, resulting in an overall decrease in TCE concentration in groundwater over time. The presence of cis-1,2-DCE and VC in 12WW24 since 2013 also indicates that biodegradation has been occurring.

The TCE trend in 12WW24 as shown in **Figure 2-3** was used to calculate a first order decay rate (**Appendix F**). Two calculations have been made; the first for the dataset through June 2011 and the second for the dataset that includes all results through December 2018. The reason for the two calculations is because drought conditions evidenced by the dry well in December 2012 later manifested in an increase in TCE concentration in this well in the December 2013, and January 2015 sampling events. However, the water levels have been increasing since the well was dry in December 2012 and an overall decrease in TCE concentration has also been observed. The first order rate constant for the period through June 2011 was 6.0 E-04 per day while the first order rate constant for the entire dataset was 2.4 E-04 per day, an implied decrease of less than one order of magnitude. Based on these rate constants, the estimated durations to achieve the MCL for TCE in 12WW24 are 15 years using the rate constant for the dataset through June 2011 and approximately 26 years using the rate constant for the entire dataset. The drought conditions along with changes in seasonal sampling have constrained the trends observed.

3.0 CONCLUSIONS AND RECOMMENDATIONS

Physical inspections and groundwater monitoring continue at LHAAP-12 in compliance with the ROD for LHAAP-12 (Shaw 2006). No damage to bollards, pads, or protective casings was observed, and limited encroachment of weeds or brush on the well pads was observed. No change in land or groundwater use has occurred at the site. Four holes were identified in the center of the landfill cap during the official 2018 physical inspection of the LHAAP-12 cap completed on 26 Jul 2018. The holes were covered with dirt in February 2019 and allowed to vegetate naturally. LUCs were verified, and the use of the site is still consistent with that mandated by the ROD. Annual LUC inspections will continue to be conducted annually.

In accordance with the Groundwater Sampling Plan, found in Appendix A of the RD Addendum for LHAAP-12 (Shaw 2007), annual sampling of wells will continue until the next Five-Year Review. Results for year 11 are documented in this report.

The groundwater gauging event from January 2019 indicates an easterly gradient similar to the most prevalent historic flow direction. The monitoring wells will continue to be gauged and inspected as part of future sampling events.

Monitoring wells 12WW20, 12WW21 and 12WW24 were sampled on 4 Dec 2018 for VOCs. No VOCs were detected in 12WW20 and 12WW21. At 12WW24, TCE was detected above its MCL of 5 µg/L at a concentration of 48 µg/L. This was a decrease since the December 2017 TCE concentration of 83 µg/L. None of the other detected compounds exceeded their respective MCLs. TCE concentrations in 12WW24 show a statistically significant decreasing trend since 2006 based on Mann-Kendall analysis (**Appendix E**).

When compared to VOC concentrations, the groundwater elevation data (**Figure 2-3**) exhibits an inverse correlation with VOC concentrations in 12WW24. The increase in TCE in the December 2013 and January 2015 sampling events in 12WW24 is thought to be associated with low water levels and sampling during drought conditions, as these were the first samples collected since the well was dry due to drought conditions in 2012. Water levels have continued to increase since 2012, and the TCE concentration from the 4 Dec 2018 sampling is the lowest value measured during the 2006-2018 period.

The Five Year review evaluations indicate the need for an additional well to the east, dowgradient of monitoring well 12WW24 to define the plume (US Army, 2019). The well will be installed at the location shown on **Figure 3-1** and sampled following development. The well will be installed in accordance with the Installation Wide Work Plan (Bhate, 2018) with the same screened interval as 12WW24 from 15.5 to 25.5 feet below ground surface (bgs) unless

different lithologies are noted during drilling. The decision to add the well to the annual monitoring list will be based on the initial sample results and consultation with the regulators.

The three wells (12WW20, 12WW21, and 12WW24) will continue to be sampled annually for VOCs. Any change in this frequency will be evaluated in the next five year review.

4.0 REFERENCES

AECOM. 2014a. *Final 2013 Five-Year Review Report*. Longhorn Army Ammunition Plant, Karnack, Texas. May.

AECOM. 2015. *Draft Final 2014 Remedial Action Operation Report*. Longhorn Army Ammunition Plant, Karnack, Texas. May.

Bhate. 2018. *Final Installation Wide Work Plan*. Longhorn Army Ammunition Plant, Karnack, Texas. March.

Shaw Environmental, Inc. (Shaw). 2006. *Final Record of Decision, Landfill 12 (LHAAP-12), Longhorn Army Ammunition Plant, Karnack, Texas*. August.

Shaw. 2007. *Final Remedial Design Addendum Landfill 12 (LHAAP-12), Longhorn Army Ammunition Plant, Karnack, Texas*. June.

Shaw. 2011. *Draft Annual Remedial Action Operation Report, Years 1 and 2, Landfill 12 (LHAAP-12), Longhorn Army Ammunition Plant, Karnack, Texas*. June.

U.S. Army, 2019. *Draft Five Year Review Report for Longhorn Army Ammunition Plant, Karnack, Texas*. February.

Contract No. W9128F-13-D-0012, Task Order No. W9128BV17F0150 • Draft Final • Rev 0 • July 2019

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Tables

Table 2-1
Groundwater Elevation Data, LHAAP-12

Well ID	Sampling Date	Top of Casing Elevation (ft)	Depth to Water from TOC (ft)	Groundwater Elevation (ft amsl)	Total Depth bgs (ft)	Screen Interval bgs (ft)
12WW01	12/1/2016	204.19	23.60	180.59	26.5	14.50 - 26.50
	3/27/2018	204.19	23.55	180.64		
	1/29/2019	204.19	22.99	181.20		
12WW02	12/1/2016	202.45	22.07	180.38	28.50	13.50 - 28.50
	3/27/2018	202.45	21.82	180.63		
	1/29/2019	202.45	20.93	181.52		
12WW05	12/1/2016	190.52	9.88	180.64	38.00	15.50 - 35.50
	3/27/2018	190.52	8.05	182.47		
	1/29/2019	190.52	6.95	183.57		
12WW20	6/15/2010	199.15	17.98	181.17	38.85	28.85 - 38.85
	6/7/2011	199.15	19.78	179.37		
	12/3/2012	199.15	22.03	177.12		
	1/8/2014	199.15	20.71	178.44		
	1/9/2015	199.15	19.34	179.81		
	12/19/2015	199.15	19.00	180.15		
	12/1/2016	199.15	18.69	180.46		
	12/4/2017	199.15	18.8	180.35		
	3/27/2018	199.15	17.6	181.55		
	12/4/2018	199.15	18.35	180.8		
1/29/2019	199.15	16.81	182.34			
12WW21	6/15/2010	202.07	21.85	180.22	41.70	31.70 - 41.70
	6/7/2011	202.07	23.35	178.72		
	12/3/2012	202.07	24.21	177.86		
	1/8/2014	202.07	24.56	177.51		
	1/9/2015	202.07	24.35	177.72		
	12/19/2015	202.07	22.95	179.12		
	12/1/2016	202.07	21.97	180.10		
	12/4/2017	202.07	21.92	180.15		
	3/27/2018	202.07	21.42	180.65		
	12/4/2018	202.07	22.2	179.87		
1/29/2019	202.07	21.00	181.07			
12WW22	6/15/2010	190.20	7.93	182.27	38.36	28.36 - 38.36
	6/7/2011	190.20	9.72	180.48		
	12/3/2012	190.20	13.46	176.74		
	1/8/2014	190.20	10.57	179.63		
	1/9/2015	190.20	9.60	180.60		
	12/1/2016	190.20	10.32	179.88		
	3/27/2018	190.20	7.81	182.39		
	1/29/2019	190.20	6.93	183.27		
12WW23	6/16/2010	196.97	18.28	178.69	25.14	15.14 - 25.14
	6/7/2011	196.97	19.40	177.57		
	12/3/2012	196.97	20.38	176.59		
	1/8/2014	196.97	20.80	176.17		
	1/9/2015	196.97	19.34	177.63		
	12/1/2016	196.97	17.23	179.74		
	3/27/2018	196.97	16.88	180.09		
1/29/2019	196.97	16.02	180.95			

Table 2-1
Groundwater Elevation Data, LHAAP-12

Well ID	Sampling Date	Top of Casing Elevation (ft)	Depth to Water from TOC (ft)	Groundwater Elevation (ft amsl)	Total Depth bgs (ft)	Screen Interval bgs (ft)
12WW24	6/16/2010	203.17	22.50	180.67	26.00	15.50 - 25.50
	6/7/2011	203.17	24.15	179.02		
	12/3/2012	203.17	Dry	Dry		
	12/9/2013	203.17	26.00	177.17		
	1/9/2015	203.17	24.22	178.95		
	12/18/2015	203.17	23.50	179.67		
	11/30/2016	203.17	22.53	180.64		
	12/4/2017	203.17	22.61	180.56		
	3/27/2018	203.17	22.35	180.82		
	12/4/2018	203.17	22.87	180.30		
	1/29/2019	203.17	21.61	181.56		

Notes:

amsl - above mean sea level

bgs - below ground surface

ft - feet

ID - identification

TOC - top of casing

Table 2-2
2018 Sample Results, LHAAP-12

			Location Code		12WW20				12WW21				12WW24	
			Sample ID		12WW20-181204		12WW21-181204		12WW21-181204-FD		12WW24-181204			
			Sample Date		12/4/2018		12/4/2018		12/4/2018		12/4/2018			
			Sample Purpose		REG		REG		FD		REG			
Parameter	Units	MCL / GW-Ind	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual				
Field Tests														
Conductivity	mS/cm	NV	1.45		5.42				1.55					
Dissolved oxygen	mg/L	NV	0.55		0.08				0.04					
Flow Rate	mL/min	NV	100		100				100					
Oxidation-Reduction Potential	mV	NV	249		202				119					
pH	STD UNIT	NV	5		5.51				5.45					
Temperature	°C	NV	17.8		16				14.68					
Total Purge	L	NV	3		3				3					
Turbidity	NTU	NV	3.7		34.4				3.9					
Water Level at Reading Time	ft btoc	NV	18.58		22.43				22.87					
VOCs														
1,1,1,2-Tetrachloroethane	µg/L	110	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,1,1-Trichloroethane	µg/L	200	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,1,2,2-Tetrachloroethane	µg/L	14	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,1,2-Trichloro-1,2,2-Trifluoroethane	µg/L	3,100,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,1,2-Trichloroethane	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,1-Dichloroethane	µg/L	10,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	< 0.5	U	0.71	J				
1,1-Dichloropropene	µg/L	2.9	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,2,3-Trichlorobenzene	µg/L	310	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,2,3-Trichloropropane	µg/L	0.041	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,2,4-Trichlorobenzene	µg/L	70	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,2,4-Trimethylbenzene	µg/L	5,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,2-Dibromo-3-chloropropane	µg/L	0.2	< 1	U	< 1	U	< 1	U	< 1	U				
1,2-Dibromoethane	µg/L	0.05	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,2-Dichlorobenzene	µg/L	600	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				
1,2-Dichloroethane	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U				

Table 2-2
2018 Sample Results, LHAAP-12

			Location Code		12WW20		12WW21			12WW24	
			Sample ID		12WW20-181204		12WW21-181204		12WW21-181204-FD	12WW24-181204	
			Sample Date		12/4/2018		12/4/2018		12/4/2018	12/4/2018	
			Sample Purpose		REG		REG		FD	REG	
Parameter	Units	MCL / GW-Ind	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	
1,2-Dichloropropane	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
1,3,5-Trimethylbenzene	µg/L	5,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
1,3-Dichlorobenzene	µg/L	3,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
1,3-Dichloropropane	µg/L	29	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
1,4-Dichlorobenzene	µg/L	75	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
2,2-Dichloropropane	µg/L	42	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
2-Butanone	µg/L	61,000	< 1	U	< 1	U	< 1	U	< 1	U	
2-Chlorotoluene	µg/L	2,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
2-Hexanone	µg/L	6,100	< 1	U	< 1	U	< 1	U	< 1	U	
4-Chlorotoluene	µg/L	2,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Acetone	µg/L	92,000	< 2	U	< 2	U	< 2	U	< 2	U	
Benzene	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Bromobenzene	µg/L	2,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Bromochloromethane	µg/L	4,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Bromodichloromethane	µg/L	4.6	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Bromoform	µg/L	36	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Bromomethane	µg/L	140	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Carbon disulfide	µg/L	10,000	< 1	U	< 1	U	< 1	U	< 1	U	
Carbon tetrachloride	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Chlorobenzene	µg/L	100	< 0.5	U	< 0.5	U	< 0.5	U	1.5	J	
Chloroethane	µg/L	41,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Chloroform	µg/L	1,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Chloromethane	µg/L	220	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
cis-1,2-Dichloroethene	µg/L	70	< 0.5	U	< 0.5	U	< 0.5	U	22		
cis-1,3-Dichloropropene	µg/L	5.3	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Dibromochloromethane	µg/L	34	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Dibromomethane	µg/L	380	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	

Table 2-2
2018 Sample Results, LHAAP-12

			Location Code		12WW20		12WW21			12WW24	
			Sample ID		12WW20-181204		12WW21-181204		12WW21-181204-FD	12WW24-181204	
			Sample Date		12/4/2018		12/4/2018		12/4/2018	12/4/2018	
			Sample Purpose		REG		REG		FD	REG	
Parameter	Units	MCL / GW-Ind	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	
Dichlorodifluoromethane	µg/L	20,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Ethylbenzene	µg/L	700	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Hexachlorobutadiene	µg/L	20	< 1	U	< 1	U	< 1	U	< 1	U	
Isopropylbenzene	µg/L	10,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
m,p-Xylenes	µg/L	10,000	< 1	U	< 1	U	< 1	U	< 1	U	
Methyl isobutyl ketone	µg/L	8,200	< 1	U	< 1	U	< 1	U	< 1	U	
Methylene chloride	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Naphthalene	µg/L	2,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
n-Butylbenzene	µg/L	4,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
n-Propylbenzene	µg/L	4,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
o-Xylene	µg/L	10,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
p-Isopropyltoluene	µg/L	10,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
sec-Butylbenzene	µg/L	4,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Styrene	µg/L	100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
tert-Butylbenzene	µg/L	4,100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Tetrachloroethene	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Toluene	µg/L	1,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
trans-1,2-Dichloroethene	µg/L	100	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
trans-1,3-Dichloropropene	µg/L	29	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Trichloroethene	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	48		
Trichlorofluoromethane	µg/L	31,000	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	

Notes:

Highlighted **bold** result indicates concentration above the MCL.

µg/L - micrograms per liter

< - less than

°C - degrees Celsius

Table 2-2
2018 Sample Results, LHAAP-12

			12WW20		12WW21			12WW24		
			12WW20-181204		12WW21-181204	12WW21-181204-FD	12WW24-181204			
			12/4/2018		12/4/2018	12/4/2018	12/4/2018			
			REG		REG	FD	REG			
Parameter	Units	MCL / GW-Ind	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual

Notes: (cont.)

FD - field duplicate

ft btoc - feet below top of casing

GW-Ind - groundwater medium-specific concentration for industrial use

ID - identification

J - estimated value

L - liter

MCL - maximum contaminant level

mg/L - milligrams per liter

mL/min - milliliters per minute

mS/cm - millisiemens per centimeter

mV - millivolts

NTU - nephelometric turbidity unit

NV - not established

REG - regular sample

STD UNIT - standard unit

U - Not detected; The analyte was analyzed for but not detected above the associated method detection limit.

Val Qual - validation qualifier

VOCs - volatile organic compounds

Table 2-3
Historic TCE Concentrations at LHAAP-12 Monitoring Wells

Sampling Date	Monitoring Wells					
	12WW20 (µg/L)	12WW21 (µg/L)	12WW22 (µg/L)	12WW23 (µg/L)	12WW24 (µg/L)	12WW25(30) ^a (µg/L)
1 Dec 2006	0.713	1 U	1 U	1 U	396	NS
1 Sep 2007	1.34	1 U	1 U	1 U	272	NS
1 Dec 2007	1.19	1 U	1 U	1 U	313	NS
1 Mar 2008	0.999 J	0.25 U	0.25 U	0.25 U	301	NS
1 Jun 2008	1.04	0.25 U	0.25 U	0.25 U	237	NS
1 Sep 2008	0.985	0.25 U	0.25 U	0.25 U	185	NS
1 Feb 2009	1.18	0.25 U	0.25 U	0.25 U	334	NS
1 Apr 2009	0.997	0.25 U	0.25 U	0.25 U	197	NS
1 Jul 2009	0.931	0.25 U	0.25 U	0.25 U	204	NS
1 Jun 2010	0.353 J	0.25 U	0.25 U	0.25 U	145	NS
1 Jun 2011	0.263 J	0.25 U	0.25 U	0.25 U	147	NS
1 Dec 2012	0.5 J	0.582 J	0.5 U	0.5 U	Dry Well	NS
1 Jan 2014	5	0.721 J	0.5 U	0.5 U	259 ^b	NS
1 Aug 2014	NS	NS	NS	NS	NS	0.317 J
1 Jan 2015	0.293 J	0.25 U	0.25 U	0.25 U	353	NS
1 Dec 2015	0.5 U	0.5 U	NS	NS	278	NS
1 Dec 2016	0.5 U	0.5 U	0.5 U	0.5 U	151	NS
1 Dec 2017	1 U	1 U	NS	NS	83	NS

Notes:

^a This was a 2014 DPT boring grab groundwater sample, no well was subsequently installed.

^b Analyte was diluted 5X

µg/L - micrograms per liter

J - estimated value

DPT - direct-push technology

NS - not sampled

TCE - trichloroethene

U - Not detected; The analyte was analyzed for but not detected above the associated method detection limit.

Table 2-4
Historic COC Concentrations at Monitoring Well 12WW24

Sampling Date	Monitoring Wells		
	Trichloroethene (µg/L)	cis-1,2-DCE (µg/L)	Vinyl Chloride (µg/L)
1 Dec 2006	396	1 U	1 U
1 Sep 2007	272	1 U	1 U
1 Dec 2007	313	1 U	1 U
1 Mar 2008	301	0.25 U	0.25 U
1 Jun 2008	237	0.25 U	0.25 U
1 Sep 2008	185	0.25 U	0.25 U
1 Feb 2009	334	0.25 U	0.25 U
1 Apr 2009	197	0.25 U	0.25 U
1 Jul 2009	204	0.25 U	0.25 U
1 Jun 2010	145	0.25 U	0.25 U
1 Jun 2011	147	0.25 U	0.25 U
11 Dec 2013	259	66.1	0.873
1 Jan 2015	353	70.8	1.25
1 Dec 2015	278	58.5	0.786
1 Dec 2016	151	36.7	0.51
1 Dec 2017	83	26	1 U
4 Dec 2018	48	22	0.5 U

Notes:

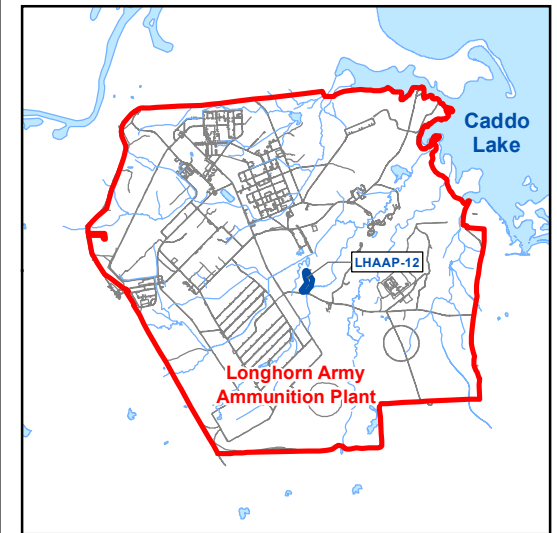
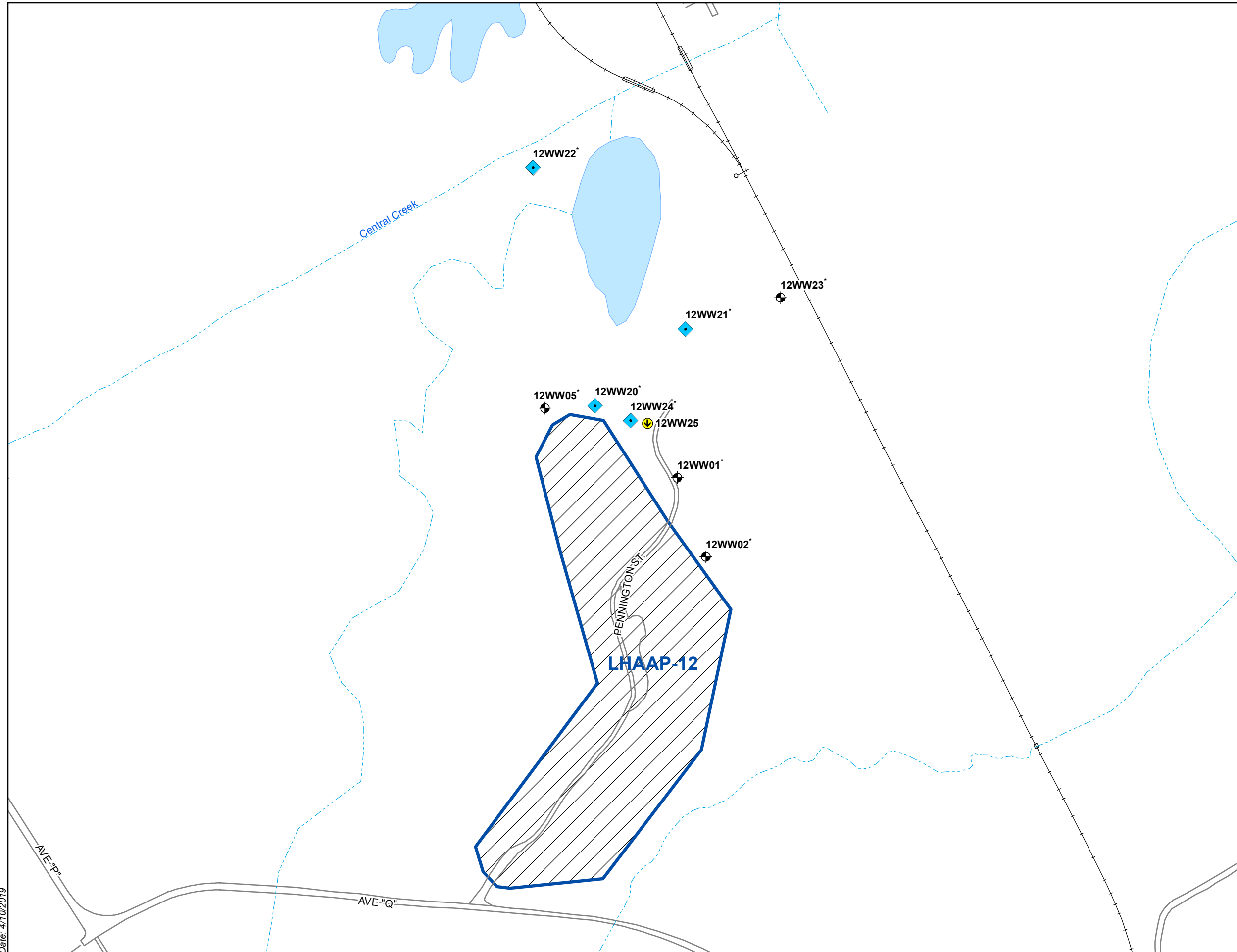
µg/L - micrograms per liter

COC - contaminant of concern

DCE - dichloroethene

U - Not detected; The analyte was analyzed for but not detected above the associated method detection limit.

Figures



- Monitoring Well
- Monitored Natural Attenuation Shallow Well
- 2014 DPT Location (This was a DPT boring grab groundwater sample, no well was subsequently installed)
- Stream
- Road
- Site Boundary
- Landfill Cap Area
- Railroad

Note:
 1. Each well has 5'x5' concrete surface monument surrounded bollards. Each well has a PVC riser. Appropriate equipment and care used around these features.
 2. * Gauged for water levels.



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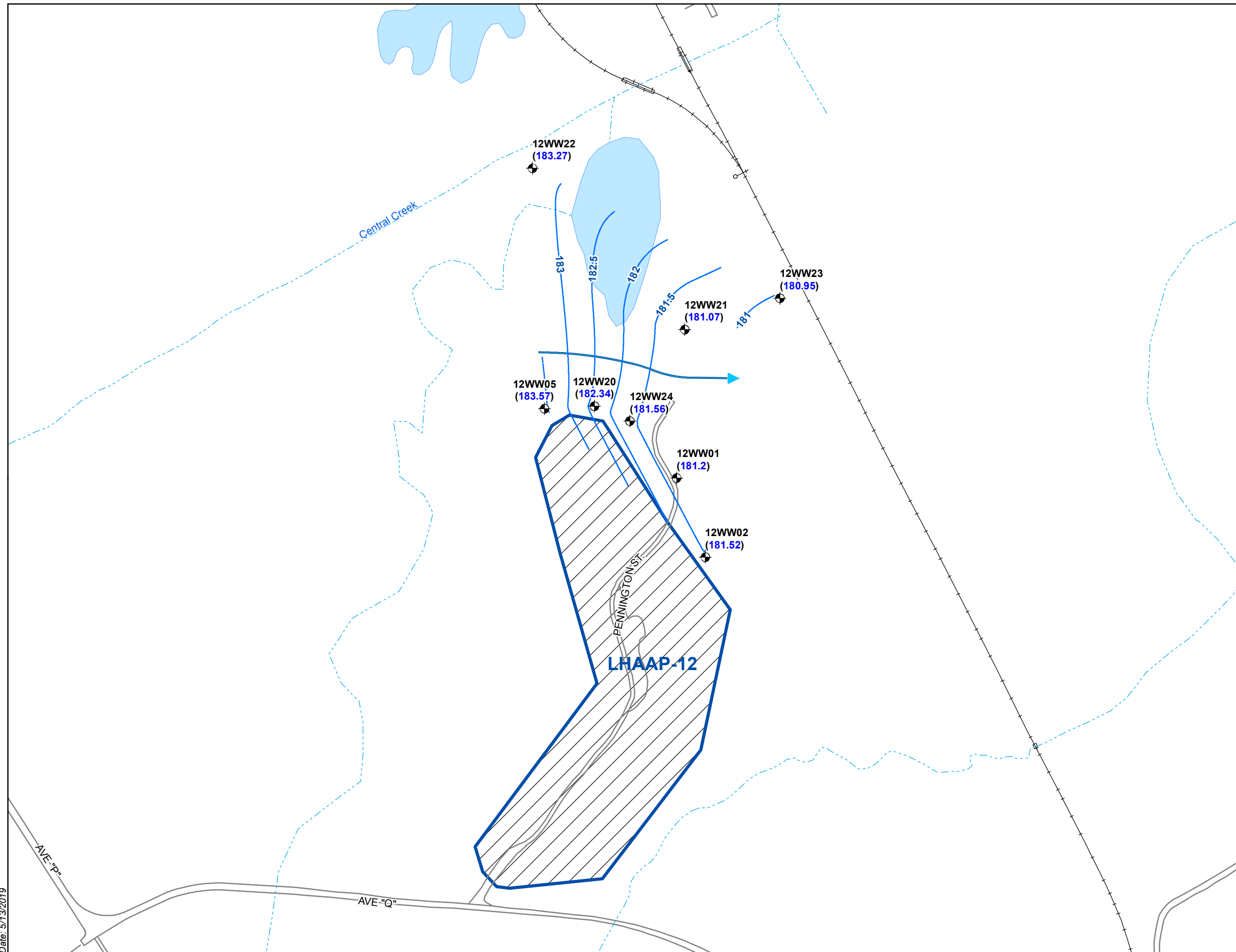


Figure 2-1

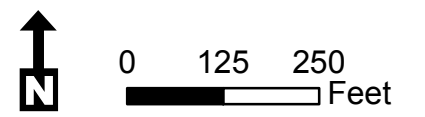
Groundwater Monitoring Well Locations
 LHAAP-12

LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS

Date: 4/10/2019



- Shallow Monitoring Well
- Groundwater Contour
- Stream
- Groundwater Flow Direction
- Road
- Site Boundary
- Landfill Cap Area
- Railroad



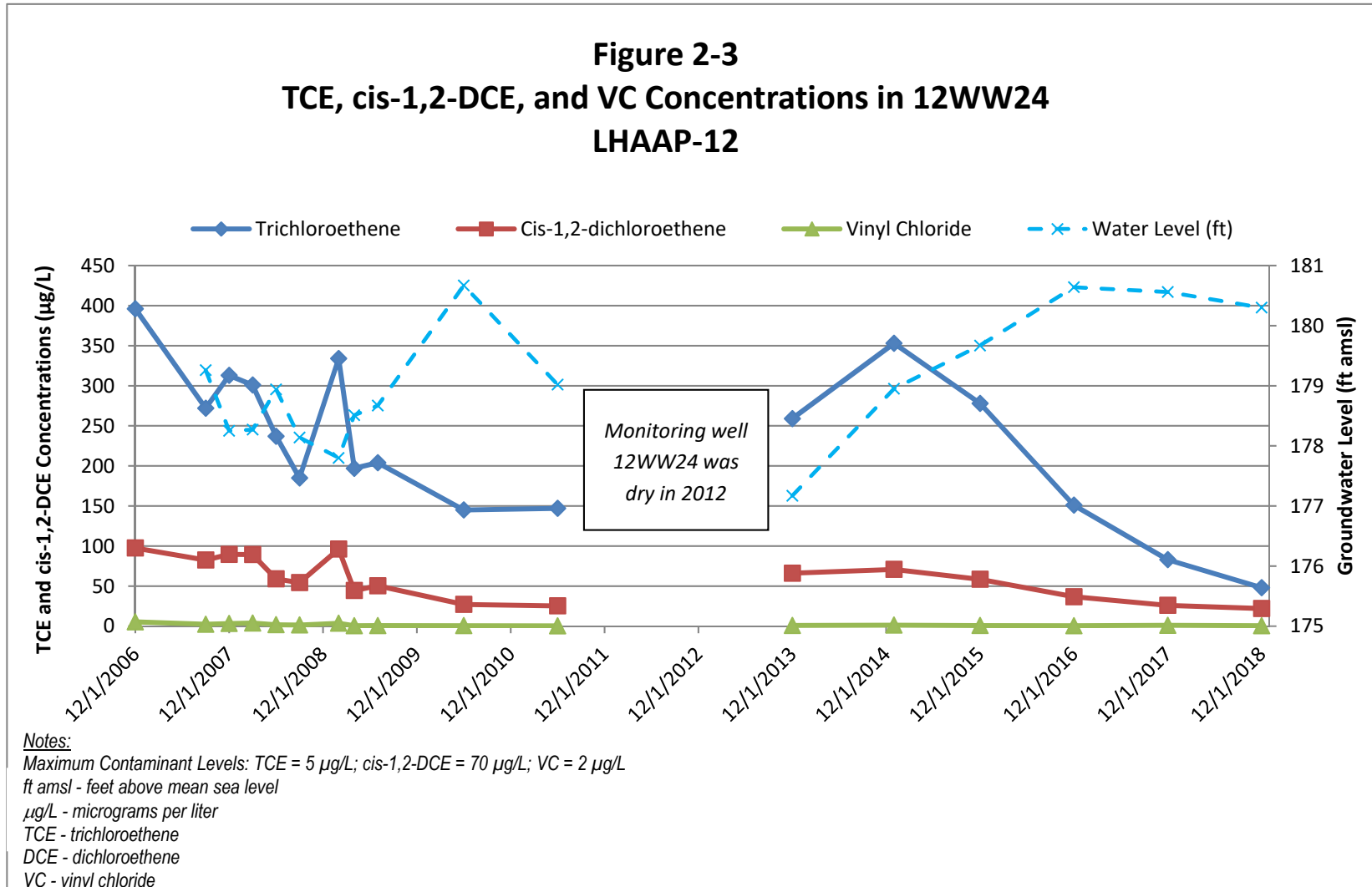
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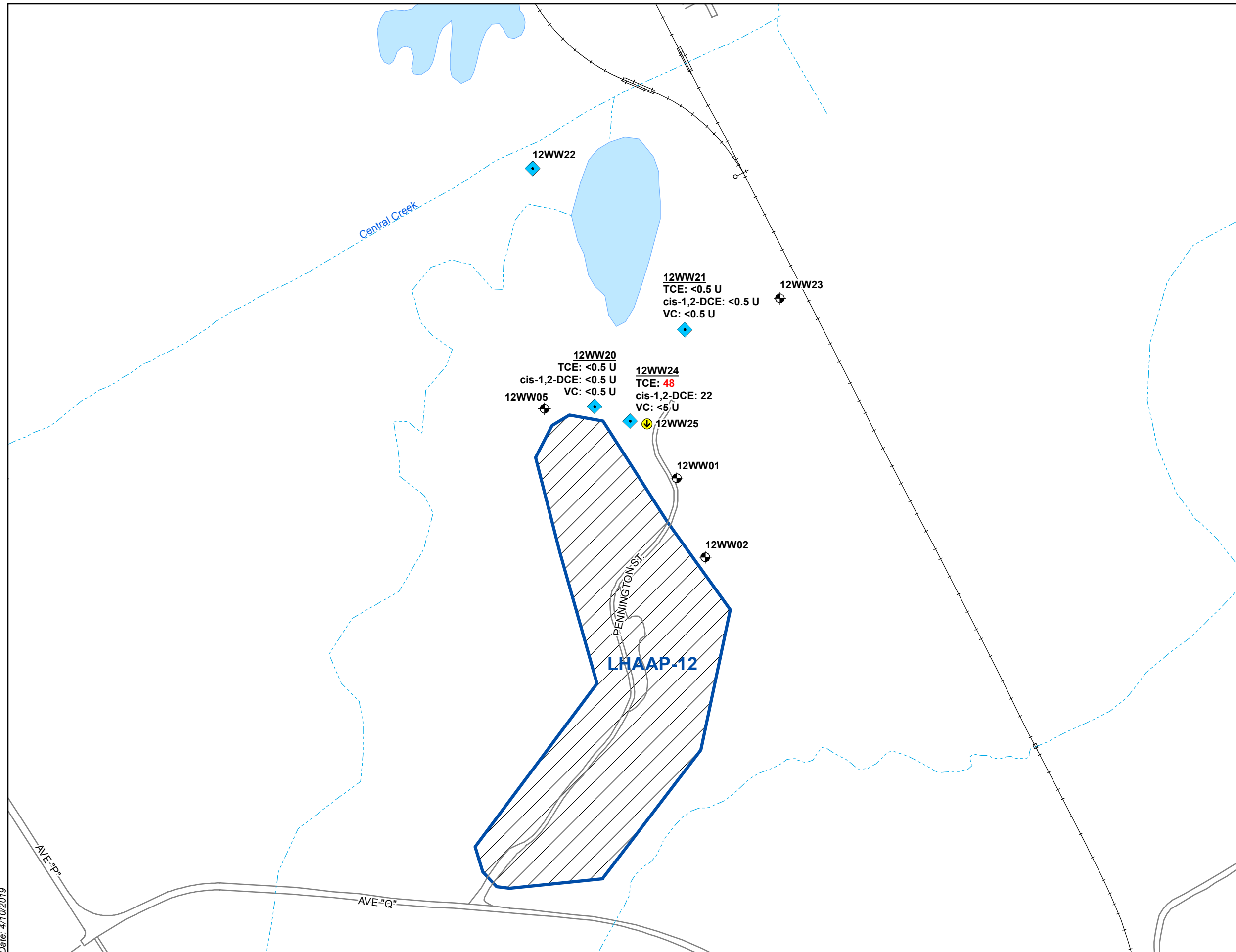


Figure 2-2
Shallow Groundwater Gradient Map
January 2019
LHAAP-12
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

Date: 5/13/2019

**Figure 2-3
TCE, cis-1,2-DCE, and VC Concentrations in 12WW24
LHAAP-12**





- Monitoring Well
- Monitored Natural Attenuation Shallow Well
- 2014 DPT Location (This was a DPT boring grab groundwater sample, no well was subsequently installed)
- Stream
- Road
- Site Boundary
- Landfill Cap Area
- Railroad

Note:

1. Each well has 5'x5' concrete surface monument surrounded bollards. Each well has a PVC riser. Appropriate equipment and care used around these features.
2. Results are in micrograms per liter (µg/L)
- U - Not Detected
- TCE - trichloroethene
- DCE - dichloroethene
- VC - Vinyl Chloride
3. Concentrations in red are above the Maximum Contaminant Limit (MCL)

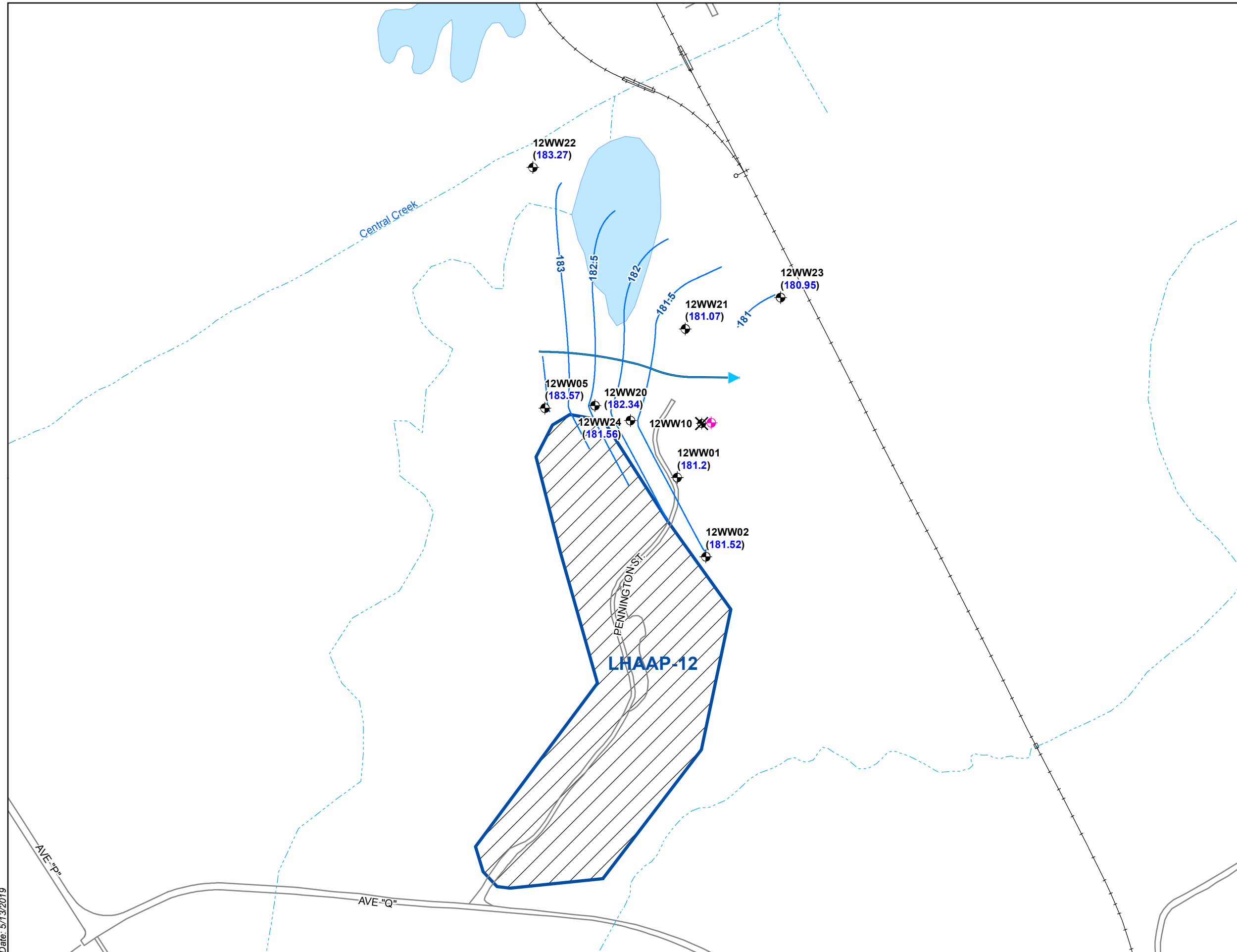


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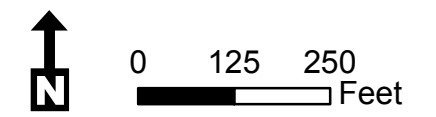


Figure 2-4
Groundwater COC Concentrations
December 2018
LHAAP-12
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

Date: 4/10/2019



- Proposed Well
- Shallow Monitoring Well
- Abandoned Well
- Groundwater Contour
- Stream
- Groundwater Flow Direction
- Road
- Site Boundary
- Landfill Cap Area
- Railroad



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Figure 3-1

Proposed Well Location
LHAAP-12

LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

Appendix A

Photo Log and Well Inspection Forms

APPENDIX A**Photo Log**

Photo No.	Date	Task and Description
1	7/25/2018	Signage Along Site in Good Condition
2	7/25/2018	Signage Along Site in Good Condition
3	1/29/2019	Monitoring Well 12WW01
4	1/29/2019	Monitoring Well 12WW02
5	1/29/2019	Monitoring Well 12WW05
6	1/29/2019	Monitoring Well 12WW20
7	1/29/2019	Monitoring Well 12WW21
8	1/29/2019	Monitoring Well 12WW22
9	1/29/2019	Monitoring Well 12WW23
10	1/29/2019	Monitoring Well 12WW24
11	2/1/2019	Repair of Hole in Landfill Cover
12	2/1/2019	Repair of Hole in Landfill Cover
13	2/1/2019	Repair of Hole in Landfill Cover
14	2/1/2019	Repair of Hole in Landfill Cover



PHOTO 1: Signage Along Site in Good Condition
DATE: July 25, 2018



PHOTO 2: Signage Along Site in Good Condition
DATE: July 25, 2018



PHOTO 3: Monitoring Well 12WW01
DATE: January 29, 2019



PHOTO 4: Monitoring Well 12WW02
DATE: January 29, 2019



PHOTO 5: Monitoring Well 12WW05
DATE: January 29, 2019



PHOTO 6: Monitoring Well 12WW20
DATE: January 29, 2019



PHOTO 7: Monitoring Well 12WW21
DATE: January 29, 2019



PHOTO 8: Monitoring Well 12WW22
DATE: January 29, 2019



PHOTO 9: Monitoring Well 12WW23

DATE: January 29, 2019



PHOTO 10: Monitoring Well 12WW24

DATE: January 29, 2019



PHOTO 11: Repair of Holes in Cover
DATE: February 1, 2019



PHOTO 12: Repair of Holes in Cover
DATE: February 1, 2019



PHOTO 13: Repair of Holes in Landfill Cover
DATE: February 1, 2019



PHOTO 14: Repair of Holes in Landfill Cover
DATE: February 1, 2019

WELL INSPECTION FORM

Job Name: LHAAP Well ID: 12ww02
 Job No.: 501032 Inspection Date: 1/29/19
 Client: Army Well Completion Depth/ Measured Depth: 31.70
 Site Name: 12 Inspector: Scott Beesinger

<input checked="" type="checkbox"/> ABOVE GROUND	
Well Accessible?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Protective casing?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Material: <u>Steel</u>	
Condition: <input checked="" type="checkbox"/> Good <input type="checkbox"/> Broken <input type="checkbox"/> Cracked <input type="checkbox"/> Need Paint	
Lid Condition: <input checked="" type="checkbox"/> Good <input type="checkbox"/> Broken <input type="checkbox"/> Cracked	
Hinge Condition: <input checked="" type="checkbox"/> Good <input type="checkbox"/> Less than 50% rusted <input type="checkbox"/> More than 50% rusted	
<input type="checkbox"/> FLUSH MOUNTED	
Well cover present?	<input type="checkbox"/> Yes <input type="checkbox"/> No
Condition: <input type="checkbox"/> Good <input type="checkbox"/> Broken <input type="checkbox"/> Cracked	
Condition of Sump: <input type="checkbox"/> Clean <input type="checkbox"/> Dry <input type="checkbox"/> Standing Water	
CONCRETE PAD:	
Sloped away from casing?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Pad Area Cleared of Vegetation?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Check any of the following features that apply:	
<input type="checkbox"/> Many Cracks <input type="checkbox"/> Gap Around Casing <input type="checkbox"/> Few Cracks <input type="checkbox"/> Ponded Water <input type="checkbox"/> No Pad Present	
WELL CONDITION:	
Inner Diameter (inches): <u>411</u>	
Condition: <input checked="" type="checkbox"/> Good <input type="checkbox"/> Broken <input type="checkbox"/> Cracked <input type="checkbox"/> Other (describe)	
Cap present?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Well lock present?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Lock functioning properly?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Are bollards present and stable?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Well ID visible?	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
WELL INTEGRITY:	
Bailer present?	<input type="checkbox"/> Yes <input checked="" type="checkbox"/> No
Visual obstruction?	<input type="checkbox"/> Yes <input checked="" type="checkbox"/> No
Is well open to completed depth? (complete when gauging)	<input checked="" type="checkbox"/> Yes <input type="checkbox"/> No
Is silt present in well? (complete when gauging)	<input type="checkbox"/> Yes <input checked="" type="checkbox"/> No
Is silt greater than 10% of well screen length? (complete when gauging)	<input type="checkbox"/> Yes <input checked="" type="checkbox"/> No
COMMENTS:	

WELL INSPECTION FORM

Job Name: LITRAP Well ID: 12WW22
 Job No.: 501032 Inspection Date: 1/29/19
 Client: Army Well Completion Depth/ Measured Depth: 38.69
 Site Name: 12 Inspector: Scott Beasinger

ABOVE GROUND

Well Accessible? Yes No
 Protective casing? Yes No
 Material: Steel
 Condition: Good Broken Cracked Need Paint
 Lid Condition: Good Broken Cracked
 Hinge Condition: Good Less than 50% rusted More than 50% rusted

FLUSH MOUNTED

NA
 Well cover present? Yes No
 Condition: Good Broken Cracked
 Condition of Sump: Clean Dry Standing Water

CONCRETE PAD:

Sloped away from casing? Yes No
 Pad Area Cleared of Vegetation? Yes No
 Check any of the following features that apply:
 Many Cracks Gap Around Casing Few Cracks Ponded Water No Pad Present

WELL CONDITION:

Inner Diameter (inches) 4"
 Condition: Good Broken Cracked Other (describe)
 Cap present? Yes No
 Well lock present? Yes No
 Lock functioning properly? Yes No
 Are bollards present and stable? Yes No
 Well ID visible? Yes No

WELL INTEGRITY:

Bailer present? Yes No
 Visual obstruction? Yes No
 Is well open to completed depth? (complete when gauging) Yes No
 Is silt present in well? (complete when gauging) Yes No
 Is silt greater than 10% of well screen length? (complete when gauging) Yes No

COMMENTS:

WELL INSPECTION FORM

Job Name: CHAAP Well ID: 12WW05
 Job No.: 501032 Inspection Date: 1/29/19
 Client: Army Well Completion Depth/ Measured Depth: 38.61
 Site Name: 12 Inspector: Scott Boesinger

ABOVE GROUND

Well Accessible? Yes No
 Protective casing? Yes No
 Material: Steel
 Condition: Good Broken Cracked Need Paint
 Lid Condition: Good Broken Cracked
 Hinge Condition: Good Less than 50% rusted More than 50% rusted

FLUSH MOUNTED

Well cover present? Yes No
 Condition: Good Broken Cracked
 Condition of Sump: Clean Dry Standing Water
NA

CONCRETE PAD:

Sloped away from casing? Yes No
 Pad Area Cleared of Vegetation? Yes No
 Check any of the following features that apply:
 Many Cracks Gap Around Casing Few Cracks Poned Water No Pad Present

WELL CONDITION:

Inner Diameter (inches) 8 4"
 Condition: Good Broken Cracked Other (describe)
 Cap present? Yes No
 Well lock present? Yes No
 Lock functioning properly? Yes No
 Are bollards present and stable? Yes No
 Well ID visible? Yes No

WELL INTEGRITY:

Bailer present? Yes No
 Visual obstruction? Yes No
 Is well open to completed depth? (complete when gauging) Yes No
 Is silt present in well? (complete when gauging) Yes No
 Is silt greater than 10% of well screen length? (complete when gauging) Yes No

COMMENTS:

WELL INSPECTION FORM

Job Name: LHAAP Well ID: 12WW20
 Job No.: 501032 Inspection Date: 1/29/19
 Client: Army Well Completion Depth/ Measured Depth: 39.33
 Site Name: 12 Inspector: Scott Bessinger

ABOVE GROUND

Well Accessible? Yes No
 Protective casing? Yes No
 Material: Steel
 Condition: Good Broken Cracked Need Paint
 Lid Condition: Good Broken Cracked
 Hinge Condition: Good Less than 50% rusted More than 50% rusted

FLUSH MOUNTED

Well cover present? Yes No
 Condition: Good Broken Cracked
 Condition of Sump: Clean Dry Standing Water
NA

CONCRETE PAD:

Sloped away from casing? Yes No
 Pad Area Cleared of Vegetation? Yes No
 Check any of the following features that apply:
 Many Cracks Gap Around Casing Few Cracks Pounded Water No Pad Present

WELL CONDITION:

Inner Diameter (inches) 44
 Condition: Good Broken Cracked Other (describe)
 Cap present? Yes No
 Well lock present? Yes No
 Lock functioning properly? Yes No
 Are bollards present and stable? Yes No
 Well ID visible? Yes No

WELL INTEGRITY:

Bailer present? Yes No
 Visual obstruction? Yes No
 Is well open to completed depth? (complete when gauging) Yes No
 Is silt present in well? (complete when gauging) Yes No
 Is silt greater than 10% of well screen length? (complete when gauging) Yes No

COMMENTS:

WELL INSPECTION FORM

Job Name: LHAAP Well ID: 12WW24
 Job No.: 501032 Inspection Date: 1/29/19
 Client: Army Well Completion Depth/ Measured Depth: 28.24
 Site Name: 12 Inspector: Scott Bessinger

ABOVE GROUND

Well Accessible? Yes No
 Protective casing? Yes No
 Material: Steel
 Condition: Good Broken Cracked Need Paint
 Lid Condition: Good Broken Cracked
 Hinge Condition: Good Less than 50% rusted More than 50% rusted

FLUSH MOUNTED

Well cover present? Yes No
 Condition: Good Broken Cracked
 Condition of Sump: Clean Dry Standing Water

CONCRETE PAD:

Sloped away from casing? Yes No
 Pad Area Cleared of Vegetation? Yes No
 Check any of the following features that apply:
 Many Cracks Gap Around Casing Few Cracks Ponded Water No Pad Present

WELL CONDITION:

Inner Diameter (inches): 41
 Condition: Good Broken Cracked Other (describe)
 Cap present? Yes No
 Well lock present? Yes No
 Lock functioning properly? Yes No
 Are bollards present and stable? Yes No
 Well ID visible? Yes No

WELL INTEGRITY:

Bailer present? Yes No
 Visual obstruction? Yes No
 Is well open to completed depth? (complete when gauging) Yes No
 Is silt present in well? (complete when gauging) Yes No
 Is silt greater than 10% of well screen length? (complete when gauging) Yes No

COMMENTS:

WELL INSPECTION FORM

Job Name: LHAAP Well ID: 12WW21
 Job No.: 501032 Inspection Date: 1/29/19
 Client: ARMY Well Completion Depth/ Measured Depth: 41.74
 Site Name: 12 Inspector: Scott Beasinger

ABOVE GROUND

Well Accessible? Yes No
 Protective casing? Yes No
 Material: Steel
 Condition: Good Broken Cracked Need Paint
 Lid Condition: Good Broken Cracked
 Hinge Condition: Good Less than 50% rusted More than 50% rusted

FLUSH MOUNTED

Well cover present? NA Yes No
 Condition: Good Broken Cracked
 Condition of Sump: Clean Dry Standing Water

CONCRETE PAD:

Sloped away from casing? Yes No
 Pad Area Cleared of Vegetation? Yes No
 Check any of the following features that apply:
 Many Cracks Gap Around Casing Few Cracks Ponded Water No Pad Present

WELL CONDITION:

Inner Diameter (inches) 411
 Condition: Good Broken Cracked Other (describe)
 Cap present? Yes No
 Well lock present? Yes No
 Lock functioning properly? Yes No
 Are bollards present and stable? Yes No
 Well ID visible? Yes No

WELL INTEGRITY:

Bailer present? Yes No
 Visual obstruction? Yes No
 Is well open to completed depth? (complete when gauging) Yes No
 Is silt present in well? (complete when gauging) Yes No
 Is silt greater than 10% of well screen length? (complete when gauging) Yes No

COMMENTS:

WELL INSPECTION FORM

Job Name: LHAAP Well ID: 12WW23
 Job No.: 501032 Inspection Date: 1/29/19
 Client: ARMY Well Completion Depth/ Measured Depth: 25.43
 Site Name: 12 Inspector: Scott Beesinger

ABOVE GROUND

Well Accessible? Yes No
 Protective casing? Yes No
 Material: steel
 Condition: Good Broken Cracked Need Paint
 Lid Condition: Good Broken Cracked
 Hinge Condition: Good Less than 50% rusted More than 50% rusted

FLUSH MOUNTED

Well cover present? NA Yes No
 Condition: Good Broken Cracked
 Condition of Sump: Clean Dry Standing Water

CONCRETE PAD:

Sloped away from casing? Yes No
 Pad Area Cleared of Vegetation? Yes No
 Check any of the following features that apply:
 Many Cracks Gap Around Casing Few Cracks Ponded Water No Pad Present

WELL CONDITION:

Inner Diameter (inches) 4 1/2
 Condition: Good Broken Cracked Other (describe)
 Cap present? Yes No
 Well lock present? Yes No
 Lock functioning properly? Yes No
 Are bollards present and stable? Yes No
 Well ID visible? Yes No

WELL INTEGRITY:

Bailer present? Yes No
 Visual obstruction? Yes No
 Is well open to completed depth? (complete when gauging) Yes No
 Is silt present in well? (complete when gauging) Yes No
 Is silt greater than 10% of well screen length? (complete when gauging) Yes No

COMMENTS:

WELL INSPECTION FORM

Job Name: LHAAP Well ID: 12WW01
 Job No.: 501032 Inspection Date: 1/29/19
 Client: Army Well Completion Depth/ Measured Depth: 29.97
 Site Name: 12 Inspector: Scott Beesinger

ABOVE GROUND

Well Accessible? Yes No
 Protective casing? Yes No
 Material: Steel
 Condition: Good Broken Cracked Need Paint
 Lid Condition: Good Broken Cracked
 Hinge Condition: Good Less than 50% rusted More than 50% rusted

FLUSH MOUNTED

Well cover present? NA Yes No
 Condition: Good Broken Cracked
 Condition of Sump: Clean Dry Standing Water

CONCRETE PAD:

Sloped away from casing? Yes No
 Pad Area Cleared of Vegetation? Yes No
 Check any of the following features that apply:
 Many Cracks Gap Around Casing Few Cracks Ponded Water No Pad Present

WELL CONDITION:

Inner Diameter (inches) 4"
 Condition: Good Broken Cracked Other (describe)
 Cap present? Yes No
 Well lock present? Yes No
 Lock functioning properly? Yes No
 Are bollards present and stable? Yes No
 Well ID visible? Yes No

WELL INTEGRITY:

Bailer present? Yes No
 Visual obstruction? Yes No
 Is well open to completed depth? (complete when gauging) Yes No
 Is silt present in well? (complete when gauging) Yes No
 Is silt greater than 10% of well screen length? (complete when gauging) Yes No

COMMENTS:

2018 Summary of Monitoring Well Inspection and Repair

Well Information		Visual Inspections of Well												Comments
Well Identification	Date	ID Plate	Lock	Gasket	Cap	Bolts	Bollards	Pad	Vegetation	Tampering/ Obstructions	Limited Access	New Wells	Subsurface Activities	
12WW01	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	
12WW02	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	
12WW05	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	
12WW20	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	
12WW21	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	
12WW22	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	
12WW23	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	
12WW24	1/29/2019	X	X	X	X	X	Present/Functioning	Good	None - Well is Clear	N	N	N	N	

Notes:

ID - identification

N - none

Well Construction Information

Well Identification	Log Date	Depth to Water (ft btoc)	Total Measured Well Depth (ft btoc)	Total Measured Well Depth (ft bgs)	Dry?	Water Column (ft)	Survey Data		Well Construction Data				Percent Siltation
							Top of Casing Elevation (ft amsl)	Ground Surface Elevation (ft amsl)	Constructed Well Depth (ft bgs)	Well Screen Beginning Depth (ft bgs)	Well Screen Ending Depth (ft bgs)	Screen Length (ft)	
12WW01	1/29/2019	22.99	29.97	27.35	No	3.51	204.19	201.57	26.5	14.50	26.50	12.00	0.0%
12WW02	1/29/2019	20.93	31.70	29.51	No	7.57	202.45	200.26	28.5	13.50	28.50	15.00	0.0%
12WW05	1/29/2019	6.95	38.61	36.46	No	31.05	190.52	188.37	38	15.50	35.50	20.00	0.0%
12WW20	1/29/2019	16.81	39.33	37.26	No	22.04	199.15	197.08	38.85	28.85	38.85	10.00	4.1%
12WW21	1/29/2019	21.00	41.74	39.77	No	20.70	202.07	200.10	41.7	31.70	41.70	10.00	4.6%
12WW22	1/29/2019	6.93	38.69	36.27	No	31.43	190.20	187.78	38.36	28.36	38.36	10.00	5.4%
12WW23	1/29/2019	16.02	25.43	23.33	No	9.12	196.97	194.87	25.14	15.14	25.14	10.00	7.2%
12WW24	1/29/2019	21.61	28.24	25.94	No	4.39	203.17	200.87	26	15.50	25.50	10.00	0.0%

Notes:

ft - feet

ft amsl - feet above mean sea level

ft bgs - feet below ground surface

ft btoc - feet below top of casing

Appendix B

Land Use Controls and Maintenance Log

LUC Inspection and Maintenance Log - LHAAP 12

Date	Inspected By	Inspection / Maintenance Activities (Yearly, at a Minimum)					Corrective Action or Repairs Required?	Repairs / Action Taken
		Protect Landfill Integrity			Prevent Human Exposure to Groundwater			
		Vegetative Cover Maintained (i.e., grass mowed at least annually)	Fence and Signage Maintained	Observance of Landfill Cover Degradation (e.g., desiccation cracks, erosion, or gulying)	Continued Compliance Verified for No Digging or Disturbance of Landfill Cover or Contents	Verified No Withdrawal of Use of Groundwater (other than environmental testing)		
7/25/2018	Beesinger, S.	Maintained. Vegetative cover is in good condition.	Fence and signage in good repair.	Cover subsidence observed and found to be in good condition. No erosion, cracks, or gulying was found. There are 4 locations in the center of the cap that have sink holes.	No evidence of animal burrowing in the cap.	Completed and verified.	Top soil needed to fill in the sink holes in center of landfill cap.	Holes were covered with dirt on 2/1/2019.

Notes:

e.g. - for example
i.e. - that is
LUC - land use control

Appendix C

Groundwater Sampling Forms



Sample Collection Log

1 of 4

Project Name: **Longhorn AAP**Location ID: **12WW20**Project No: **501032**Sampler(s): **Scott Beesinger**

FIELD CONDITIONS [Clear](#)

SAMPLING INFORMATION

Sample No: **12WW20-181204**DATE/TIME: **12/4/2018 / 12:05**Pump Inlet Depth (ft): **34.30**Sampling Method: **Low Flow**Sample Purpose: **REG**Sample Matrix: **GW**

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
12-DEC2018-GW-ALSHT-181204	None	VOLATILES	SW8260

Sampler:	SCOTT BEESINGER / BITE	Scott Beesinger
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QC'ed By:	<i>Jacoma</i>		
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WELL AND PURGING INFORMATION

Measuring Point : Top of CasingPurging Method/Equipment: Bladder PumpCasing ID (in.): 4Purge Start Date/Time: 12/4/2018 / 11:35Depth to Water - Initial (DTWi) (ft) 18.35Purge End Date/Time: 12/4/2018 / 12:05Measured Depth of Well (ft): 39.6PID Reading: N/AScreen Interval (ft): 29.30 - 39.30Pump Start Time: 12/4/2018 / 11:35



Sample Collection Log

Location ID: 12WW20 Sample No: 12WW20-181204										
Date of Reading	Time of Reading	Purge Rate	Total Purge	DTW	Cond.	Temp.	pH	Turbidity	ORP	DO
		(ml/min)	(L)	(ft)	(mS/cm)	(°C)		(NTU)	(mV)	(mg/L)
	Purge Stabilization Criteria	-	-	Drawdown limit 0.3 ft	±10%		±0.1 units	No criteria		± 10% or 0.2 mg/L
12/4/2018	11:40	100	0.5	18.42	1.47	16.07	5.45	6.6	164	0.95
12/4/2018	11:45	100	1.0	18.48	1.46	17.33	5.10	6.0	223	0.63
12/4/2018	11:50	100	1.5	18.52	1.45	17.61	5.01	5.5	246	0.57
12/4/2018	11:55	100	2.0	18.55	1.45	17.69	5.0	4.9	247	0.56
12/4/2018	12:00	100	2.5	18.57	1.45	17.74	5.0	4.1	248	0.56
12/4/2018	12:05	100	3.0	18.58	1.45	17.80	5.0	3.7	249	0.55



Sample Collection Log

1 of 4

Project Name: **Longhorn AAP**Location ID: **12WW21**Project No: **501032**Sampler(s): **Scott Beesinger**

FIELD CONDITIONS [Clear](#)

SAMPLING INFORMATION

Sample No: **12WW21-181204**DATE/TIME: **12/4/2018 / 10:05**Pump Inlet Depth (ft): **36.80**Sampling Method: **Low Flow**Sample Purpose: **REG**Sample Matrix: **GW**

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
12-DEC2018-GW-ALSHT-181204	None	VOLATILES	SW8260

Sampler:	SCOTT BEESINGER / BITITE Scott Beesinger
----------	--

QC'ed By:	
-----------	--

WELL AND PURGING INFORMATION

Measuring Point : Top of CasingPurging Method/Equipment: Bladder PumpCasing ID (in.): 4Purge Start Date/Time: 12/4/2018 / 09:35Depth to Water - Initial (DTWi) (ft) 22.2Purge End Date/Time: 12/4/2018 / 10:05Measured Depth of Well (ft): 42.15PID Reading: N/AScreen Interval (ft): 31.80 - 41.80Pump Start Time: 12/4/2018 / 09:35



Sample Collection Log

3 of 4

Location ID: 12WW21 Sample No: 12WW21-181204										
Date of Reading	Time of Reading	Purge Rate	Total Purge	DTW	Cond.	Temp.	pH	Turbidity	ORP	DO
		(ml/min)	(L)	(ft)	(mS/cm)	(°C)		(NTU)	(mV)	(mg/L)
	Purge Stabilization Criteria	-	-	Drawdown limit 0.3 ft	±10%		±0.1 units	No criteria		± 10% or 0.2 mg/L
12/4/2018	09:40	100	0.5	22.27	4.94	14.83	6.68	51.3	162	0.73
12/4/2018	09:45	100	1.0	22.34	5.40	15.67	5.72	39.5	187	0.25
12/4/2018	09:50	100	1.5	22.38	5.41	15.89	5.53	35.9	200	0.09
12/4/2018	09:55	100	2.0	22.40	5.42	15.93	5.52	35.2	201	0.09
12/4/2018	10:00	100	2.5	22.42	5.42	15.97	5.51	34.8	201	0.08
12/4/2018	10:05	100	3.0	22.43	5.42	16.0	5.51	34.4	202	0.08



Sample Collection Log

Project Name: Longhorn AAP

Location ID: 12WW24

Project No: 501032

Sampler(s): Scott Beesinger

FIELD CONDITIONS [Clear](#)

SAMPLING INFORMATION

Sample No: 12WW24-181204

DATE/TIME: 12/4/2018 / 11:10

Pump Inlet Depth (ft): 24.0

Sampling Method: Low Flow

Sample Purpose: REG

Sample Matrix: GW

Sample Notes:

Chain of Custody	COC Notes	Analysis Group	Analytic Method
12-DEC2018-GW-ALSHT-181204	None	VOLATILES	SW8260

Sampler:	<u>Scott Beesinger / BITE</u>	Scott Beesinger
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QC'ed By:	<u>Joanna</u>		
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WELL AND PURGING INFORMATION

Measuring Point : Top of Casing

Purging Method/Equipment: Bladder Pump

Casing ID (in.): 4

Purge Start Date/Time: 12/4/2018 / 10:40

Depth to Water - Initial (DTWi) (ft) 22.87

Purge End Date/Time: 12/4/2018 / 11:10

Measured Depth of Well (ft): 28.25

PID Reading: N/A

Screen Interval (ft): 15.50 - 25.50

Pump Start Time: 12/4/2018 / 10:40

Location ID: 12WW24 Sample No: 12WW24-181204										
Date of Reading	Time of Reading	Purge Rate	Total Purge	DTW	Cond.	Temp.	pH	Turbidity	ORP	DO
		(ml/min)	(L)	(ft)	(mS/cm)	(°C)		(NTU)	(mV)	(mg/L)
	Purge Stabilization Criteria	-	-	Drawdown limit 0.3 ft	±10%		±0.1 units	No criteria		± 10% or 0.2 mg/L
12/4/2018	10:45	100	0.5	22.95	1.59	13.85	5.49	9.4	134	0.42
12/4/2018	10:50	100	1.0	23.01	1.55	14.53	5.48	4.4	123	0.22
12/4/2018	10:55	100	1.5	23.05	1.55	14.57	5.46	4.2	120	0.05
12/4/2018	11:00	100	2.0	23.08	1.55	14.60	5.45	4.1	120	0.05
12/4/2018	11:05	100	2.5	23.10	1.55	14.64	5.45	4.0	119	0.05
12/4/2018	11:10	100	3.0	23.12	1.55	14.68	5.45	3.9	119	0.04

Appendix D

Laboratory Analytical Data Package

HS18120278 Cover Page

ALS WO HS18120278





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Dates Report
QC
Acronyms
Certifications
SampleReceiptCheckList
COC
HS18120278 8260 Raw Data





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WorkOrder: HS18120278

LHAARP - 12

Aptim Environmental & Infrastructure, Inc.

Susan Huang
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28-Jan-2019



HS18120278 LHAARP 12 Final

ALS WO HS18120278





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December 19, 2018

Susan Huang
Aptim Environmental & Infrastructure, Inc.
2500 City West Blvd., Suite 1700
Houston, TX 77042

Work Order: **HS18120278**

Laboratory Results for: **LHAARP - 12**

Dear Susan,

ALS Environmental received 5 sample(s) on Dec 05, 2018 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read "Raj. P. Modashia", enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
Work Order: HS18120278

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS18120278-01	12WW21-181204	Groundwater		04-Dec-2018 10:05	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-02	12WW21-181204 FD	Groundwater		04-Dec-2018 10:05	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-03	12WW24-181204	Groundwater		04-Dec-2018 13:10	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-04	12WW20-181204	Groundwater		04-Dec-2018 12:05	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-05	Trip Blank ALS-071918-78	Water		04-Dec-2018 00:00	05-Dec-2018 09:30	<input type="checkbox"/>



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
Work Order: HS18120278

CASE NARRATIVE**GCMS Volatiles by Method SW8260****Batch ID: R329535****Sample ID: VLCSW-1812018**

- 1,4_Dichlorobenzene and Hexachlorobutadiene exceeded QC limits for LCS. CCV is OK.

Sample ID: VSTD050

- cis-1,3-Dichloropropene exceeded %D limits for CCV. Samples are ND for this compound.

Sample ID: 12WW24-181204 (HS18120278-03MS)

- MS/MSD recovered outside the control limits for multiple compounds
-

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-01
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 16:27	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-01
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:27	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 16:27	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Surr: 1,2-Dichloroethane-d4	91.4			0	70-126	%REC	1	18-Dec-2018 16:27	
Surr: 4-Bromofluorobenzene	99.2			0	81-113	%REC	1	18-Dec-2018 16:27	
Surr: Dibromofluoromethane	87.4			0	77-123	%REC	1	18-Dec-2018 16:27	
Surr: Toluene-d8	106			0	82-127	%REC	1	18-Dec-2018 16:27	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204 FD
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-02
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 15:13	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204 FD
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT

WorkOrder:HS18120278
 Lab ID:HS18120278-02
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260		Analyst: PC					
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:13	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 15:13	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Surr: 1,2-Dichloroethane-d4	90.5			0	70-126	%REC	1	18-Dec-2018 15:13	
Surr: 4-Bromofluorobenzene	97.8			0	81-113	%REC	1	18-Dec-2018 15:13	
Surr: Dibromofluoromethane	88.4			0	77-123	%REC	1	18-Dec-2018 15:13	
Surr: Toluene-d8	105			0	82-127	%REC	1	18-Dec-2018 15:13	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW24-181204
 Collection Date: 04-Dec-2018 13:10

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-03
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1-Dichloroethene	0.71	J	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 16:02	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Chlorobenzene	1.5		0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW24-181204
 Collection Date: 04-Dec-2018 13:10

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-03
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
cis-1,2-Dichloroethene	22		0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:02	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 16:02	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Trichloroethene	48		0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>91.4</i>			0	<i>70-126</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>97.6</i>			0	<i>81-113</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	
<i>Surr: Dibromofluoromethane</i>	<i>89.1</i>			0	<i>77-123</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	
<i>Surr: Toluene-d8</i>	<i>105</i>			0	<i>82-127</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW20-181204
 Collection Date: 04-Dec-2018 12:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-04
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 15:37	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW20-181204
 Collection Date: 04-Dec-2018 12:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-04
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:37	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 15:37	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Surr: 1,2-Dichloroethane-d4	91.4			0	70-126	%REC	1	18-Dec-2018 15:37	
Surr: 4-Bromofluorobenzene	96.8			0	81-113	%REC	1	18-Dec-2018 15:37	
Surr: Dibromofluoromethane	88.0			0	77-123	%REC	1	18-Dec-2018 15:37	
Surr: Toluene-d8	105			0	82-127	%REC	1	18-Dec-2018 15:37	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: Trip Blank ALS-071918-78
 Collection Date: 04-Dec-2018 00:00

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-05
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
Acetone	2.7		2.0	2.0	2.0	ug/L	1	18-Dec-2018 14:48	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: Trip Blank ALS-071918-78
 Collection Date: 04-Dec-2018 00:00

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-05
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 14:48	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 14:48	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Surr: 1,2-Dichloroethane-d4	89.5			0	70-126	%REC	1	18-Dec-2018 14:48	
Surr: 4-Bromofluorobenzene	98.2			0	81-113	%REC	1	18-Dec-2018 14:48	
Surr: Dibromofluoromethane	87.9			0	77-123	%REC	1	18-Dec-2018 14:48	
Surr: Toluene-d8	106			0	82-127	%REC	1	18-Dec-2018 14:48	

Note: See Qualifiers Page for a list of qualifiers and their explanation.



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R329535	Test Name : LOW LEVEL VOLATILES BY SW8260C			Matrix: Water		
HS18120278-05	Trip Blank ALS-071918-78	04 Dec 2018 00:00			18 Dec 2018 14:48	1
Batch ID R329535	Test Name : LOW LEVEL VOLATILES BY SW8260C			Matrix: Groundwater		
HS18120278-01	12WW21-181204	04 Dec 2018 10:05			18 Dec 2018 16:27	1
HS18120278-02	12WW21-181204 FD	04 Dec 2018 10:05			18 Dec 2018 15:13	1
HS18120278-03	12WW24-181204	04 Dec 2018 13:10			18 Dec 2018 16:02	1
HS18120278-04	12WW20-181204	04 Dec 2018 12:05			18 Dec 2018 15:37	1



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MBLK	Sample ID: VBLKW-181218	Units: ug/L			Analysis Date: 18-Dec-2018 13:34					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872159	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	0.50	1.0								U
1,1,1-Trichloroethane	0.50	1.0								U
1,1,2,2-Tetrachloroethane	0.50	1.0								U
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	1.0								U
1,1,2-Trichloroethane	0.50	1.0								U
1,1-Dichloroethane	0.50	1.0								U
1,1-Dichloroethene	0.50	1.0								U
1,1-Dichloropropene	0.50	1.0								U
1,2,3-Trichlorobenzene	0.50	1.0								U
1,2,3-Trichloropropane	0.50	1.0								U
1,2,4-Trichlorobenzene	0.50	1.0								U
1,2,4-Trimethylbenzene	0.50	1.0								U
1,2-Dibromo-3-chloropropane	1.0	1.0								U
1,2-Dibromoethane	0.50	1.0								U
1,2-Dichlorobenzene	0.50	1.0								U
1,2-Dichloroethane	0.50	1.0								U
1,2-Dichloropropane	0.50	1.0								U
1,3,5-Trimethylbenzene	0.50	1.0								U
1,3-Dichlorobenzene	0.50	1.0								U
1,3-Dichloropropane	0.50	1.0								U
1,4-Dichlorobenzene	0.50	1.0								U
2,2-Dichloropropane	0.50	1.0								U
2-Butanone	1.0	2.0								U
2-Chlorotoluene	0.50	1.0								U
2-Hexanone	1.0	2.0								U
4-Chlorotoluene	0.50	1.0								U
4-Isopropyltoluene	0.50	1.0								U
4-Methyl-2-pentanone	1.0	2.0								U
Acetone	2.0	2.0								U
Benzene	0.50	1.0								U
Bromobenzene	0.50	1.0								U
Bromochloromethane	0.50	1.0								U
Bromodichloromethane	0.50	1.0								U
Bromoform	0.50	1.0								U



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MBLK	Sample ID: VBLKW-181218	Units: ug/L			Analysis Date: 18-Dec-2018 13:34					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872159	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Bromomethane	0.50	1.0								U
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>45.19</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>90.4</i>	<i>70 - 123</i>				
<i>Surr: 4-Bromofluorobenzene</i>	<i>48.92</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>97.8</i>	<i>82 - 115</i>				
<i>Surr: Dibromofluoromethane</i>	<i>43.69</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>87.4</i>	<i>73 - 126</i>				



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MBLK	Sample ID: VBLKW-181218	Units: ug/L			Analysis Date: 18-Dec-2018 13:34					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872159		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	52.1	1.0	50	0	104	81 - 120				



ALS Houston, US

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Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
LCS	Sample ID: VLCSW-1812018	Units: ug/L			Analysis Date: 18-Dec-2018 13:58					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872160		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	21.7	1.0	20	0	109	77 - 118				
1,1,1-Trichloroethane	20.06	1.0	20	0	100	70 - 130				
1,1,2,2-Tetrachloroethane	21.61	1.0	20	0	108	70 - 120				
1,1,2-Trichlor-1,2,2-trifluoroethane	21.66	1.0	20	0	108	70 - 130				
1,1,2-Trichloroethane	21.9	1.0	20	0	109	77 - 113				
1,1-Dichloroethane	20.96	1.0	20	0	105	71 - 122				
1,1-Dichloroethene	20.63	1.0	20	0	103	70 - 130				
1,1-Dichloropropene	22.62	1.0	20	0	113	78 - 118				
1,2,3-Trichlorobenzene	22.19	1.0	20	0	111	70 - 130				
1,2,3-Trichloropropane	21.99	1.0	20	0	110	70 - 127				
1,2,4-Trichlorobenzene	21.92	1.0	20	0	110	77 - 126				
1,2,4-Trimethylbenzene	23.08	1.0	20	0	115	73 - 121				
1,2-Dibromo-3-chloropropane	19.69	1.0	20	0	98.4	70 - 130				
1,2-Dibromoethane	22.19	1.0	20	0	111	76 - 123				
1,2-Dichlorobenzene	21.19	1.0	20	0	106	77 - 113				
1,2-Dichloroethane	20.79	1.0	20	0	104	70 - 124				
1,2-Dichloropropane	23.16	1.0	20	0	116	72 - 119				
1,3,5-Trimethylbenzene	23.2	1.0	20	0	116	75 - 118				
1,3-Dichlorobenzene	21.7	1.0	20	0	109	78 - 118				
1,3-Dichloropropane	22.28	1.0	20	0	111	75 - 116				
1,4-Dichlorobenzene	22.92	1.0	20	0	115	79 - 113				S
2,2-Dichloropropane	21.28	1.0	20	0	106	70 - 130				
2-Butanone	42.42	2.0	40	0	106	70 - 130				
2-Chlorotoluene	22.49	1.0	20	0	112	70 - 128				
2-Hexanone	46.58	2.0	40	0	116	70 - 130				
4-Chlorotoluene	22.67	1.0	20	0	113	74 - 126				
4-Isopropyltoluene	23.85	1.0	20	0	119	74 - 126				
4-Methyl-2-pentanone	46.52	2.0	40	0	116	70 - 130				
Acetone	42.82	2.0	40	0	107	70 - 130				
Benzene	22.64	1.0	20	0	113	74 - 120				
Bromobenzene	21.59	1.0	20	0	108	78 - 113				
Bromochloromethane	22.21	1.0	20	0	111	76 - 124				
Bromodichloromethane	20.86	1.0	20	0	104	74 - 122				
Bromoform	19.3	1.0	20	0	96.5	73 - 128				



ALS Houston, US

Date: 19-Dec-18

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Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
LCS	Sample ID: VLCSW-1812018	Units: ug/L			Analysis Date: 18-Dec-2018 13:58					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872160		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Bromomethane	23.28	1.0	20	0	116	70 - 130				
Carbon disulfide	43.3	2.0	40	0	108	70 - 130				
Carbon tetrachloride	21.74	1.0	20	0	109	71 - 125				
Chlorobenzene	21.78	1.0	20	0	109	76 - 113				
Chloroethane	23.06	1.0	20	0	115	70 - 130				
Chloroform	20	1.0	20	0	100	71 - 121				
Chloromethane	24.03	1.0	20	0	120	70 - 129				
cis-1,2-Dichloroethene	20.82	1.0	20	0	104	75 - 122				
cis-1,3-Dichloropropene	22.79	1.0	20	0	114	73 - 127				
Dibromochloromethane	21.85	1.0	20	0	109	77 - 122				
Dibromomethane	21.43	1.0	20	0	107	78 - 121				
Dichlorodifluoromethane	21.72	1.0	20	0	109	70 - 130				
Ethylbenzene	22.49	1.0	20	0	112	77 - 117				
Hexachlorobutadiene	26.61	1.0	20	0	133	70 - 130				S
Isopropylbenzene	23.13	1.0	20	0	116	73 - 127				
m,p-Xylene	45.63	2.0	40	0	114	77 - 122				
Methylene chloride	22.57	2.0	20	0	113	70 - 127				
Naphthalene	22.83	1.0	20	0	114	70 - 130				
n-Butylbenzene	24.18	1.0	20	0	121	72 - 130				
n-Propylbenzene	23.41	1.0	20	0	117	73 - 124				
o-Xylene	22.9	1.0	20	0	114	75 - 119				
sec-Butylbenzene	23.6	1.0	20	0	118	73 - 128				
Styrene	23.39	1.0	20	0	117	72 - 126				
tert-Butylbenzene	23.13	1.0	20	0	116	73 - 124				
Tetrachloroethene	22.36	1.0	20	0	112	76 - 119				
Toluene	22.56	1.0	20	0	113	77 - 118				
trans-1,2-Dichloroethene	20.73	1.0	20	0	104	72 - 127				
trans-1,3-Dichloropropene	19.97	1.0	20	0	99.9	77 - 119				
Trichloroethene	21.82	1.0	20	0	109	77 - 121				
Trichlorofluoromethane	20.41	1.0	20	0	102	70 - 130				
Vinyl chloride	23.07	1.0	20	0	115	70 - 130				
Surr: 1,2-Dichloroethane-d4	44.22	1.0	50	0	88.4	70 - 130				
Surr: 4-Bromofluorobenzene	51.18	1.0	50	0	102	82 - 115				
Surr: Dibromofluoromethane	44.56	1.0	50	0	89.1	73 - 126				



ALS Houston, US

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Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
LCS	Sample ID: VLCSW-1812018	Units: ug/L		Analysis Date: 18-Dec-2018 13:58						
Client ID:	Run ID: VOA9_329535	SeqNo: 4872160		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	52.7	1.0	50	0	105	81 - 120				



ALS Houston, US

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Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MS	Sample ID: HS18120278-03MS	Units: ug/L			Analysis Date: 18-Dec-2018 17:41					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872166	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	23.88	1.0	20	0	119	70 - 120				
1,1,1-Trichloroethane	22.68	1.0	20	0	113	70 - 130				
1,1,2,2-Tetrachloroethane	23.94	1.0	20	0	120	70 - 123				
1,1,2-Trichlor-1,2,2-trifluoroethane	22.2	1.0	20	0	111	70 - 130				
1,1,2-Trichloroethane	24.23	1.0	20	0	121	70 - 117				S
1,1-Dichloroethane	23.28	1.0	20	0	116	70 - 127				
1,1-Dichloroethene	23.18	1.0	20	0.7106	112	70 - 130				
1,1-Dichloropropene	25.8	1.0	20	0	129	70 - 129				S
1,2,3-Trichlorobenzene	24.51	1.0	20	0	123	70 - 130				
1,2,3-Trichloropropane	23.25	1.0	20	0	116	70 - 130				
1,2,4-Trichlorobenzene	23.82	1.0	20	0	119	70 - 125				
1,2,4-Trimethylbenzene	25.68	1.0	20	0	128	70 - 125				S
1,2-Dibromo-3-chloropropane	21.8	1.0	20	0	109	70 - 130				
1,2-Dibromoethane	24.17	1.0	20	0	121	70 - 124				
1,2-Dichlorobenzene	23.25	1.0	20	0	116	70 - 115				S
1,2-Dichloroethane	22.65	1.0	20	0	113	70 - 127				
1,2-Dichloropropane	25.31	1.0	20	0	127	70 - 122				S
1,3,5-Trimethylbenzene	25.78	1.0	20	0	129	70 - 126				S
1,3-Dichlorobenzene	23.52	1.0	20	0	118	70 - 119				
1,3-Dichloropropane	24.4	1.0	20	0	122	70 - 121				S
1,4-Dichlorobenzene	25.75	1.0	20	0	129	70 - 114				S
2,2-Dichloropropane	21.93	1.0	20	0	110	70 - 130				
2-Butanone	45.49	2.0	40	0	114	70 - 130				
2-Chlorotoluene	24.73	1.0	20	0	124	70 - 130				
2-Hexanone	51.07	2.0	40	0	128	70 - 130				
4-Chlorotoluene	25.15	1.0	20	0	126	70 - 130				
4-Isopropyltoluene	26.9	1.0	20	0	134	70 - 130				S
4-Methyl-2-pentanone	50.44	2.0	40	0	126	70 - 130				
Acetone	46	2.0	40	0	115	70 - 130				
Benzene	25.3	1.0	20	0	126	70 - 127				
Bromobenzene	23.78	1.0	20	0	119	70 - 115				S
Bromochloromethane	24.54	1.0	20	0	123	70 - 127				
Bromodichloromethane	22.91	1.0	20	0	115	70 - 124				
Bromoform	20.79	1.0	20	0	104	70 - 129				



ALS Houston, US

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Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MS	Sample ID: HS18120278-03MS	Units: ug/L			Analysis Date: 18-Dec-2018 17:41					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872166	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Bromomethane	12.09	1.0	20	0	60.5	70 - 130				S
Carbon disulfide	44.81	2.0	40	0	112	70 - 130				
Carbon tetrachloride	24.05	1.0	20	0	120	70 - 130				
Chlorobenzene	25.62	1.0	20	1.542	120	70 - 114				S
Chloroethane	24.72	1.0	20	0	124	70 - 130				
Chloroform	22.07	1.0	20	0	110	70 - 125				
Chloromethane	15.68	1.0	20	0	78.4	70 - 130				
cis-1,2-Dichloroethene	44.9	1.0	20	21.65	116	70 - 128				
cis-1,3-Dichloropropene	24.3	1.0	20	0	121	70 - 125				
Dibromochloromethane	23.74	1.0	20	0	119	70 - 124				
Dibromomethane	22.87	1.0	20	0	114	70 - 124				
Dichlorodifluoromethane	10.42	1.0	20	0	52.1	70 - 130				S
Ethylbenzene	25.15	1.0	20	0	126	70 - 124				S
Hexachlorobutadiene	28.15	1.0	20	0	141	70 - 130				S
Isopropylbenzene	26.04	1.0	20	0	130	70 - 130				S
m,p-Xylene	50.97	2.0	40	0	127	70 - 130				
Methylene chloride	24.38	2.0	20	0	122	70 - 128				
Naphthalene	25.37	1.0	20	0	127	70 - 130				
n-Butylbenzene	26.87	1.0	20	0	134	70 - 130				S
n-Propylbenzene	26.34	1.0	20	0	132	70 - 130				S
o-Xylene	25.44	1.0	20	0	127	70 - 124				S
sec-Butylbenzene	26.76	1.0	20	0	134	70 - 130				S
Styrene	25.73	1.0	20	0	129	70 - 130				
tert-Butylbenzene	26.06	1.0	20	0	130	70 - 130				S
Tetrachloroethene	25.53	1.0	20	0	128	70 - 130				
Toluene	25.16	1.0	20	0	126	70 - 123				S
trans-1,2-Dichloroethene	23.47	1.0	20	0	117	70 - 130				
trans-1,3-Dichloropropene	21.37	1.0	20	0	107	70 - 121				
Trichloroethene	72.45	1.0	20	47.56	124	70 - 129				
Trichlorofluoromethane	20.92	1.0	20	0	105	70 - 130				
Vinyl chloride	20.18	1.0	20	0	101	70 - 130				
Surr: 1,2-Dichloroethane-d4	44.17	1.0	50	0	88.3	70 - 126				
Surr: 4-Bromofluorobenzene	51.45	1.0	50	0	103	81 - 113				
Surr: Dibromofluoromethane	45.03	1.0	50	0	90.1	77 - 123				



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MS	Sample ID: HS18120278-03MS	Units: ug/L			Analysis Date: 18-Dec-2018 17:41					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872166		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	53.03	1.0	50	0	106	82 - 127				



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MSD	Sample ID: HS18120278-03MSD	Units: ug/L			Analysis Date: 18-Dec-2018 18:06					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872167	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	22.94	1.0	20	0	115	70 - 120	23.88	3.99	20	
1,1,1-Trichloroethane	21.46	1.0	20	0	107	70 - 130	22.68	5.56	20	
1,1,2,2-Tetrachloroethane	23.01	1.0	20	0	115	70 - 123	23.94	3.93	20	
1,1,2-Trichlor-1,2,2-trifluoroethane	20.91	1.0	20	0	105	70 - 130	22.2	6	20	
1,1,2-Trichloroethane	23.16	1.0	20	0	116	70 - 117	24.23	4.52	20	
1,1-Dichloroethane	22.19	1.0	20	0	111	70 - 127	23.28	4.8	20	
1,1-Dichloroethene	21.54	1.0	20	0.7106	104	70 - 130	23.18	7.34	20	
1,1-Dichloropropene	24.29	1.0	20	0	121	70 - 129	25.8	6.04	20	
1,2,3-Trichlorobenzene	22.99	1.0	20	0	115	70 - 130	24.51	6.4	20	
1,2,3-Trichloropropane	21.98	1.0	20	0	110	70 - 130	23.25	5.62	20	
1,2,4-Trichlorobenzene	22.88	1.0	20	0	114	70 - 125	23.82	4.03	20	
1,2,4-Trimethylbenzene	24.38	1.0	20	0	122	70 - 125	25.68	5.16	20	
1,2-Dibromo-3-chloropropane	20.49	1.0	20	0	102	70 - 130	21.8	6.15	20	
1,2-Dibromoethane	23.25	1.0	20	0	116	70 - 124	24.17	3.85	20	
1,2-Dichlorobenzene	22.08	1.0	20	0	110	70 - 115	23.25	5.16	20	
1,2-Dichloroethane	21.98	1.0	20	0	110	70 - 127	22.65	3	20	
1,2-Dichloropropane	24.76	1.0	20	0	124	70 - 122	25.31	2.19	20	S
1,3,5-Trimethylbenzene	24.4	1.0	20	0	122	70 - 126	25.78	5.52	20	
1,3-Dichlorobenzene	22.5	1.0	20	0	112	70 - 119	23.52	4.46	20	
1,3-Dichloropropane	23.35	1.0	20	0	117	70 - 121	24.4	4.39	20	
1,4-Dichlorobenzene	24.57	1.0	20	0	123	70 - 114	25.75	4.7	20	S
2,2-Dichloropropane	20.71	1.0	20	0	104	70 - 130	21.93	5.72	20	
2-Butanone	45.19	2.0	40	0	113	70 - 130	45.49	0.66	20	
2-Chlorotoluene	23.65	1.0	20	0	118	70 - 130	24.73	4.47	20	
2-Hexanone	49.4	2.0	40	0	124	70 - 130	51.07	3.32	20	
4-Chlorotoluene	23.94	1.0	20	0	120	70 - 130	25.15	4.92	20	
4-Isopropyltoluene	25.24	1.0	20	0	126	70 - 130	26.9	6.37	20	
4-Methyl-2-pentanone	48.64	2.0	40	0	122	70 - 130	50.44	3.64	20	
Acetone	43.54	2.0	40	0	109	70 - 130	46	5.49	20	
Benzene	24.46	1.0	20	0	122	70 - 127	25.3	3.38	20	
Bromobenzene	22.37	1.0	20	0	112	70 - 115	23.78	6.13	20	
Bromochloromethane	23.65	1.0	20	0	118	70 - 127	24.54	3.69	20	
Bromodichloromethane	22.31	1.0	20	0	112	70 - 124	22.91	2.64	20	
Bromoform	20.15	1.0	20	0	101	70 - 129	20.79	3.11	20	



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260							
MSD	Sample ID: HS18120278-03MSD	Units: ug/L			Analysis Date: 18-Dec-2018 18:06						
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872167	PrepDate:	DF: 1							
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual	
Bromomethane	13.33	1.0	20	0	66.6	70 - 130	12.09	9.71	20	S	
Carbon disulfide	42.21	2.0	40	0	106	70 - 130	44.81	5.98	20		
Carbon tetrachloride	23.31	1.0	20	0	117	70 - 130	24.05	3.09	20		
Chlorobenzene	24.33	1.0	20	1.542	114	70 - 114	25.62	5.14	20		
Chloroethane	20.67	1.0	20	0	103	70 - 130	24.72	17.8	20		
Chloroform	21.16	1.0	20	0	106	70 - 125	22.07	4.18	20		
Chloromethane	14.94	1.0	20	0	74.7	70 - 130	15.68	4.84	20		
cis-1,2-Dichloroethene	42.81	1.0	20	21.65	106	70 - 128	44.9	4.77	20		
cis-1,3-Dichloropropene	23.85	1.0	20	0	119	70 - 125	24.3	1.86	20		
Dibromochloromethane	23.1	1.0	20	0	115	70 - 124	23.74	2.73	20		
Dibromomethane	22.89	1.0	20	0	114	70 - 124	22.87	0.084	20		
Dichlorodifluoromethane	9.87	1.0	20	0	49.3	70 - 130	10.42	5.41	20	S	
Ethylbenzene	23.86	1.0	20	0	119	70 - 124	25.15	5.24	20		
Hexachlorobutadiene	26.61	1.0	20	0	133	70 - 130	28.15	5.63	20	S	
Isopropylbenzene	24.62	1.0	20	0	123	70 - 130	26.04	5.58	20		
m,p-Xylene	48.45	2.0	40	0	121	70 - 130	50.97	5.06	20		
Methylene chloride	22.91	2.0	20	0	115	70 - 128	24.38	6.23	20		
Naphthalene	24.2	1.0	20	0	121	70 - 130	25.37	4.71	20		
n-Butylbenzene	25.57	1.0	20	0	128	70 - 130	26.87	4.97	20		
n-Propylbenzene	24.76	1.0	20	0	124	70 - 130	26.34	6.18	20		
o-Xylene	24	1.0	20	0	120	70 - 124	25.44	5.82	20		
sec-Butylbenzene	25.1	1.0	20	0	125	70 - 130	26.76	6.41	20		
Styrene	24.44	1.0	20	0	122	70 - 130	25.73	5.15	20		
tert-Butylbenzene	24.73	1.0	20	0	124	70 - 130	26.06	5.27	20		
Tetrachloroethene	24.06	1.0	20	0	120	70 - 130	25.53	5.9	20		
Toluene	23.77	1.0	20	0	119	70 - 123	25.16	5.68	20		
trans-1,2-Dichloroethene	21.99	1.0	20	0	110	70 - 130	23.47	6.52	20		
trans-1,3-Dichloropropene	20.89	1.0	20	0	104	70 - 121	21.37	2.3	20		
Trichloroethene	68.98	1.0	20	47.56	107	70 - 129	72.45	4.9	20		
Trichlorofluoromethane	19.31	1.0	20	0	96.6	70 - 130	20.92	8.01	20		
Vinyl chloride	18.76	1.0	20	0	93.8	70 - 130	20.18	7.31	20		
Surr: 1,2-Dichloroethane-d4	43.56	1.0	50	0	87.1	70 - 126	44.17	1.38	20		
Surr: 4-Bromofluorobenzene	51.09	1.0	50	0	102	81 - 113	51.45	0.704	20		
Surr: Dibromofluoromethane	44.93	1.0	50	0	89.9	77 - 123	45.03	0.218	20		



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.

Project: LHAARP - 12

WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MSD	Sample ID: HS18120278-03MSD	Units: ug/L		Analysis Date: 18-Dec-2018 18:06						
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872167		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	52.59	1.0	50	0	105	82 - 127	53.03	0.822	20	

The following samples were analyzed in this batch:

HS18120278-01	HS18120278-02	HS18120278-03	HS18120278-04
HS18120278-05			



ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program



CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
North Carolina	624-2018	31-Dec-2018
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	22-Dec-2018
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019



Sample Receipt Checklist

Client Name: CBI-Houston
 Work Order: HS18120278

Date/Time Received: **05-Dec-2018 09:30**
 Received by: **RPG**

Checklist completed by: Raegen Giga 6-Dec-2018 Reviewed by: Sonia West 10-Dec-2018
 eSignature Date eSignature Date

Matrices: **GW** Carrier name: **ALS.HS**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- Chain of custody present? Yes No
- Chain of custody signed when relinquished and received? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- TX1005 solids received in hermetically sealed vials? Yes No N/A
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.8c/2.1c uc/c IR 25
 Cooler(s)/Kit(s): 25741
 Date/Time sample(s) sent to storage: 12/05/2018 19:55

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

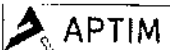
Client Contacted: Date Contacted: Person Contacted:

Contacted By: Regarding:

Comments:

Corrective Action:






COC ID: **LHAAP12-DEC2018-ALSHT-1812-** TURNAROUND TIME: normal RUSH: Page 1 of

PROJECT/CLIENT INFO				LABORATORY				OTHER INFO			
Facility Name	Longhorn AAP			Lab Name	ALS Laboratories			Email Invoice To	FedInvoices@aptim.com		
Project Number	501032			Lab Contact	RJ Modashia			Email Report To	Susan.Huang@aptim.com		
Address	LHAAP-12 1203-B East Grand Avenue PMB 202			Email	RJ.Modashia@alsglobal.com			Mail Reports To	Susan Huang		
City	Marshall	State	TX	Address	10450 Stauchiff Rd., Suite 210			Address	4005 Port Chicago Highway, Suit 200		
Postal Code	75670	Country	USA	City	Houston	State	TX	City	Concord	State	CA
Phone Number	713.243.7264			Postal Code	77099	Country	USA	Postal Code	94520	Country	USA
Project Manager	Praveen Srivastav			Phone Number	281.575.2279 or 281.530.5656			Shipping Company			

SAMPLE DETAILS										ANALYSIS REQUESTED	
Sample ID	Location	Start Depth	End Depth	Depth Unit	Field Matrix	Date	Time (24hr)	# Of Cont.	ANALYSIS	Yes by 8260B (3-40mm Von Vriest-w/TCU)	
12ww21-181204	LHAAP12	22.20	22.43		WG	12/4/18	1005	3	X		 HS18120278 Aptim Environmental & Infrastructure, Inc. LHAARP - 12
12ww21-181204-FD	LHAAP12	22.20	22.43		WG	12/4/18	1005	3	X		
12ww24-181204	LHAAP12	22.87	23.12		WG	12/4/18	110	3	X		
12ww24-181204-MS	LHAAP12	22.87	23.12		WG	12/4/18	1110	3	X		
12ww24-181204-MSD	LHAAP12	22.87	23.12		WG	12/4/18	1110	3	X		
12ww20-181204	LHAAP12	18.35	18.58		WG	12/4/18	1205	3	X		
TRIP BLANK					W	12/4/18		2	X		

ADDITIONAL COMMENTS/SPECIAL INSTRUCTIONS	RELINQUISHED BY/AFFILIATION	DATE/TIME	ACCEPTED BY/AFFILIATION	DATE/TIME
	Srinivasan / BHATIA	12/4/18 1400	R Ciga	12/5/18 09:30am
				COOL # 25741
				TEMP - 1.8c
				IR 25 CF +0.3c



 ALS 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CUSTODY SEAL		Seal Broken By: <i>SM</i>
	Date: <i>12/11/18</i>	Time: <i>14:00</i>	Date: <i>12/05/18</i>
	Name: <i>Scott Beesinger</i>		Comp any: <i>STATE</i>

25741 DEC 05 2018



Must Deliver Next Business Day
Time and Tempature Sensitive!

25741

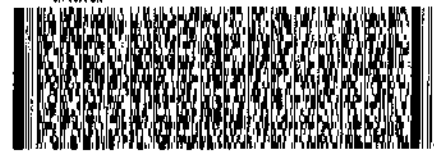
ORIGIN ID: SGRA (903) 930-6193
 ATT: SCOTT BEESINGER
 APTM ENVIRONMENTAL & INFRASTR. INC
 1203-B EAST GRAND AVE PMB202
 MARSHALL, TX 75670
 UNITED STATES US

SHIP DATE: 28NOV18
 ACTNGT: 1.00 LB MAN
 CAD: 300130/CAF3211
 DIMS: 19x15x13 IN

TO CLIENT SERVICES
 ALS LABORATORY GROUP
 10450 STANCLIFF ROAD
 SUITE 210
 HOUSTON TX 77099

(281) 530-6656
 REF: LHAAP-BO 62490-RJ

RMA: III III III

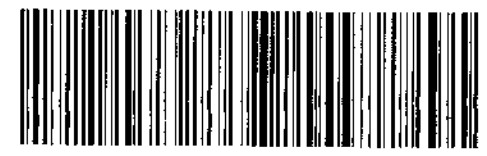


FedEx
 TRK#
 0221 4380 9535 0865

WED - 05 DEC 10:30A
 PRIORITY OVERNIGHT

AB SGRA

77099
 TX-US
 IAH



FID 162705 04DEC18 666A 553C1/71FF/BC0A



HS18120278 8260 Raw Data

ALS WO HS18120278



MSVOA09 -Logbook

Batch: 33625
 Date: 11-13-2018
 Method: 8260
 Comments:

Analyst: Presenta Cabascango
 Reviewer: Anjana Poluri
 Laboratory: Houston

#	Samp ID	Type	Analyzed	DF	Init Wt/Vol	Final Vol	File ID	Matrix	Status	pH
1	BFB	TUNE	11-13-2018 10:58 am	1.00	50 mL	50 mL	U111301.D	Liquid	Y	NA
	<i>Auto find/purged</i>									
2	VSTD000.25	ICAL1	11-13-2018 11:23 am	1.00	50 mL	50 mL	U111302.D	Liquid	Y	NA
	<i>0.25 uL cal std/250 mL DI</i>									
3	VSTD000.5	ICAL2	11-13-2018 11:47 am	1.00	50 mL	50 mL	U111303.D	Liquid	Y	NA
	<i>0.5 uL cal std/250 mL DI</i>									
4	VSTD001	ICAL3	11-13-2018 12:36 pm	1.00	50 mL	50 mL	U111304.D	Liquid	Y	NA
	<i>1 uL cal std/250 mL DI</i>									
5	VSTD002	ICAL4	11-13-2018 01:01 pm	1.00	50 mL	50 mL	U111305.D	Liquid	Y	NA
	<i>2 uL cal std/250 mL DI</i>									
6	VSTD005	ICAL5	11-13-2018 01:25 pm	1.00	50 mL	50 mL	U111306.D	Liquid	Y	NA
	<i>5 uL cal std/250 mL DI</i>									
7	VSTD020	ICAL6	11-13-2018 01:50 pm	1.00	50 mL	50 mL	U111307.D	Liquid	Y	NA
	<i>4 uL cal std/50 mL DI</i>									
8	VSTD050	ICAL7	11-13-2018 02:15 pm	1.00	50 mL	50 mL	U111308.D	Liquid	Y	NA
	<i>10 uL cal std/50 mL DI</i>									
9	VSTD100	ICAL8	11-13-2018 02:39 pm	1.00	50 mL	50 mL	U111309.D	Liquid	Y	NA
	<i>20 uL cal std/50 mL DI</i>									
10	VSTD150	ICAL9	11-13-2018 03:04 pm	1.00	50 mL	50 mL	U111310.D	Liquid	Y	NA
	<i>30 uL cal std/50 mL DI</i>									
11	VSTD200	ICAL	11-13-2018 03:28 pm	1.00	50 mL	50 mL	U111311.D	Liquid	Y	NA
	<i>40 uL cal std/50 mL DI</i>									
12	BLANK	SAMP	11-13-2018 03:53 pm	1.00	50 mL	50 mL	U111312.D	Liquid	Y	NA
	<i>Cleanup blk</i>									
13	VSTD050	ICV	11-13-2018 04:18 pm	1.00	50 mL	50 mL	U111313.D	Liquid	Y	NA
	<i>10 uL ICV std/50 mL DI</i>									
14	BFB	TUNE	11-13-2018 05:07 pm	1.00	50 mL	50 mL	V111301.D	Liquid	Y	NA
	<i>Auto find/purged</i>									
15	VSTD050	CCV	11-13-2018 05:32 pm	1.00	50 mL	50 mL	V111302.D	Liquid	Y	NA
	<i>10 uL cal std/50 mL DI</i>									
16	VSTD050	CCV	11-13-2018 05:56 pm	1.00	50 mL	50 mL	V111303.D	Liquid	Y	NA
	<i>10 uL cal std/50 mL DI</i>									
17	VLCSW-181113	LCS	11-13-2018 06:21 pm	1.00	50 mL	50 mL	V111304.D	Liquid	Y	NA
	<i>4 uL cal std/50 mL DI</i>									
18	BLANK	SAMP	11-13-2018 06:46 pm	1.00	50 mL	50 mL	V111305.D	Liquid	Y	NA
19	VBLKW-181113	MBLK	11-13-2018 07:10 pm	1.00	50 mL	50 mL	V111306.D	Liquid	Y	NA
20	HS18110428-01	SAMP	11-13-2018 07:35 pm	1.00	50 mL	50 mL	V111307.D	Liquid	Y	<2
21	HS18110496-01	SAMP	11-13-2018 08:00 pm	1.00	50 mL	50 mL	V111308.D	Liquid	Y	<2
22	HS18110428-02	SAMP	11-13-2018 08:25 pm	1.00	50 mL	50 mL	V111309.D	Liquid	Y	<2
23	HS18110428-03	SAMP	11-13-2018 08:49 pm	1.00	50 mL	50 mL	V111310.D	Liquid	Y	<2
24	HS18110428-04	SAMP	11-13-2018 09:14 pm	1.00	50 mL	50 mL	V111311.D	Liquid	Y	<2
25	HS18110496-04	SAMP	11-13-2018 09:39 pm	1.00	50 mL	50 mL	V111312.D	Liquid	Y	<2
26	HS18110496-05	SAMP	11-13-2018 10:04 pm	1.00	50 mL	50 mL	V111313.D	Liquid	Y	<2
27	HS18110496-06	SAMP	11-13-2018 10:28 pm	1.00	50 mL	50 mL	V111314.D	Liquid	Y	<2
28	HS18110428-08	SAMP	11-13-2018 10:53 pm	1.00	50 mL	50 mL	V111315.D	Liquid	Y	<2
29	HS18110428-09	SAMP	11-13-2018 11:18 pm	1.00	50 mL	50 mL	V111316.D	Liquid	Y	<2
30	HS18110496-04MS	MS	11-13-2018 11:43 pm	1.00	50 mL	50 mL	V111317.D	Liquid	Y	<2
	<i>8 uL cal std/40 mL Sample</i>									
31	HS18110496-04MSD	MSD	11-14-2018 12:07 am	1.00	50 mL	50 mL	V111318.D	Liquid	Y	<2
	<i>8 uL cal std/40 mL Sample</i>									



MSVOA09 -Logbook

#	<u>Samp ID</u>	<u>Type</u>	<u>Analyzed</u>	<u>DF</u>	<u>Init Wt/Vol</u>	<u>Final Vol</u>	<u>File ID</u>	<u>Matrix</u>	<u>Status</u>	<u>pH</u>
32	HS18110650-09	SAMP	11-14-2018 12:32 am	1.00	50 mL	50 mL	V111319.D	Liquid	Y	<2
33	HS18110428-05	SAMP	11-14-2018 12:57 am	5.00	10 mL	50 mL	V111320.D	Liquid	Y	<2
34	HS18110428-05	SAMP	11-14-2018 01:22 am	25.00	2 mL	50 mL	V111321.D	Liquid	Y	<2
35	HS18110428-06	SAMP	11-14-2018 01:46 am	5.00	10 mL	50 mL	V111322.D	Liquid	Y	<2
36	HS18110428-06	SAMP	11-14-2018 02:11 am	25.00	2 mL	50 mL	V111323.D	Liquid	Y	<2
37	HS18110428-07	SAMP	11-14-2018 02:36 am	100.00	500 µL	50 mL	V111324.D	Liquid	Y	<2
38	HS18110428-07	SAMP	11-14-2018 03:01 am	10.00	5 mL	50 mL	V111325.D	Liquid	Y	<2
39	HS18110650-05	SAMP	11-14-2018 03:25 am	1.00	50 mL	50 mL	V111326.D	Liquid	Y	<2
40	HS18110650-06	SAMP	11-14-2018 03:50 am	1.00	50 mL	50 mL	V111327.D	Liquid	Y	<2
41	VSTD050-END	CCV	11-14-2018 04:15 am	1.00	50 mL	50 mL	V111328.D	Liquid	Y	NA

10 uL cal std/50 mL DI

Chemical	Value
SURR SPK ID	30502-51-03
IS ID	30502-51-03
ICV STD ID	30603-41-01
LCS/MS ID	30502-51-01/02
CAL STD ID	30502-51-01/02
BFB ID	30502-51-03



FORM 3
WATER VOLATILE METHOD SPIKE RECOVERY

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Matrix Spike - Sample No.: VICV050

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE AMOUNT ()	% REC #	QC. LIMITS REC.
===== cis-1,3-Dichloropropene	50.00	53.22	106	80-120
trans-1,3-Dichloropropene	50.00	47.39	95	80-120
1,3-Dichlorobenzene	50.00	45.55	91	80-120
2,2-Dichloropropane	50.00	51.21	102	80-120
1,1-Dichloropropene	50.00	46.61	93	80-120
Dibromomethane	50.00	47.16	94	80-120
1,2-Dibromoethane	50.00	48.42	97	80-120
trans-1,2-Dichloroethene	50.00	46.36	93	80-120
1,1,1,2-Tetrachloroethane	50.00	49.98	100	80-120
1,1,1-Trichloroethane	50.00	48.15	96	80-120
1,1,2,2-Tetrachloroethane	50.00	45.96	92	80-120
Freon TF	50.00	46.03	92	80-120
1,1,2-Trichloroethane	50.00	45.83	92	80-120
1,1-Dichloroethane	50.00	45.57	91	80-120
1,1-Dichloroethene	50.00	45.66	91	80-120
Trichlorofluoromethane	50.00	46.76	94	80-120
1,2,3-Trichlorobenzene	50.00	48.30	97	80-120
Toluene	50.00	46.16	92	80-120
1,2,4-Trichlorobenzene	50.00	47.47	95	80-120
1,2,4-Trimethylbenzene	50.00	47.71	95	80-120
Tetrachloroethene	50.00	45.53	91	80-120
Trichloroethene	50.00	46.38	93	80-120
1,2-Dichlorobenzene	50.00	44.93	90	80-120
1,2-Dichloroethane	50.00	45.18	90	80-120
1,2-Dichloropropane	50.00	47.27	94	80-120
1,3,5-Trimethylbenzene	50.00	47.70	95	80-120
1,3-Dichloropropane	50.00	46.16	92	80-120
1,4-Dichlorobenzene	50.00	48.32	97	80-120

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits

COMMENTS: _____

FORM III VOA



FORM 3
WATER VOLATILE METHOD SPIKE RECOVERY

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No. : HS18120278

Matrix Spike - Sample No.: VICV050

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE AMOUNT ()	% REC #	QC. LIMITS REC.
2-Butanone	100.00	92.32	92	80-120
2-Chlorotoluene	50.00	45.71	91	80-120
2-Hexanone	100.00	99.26	99	80-120
4-Chlorotoluene	50.00	46.84	94	80-120
tert-Butylbenzene	50.00	47.82	96	80-120
4-Methyl-2-Pentanone	100.00	96.36	96	80-120
Acetone	100.00	95.46	95	80-120
Benzene	50.00	46.32	93	80-120
Bromobenzene	50.00	46.19	92	80-120
Bromochloromethane	50.00	50.90	102	80-120
Bromodichloromethane	50.00	48.53	97	80-120
Bromoform	50.00	47.71	95	80-120
Bromomethane	50.00	50.93	102	80-120
Carbon Disulfide	100.00	98.32	98	80-120
Carbon Tetrachloride	50.00	49.39	99	80-120
Chlorobenzene	50.00	45.35	91	80-120
Chloroethane	50.00	44.97	90	80-120
Chloroform	50.00	45.67	91	80-120
Chloromethane	50.00	46.70	93	80-120
cis-1,2-Dichloroethene	50.00	45.95	92	80-120
Dibromochloromethane	50.00	52.11	104	80-120
Dichlorodifluoromethane	50.00	47.87	96	80-120
Ethylbenzene	50.00	46.81	94	80-120
Hexachlorobutadiene	50.00	55.07	110	80-120
Isopropylbenzene	50.00	47.92	96	80-120
m,p-Xylenes	100.00	94.65	95	80-120
Methylene Chloride	50.00	48.07	96	80-120
n-Butylbenzene	50.00	49.50	99	80-120

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits

COMMENTS:

FORM III VOA



FORM 3
WATER VOLATILE METHOD SPIKE RECOVERY

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Matrix Spike - Sample No.: VICV050

COMPOUND	SPIKE ADDED (ug/L)	SAMPLE AMOUNT ()	% REC #	QC. LIMITS REC.
n-Propylbenzene	50.00	47.57	95	80-120
Naphthalene	50.00	51.20	102	80-120
o-Xylene	50.00	47.39	95	80-120
sec-Butylbenzene	50.00	48.40	97	80-120
Styrene	50.00	49.81	100	80-120
Vinyl Chloride	50.00	47.66	95	80-120
1,2,3-Trichloropropane	50.00	49.28	98	80-120
p-Isopropyltoluene	50.00	49.15	98	80-120
1,2-Dibromo-3-Chloropro	50.00	47.82	96	80-120

Column to be used to flag recovery and RPD values with an asterisk

* Values outside of QC limits

COMMENTS:

FORM III VOA



FORM 5
VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Lab File ID: U111301

BFB Injection Date: 11/13/18

Instrument ID: VOA9

BFB Injection Time: 1058

GC Column: DB624

ID: 0.25 (mm)

Heated Purge: (Y/N) N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	20.9
75	30.0 - 60.0% of mass 95	53.3
95	Base Peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	7.1
173	Less than 2.0% of mass 174	0.6 (1.0)1
174	Greater than 50.0% of mass 95	67.7
175	5.0 - 9.0% of mass 174	5.2 (7.7)1
176	95.0 - 101.0% of mass 174	64.6 (95.4)1
177	5.0 - 9.0% of mass 176	4.8 (7.4)2

1-Value is % mass 174

2-Value is % mass 176

THIS CHECK APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

	EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
01	VSTD000.25	VSTD000.25	U111302	11/13/18	1123
02	VSTD000.5	VSTD000.5	U111303	11/13/18	1147
03	VSTD001	VSTD001	U111304	11/13/18	1236
04	VSTD002	VSTD002	U111305	11/13/18	1301
05	VSTD005	VSTD005	U111306	11/13/18	1325
06	VSTD020	VSTD020	U111307	11/13/18	1350
07	VSTD050	VSTD050	U111308	11/13/18	1415
08	VSTD100	VSTD100	U111309	11/13/18	1439
09	VSTD150	VSTD150	U111310	11/13/18	1504
10	VSTD200	VSTD200	U111311	11/13/18	1528
11	VICV050	VICV050	U111313	11/13/18	1618
12					
13					
14					
15					
16					
17					
18					
19					
20					
21					
22					

page 1 of 1

FORM V VOA



FORM 6
VOLATILE INITIAL CALIBRATION DATA

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Instrument ID: VOA9

Calibration Date(s): 11/13/18 11/13/18

Column: DB624

ID: 0.18 (mm)

Calibration Time(s): 1123

1528

LAB FILE ID:

RF0.25: U111302

RF0.5: U111303

RF1: U111304

RF2: U111305

RF5: U111306

RF20: U111307

COMPOUND	RF0.25	RF0.5	RF1	RF2	RF5	RF20
===== cis-1,3-Dichloropropene		0.514	0.452	0.424	0.458	0.523
trans-1,3-Dichloropropene		3218	5482	10881	30108	147063
1,3-Dichlorobenzene		2.016	1.635	1.560	1.502	1.485
2,2-Dichloropropane		0.895	0.675	0.644	0.597	0.645
1,1-Dichloropropene		0.589	0.484	0.455	0.407	0.432
Dibromomethane		0.272	0.224	0.217	0.212	0.216
1,2-Dibromoethane		0.438	0.364	0.337	0.365	0.365
trans-1,2-Dichloroethene		0.776	0.676	0.607	0.562	0.575
1,1,1,2-Tetrachloroethane		0.357	0.286	0.272	0.294	0.309
1,1,1-Trichloroethane		1.047	0.838	0.848	0.741	0.818
1,1,2,2-Tetrachloroethane		1.478	1.172	1.145	1.215	1.150
Freon TF		0.612	0.528	0.532	0.409	0.460
1,1,2-Trichloroethane		0.411	0.336	0.304	0.314	0.307
1,1-Dichloroethane		1.465	1.215	1.128	1.077	1.065
1,1-Dichloroethene		0.736	0.626	0.565	0.474	0.526
Trichlorofluoromethane		1.241	1.099	1.043	0.817	0.913
1,2,3-Trichlorobenzene		1.196	0.776	0.891	0.911	0.922
Toluene		2.078	1.628	1.576	1.529	1.532
1,2,4-Trichlorobenzene		1.260	0.914	0.912	0.902	0.924
1,2,4-Trimethylbenzene		3.320	2.638	2.610	2.690	2.770
Tetrachloroethene		0.336	0.285	0.263	0.234	0.246
Trichloroethene		0.440	0.361	0.330	0.306	0.325
1,2-Dichlorobenzene		2.072	1.662	1.534	1.584	1.503
1,2-Dichloroethane		0.688	0.543	0.533	0.514	0.502
1,2-Dichloropropane		0.420	0.368	0.338	0.340	0.346
1,3,5-Trimethylbenzene		3.173	2.583	2.523	2.530	2.676
1,3-Dichloropropane		0.875	0.710	0.656	0.685	0.660
1,4-Dichlorobenzene		7995	11659	22453	55298	228900
2-Butanone		0.438	0.357	0.328	0.355	0.335
2-Chlorotoluene		3.108	2.512	2.344	2.371	2.366
2-Hexanone		0.309	0.250	0.264	0.296	0.296
4-Chlorotoluene		3.515	2.726	2.736	2.786	2.768
tert-Butylbenzene		2.788	2.149	2.208	2.094	2.254
4-Methyl-2-Pentanone		0.448	0.409	0.382	0.418	0.418
Acetone		5795	7657	12023	24249	87797
Benzene		1.791	1.472	1.332	1.296	1.322
Bromobenzene		1.057	0.812	0.769	0.798	0.781
Bromochloromethane		0.288	0.289	0.307	0.306	0.304
Bromodichloromethane		0.499	0.390	0.373	0.378	0.400
Bromoform		1478	2470	4530	13391	60993
Bromomethane		2224	4167	7606	17760	65394
Carbon Disulfide		2.186	1.775	1.666	1.469	1.640
Carbon Tetrachloride		0.413	0.348	0.330	0.290	0.344
Chlorobenzene		1.390	1.114	0.980	0.979	0.984

FORM VI VOA



FORM 6
VOLATILE INITIAL CALIBRATION DATA

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS1812027

Instrument ID: VOA9

Calibration Date(s): 11/13/18 11/13/18

Column: DB624

ID: 0.18 (mm)

Calibration Time(s): 1123

1528

LAB FILE ID:
RF2: U111305RF0.25: U111302
RF5: U111306RF0.5: U111303
RF20: U111307

RF1: U111304

COMPOUND	RF0.25	RF0.5	RF1	RF2	RF5	RF20
Chloroethane		0.691	0.561	0.547	0.455	0.484
Chloroform		1.493	1.230	1.118	1.087	1.079
Chloromethane		4145	6506	12000	26431	114171
cis-1,2-Dichloroethene		0.890	0.758	0.706	0.655	0.674
Dibromochloromethane		0.340	0.262	0.256	0.287	0.314
Dichlorodifluoromethane		4116	6358	12615	24157	119108
Ethylbenzene		0.660	0.543	0.490	0.475	0.502
Hexachlorobutadiene		2273	2210	4667	8782	45235
Isopropylbenzene		1.845	1.478	1.459	1.431	1.486
m,p-Xylenes		0.830	0.625	0.602	0.602	0.634
Methylene Chloride		5188	7766	13586	29126	113894
n-Butylbenzene		11934	17894	36189	79206	381141
n-Propylbenzene		4.808	3.945	3.820	3.699	3.871
Naphthalene		2.957	2.577	2.535	2.887	3.092
o-Xylene		0.725	0.632	0.608	0.619	0.643
sec-Butylbenzene		4.052	3.193	3.096	2.852	3.218
Styrene		1.201	0.990	1.012	1.047	1.097
Vinyl Chloride		4801	7374	13465	29718	136290
1,2,3-Trichloropropane		1.381	1.095	1.101	1.220	1.204
p-Isopropyltoluene		3.227	2.386	2.598	2.465	2.727
1,2-Dibromo-3-Chloropropane		479	955	1762	5351	24969
1,2-Dichloroethane-d4		8937	11059	18531	38984	138453
Dibromofluoromethane		5664	7705	12925	27004	102630
Toluene-d8		23871	30489	51591	112942	412445
4-Bromofluorobenzene		8554	10512	18827	41214	154498

FORM VI VOA



FORM 6
VOLATILE INITIAL CALIBRATION DATA

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS1812027

Instrument ID: VOA9

Calibration Date(s): 11/13/18 11/13/18

Column: DB624

ID: 0.18 (mm)

Calibration Time(s): 1123 1528

LAB FILE ID: RF50: U111308 RF100: U111309 RF150: U111310
RF200: U111311

COMPOUND	RF50	RF100	RF150	RF200
cis-1,3-Dichloropropene	0.547	0.595	0.608	0.588
trans-1,3-Dichloropropene	407060	912116	1439550	1983766
1,3-Dichlorobenzene	1.441	1.512	1.522	1.494
2,2-Dichloropropane	0.668	0.740	0.784	0.738
1,1-Dichloropropene	0.414	0.457	0.468	0.451
Dibromomethane	0.214	0.221	0.220	0.212
1,2-Dibromoethane	0.363	0.366	0.360	0.353
trans-1,2-Dichloroethene	0.578	0.598	0.615	0.580
1,1,1,2-Tetrachloroethane	0.320	0.333	0.335	0.332
1,1,1-Trichloroethane	0.837	0.918	0.956	0.902
1,1,2,2-Tetrachloroethane	1.122	1.135	1.138	1.104
Freon TF	0.433	0.506	0.529	0.498
1,1,2-Trichloroethane	0.296	0.302	0.299	0.291
1,1-Dichloroethane	1.077	1.100	1.127	1.057
1,1-Dichloroethene	0.507	0.543	0.568	0.534
Trichlorofluoromethane	0.888	0.996	1.035	0.964
1,2,3-Trichlorobenzene	0.895	0.964	0.981	0.959
Toluene	1.480	1.511	1.507	1.445
1,2,4-Trichlorobenzene	0.918	0.997	1.018	1.004
1,2,4-Trimethylbenzene	2.646	2.808	2.798	2.714
Tetrachloroethene	0.231	0.253	0.257	0.253
Trichloroethene	0.310	0.331	0.336	0.327
1,2-Dichlorobenzene	1.470	1.530	1.529	1.486
1,2-Dichloroethane	0.482	0.494	0.487	0.468
1,2-Dichloropropane	0.339	0.355	0.356	0.342
1,3,5-Trimethylbenzene	2.552	2.731	2.750	2.658
1,3-Dichloropropane	0.642	0.647	0.644	0.623
1,4-Dichlorobenzene	576011	1240019	1907496	2606334
2-Butanone	0.340	0.348	0.343	0.318
2-Chlorotoluene	2.278	2.377	2.372	2.306
2-Hexanone	0.297	0.307	0.305	0.292
4-Chlorotoluene	2.640	2.769	2.781	2.705
tert-Butylbenzene	2.136	2.334	2.363	2.278
4-Methyl-2-Pentanone	0.416	0.431	0.421	0.410
Acetone	201617	430350	636098	869596
Benzene	1.290	1.330	1.333	1.271
Bromobenzene	0.757	0.802	0.810	0.796
Bromochloromethane	0.308	0.299	0.285	0.266
Bromodichloromethane	0.406	0.432	0.438	0.424
Bromoform	170886	383125	610635	844522
Bromomethane	154362	367005	609177	
Carbon Disulfide	1.661	1.759	1.826	1.705
Carbon Tetrachloride	0.337	0.388	0.405	0.392
Chlorobenzene	0.950	0.978	0.966	0.946

FORM VI VOA



FORM 6
VOLATILE INITIAL CALIBRATION DATA

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS1812027

Instrument ID: VOA9

Calibration Date(s): 11/13/18 11/13/18

Column: DB624

ID: 0.18 (mm)

Calibration Time(s): 1123

1528

LAB FILE ID:

RF50: U111308

RF100: U111309

RF150: U111310

RF200: U111311

COMPOUND	RF50	RF100	RF150	RF200
Chloroethane	0.482	0.493	0.509	0.472
Chloroform	1.082	1.104	1.121	1.047
Chloromethane	285521	595590	963160	1334230
cis-1,2-Dichloroethene	0.664	0.681	0.694	0.654
Dibromochloromethane	0.333	0.351	0.354	0.351
Dichlorodifluoromethane	293949	680233	1086406	1483355
Ethylbenzene	0.492	0.514	0.517	0.504
Hexachlorobutadiene	110558	270039	426795	
Isopropylbenzene	1.427	1.543	1.535	1.471
m,p-Xylenes	0.609	0.638	0.632	0.615
Methylene Chloride	276248	577262	911224	1246871
n-Butylbenzene	954951	2203969	3347969	
n-Propylbenzene	3.668	3.966	3.962	3.771
Naphthalene	3.062	3.229	3.212	3.095
o-Xylene	0.629	0.652	0.649	0.635
sec-Butylbenzene	3.062	3.389	3.420	3.284
Styrene	1.100	1.130	1.117	1.082
Vinyl Chloride	335638	744825	1189386	1622522
1,2,3-Trichloropropane	1.240	1.296	1.309	1.275
p-Isopropyltoluene	2.627	2.886	2.907	2.820
1,2-Dibromo-3-Chloropropane	66581	149236	234820	321770
1,2-Dichloroethane-d4	335901	696595	1062364	1441844
Dibromofluoromethane	255826	526814	809097	1112100
Toluene-d8	1045096	2142731	3238162	4386438
4-Bromofluorobenzene	396604	823996	1261578	1723103

FORM VI VOA



FORM 6
VOLATILE INITIAL CALIBRATION DATA

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS1812027

Instrument ID: VOA9

Calibration Date(s): 11/13/18 11/13/18

Column: DB624

ID: 0.18 (mm)

Calibration Time(s): 1123

1528

COMPOUND	CURVE	COEFFICIENTS			%RSD OR R ²	MAX %RSD OR R ²
		A0	A1	A2		
===== cis-1,3-Dichloropropene	AVRG		0.52317106		12.886	20.000
trans-1,3-Dichloropropene	LINR	2.955e-002	1.87475164		0.9992282	0.9900000
1,3-Dichlorobenzene	AVRG		1.57423339		11.066	20.000
2,2-Dichloropropane	AVRG		0.70955952		12.773	20.000
1,1-Dichloropropene	AVRG		0.46178698		11.618	20.000
Dibromomethane	AVRG		0.22310340		8.428	20.000
1,2-Dibromoethane	AVRG		0.36809182		7.515	20.000
trans-1,2-Dichloroethene	AVRG		0.61854084		10.984	20.000
1,1,1,2-Tetrachloroethane	AVRG		0.31538401		8.665	20.000
1,1,1-Trichloroethane	AVRG		0.87828248		10.125	20.000
1,1,2,2-Tetrachloroethane	AVRG		1.18433615		9.680	20.000
Freon TF	AVRG		0.50088227		12.161	20.000
1,1,2-Trichloroethane	AVRG		0.31787570		11.701	20.000
1,1-Dichloroethane	AVRG		1.14578561		11.273	20.000
1,1-Dichloroethene	AVRG		0.56438078		13.689	20.000
Trichlorofluoromethane	AVRG		0.99961564		12.536	20.000
1,2,3-Trichlorobenzene	AVRG		0.94398267		11.868	20.000
Toluene	AVRG		1.58742783		12.041	20.000
1,2,4-Trichlorobenzene	AVRG		0.98325969		11.514	20.000
1,2,4-Trimethylbenzene	AVRG		2.77717297		7.772	20.000
Tetrachloroethene	AVRG		0.26191977		12.290	20.000
Trichloroethene	AVRG		0.34069901		11.908	20.000
1,2-Dichlorobenzene	AVRG		1.59672308		11.731	20.000
1,2-Dichloroethane	AVRG		0.52321079		12.632	20.000
1,2-Dichloropropane	AVRG		0.35598971		7.249	20.000
1,3,5-Trimethylbenzene	AVRG		2.68643867		7.489	20.000
1,3-Dichloropropane	AVRG		0.68254990		11.227	20.000
1,4-Dichlorobenzene	LINR	-8.51e-004	0.64997268		0.9997856	0.9900000
2-Butanone	AVRG		0.35147036		9.867	20.000
2-Chlorotoluene	AVRG		2.44838115		10.438	20.000
2-Hexanone	AVRG		0.29065206		6.972	20.000
4-Chlorotoluene	AVRG		2.82514687		9.297	20.000
tert-Butylbenzene	AVRG		2.28952823		9.083	20.000
4-Methyl-2-Pentanone	AVRG		0.41710960		4.264	20.000
Acetone	LINR	-8.86e-002	4.54997798		0.9976530	0.9900000
Benzene	AVRG		1.38203708		11.842	20.000
Bromobenzene	AVRG		0.82019754		11.060	20.000
Bromochloromethane	AVRG		0.29476503		4.761	20.000
Bromodichloromethane	AVRG		0.41544996		9.300	20.000
Bromoform	2ORDR	2.367e-002	4.32863402	-0.1428982	0.9996117	0.9900000
Bromomethane	LINR	3.806e-002	2.31863844		0.9951529	0.9900000
Carbon Disulfide	AVRG		1.74293069		11.174	20.000
Carbon Tetrachloride	AVRG		0.36075188		11.334	20.000
Chlorobenzene	AVRG		1.03196120		13.892	20.000

FORM VI VOA



FORM 6
VOLATILE INITIAL CALIBRATION DATA

Lab Name: ALS LABORATORY GROUP Contract:
 Lab Code: ALS-HS Case No.: SAS No.: SDG No.: HS1812027
 Instrument ID: VOA9 Calibration Date(s): 11/13/18 11/13/18
 Column: DB624 ID: 0.18 (mm) Calibration Time(s): 1123 1528

COMPOUND	CURVE	COEFFICENTS			%RSD	MAX %RSD
		A0	A1	A2	OR R ²	OR R ²
Chloroethane	AVRG		0.52169853		13.844	20.000
Chloroform	AVRG		1.15119170		11.977	20.000
Chloromethane	LINR	4.59e-003	1.49116761		0.9990790	0.9900000
cis-1,2-Dichloroethene	AVRG		0.70849189		10.610	20.000
Dibromochloromethane	AVRG		0.31638584		12.330	20.000
Dichlorodifluoromethane	LINR	2.002e-002	1.32602499		0.9979384	0.9900000
Ethylbenzene	AVRG		0.52179627		10.571	20.000
Hexachlorobutadiene	2ORDR	5.037e-003	3.37955533	-0.5103543	0.9988978	0.9900000
Isopropylbenzene	AVRG		1.51953048		8.449	20.000
m,p-Xylenes	AVRG		0.64297054		11.093	20.000
Methylene Chloride	LINR	-1.4e-002	1.58950727		0.9987446	0.9900000
n-Butylbenzene	2ORDR	7.523e-003	0.38680994	-2.96e-003	0.9992256	0.9900000
n-Propylbenzene	AVRG		3.94550864		8.657	20.000
Naphthalene	AVRG		2.96078148		8.567	20.000
o-Xylene	AVRG		0.64354428		5.210	20.000
sec-Butylbenzene	AVRG		3.28524136		10.228	20.000
Styrene	AVRG		1.08640200		5.880	20.000
Vinyl Chloride	LINR	1.016e-002	1.21438330		0.9982930	0.9900000
1,2,3-Trichloropropane	AVRG		1.23587941		7.613	20.000
p-Isopropyltoluene	AVRG		2.73812048		9.374	20.000
1,2-Dibromo-3-Chloropropane	2ORDR	2.08e-002	5.38876079	-0.2033476	0.9995316	0.9900000
1,2-Dichloroethane-d4	LINR	-3.55e-002	1.36996396		0.9980055	0.9900000
Dibromofluoromethane	LINR	-2.51e-002	1.78351549		0.9987270	0.9900000
Toluene-d8	LINR	-3.42e-002	0.80924260		0.9991601	0.9900000
4-Bromofluorobenzene	LINR	-2.13e-002	2.06564990		0.9996344	0.9900000

FORM VI VOA



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111301.D

Page 1

Date : 13-NOV-2018 10:58

Client ID: BFB

Instrument: VOA9.i

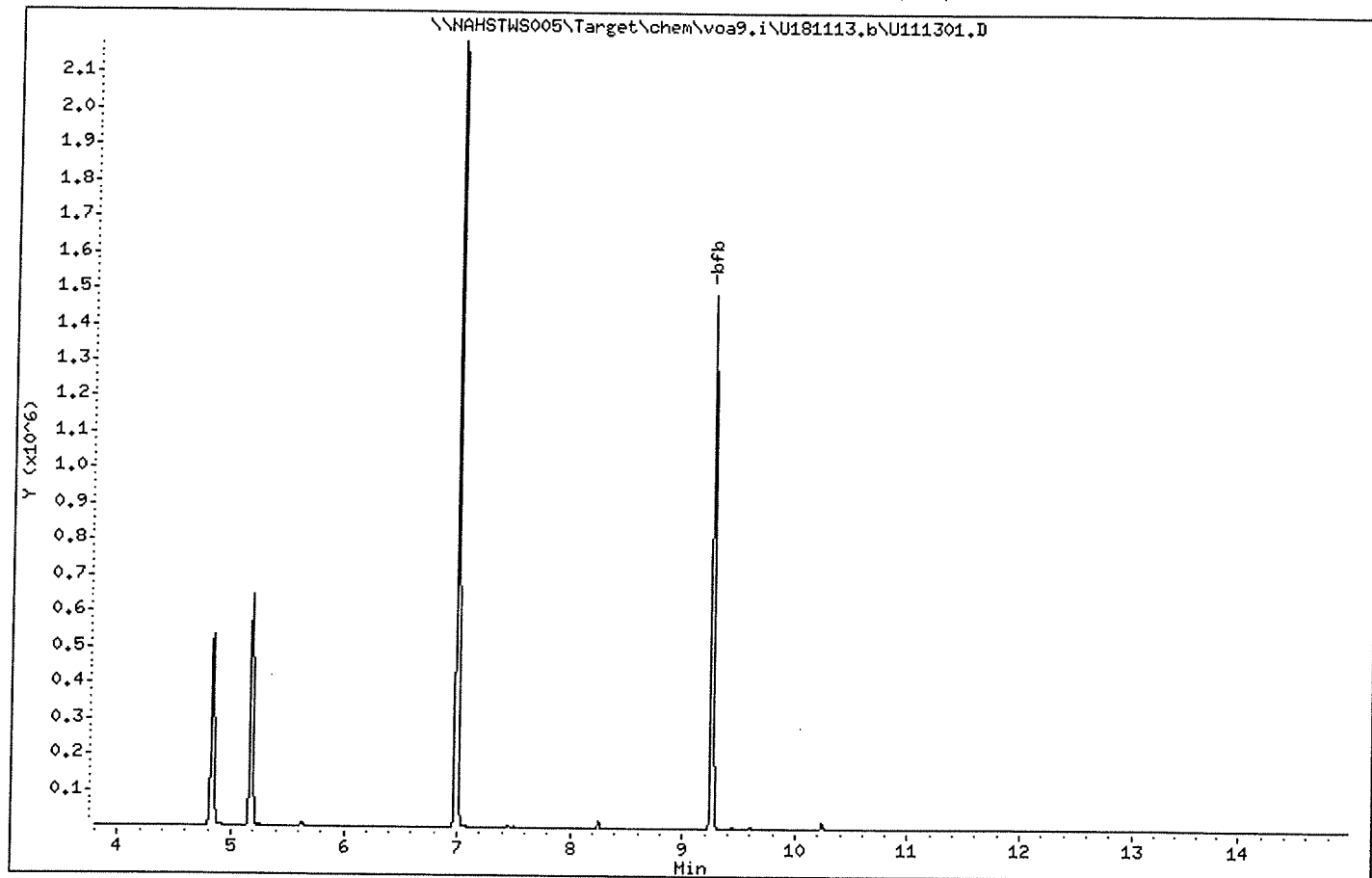
Sample Info: BFB;BFB;3;;BFB

Volume Injected (uL): 2.0

Operator: PC

Column phase: DB624

Column diameter: 0.25



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111301.D

Page 2

Date : 13-NOV-2018 10:58

Client ID: BFB

Instrument: VOA9.i

Sample Info: BFB;BFB;3;;BFB

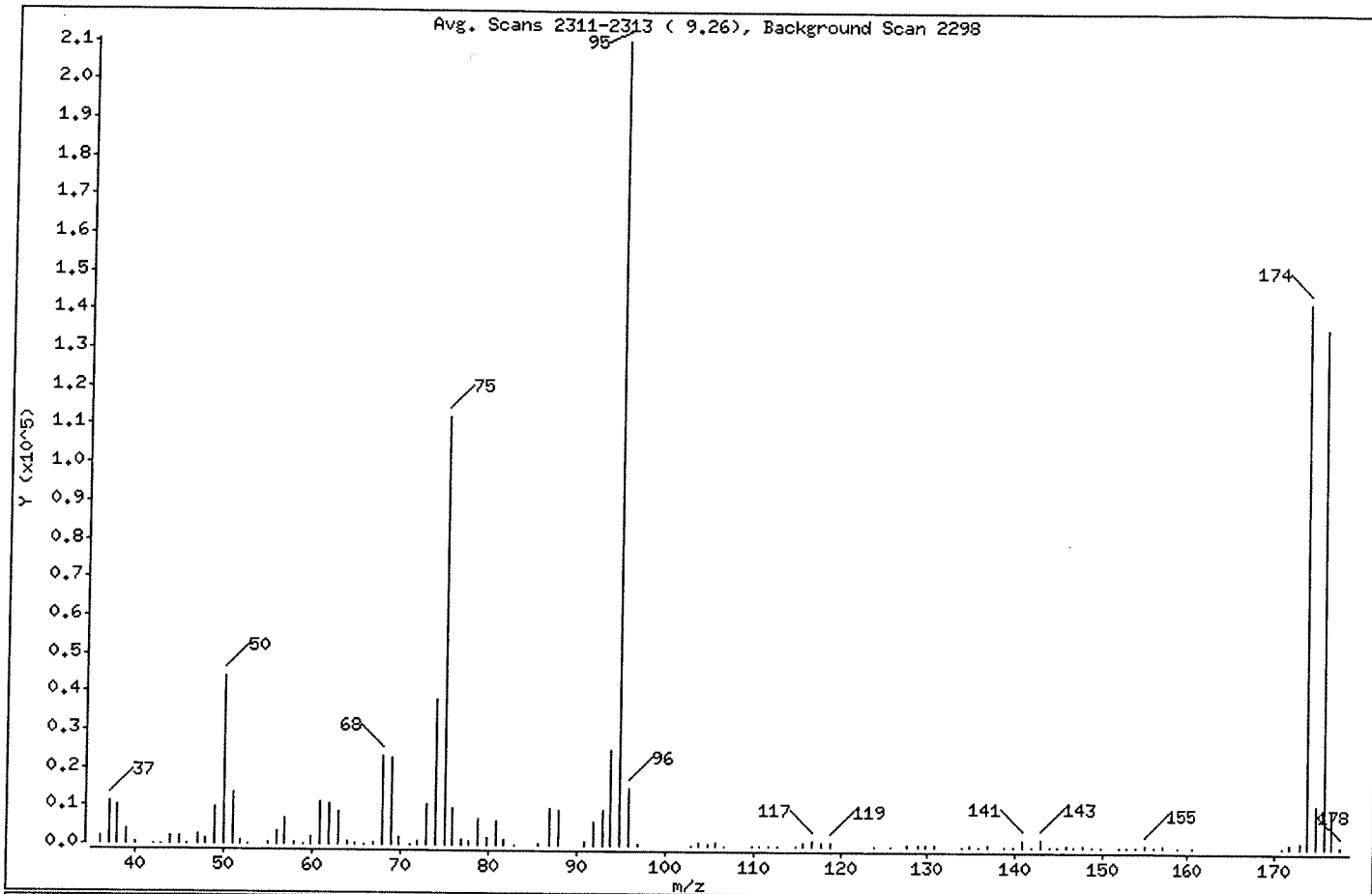
Volume Injected (uL): 2.0

Operator: PC

Column phase: DB624

Column diameter: 0.25

1 bfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
95	Base Peak, 100% relative abundance	100.00
50	15.00 - 40.00% of mass 95	20.93
75	30.00 - 60.00% of mass 95	53.26
96	5.00 - 9.00% of mass 95	7.05
173	Less than 2.00% of mass 174	0.64 (0.95)
174	Greater than 50.00% of mass 95	67.71
175	5.00 - 9.00% of mass 174	5.22 (7.70)
176	95.00 - 101.00% of mass 174	64.61 (95.42)
177	5.00 - 9.00% of mass 176	4.79 (7.41)



Data File: \\NAHSTMS005\Target\chem\voa9.i\U181113.b\U111301.D

Page 3

Date : 13-NOV-2018 10:58

Client ID: BFB

Instrument: VOA9.i

Sample Info: BFB;BFB;3;BFB

Volume Injected (uL): 2.0

Operator: PC

Column phase: DB624

Column diameter: 0.25

Data File: U111301.D
 Spectrum: Avg. Scans 2311-2313 (9.26), Background Scan 2298
 Location of Maximum: 94.95
 Number of points: 108

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	2069	65.90	90	102.90	106	142.80	1933
37.00	11082	66.90	610	103.80	947	143.90	71
38.00	10274	67.90	23360	104.80	338	144.70	179
39.00	3978	69.00	22944	105.80	973	145.70	325
40.00	381	69.90	1893	106.80	216	146.60	113
42.00	60	71.00	66	109.80	116	147.70	442
42.90	178	71.90	1065	110.70	166	148.60	101
43.90	1871	72.90	10562	111.80	126	149.70	180
45.00	2116	73.90	38432	112.70	189	151.80	82
45.90	123	75.00	112016	114.80	219	152.70	121
47.00	2495	75.90	9870	115.70	740	153.70	104
48.00	1426	76.90	1360	116.70	1240	154.70	552
49.00	9563	77.80	949	117.80	730	155.80	58
50.00	44016	78.80	6592	118.80	907	156.70	431
51.00	13714	79.80	1796	123.80	120	158.70	192
51.90	746	80.80	6503	125.70	57	160.60	164
52.70	59	81.80	1400	127.70	639	170.70	72
55.00	590	82.90	197	128.90	349	171.70	746
56.00	3508	85.70	252	129.70	626	172.80	1355
57.00	6688	86.90	9661	130.80	249	173.70	142400
57.90	351	87.90	8961	133.90	56	174.70	10971
58.90	116	90.80	826	134.70	394	175.70	135872
59.90	2009	91.90	6125	135.70	82	176.70	10067
60.90	11206	92.90	9048	136.70	337	177.70	296
62.00	10418	93.90	25168	138.70	60		
63.00	8677	94.90	210304	139.60	79		
63.90	848	95.90	14829	140.70	1870		
64.90	610	96.90	545	141.80	179		



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111302.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111302.D
 Lab Smp Id: VSTD000.25 Client Smp ID: VSTD000.25
 Inj Date : 13-NOV-2018 11:23
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD000.25;VSTD000.25;1;1;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 11:23 Cal File: U111302.D
 Als bottle: 2 Calibration Sample, Level: 1
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG				AMOUNTS		
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	421654	50.0000	
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	823585	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	757808	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	346489	50.0000	
\$ 30 Dibromofluoromethane	113	4.830	4.830	(0.987)	4873	0.25000	(a)
\$ 35 1,2-Dichloroethane-d4	65	5.171	5.171	(1.057)	7659	0.25000	(a)
\$ 48 Toluene-d8	98	6.989	6.989	(0.847)	19596	0.25000	(a)
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	7125	0.25000	(a)

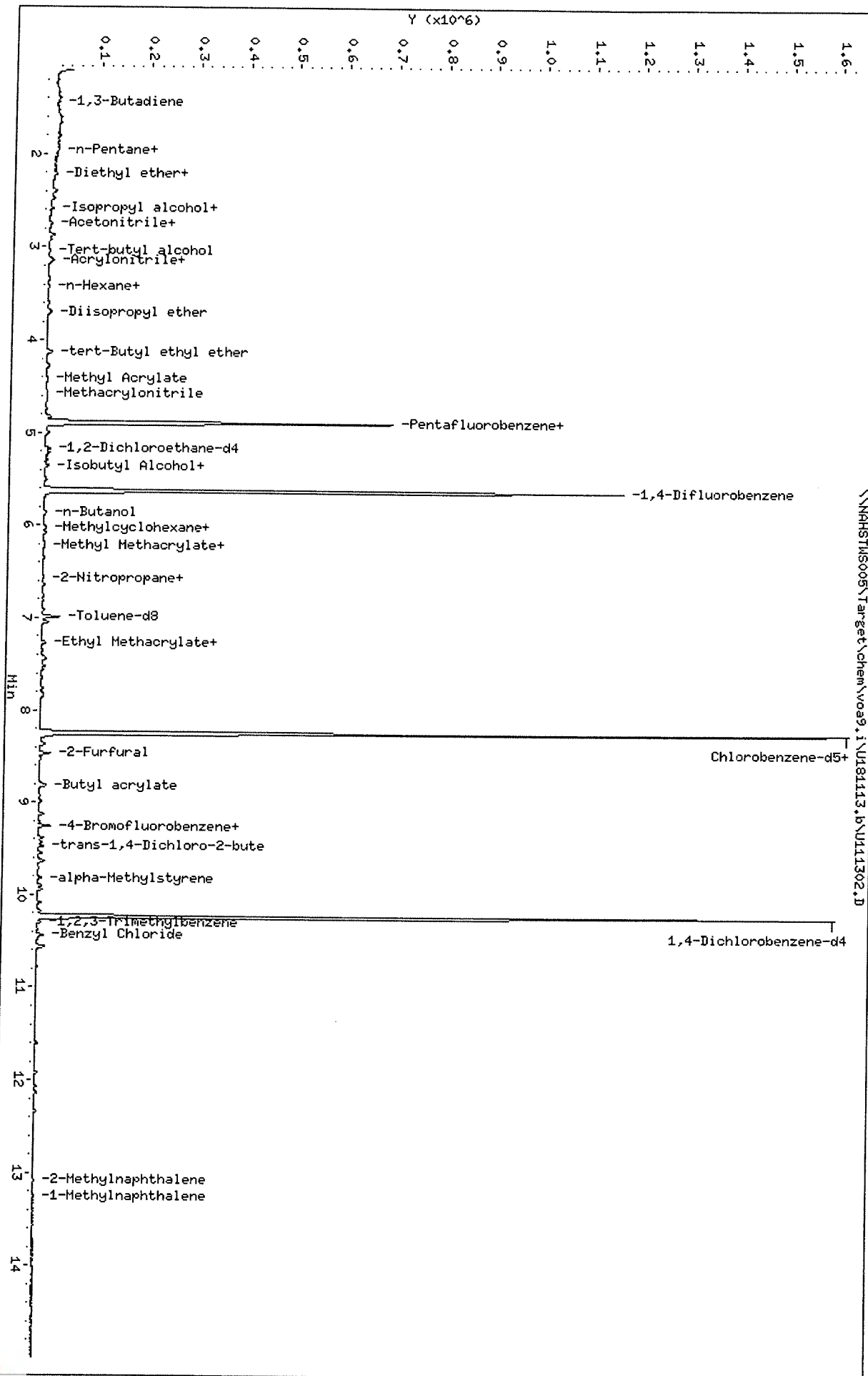
QC Flag Legend

a - Target compound detected but, quantitated amount
 Below Limit Of Quantitation(BLOQ).



Data File: \\NAHSTMS005\Target\chem\voa9.i\U18113.b\U111302.D
 Date : 13-NOV-2018 11:23
 Client ID: VSTD000.25
 Sample Info: VSTD000.25;VSTD000.25;1:1;
 Purge Volume: 5.0
 Column phase: DB624

Instrument: W0A9.i
 Operator: PC
 Column diameter: 0.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111303.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111303.D
 Lab Smp Id: VSTD000.5 Client Smp ID: VSTD000.5
 Inj Date : 13-NOV-2018 11:47
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD000.5;VSTD000.5;1;2;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 11:23 Cal File: U111302.D
 Als bottle: 3 Calibration Sample, Level: 2
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG				AMOUNTS		
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.898	(1.000)	421285	50.0000	
* 36 1,4-Difluorobenzene	114	5.629	5.629	(1.000)	817288	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	753573	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	345123	50.0000	
\$ 30 Dibromofluoromethane	113	4.834	4.834	(0.987)	5664	0.50000	(a)
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.179	(1.057)	8937	0.50000	(a)
\$ 48 Toluene-d8	98	6.990	6.990	(0.847)	23871	0.50000	(a)
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	8554	0.50000	0.10(a)
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	2690	0.50000	0.56(a)
31 1,1,1-Trichloroethane	97	4.830	4.830	(0.986)	4410	0.50000	0.59(a)
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	5102	0.50000	0.62(aM)
138 Freon TF	101	2.409	2.409	(0.492)	2580	0.50000	0.61(a)
53 1,1,2-Trichloroethane	83	7.421	7.421	(0.900)	3095	0.50000	0.64(a)
22 1,1-Dichloroethane	63	3.608	3.608	(0.737)	6173	0.50000	0.63(a)
11 1,1-Dichloroethene	96	2.409	2.409	(0.492)	3102	0.50000	0.65(a)
32 1,1-Dichloropropene	75	5.006	5.006	(0.889)	4812	0.50000	0.63(a)
93 1,2,3-Trichlorobenzene	180	12.339	12.339	(1.205)	4127	0.50000	0.63(a)
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	4767	0.50000	0.55(a)
90 1,2,4-Trichlorobenzene	180	11.927	11.927	(1.165)	4348	0.50000	0.64(a)
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	11460	0.50000	0.59(a)
89 1,2-Dibromo-3-Chloropropane	155	11.237	11.237	(1.098)	479	0.50000	1.41(aM)
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	3298	0.50000	0.59(a)



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111303.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
=====	=====	=====	=====	=====	=====	=====	=====
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	7152	0.50000	0.64(a)
33 1,2-Dichloroethane	62	5.258	5.258	(0.934)	5619	0.50000	0.65(aM)
42 1,2-Dichloropropane	63	6.082	6.082	(1.081)	3429	0.50000	0.58(a)
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	10952	0.50000	0.59(a)
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	6957	0.50000	0.64(a)
54 1,3-Dichloropropane	76	7.567	7.567	(0.917)	6595	0.50000	0.64(a)
84 1,4-Dichlorobenzene	146	10.258	10.258	(1.002)	7995	0.50000	0.71(a)
26 2,2-Dichloropropane	77	4.279	4.279	(0.874)	3772	0.50000	0.63(a)
24 2-Butanone	43	4.354	4.354	(0.889)	3689	1.00000	1.24(a)
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	10726	0.50000	0.63(a)
52 2-Hexanone	43	7.649	7.649	(0.927)	4658	1.00000	1.06(a)
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	12130	0.50000	0.62(a)
82 p-Isopropyltoluene	119	10.213	10.213	(0.998)	11138	0.50000	0.58(a)
45 4-Methyl-2-Pentanone	43	6.915	6.915	(0.838)	6749	1.00000	1.07(a)
10 Acetone	43	2.487	2.487	(0.508)	5795	1.00000	(a)
37 Benzene	78	5.220	5.220	(0.927)	14635	0.50000	0.64(a)
74 Bromobenzene	156	9.381	9.381	(0.917)	3648	0.50000	0.64(a)
29 Bromochloromethane	128	4.560	4.560	(0.931)	1212	0.50000	0.48(a)
39 Bromodichloromethane	83	6.348	6.348	(1.128)	4075	0.50000	0.60(aM)
66 Bromoform	173	8.988	8.988	(1.090)	1478	0.50000	1.60(Ta)
6 Bromomethane	94	1.678	1.678	(0.343)	2224	0.50000	2.51(a)
19 Carbon Disulfide	76	2.596	2.596	(0.530)	18415	1.00000	1.25(a)
34 Carbon Tetrachloride	117	4.995	4.995	(0.887)	3377	0.50000	0.57(a)
59 Chlorobenzene	112	8.275	8.275	(1.003)	10477	0.50000	0.67(a)
7 Chloroethane	64	1.760	1.760	(0.359)	2912	0.50000	0.66(aM)
28 Chloroform	83	4.662	4.662	(0.952)	6289	0.50000	0.64(a)
3 Chloromethane	50	1.348	1.348	(0.275)	4145	0.50000	0.96(a)
27 cis-1,2-Dichloroethene	96	4.294	4.294	(0.877)	3748	0.50000	0.62(a)
46 cis-1,3-Dichloropropene	75	6.761	6.761	(1.201)	4205	0.50000	0.49(a)
55 Dibromochloromethane	129	7.758	7.758	(0.940)	2561	0.50000	0.53(a)
44 Dibromomethane	93	6.191	6.191	(1.100)	2223	0.50000	0.60(aM)
2 Dichlorodifluoromethane	85	1.213	1.213	(0.248)	4116	0.50000	1.64(a)
61 Ethylbenzene	106	8.373	8.373	(1.015)	4972	0.50000	0.63(a)
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	2273	0.50000	1.36(a)
67 Isopropylbenzene	105	9.126	9.126	(1.106)	13902	0.50000	0.60(a)
62 m,p-Xylenes	106	8.474	8.474	(1.027)	12504	1.00000	1.29(a)
17 Methylene Chloride	84	2.877	2.877	(0.587)	5188	0.50000	0.27(a)
87 n-Butylbenzene	91	10.558	10.558	(1.031)	11934	0.50000	1.04(a)
73 n-Propylbenzene	91	9.475	9.475	(0.926)	16592	0.50000	0.60(a)
92 Naphthalene	128	12.133	12.133	(1.185)	10205	0.50000	0.49(a)
63 o-Xylene	106	8.811	8.811	(1.068)	5464	0.50000	0.56(aH)
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	13984	0.50000	0.61(a)
64 Styrene	104	8.826	8.826	(1.070)	9048	0.50000	0.55(a)
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	9624	0.50000	0.60(a)
56 Tetrachloroethene	164	7.526	7.526	(0.912)	2536	0.50000	0.64(a)
50 Toluene	91	7.049	7.049	(0.855)	15656	0.50000	0.65(a)
20 trans-1,2-Dichloroethene	96	3.143	3.143	(0.642)	3270	0.50000	0.62(a)
51 trans-1,3-Dichloropropene	75	7.263	7.263	(1.290)	3218	0.50000	1.84(a)
38 Trichloroethene	130	5.865	5.865	(1.042)	3600	0.50000	0.64(a)
8 Trichlorofluoromethane	101	1.963	1.963	(0.401)	5229	0.50000	0.62(a)
5 Vinyl Chloride	62	1.426	1.426	(0.291)	4801	0.50000	1.20(a)



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111303.D Page 3
Report Date: 24-Jan-2019 18:55

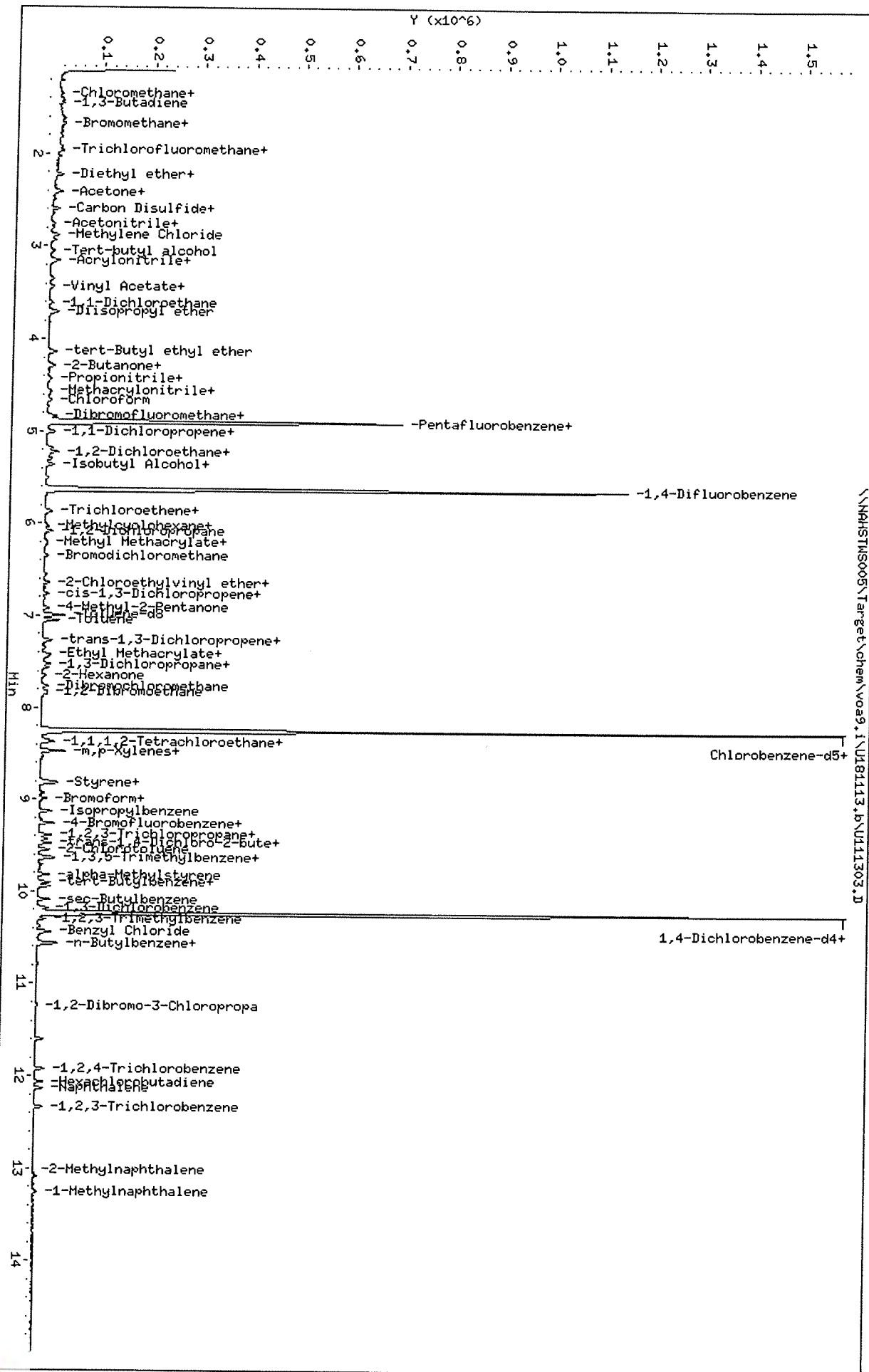
QC Flag Legend

- T - Target compound detected outside RT window.
- a - Target compound detected but, quantitated amount
Below Limit Of Quantitation(BLOQ).
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



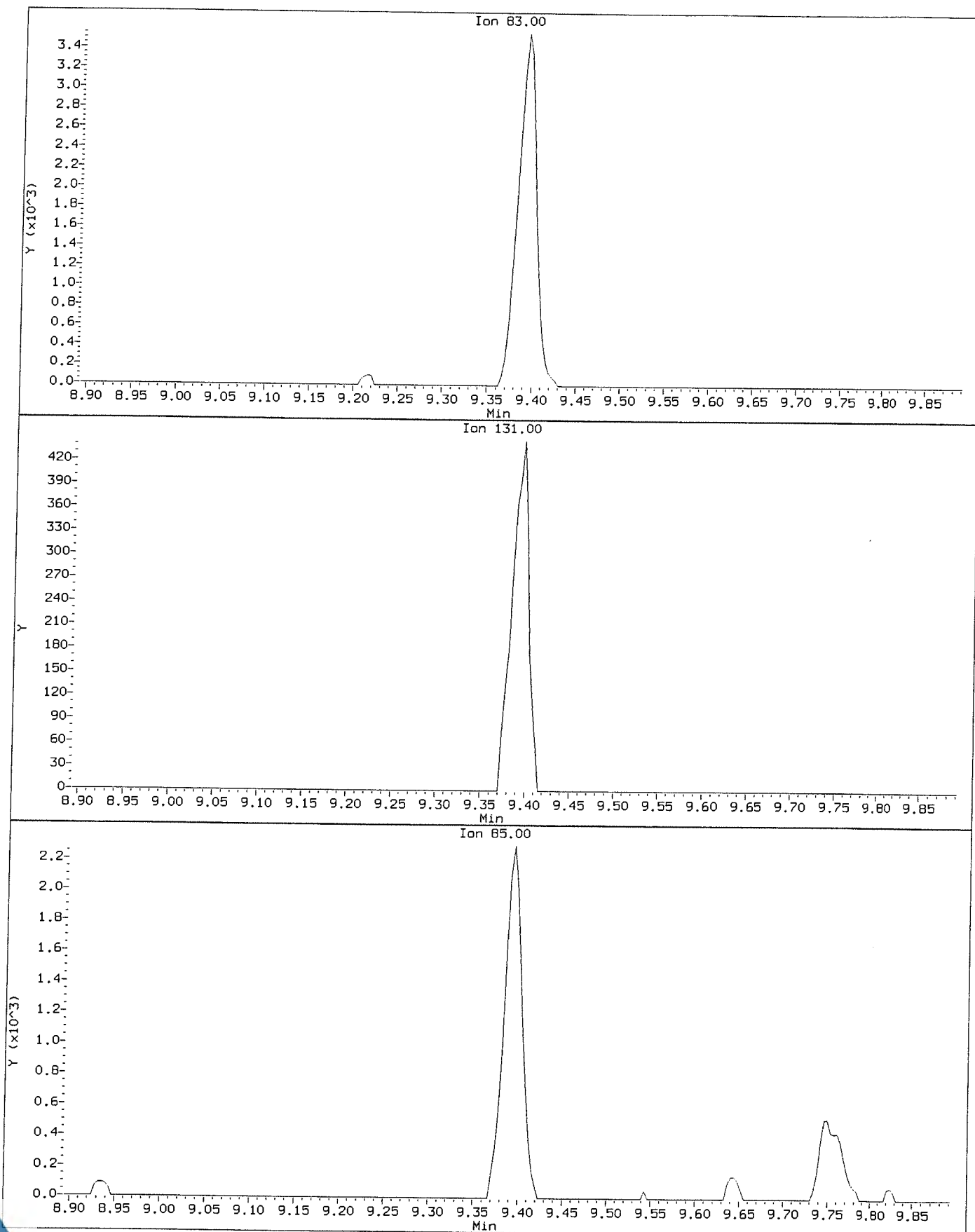
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 Sample Info: VSTD000.5;VSTD000.5;1;2;
 Purge Volume: 5.0
 Column phase: DB624

Instrument: VOA9.i
 Operator: PC
 Column diameter: 0.18



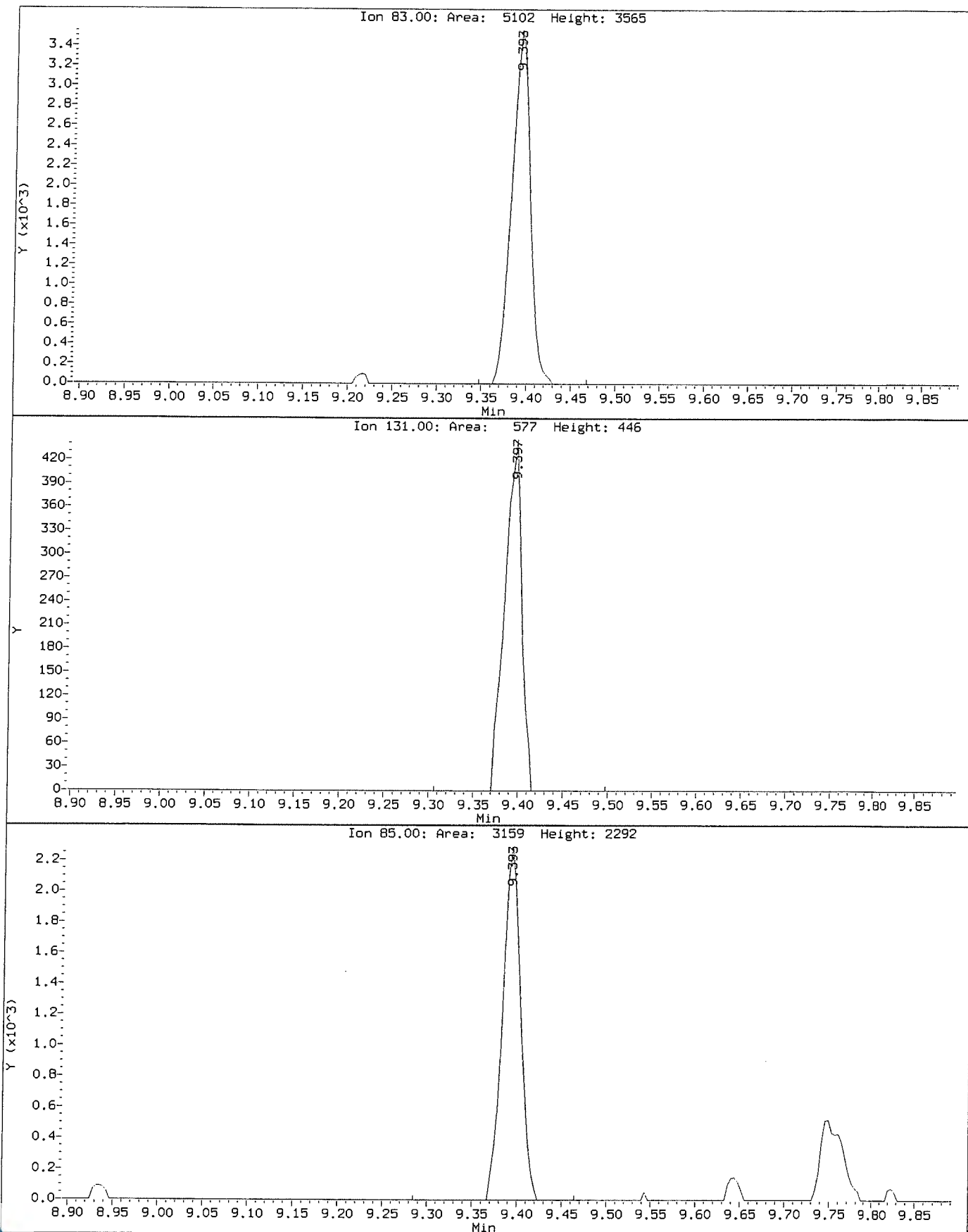
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Instrument: VOA9.1
Client Sample ID: VSTD000.5

Compound: 1,1,2,2-Tetrachloroethane
CAS Number: 79-34-5



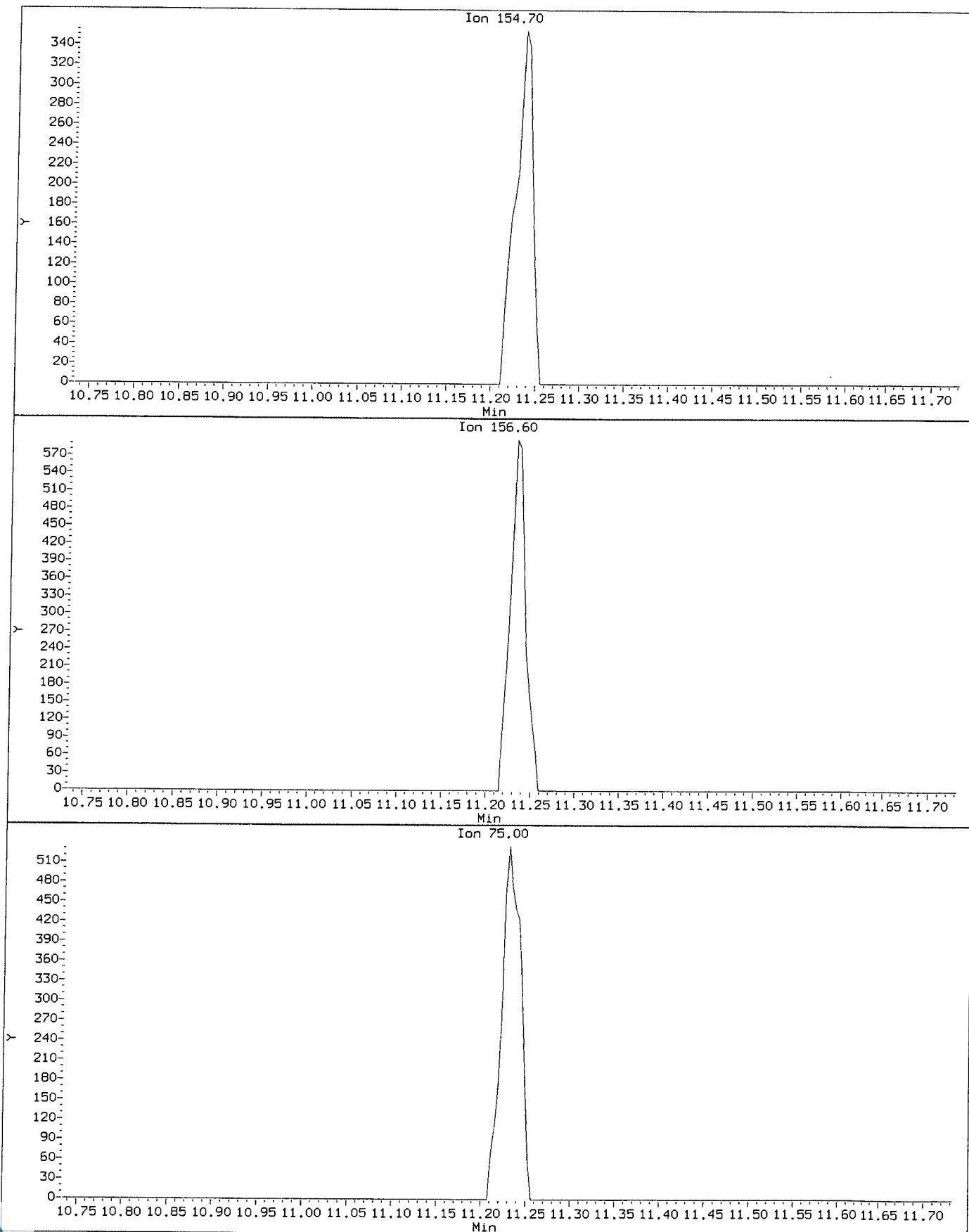
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Compound: 1,1,2,2-Tetrachloroethane
CAS Number: 79-34-5



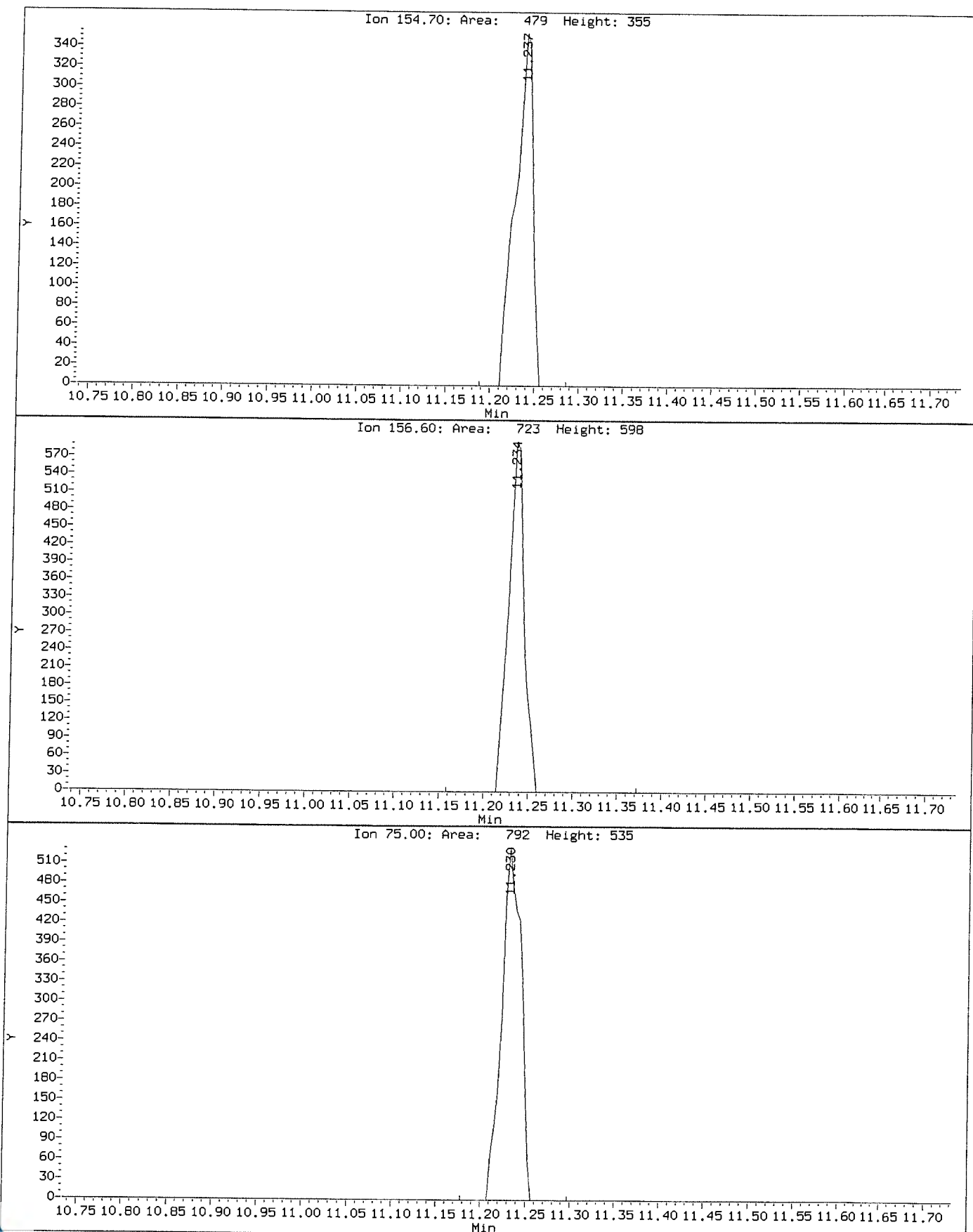
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Client Sample ID: VSTD000.5

Compound: 1,2-Dibromo-3-Chloropropane
CAS Number: 96-12-8



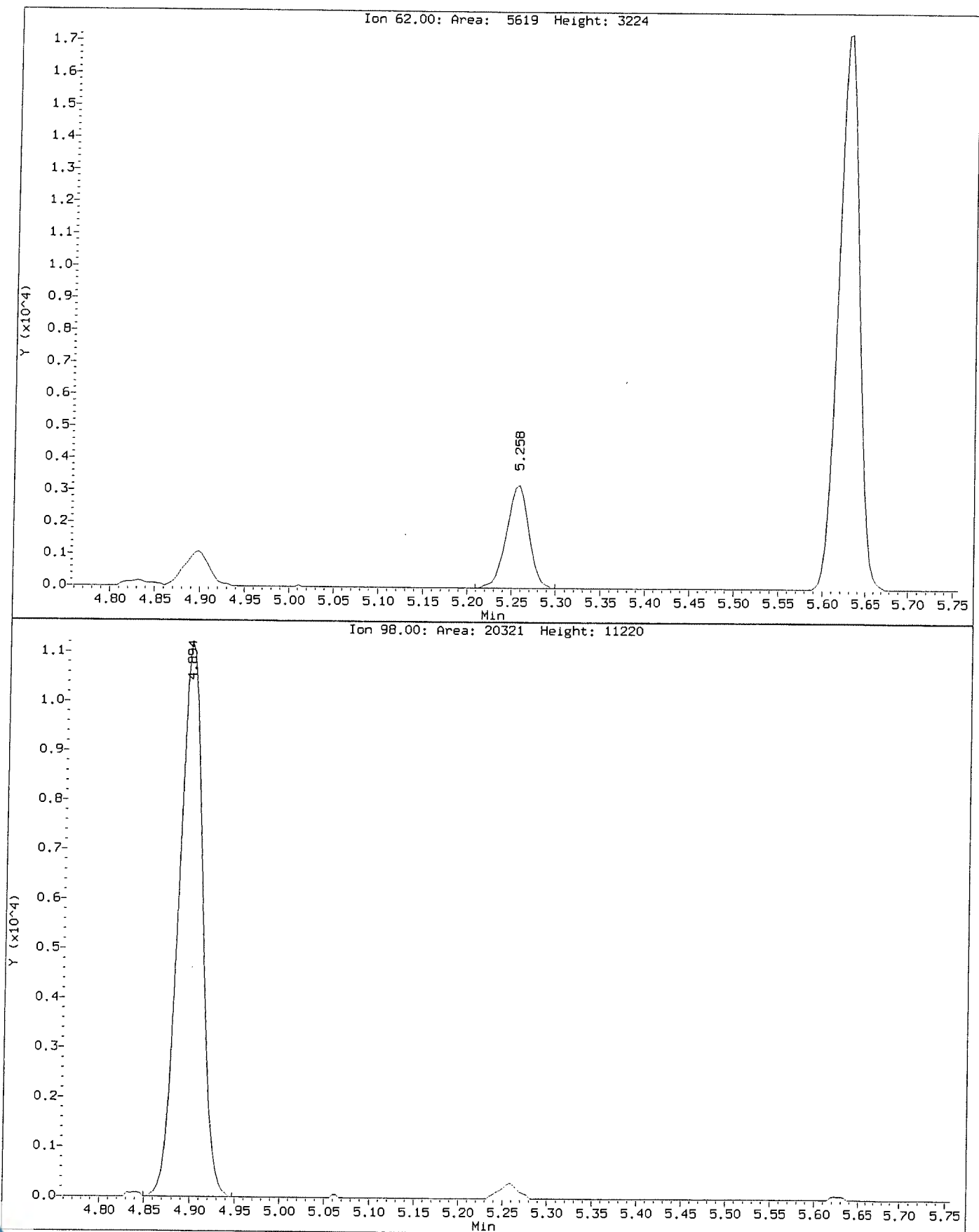
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Compound: 1,2-Dibromo-3-Chloropropane
CAS Number: 96-12-8



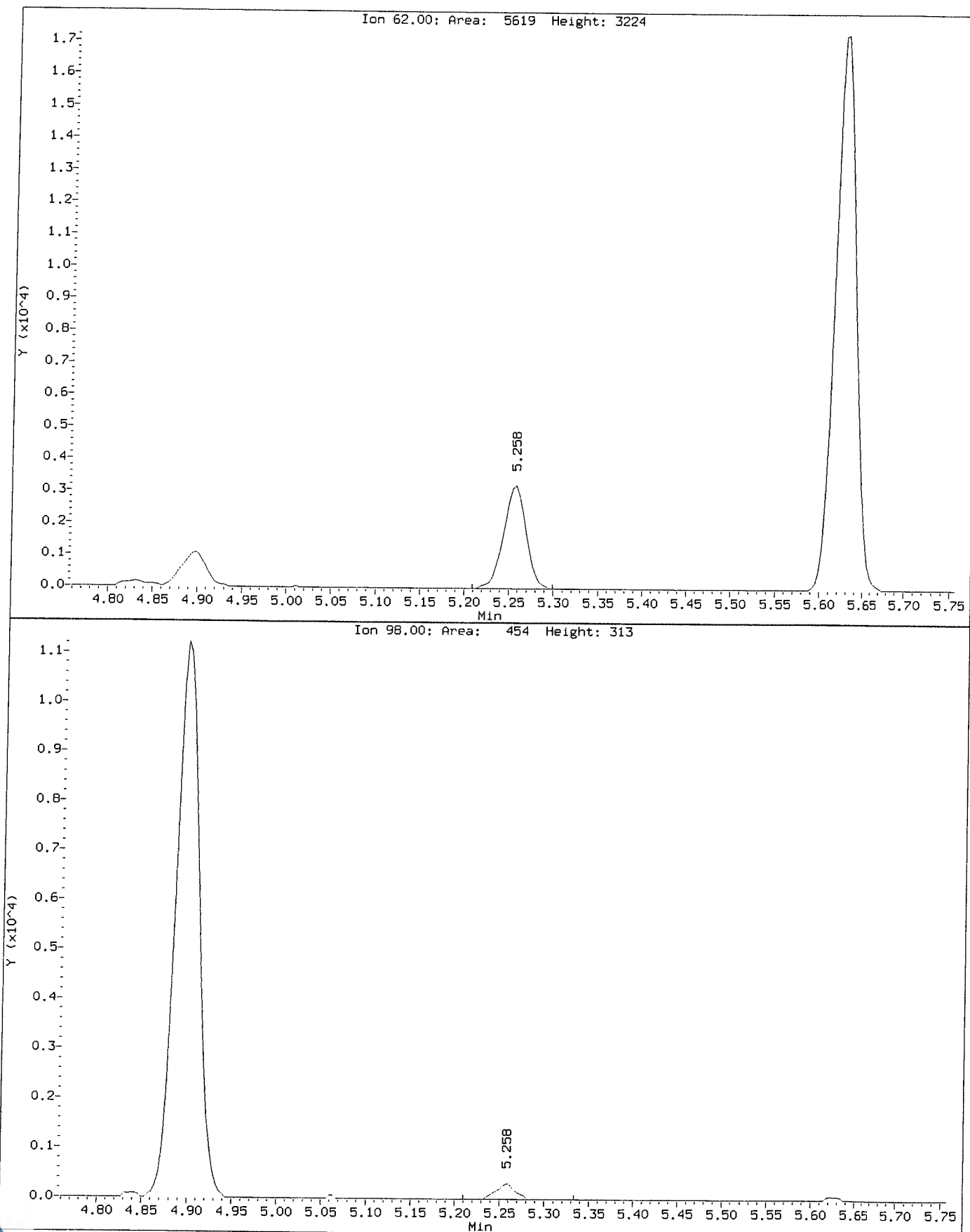
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Client Sample ID: VSTD000.5

Compound: 1,2-Dichloroethane
CAS Number: 107-06-2



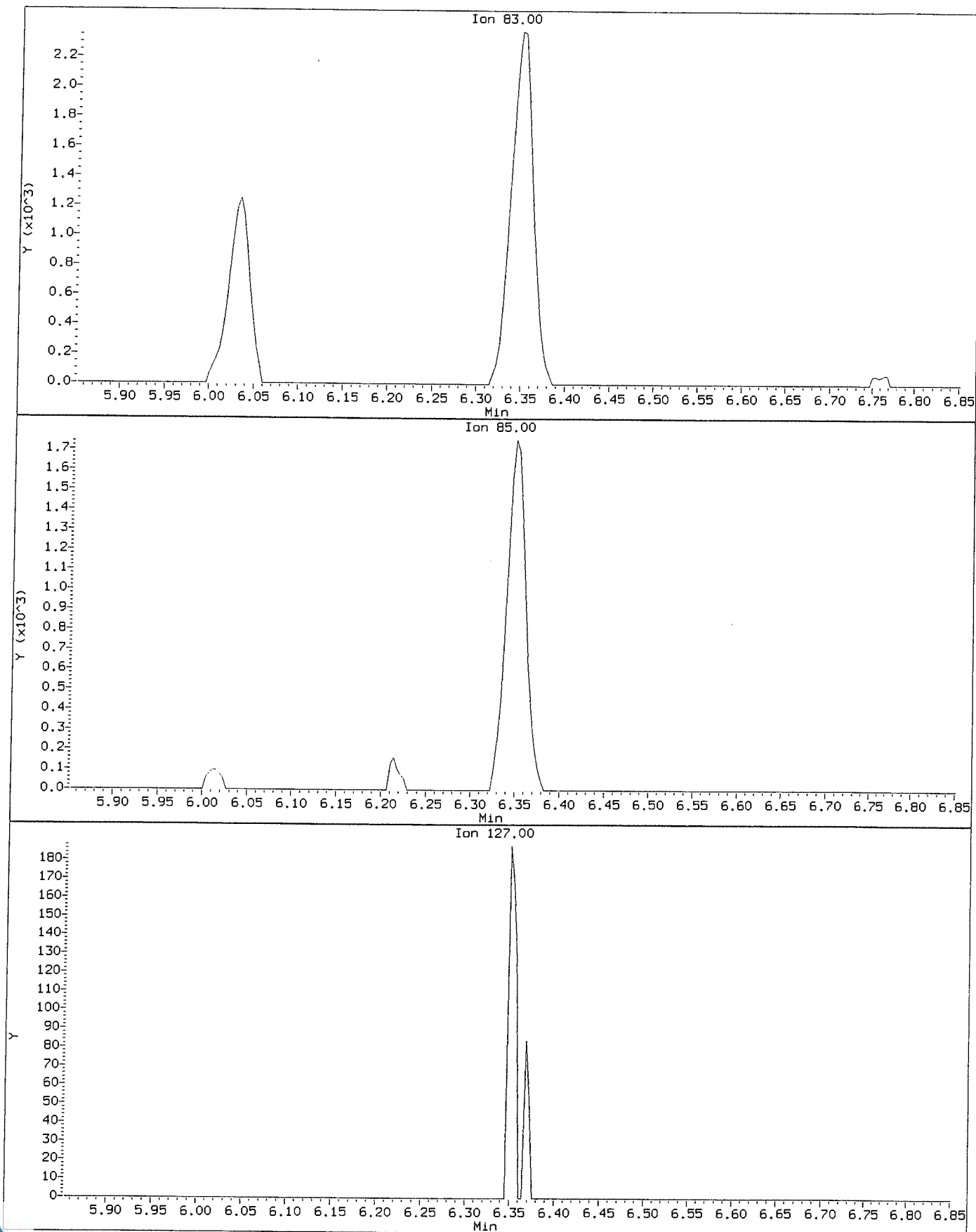
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Client Sample ID: VSTD000.5

Compound: 1,2-Dichloroethane
CAS Number: 107-06-2



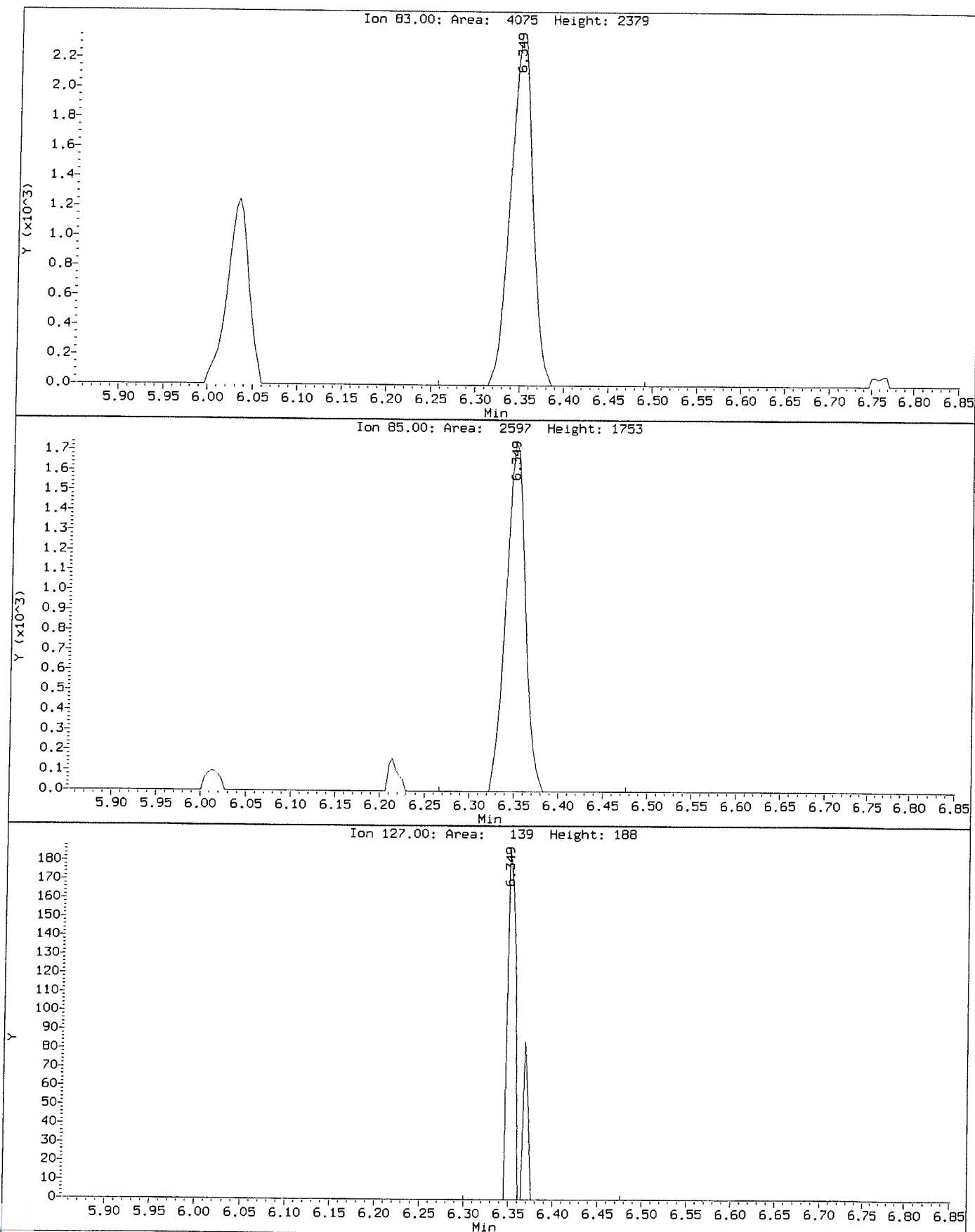
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Instrument: VOA9.i
Client Sample ID: VSTD000.5

Compound: Bromodichloromethane
CAS Number: 75-27-4



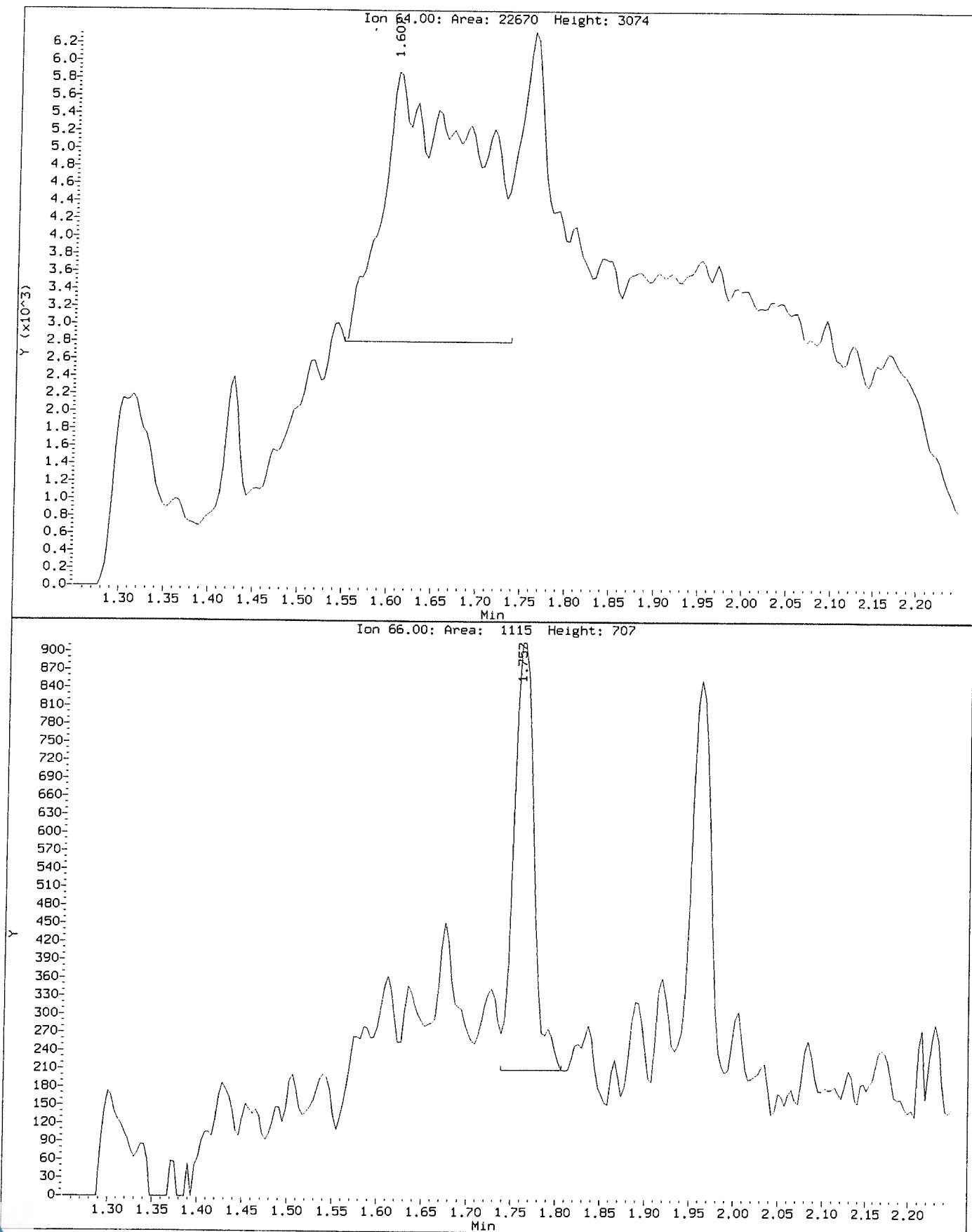
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Client Sample ID: VSTD000.5

Compound: Bromodichloromethane
CAS Number: 75-27-4



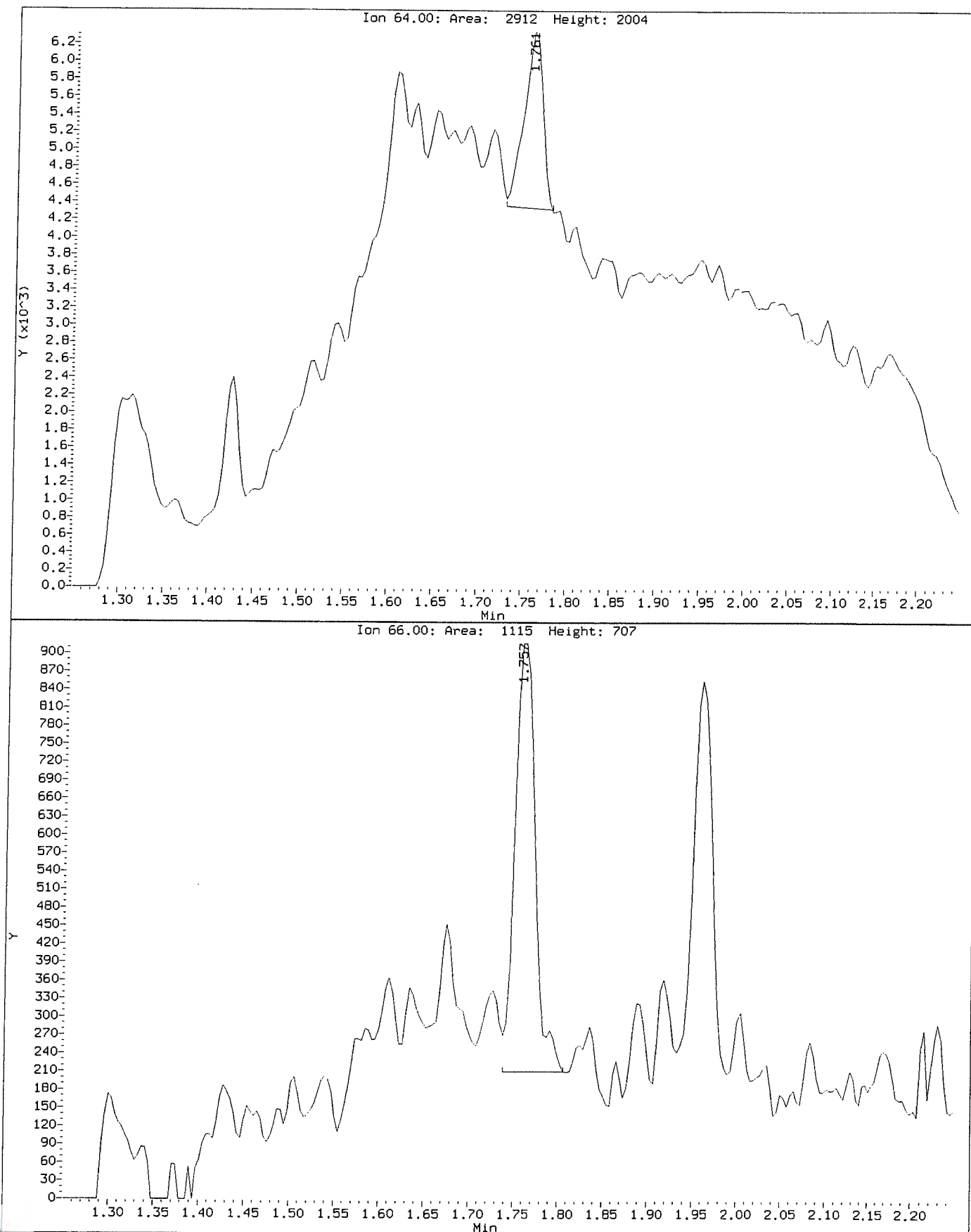
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Instrument: VOA9.1
Client Sample ID: VSTD000.5

Compound: Chloroethane
CAS Number: 75-00-3



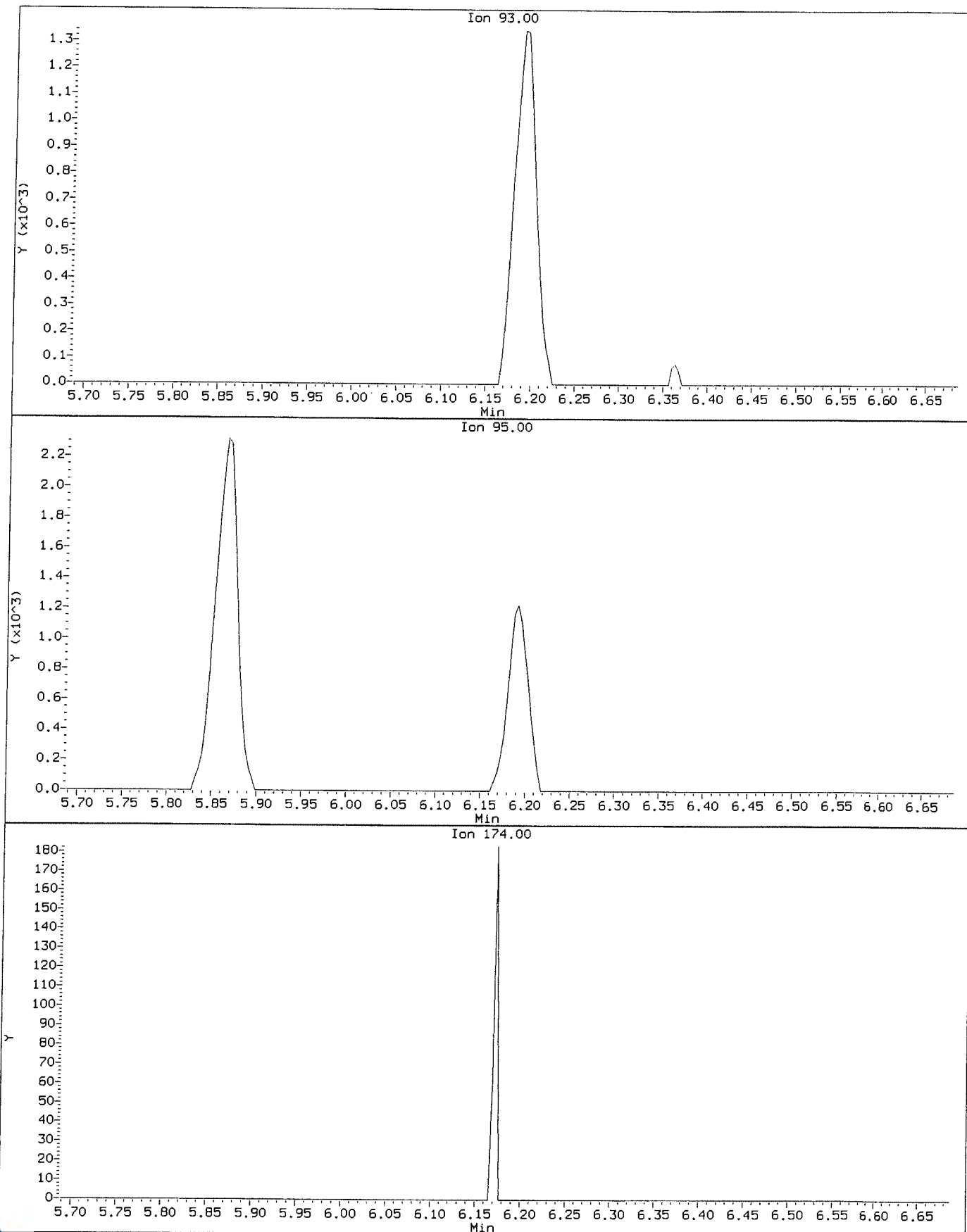
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Instrument: VOA9.i
Client Sample ID: VSTD000.5

Compound: Chloroethane
CAS Number: 75-00-3



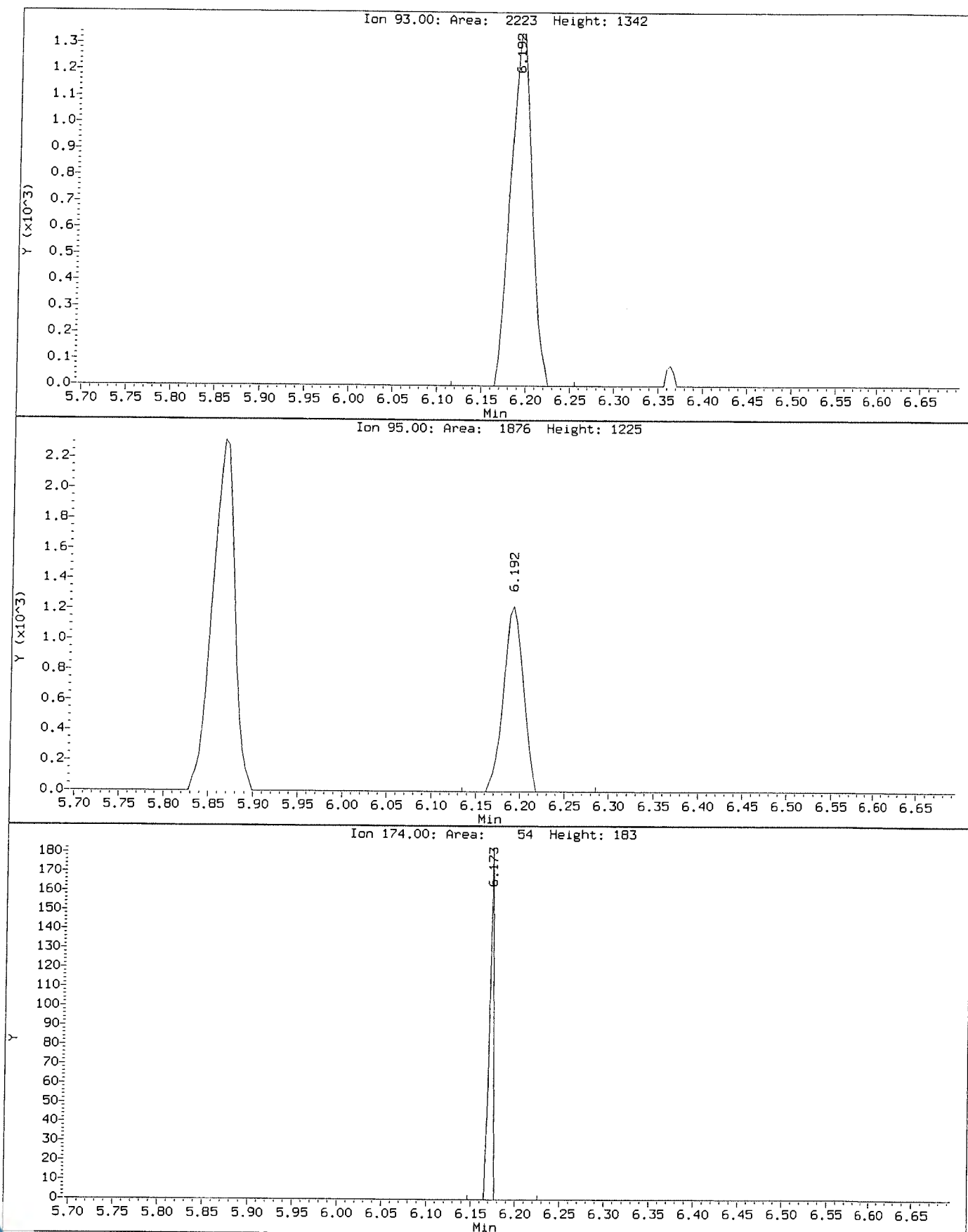
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Instrument: VOA9.1
Client Sample ID: VSTD000.5

Compound: Dibromomethane
CAS Number: 74-95-3



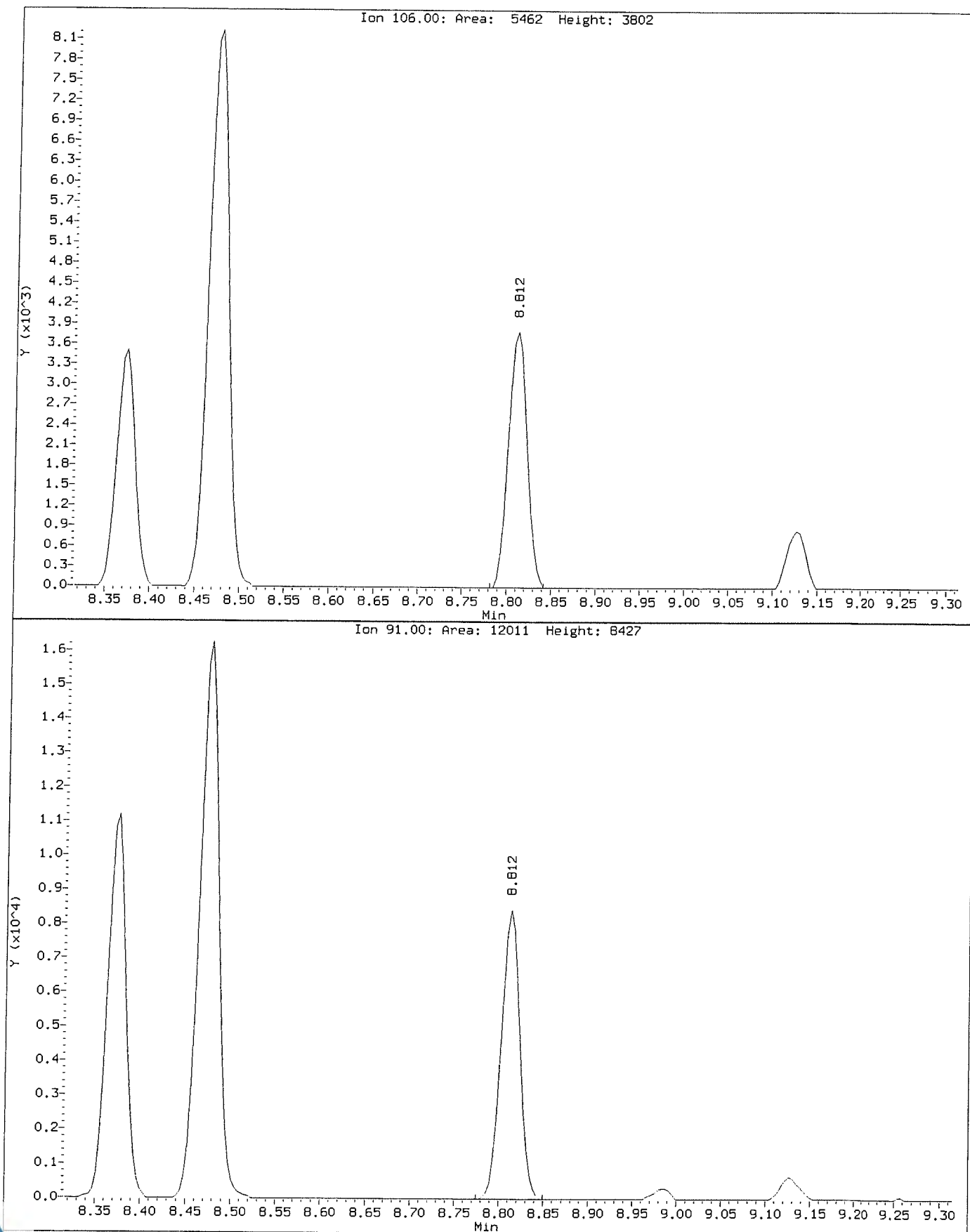
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Client Sample ID: VSTD000.5

Compound: Dibromomethane
CAS Number: 74-95-3



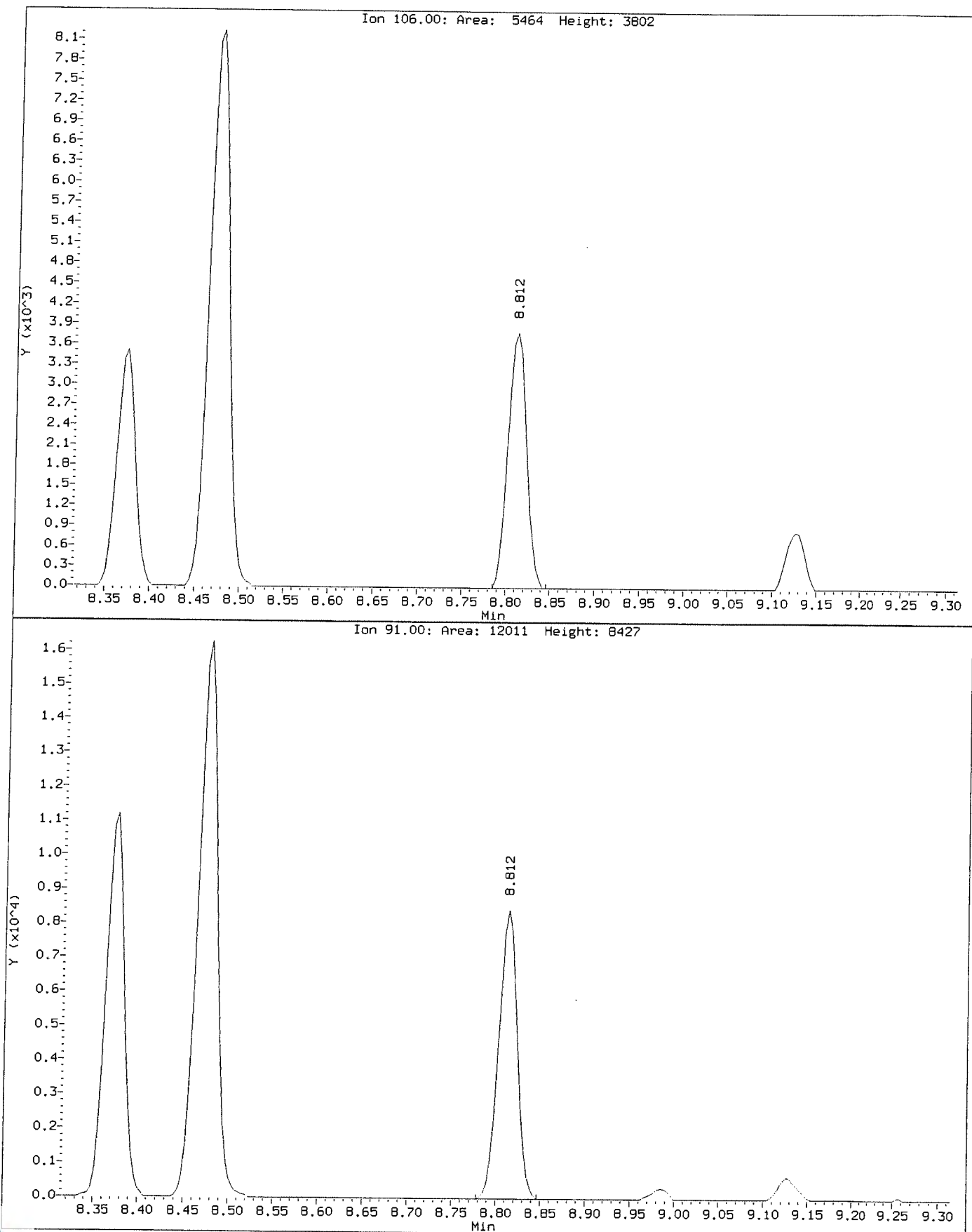
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Instrument: VOA9.i
Client Sample ID: VSTD000.5

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.1\U181113.b\U111303.D
Injection Date: 13-NOV-2018 11:47
Instrument: VDA9.i
Client Sample ID: VSTD000.5

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111304.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111304.D
 Lab Smp Id: VSTD001 Client Smp ID: VSTD001
 Inj Date : 13-NOV-2018 12:36
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD001;VSTD001;1;3;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 11:47 Cal File: U111303.D
 Als bottle: 5 Calibration Sample, Level: 3
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG						AMOUNTS	
		MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.898	(1.000)	405565	50.0000		
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	804218	50.0000		
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	748347	50.0000		
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	345100	50.0000		
\$ 30 Dibromofluoromethane	113	4.834	4.834	(0.987)	7705	1.00000	0.43(a)	
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.179	(1.057)	11059	1.00000	0.09(a)	
\$ 48 Toluene-d8	98	6.989	6.989	(0.847)	30489	1.00000	(a)	
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	10512	1.00000	0.38(a)	
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	4277	1.00000	0.90(a)	
31 1,1,1-Trichloroethane	97	4.826	4.826	(0.985)	6795	1.00000	0.95(a)	
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	8088	1.00000	0.98(a)	
138 Freon TF	101	2.405	2.405	(0.491)	4282	1.00000	1.05(a)	
53 1,1,2-Trichloroethane	83	7.421	7.421	(0.900)	5034	1.00000	1.05(a)	
22 1,1-Dichloroethane	63	3.608	3.608	(0.737)	9857	1.00000	1.06(a)	
11 1,1-Dichloroethene	96	2.405	2.405	(0.491)	5079	1.00000	1.10(a)	
32 1,1-Dichloropropene	75	5.006	5.006	(0.890)	7778	1.00000	1.04(a)	
93 1,2,3-Trichlorobenzene	180	12.339	12.339	(1.205)	5356	1.00000	0.82(a)	
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	7560	1.00000	0.88(a)	
90 1,2,4-Trichlorobenzene	180	11.926	11.926	(1.165)	6307	1.00000	0.92(a)	
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	18207	1.00000	0.94(a)	
89 1,2-Dibromo-3-Chloropropane	155	11.237	11.237	(1.098)	955	1.00000	1.78(a)	
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	5452	1.00000	0.98(a)	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111304.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.569	10.569	(1.033)	11469	1.00000	1.04 (a)
33 1,2-Dichloroethane	62	5.254	5.254	(0.934)	8729	1.00000	1.03 (a)
42 1,2-Dichloropropane	63	6.082	6.082	(1.081)	5924	1.00000	1.03 (a)
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	17829	1.00000	0.96 (a)
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	11288	1.00000	1.03 (a)
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	10633	1.00000	1.04 (a)
84 1,4-Dichlorobenzene	146	10.255	10.255	(1.002)	11659	1.00000	1.05 (a)
26 2,2-Dichloropropane	77	4.275	4.275	(0.873)	5473	1.00000	0.95 (a)
24 2-Butanone	43	4.343	4.343	(0.887)	5794	2.00000	2.03 (a)
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	17337	1.00000	1.02 (a)
52 2-Hexanone	43	7.653	7.653	(0.928)	7477	2.00000	1.71 (a)
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	18812	1.00000	0.96 (a)
82 p-Isopropyltoluene	119	10.210	10.210	(0.997)	16467	1.00000	0.87 (a)
45 4-Methyl-2-Pentanone	43	6.914	6.914	(0.838)	12236	2.00000	1.96 (a)
10 Acetone	43	2.484	2.484	(0.507)	7657	2.00000	(a)
37 Benzene	78	5.220	5.220	(0.928)	23677	1.00000	1.06 (a)
74 Bromobenzene	156	9.385	9.385	(0.917)	5602	1.00000	0.98 (a)
29 Bromochloromethane	128	4.557	4.557	(0.930)	2347	1.00000	0.98 (aM)
39 Bromodichloromethane	83	6.348	6.348	(1.129)	6268	1.00000	0.93 (aM)
66 Bromoform	173	8.984	8.984	(1.089)	2470	1.00000	1.89 (Ta)
6 Bromomethane	94	1.674	1.674	(0.342)	4167	1.00000	3.09 (a)
19 Carbon Disulfide	76	2.592	2.592	(0.529)	28789	2.00000	2.03 (a)
34 Carbon Tetrachloride	117	4.999	4.999	(0.889)	5592	1.00000	0.96 (a)
59 Chlorobenzene	112	8.275	8.275	(1.003)	16679	1.00000	1.07 (a)
7 Chloroethane	64	1.756	1.756	(0.359)	4550	1.00000	1.07 (aM)
28 Chloroform	83	4.658	4.658	(0.951)	9975	1.00000	1.06 (a)
3 Chloromethane	50	1.344	1.344	(0.274)	6506	1.00000	1.42 (a)
27 cis-1,2-Dichloroethene	96	4.290	4.290	(0.876)	6147	1.00000	1.06 (a)
46 cis-1,3-Dichloropropene	75	6.757	6.757	(1.201)	7263	1.00000	0.86 (a)
55 Dibromochloromethane	129	7.758	7.758	(0.940)	3914	1.00000	0.82 (a)
44 Dibromomethane	93	6.191	6.191	(1.101)	3612	1.00000	1.00 (aM)
2 Dichlorodifluoromethane	85	1.209	1.209	(0.247)	6358	1.00000	2.04 (a)
61 Ethylbenzene	106	8.373	8.373	(1.015)	8122	1.00000	1.03 (a)
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	2210	1.00000	1.33 (a)
67 Isopropylbenzene	105	9.126	9.126	(1.106)	22125	1.00000	0.97 (a)
62 m,p-Xylenes	106	8.474	8.474	(1.027)	18714	2.00000	1.94 (a)
17 Methylene Chloride	84	2.877	2.877	(0.587)	7766	1.00000	0.81 (a)
87 n-Butylbenzene	91	10.558	10.558	(1.031)	17894	1.00000	1.37 (a)
73 n-Propylbenzene	91	9.475	9.475	(0.926)	27227	1.00000	0.99 (a)
92 Naphthalene	128	12.133	12.133	(1.185)	17787	1.00000	0.87 (a)
63 o-Xylene	106	8.811	8.811	(1.068)	9461	1.00000	0.98 (aH)
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	22039	1.00000	0.97 (a)
64 Styrene	104	8.826	8.826	(1.070)	14819	1.00000	0.91 (a)
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	14832	1.00000	0.93 (a)
56 Tetrachloroethene	164	7.525	7.525	(0.912)	4264	1.00000	1.08 (a)
50 Toluene	91	7.049	7.049	(0.855)	24363	1.00000	1.02 (a)
20 trans-1,2-Dichloroethene	96	3.147	3.147	(0.643)	5480	1.00000	1.09 (a)
51 trans-1,3-Dichloropropene	75	7.263	7.263	(1.291)	5482	1.00000	2.11 (a)
38 Trichloroethene	130	5.861	5.861	(1.042)	5801	1.00000	1.05 (a)
8 Trichlorofluoromethane	101	1.959	1.959	(0.400)	8913	1.00000	1.09 (a)
5 Vinyl Chloride	62	1.423	1.423	(0.291)	7374	1.00000	1.61 (a)



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111304.D Page 3
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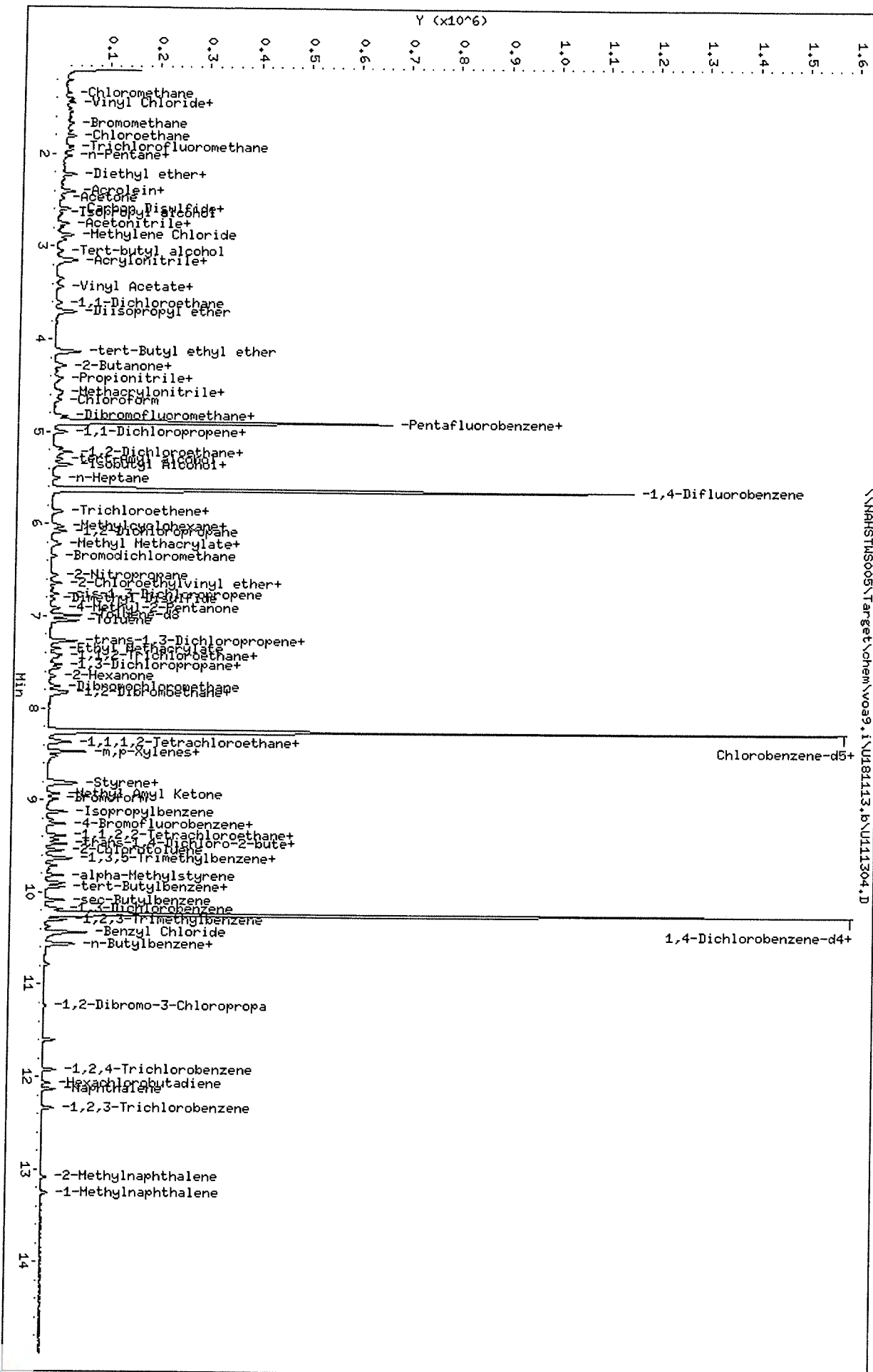
QC Flag Legend

- T - Target compound detected outside RT window.
- a - Target compound detected but, quantitated amount
Below Limit Of Quantitation(BLOQ).
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



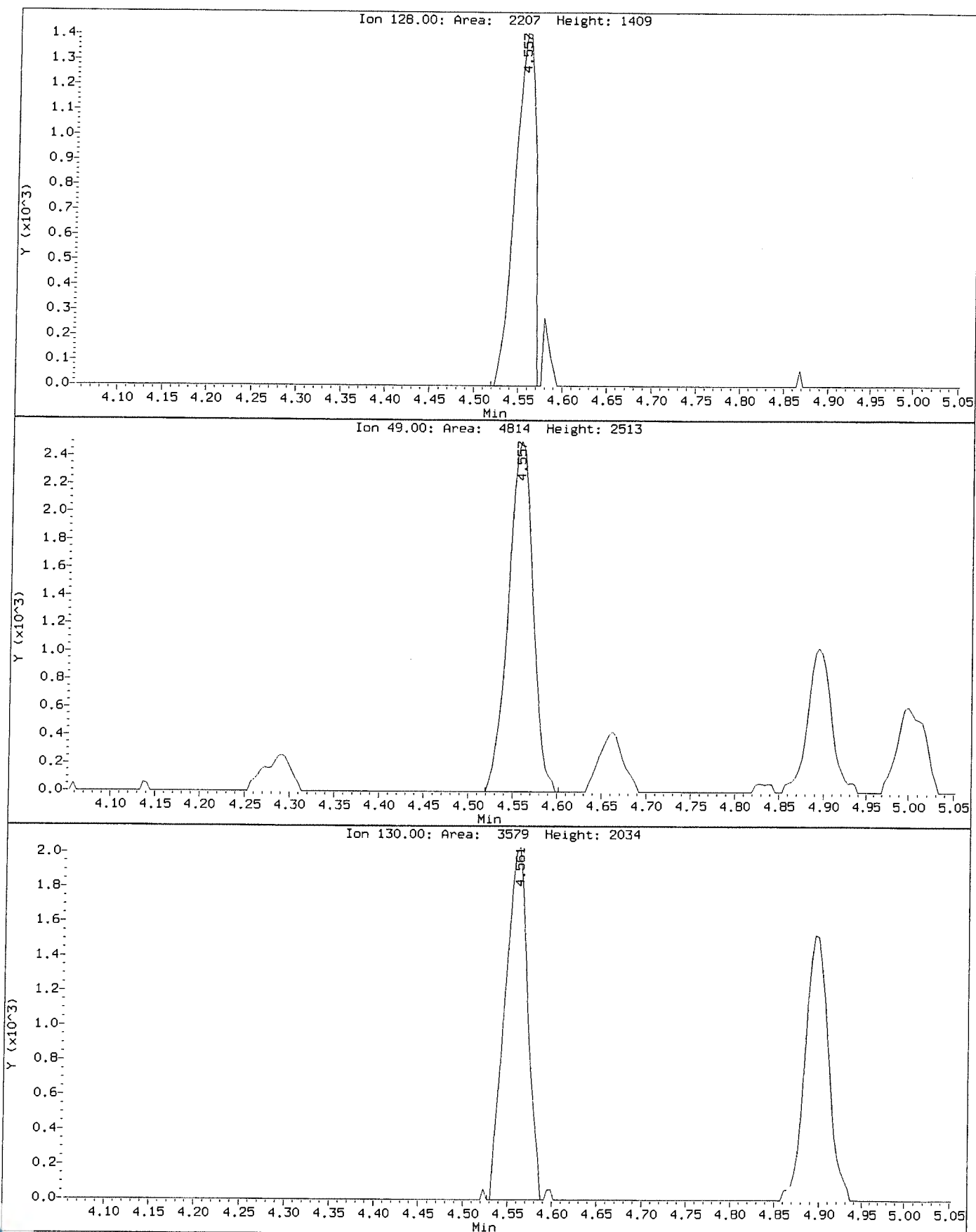
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 Column phase: DB624

Instrument: W0A9.i
 Operator: PC
 Column diameter: 0.18



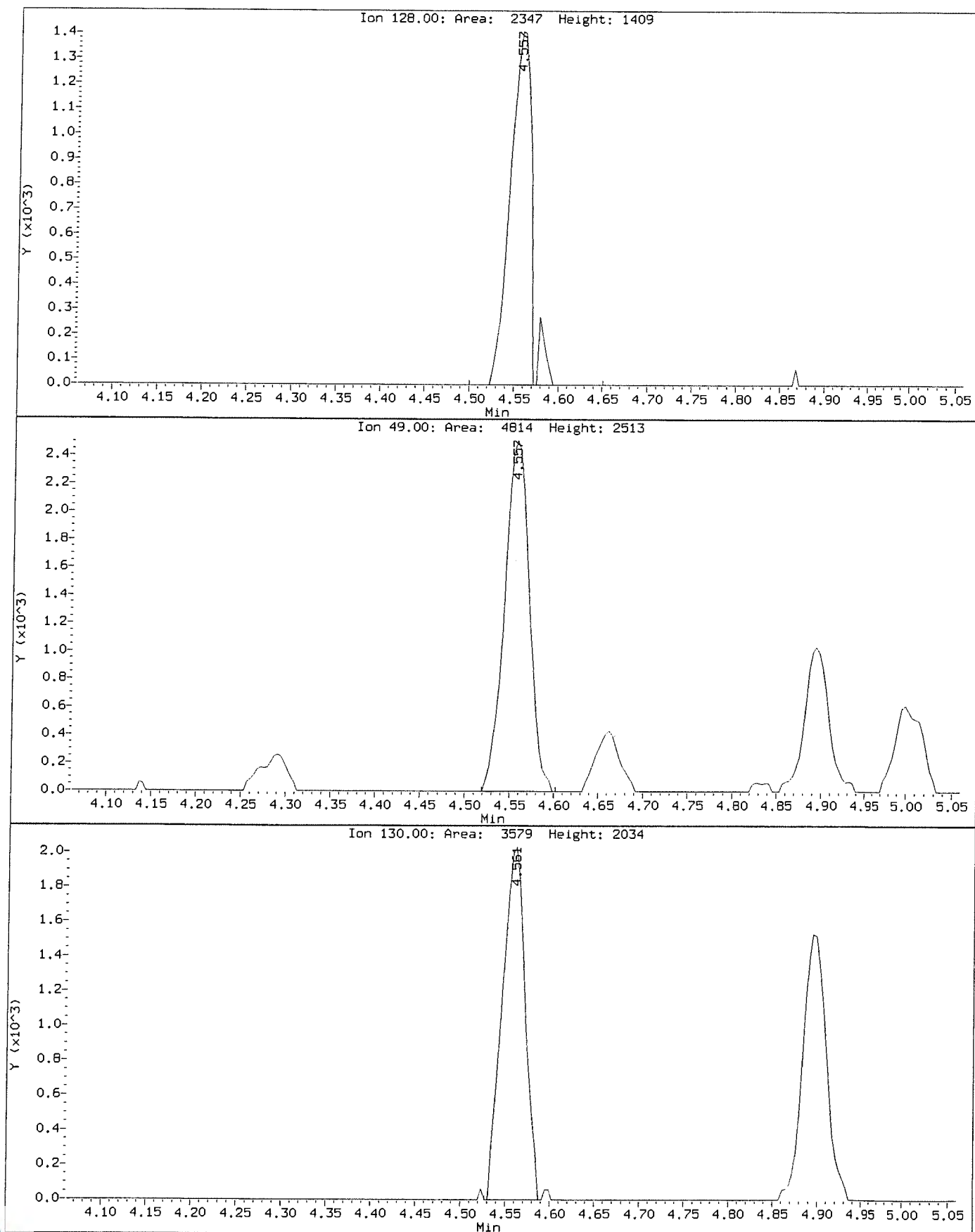
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Client Sample ID: VSTD001

Compound: Bromochloromethane
CAS Number: 74-97-5



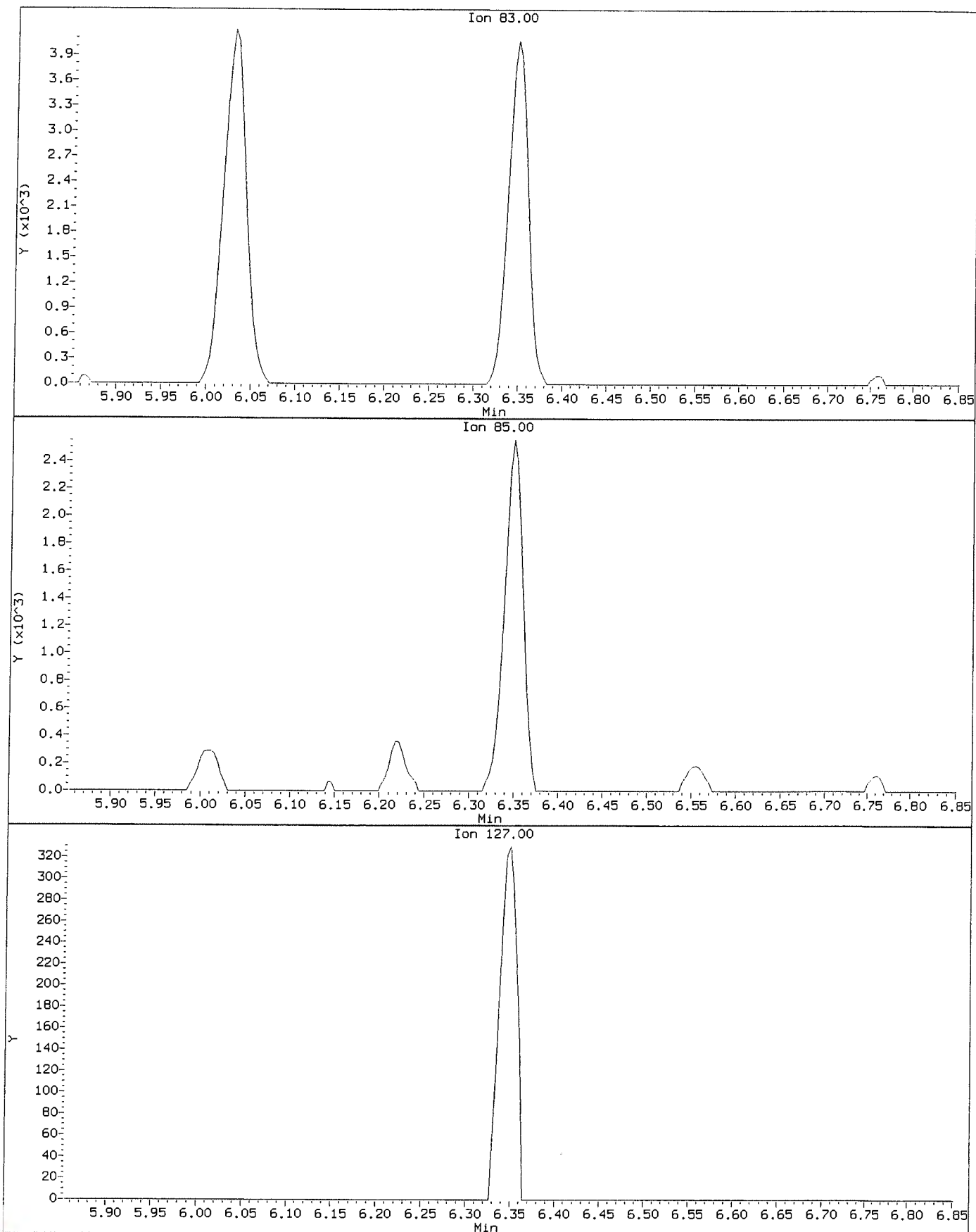
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Client Sample ID: VSTD001

Compound: Bromochloromethane
CAS Number: 74-97-5



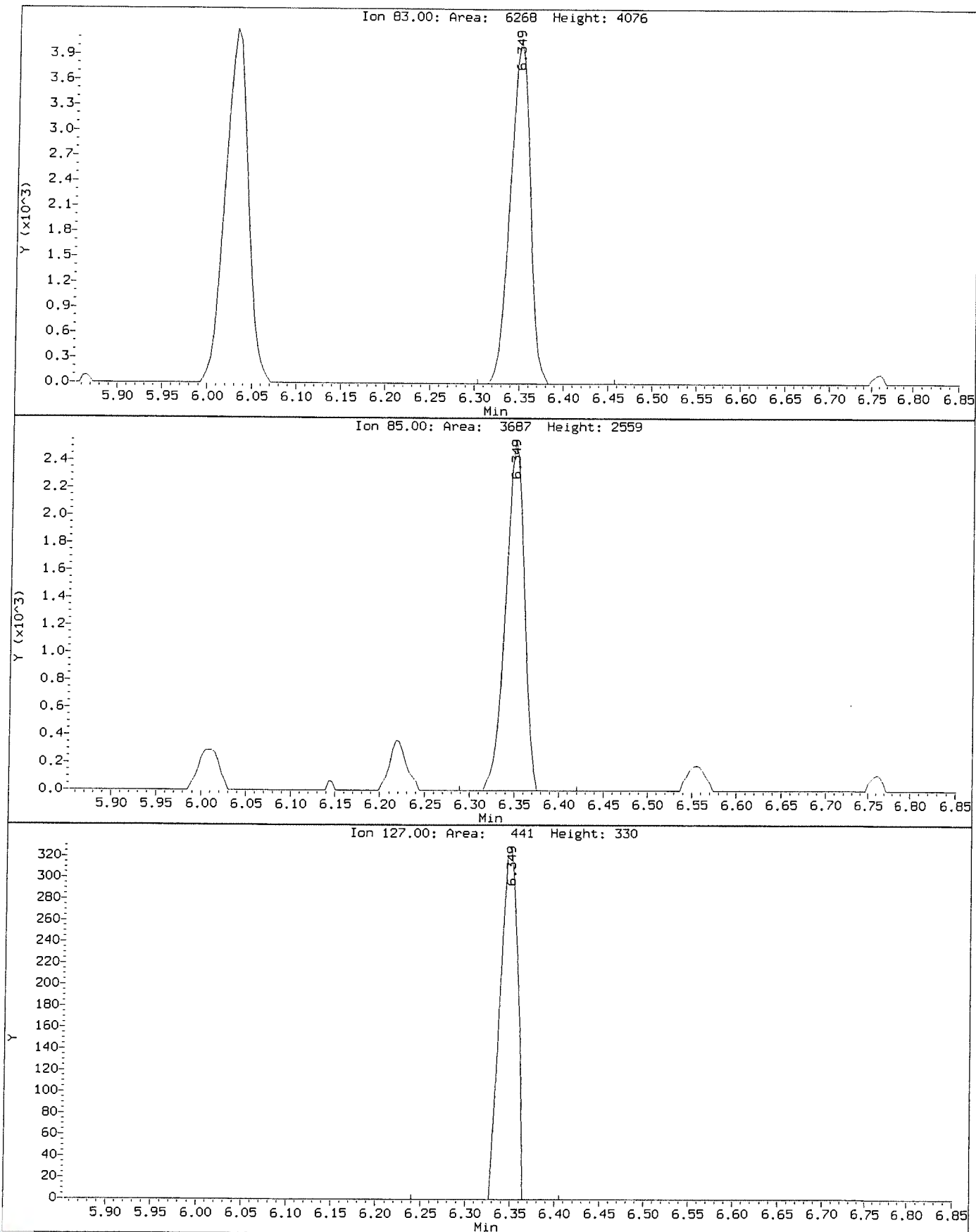
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Instrument: VOA9.1
Client Sample ID: VSTD001

Compound: Bromodichloromethane
CAS Number: 75-27-4



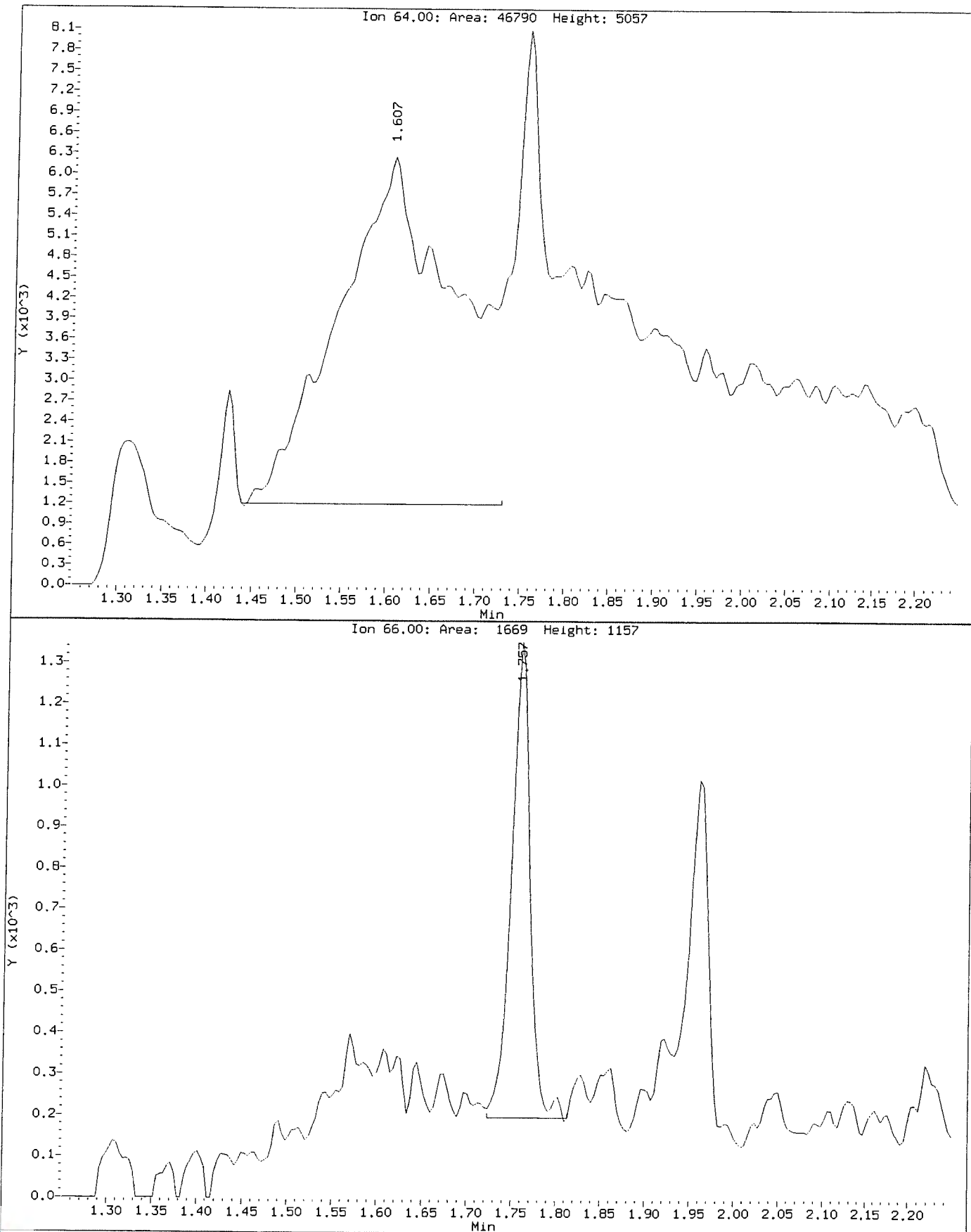
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Instrument: VOA9.1
Client Sample ID: VSTD001

Compound: Bromodichloromethane
CAS Number: 75-27-4



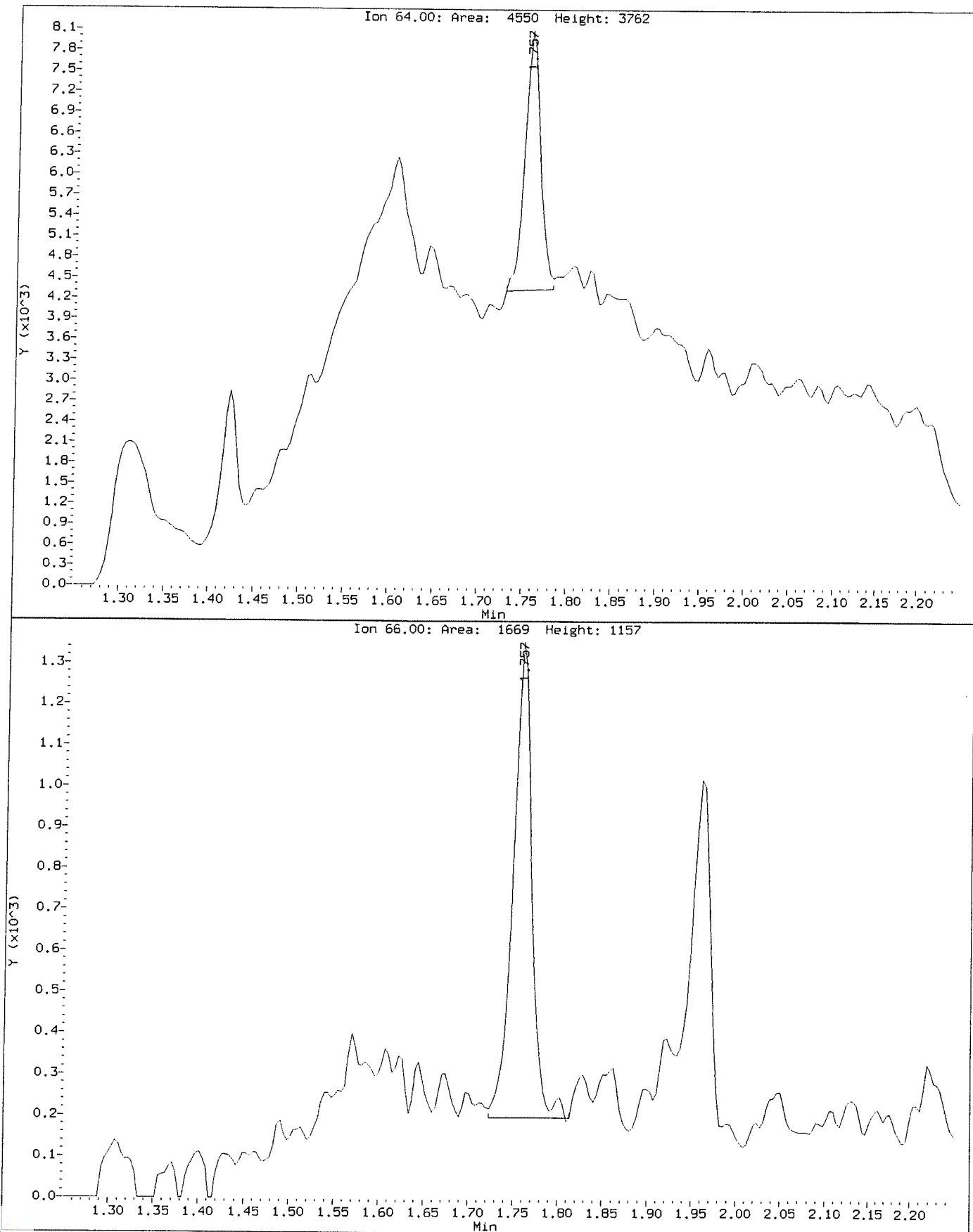
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Instrument: VOA9.i
Client Sample ID: VSTD001

Compound: Chloroethane
CAS Number: 75-00-3



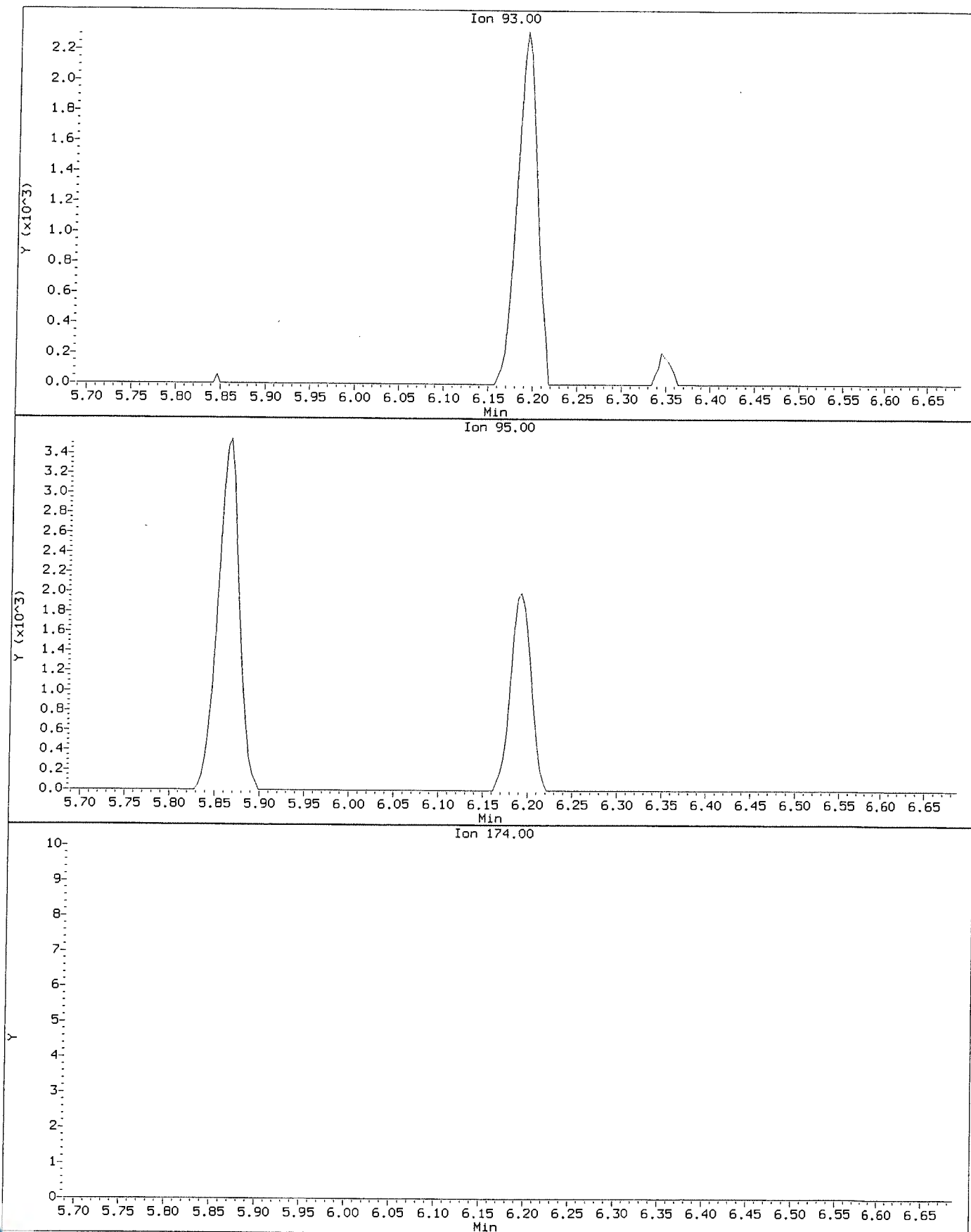
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Injection Date: 13-NOV-2018 12:36
Instrument: VOA9.1
Client Sample ID: VSTD001

Compound: Chloroethane
CAS Number: 75-00-3



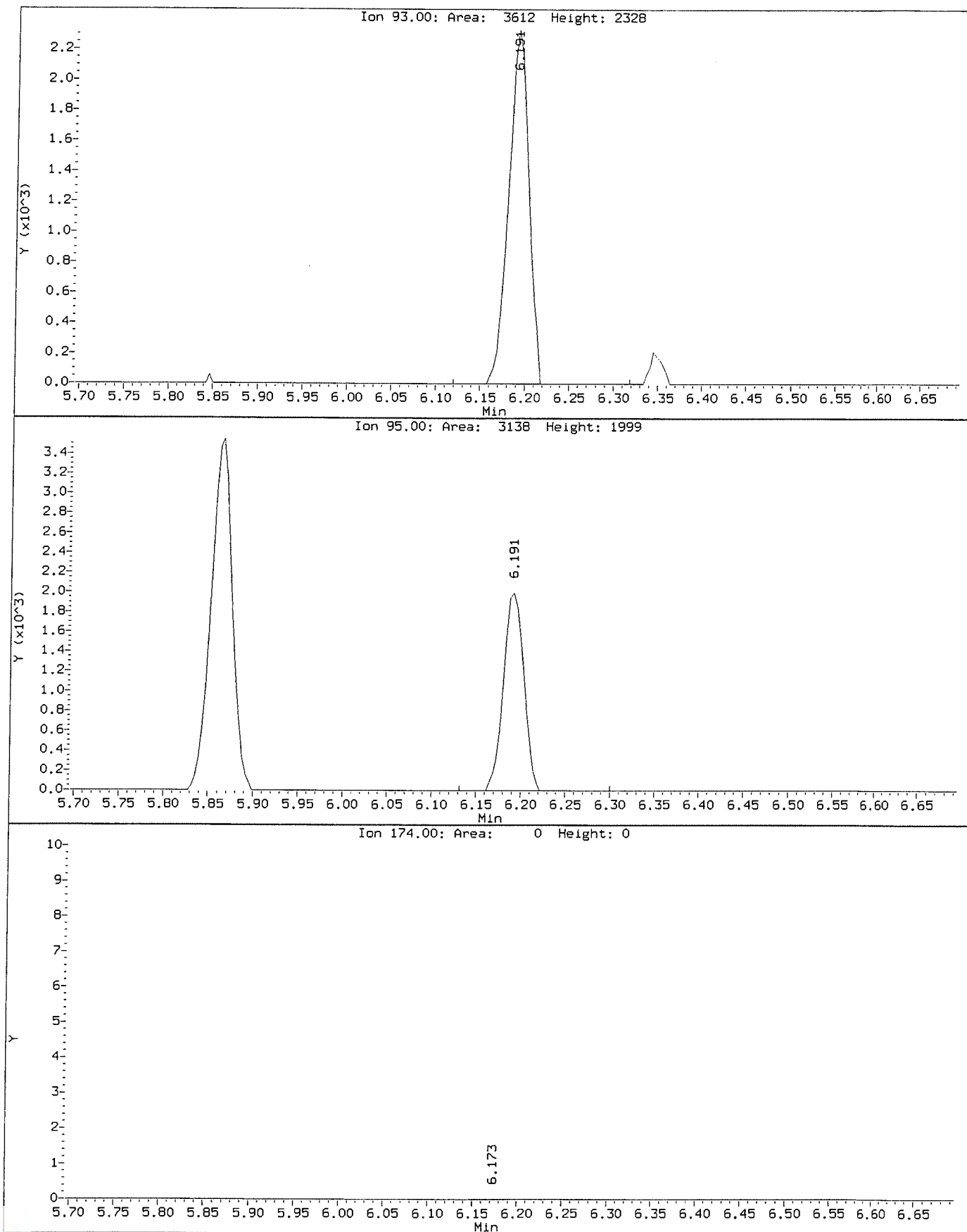
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Injection Date: 13-NOV-2018 12:36
Instrument: VOA9.i
Client Sample ID: VSTD001

Compound: Dibromomethane
CAS Number: 74-95-3



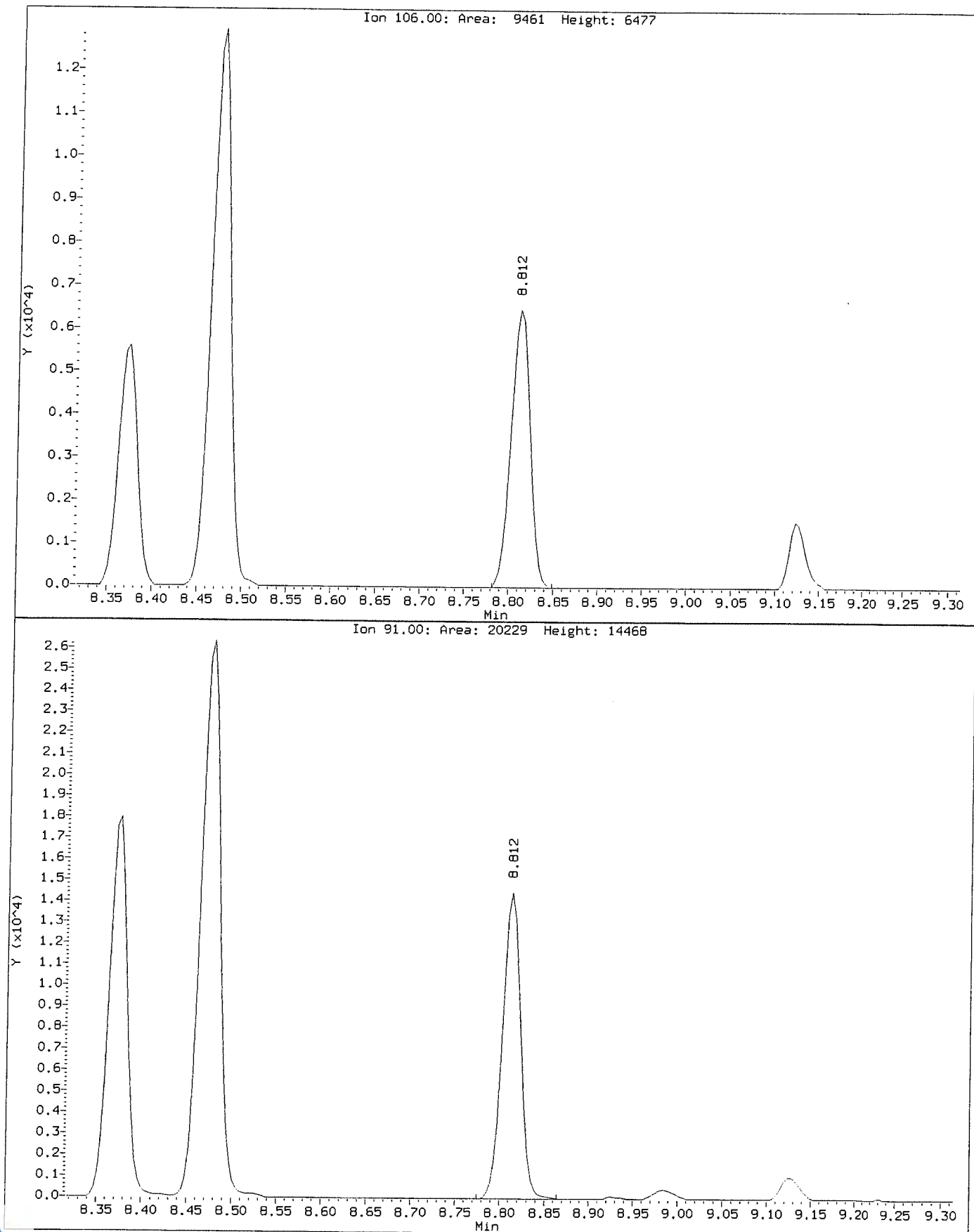
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Instrument: VOA9.1
Client Sample ID: VSTD001

Compound: Dibromomethane
CAS Number: 74-95-3



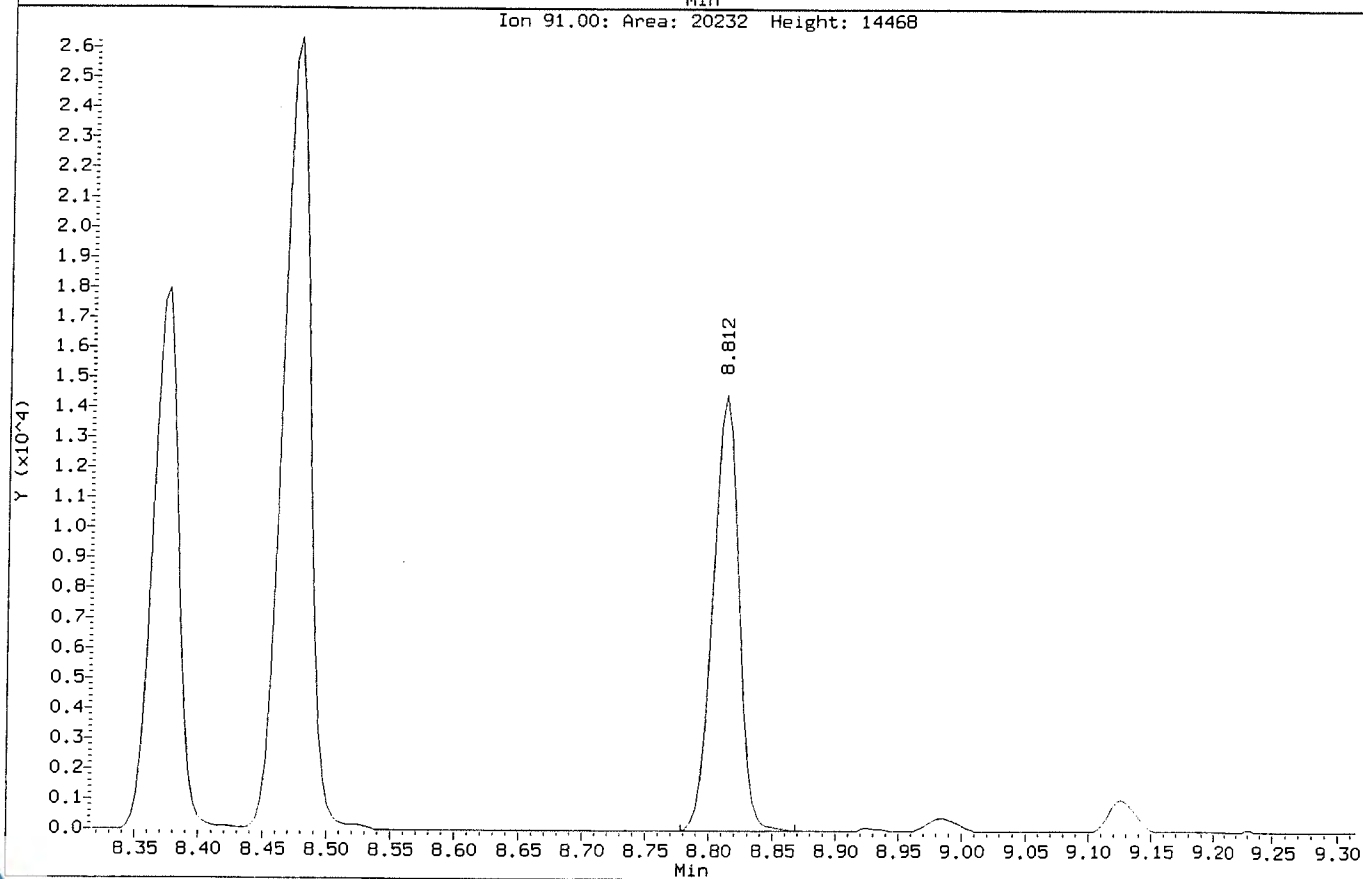
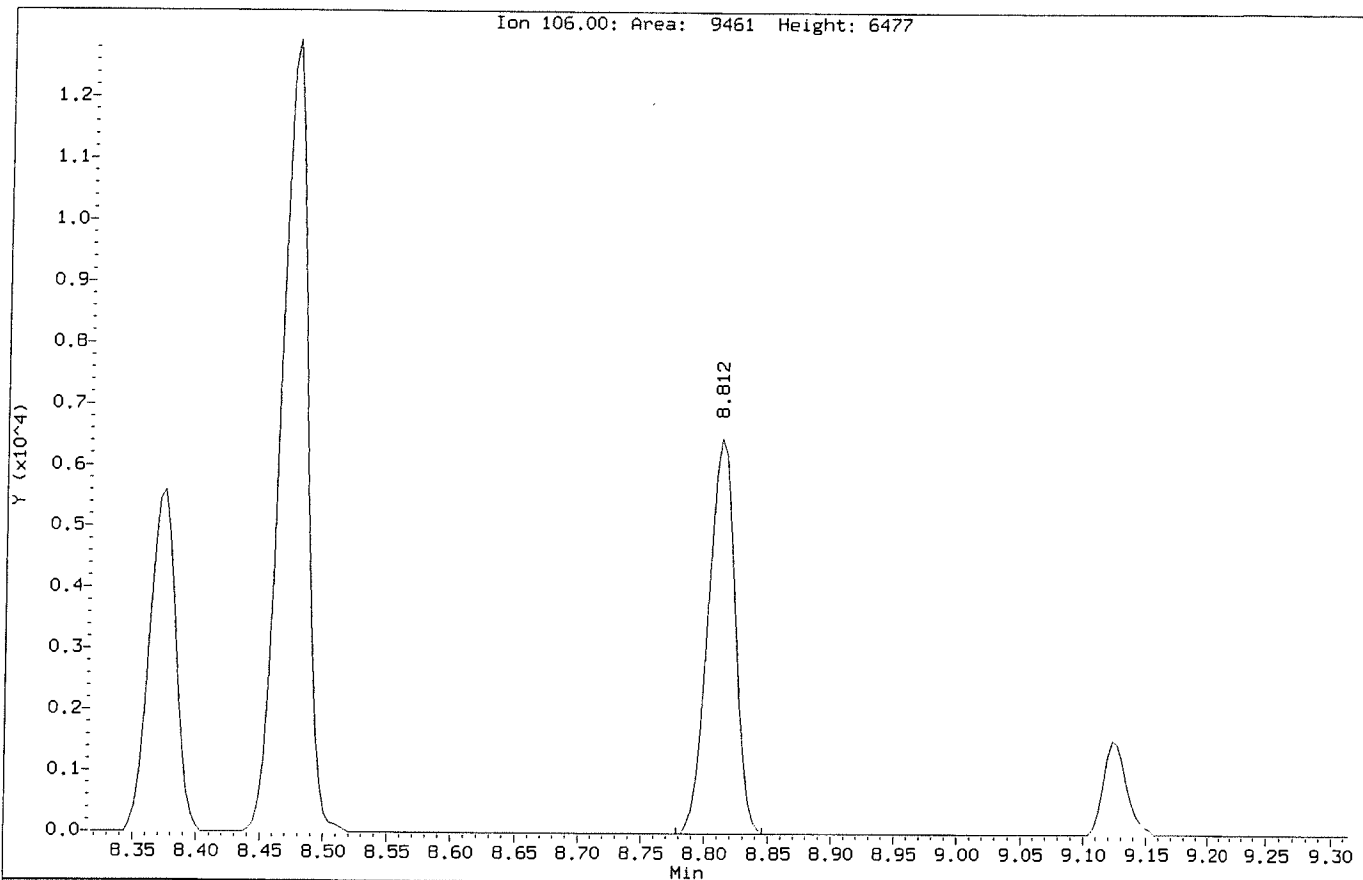
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Injection Date: 13-NOV-2018 12:36
Instrument: VDA9.i
Client Sample ID: VSTD001

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.1\U181113.b\U111304.D
Injection Date: 13-NOV-2018 12:36
Instrument: VOA9.i
Client Sample ID: VSTD001

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111305.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111305.D
 Lab Smp Id: VSTD002 Client Smp ID: VSTD002
 Inj Date : 13-NOV-2018 13:01
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD002;VSTD002;1;4;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 12:36 Cal File: U111304.D
 Als bottle: 6 Calibration Sample, Level: 4
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	AMOUNTS				CAL-AMT (ug/l)	ON-COL (ug/l)
			MASS	RT	EXP RT	REL RT		
* 1 Pentafluorobenzene	168	====	4.894	4.894	(1.000)	414744	50.0000	
* 36 1,4-Difluorobenzene	114		5.625	5.625	(1.000)	809284	50.0000	
* 47 Chlorobenzene-d5	117		8.249	8.249	(1.000)	755385	50.0000	
* 70 1,4-Dichlorobenzene-d4	152		10.236	10.236	(1.000)	352080	50.0000	
\$ 30 Dibromofluoromethane	113		4.830	4.830	(0.987)	12925	2.00000	1.52(a)
\$ 35 1,2-Dichloroethane-d4	65		5.175	5.175	(1.057)	18531	2.00000	1.28(a)
\$ 48 Toluene-d8	98		6.990	6.990	(0.847)	51591	2.00000	1.05(a)
\$ 69 4-Bromofluorobenzene	95		9.257	9.257	(1.122)	18827	2.00000	1.50(a)
60 1,1,1,2-Tetrachloroethane	131		8.350	8.350	(1.012)	8223	2.00000	1.72(a)
31 1,1,1-Trichloroethane	97		4.827	4.827	(0.986)	14071	2.00000	1.93(a)
68 1,1,2,2-Tetrachloroethane	83		9.392	9.392	(0.918)	16127	2.00000	1.93(a)
138 Freon TF	101		2.401	2.401	(0.491)	8834	2.00000	2.12(a)
53 1,1,2-Trichloroethane	83		7.421	7.421	(0.900)	9196	2.00000	1.91(a)
22 1,1-Dichloroethane	63		3.604	3.604	(0.737)	18719	2.00000	1.96(a)
11 1,1-Dichloroethene	96		2.397	2.397	(0.490)	9368	2.00000	2.00(a)
32 1,1-Dichloropropene	75		5.003	5.003	(0.889)	14736	2.00000	1.97(a)
93 1,2,3-Trichlorobenzene	180		12.335	12.335	(1.205)	12553	2.00000	1.88(a)
71 1,2,3-Trichloropropane	75		9.426	9.426	(0.921)	15506	2.00000	1.78(a)
90 1,2,4-Trichlorobenzene	180		11.927	11.927	(1.165)	12849	2.00000	1.85(a)
79 1,2,4-Trimethylbenzene	105		9.943	9.943	(0.971)	36765	2.00000	1.88(a)
89 1,2-Dibromo-3-Chloropropane	155		11.233	11.233	(1.097)	1762	2.00000	2.38(a)
57 1,2-Dibromoethane	107		7.852	7.852	(0.952)	10196	2.00000	1.83(a)



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111305.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG				AMOUNTS		
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	21603	2.00000	1.92 (a)
33 1,2-Dichloroethane	62	5.250	5.250	(0.933)	17243	2.00000	2.03 (a)
42 1,2-Dichloropropane	63	6.082	6.082	(1.081)	10957	2.00000	1.90 (a)
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	35537	2.00000	1.87 (a)
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	21971	2.00000	1.98 (a)
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	19816	2.00000	1.92 (a)
84 1,4-Dichlorobenzene	146	10.255	10.255	(1.002)	22453	2.00000	2.02 (a)
26 2,2-Dichloropropane	77	4.275	4.275	(0.874)	10687	2.00000	1.81 (a)
24 2-Butanone	43	4.343	4.343	(0.887)	10886	4.00000	3.73 (a)
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	33012	2.00000	1.91 (a)
52 2-Hexanone	43	7.649	7.649	(0.927)	15967	4.00000	3.63 (a)
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	38540	2.00000	1.93 (a)
82 p-Isopropyltoluene	119	10.210	10.210	(0.997)	36591	2.00000	1.89 (a)
45 4-Methyl-2-Pentanone	43	6.915	6.915	(0.838)	23062	4.00000	3.65 (a)
10 Acetone	43	2.480	2.480	(0.507)	12023	4.00000	2.16 (a)
37 Benzene	78	5.216	5.216	(0.927)	43114	2.00000	1.92 (a)
74 Bromobenzene	156	9.381	9.381	(0.917)	10831	2.00000	1.87 (a)
29 Bromochloromethane	128	4.553	4.553	(0.930)	5091	2.00000	2.08 (a)
39 Bromodichloromethane	83	6.348	6.348	(1.129)	12072	2.00000	1.79 (a)
66 Bromoform	173	8.984	8.984	(1.089)	4530	2.00000	2.48 (Ta)
6 Bromomethane	94	1.670	1.670	(0.341)	7606	2.00000	4.02 (a)
19 Carbon Disulfide	76	2.589	2.589	(0.529)	55277	4.00000	3.82 (a)
34 Carbon Tetrachloride	117	4.995	4.995	(0.888)	10688	2.00000	1.83 (a)
59 Chlorobenzene	112	8.275	8.275	(1.003)	29617	2.00000	1.89 (a)
7 Chloroethane	64	1.749	1.749	(0.357)	9074	2.00000	2.09 (aM)
28 Chloroform	83	4.658	4.658	(0.952)	18550	2.00000	1.94 (a)
3 Chloromethane	50	1.336	1.336	(0.273)	12000	2.00000	2.38 (a)
27 cis-1,2-Dichloroethene	96	4.283	4.283	(0.875)	11719	2.00000	1.99 (a)
46 cis-1,3-Dichloropropene	75	6.757	6.757	(1.201)	13722	2.00000	1.62 (a)
55 Dibromochloromethane	129	7.758	7.758	(0.940)	7733	2.00000	1.61 (a)
44 Dibromomethane	93	6.191	6.191	(1.101)	7013	2.00000	1.94 (aM)
2 Dichlorodifluoromethane	85	1.202	1.202	(0.246)	12615	2.00000	3.01 (a)
61 Ethylbenzene	106	8.373	8.373	(1.015)	14817	2.00000	1.87 (a)
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	4667	2.00000	2.48 (a)
67 Isopropylbenzene	105	9.126	9.126	(1.106)	44094	2.00000	1.92 (a)
62 m,p-Xylenes	106	8.474	8.474	(1.027)	36359	4.00000	3.74 (a)
17 Methylene Chloride	84	2.870	2.870	(0.586)	13586	2.00000	1.90 (a)
87 n-Butylbenzene	91	10.558	10.558	(1.031)	36189	2.00000	2.36 (a)
73 n-Propylbenzene	91	9.475	9.475	(0.926)	53805	2.00000	1.93 (a)
92 Naphthalene	128	12.133	12.133	(1.185)	35700	2.00000	1.71 (a)
63 o-Xylene	106	8.811	8.811	(1.068)	18379	2.00000	1.89 (aH)
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	43610	2.00000	1.88 (a)
64 Styrene	104	8.826	8.826	(1.070)	30570	2.00000	1.86 (a)
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	31095	2.00000	1.92 (a)
56 Tetrachloroethene	164	7.526	7.526	(0.912)	7936	2.00000	2.00 (a)
50 Toluene	91	7.046	7.046	(0.854)	47615	2.00000	1.98 (a)
20 trans-1,2-Dichloroethene	96	3.143	3.143	(0.642)	10067	2.00000	1.96 (a)
51 trans-1,3-Dichloropropene	75	7.263	7.263	(1.291)	10881	2.00000	2.73 (a)
38 Trichloroethene	130	5.861	5.861	(1.042)	10685	2.00000	1.93 (a)
8 Trichlorofluoromethane	101	1.951	1.951	(0.399)	17301	2.00000	2.08 (a)
5 Vinyl Chloride	62	1.415	1.415	(0.289)	13465	2.00000	2.47 (a)



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111305.D Page 3
Report Date: 24-Jan-2019 18:55

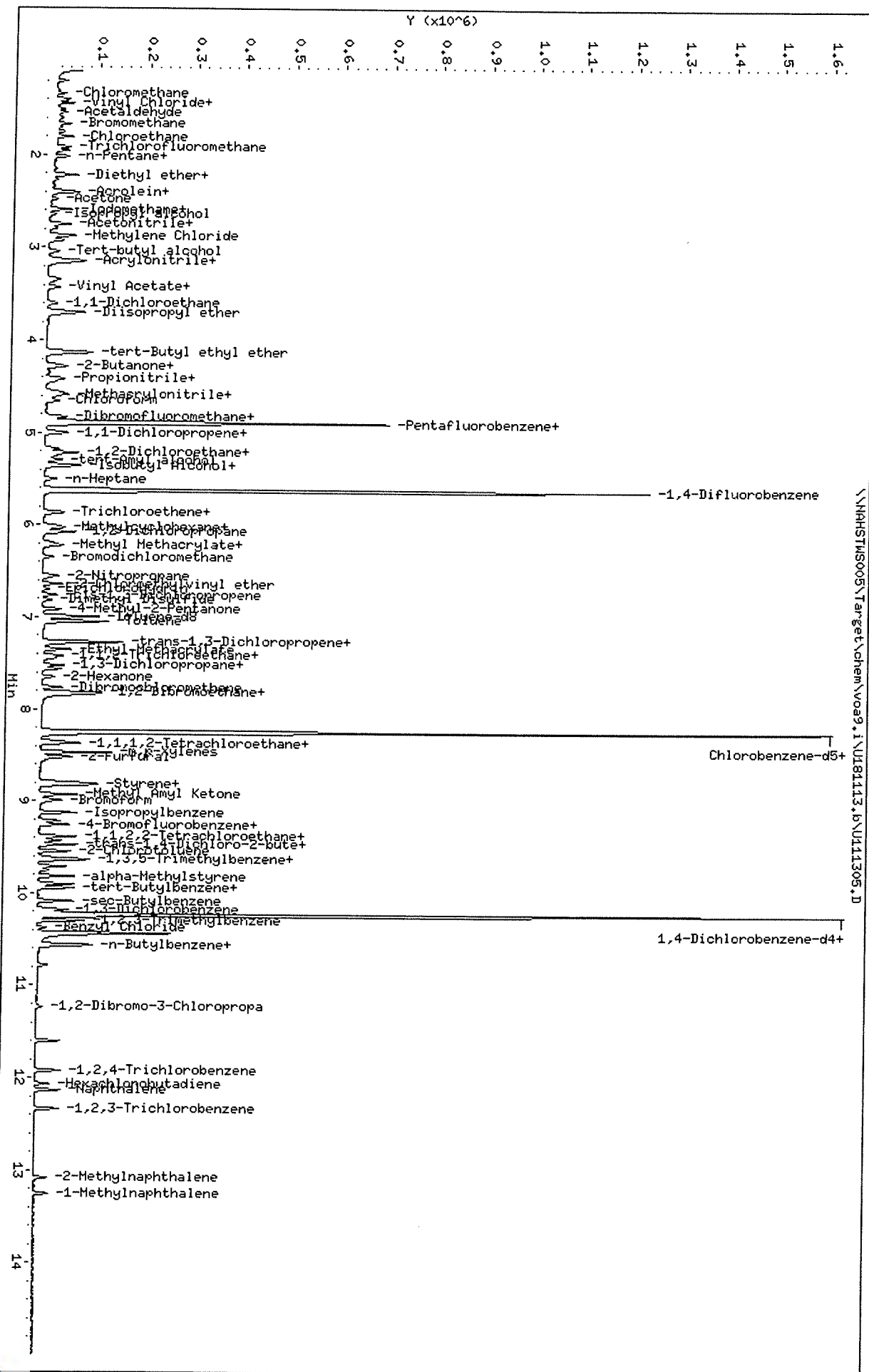
QC Flag Legend

- T - Target compound detected outside RT window.
- a - Target compound detected but, quantitated amount
Below Limit Of Quantitation(BLOQ).
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



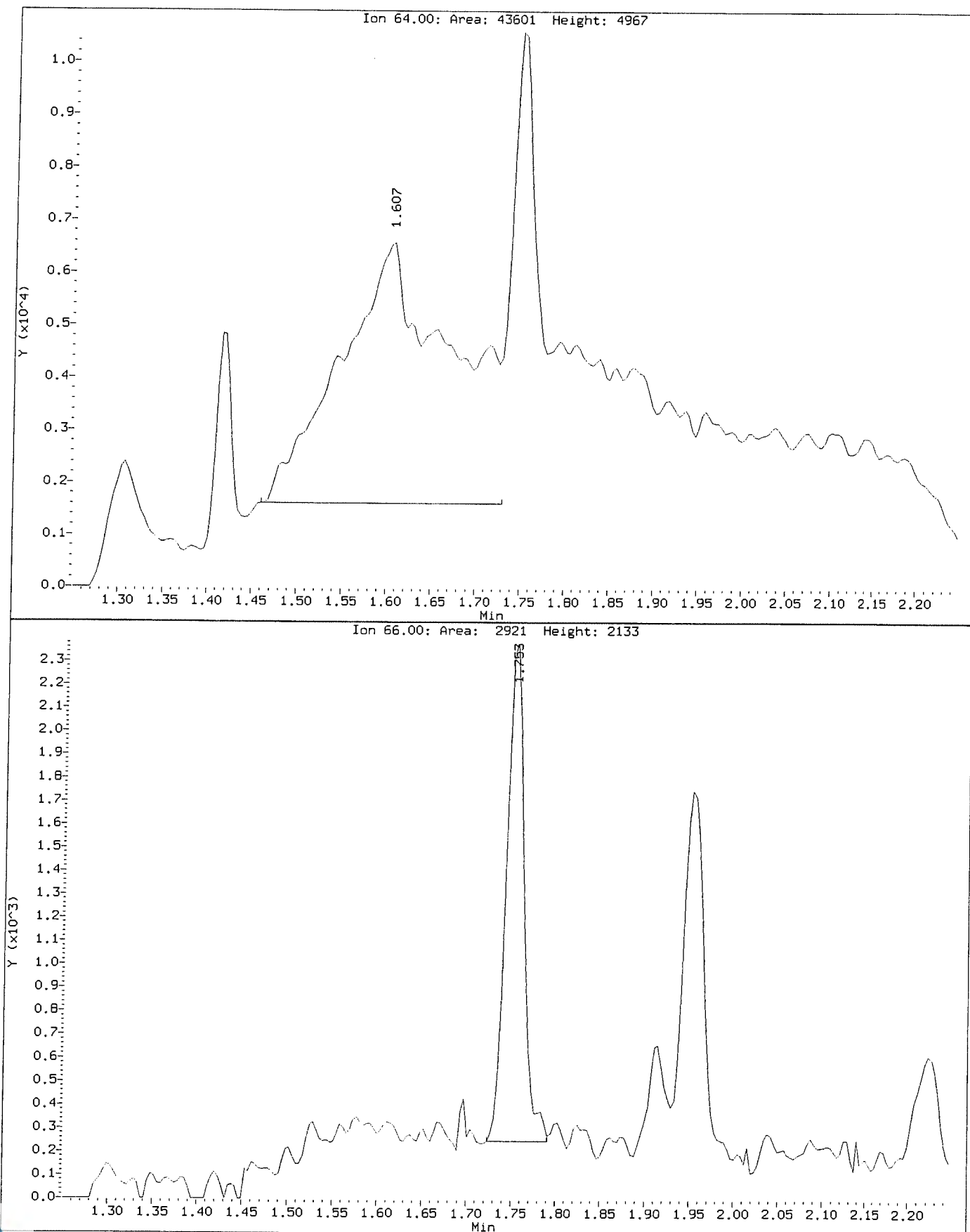
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Date : 13-NOV-2018 13:01
Client ID: VSTD002
Sample Info: VSTD002;VSTD002;1;4;
Purge Volume: 5.0
Column phase: DB624

Instrument: VOA9.i
Operator: PC
Column diameter: 0.18



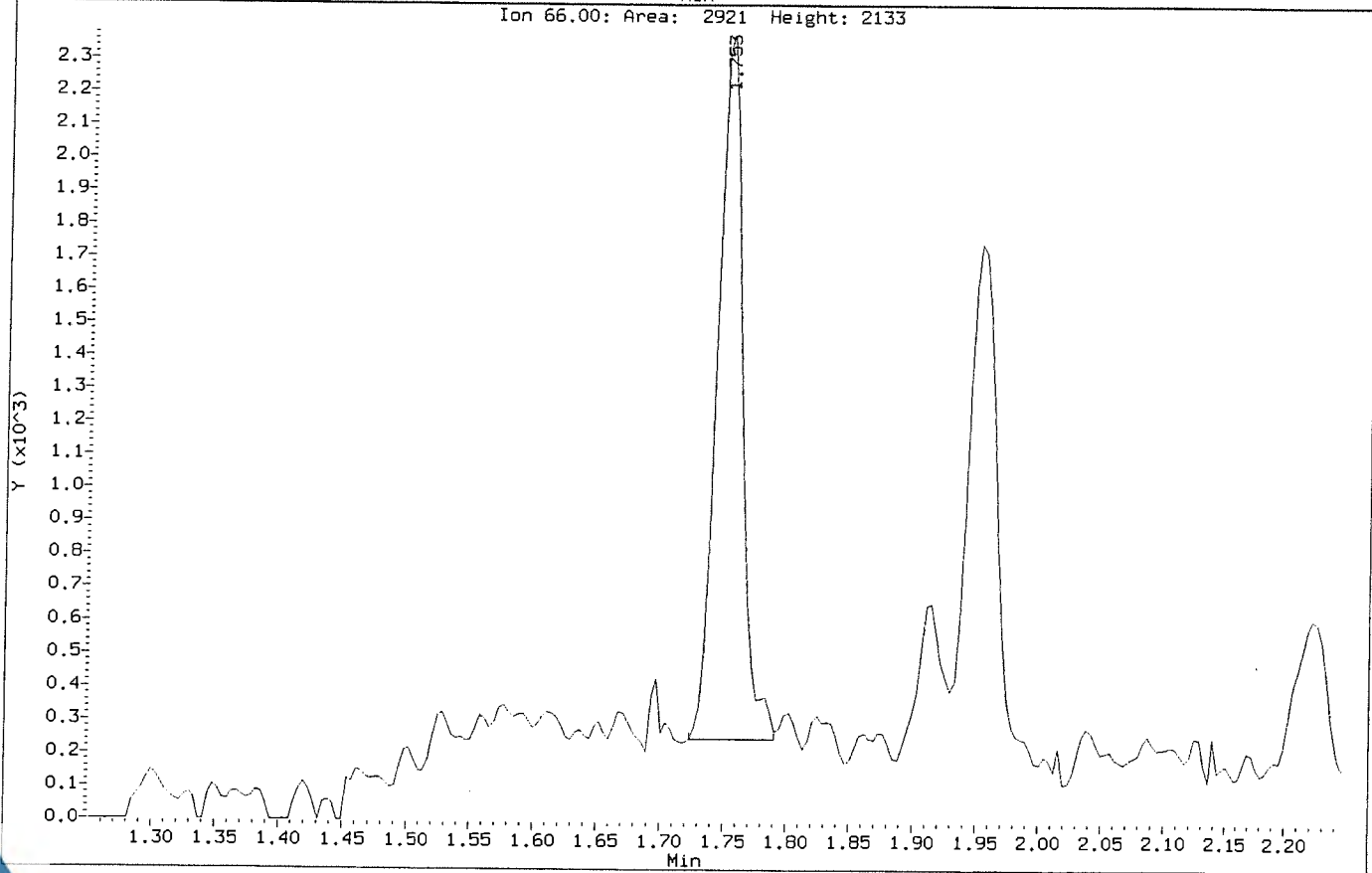
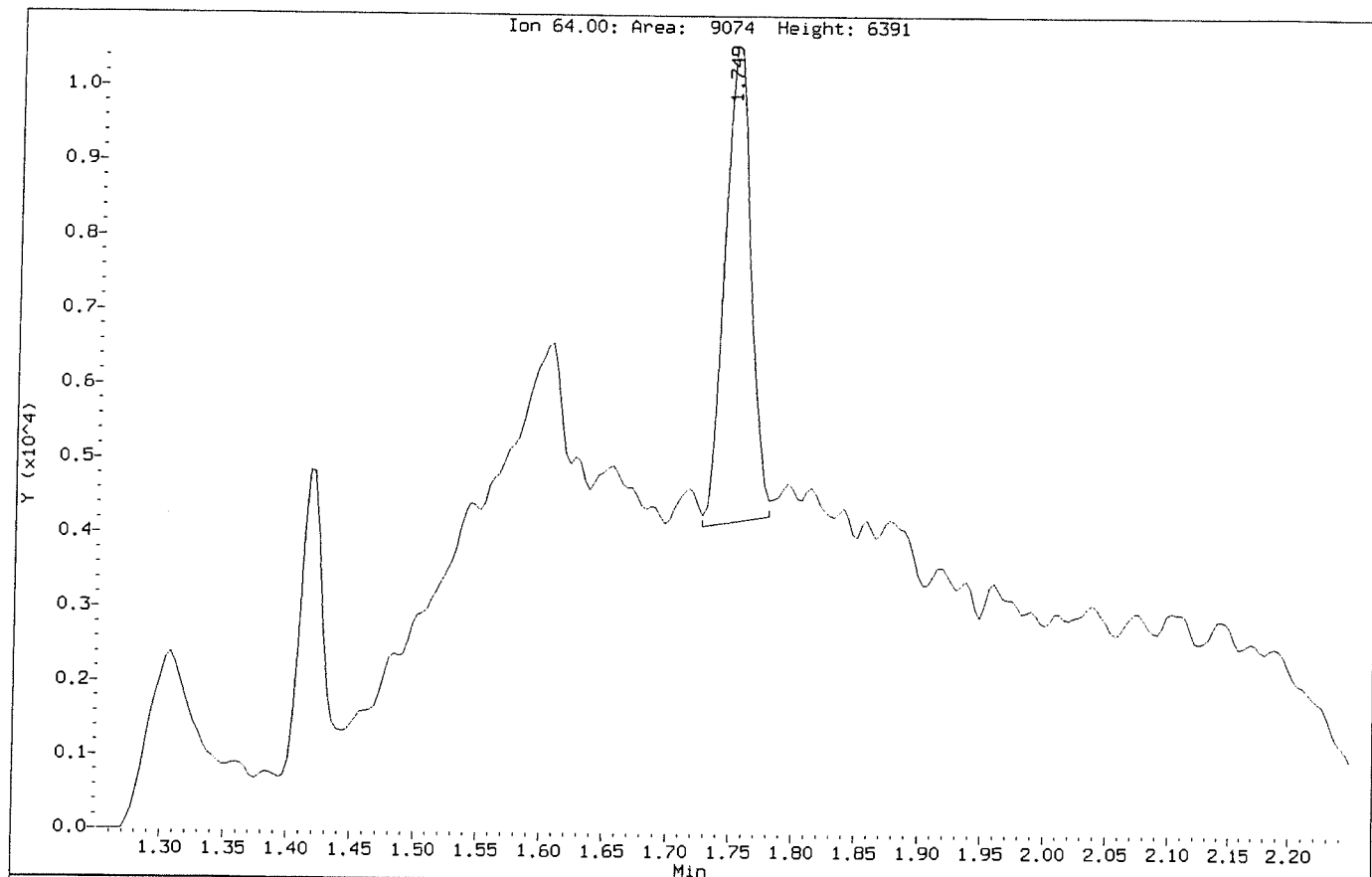
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Injection Date: 13-NOV-2018 13:01
Instrument: VOA9.1
Client Sample ID: VSTD002

Compound: Chloroethane
CAS Number: 75-00-3



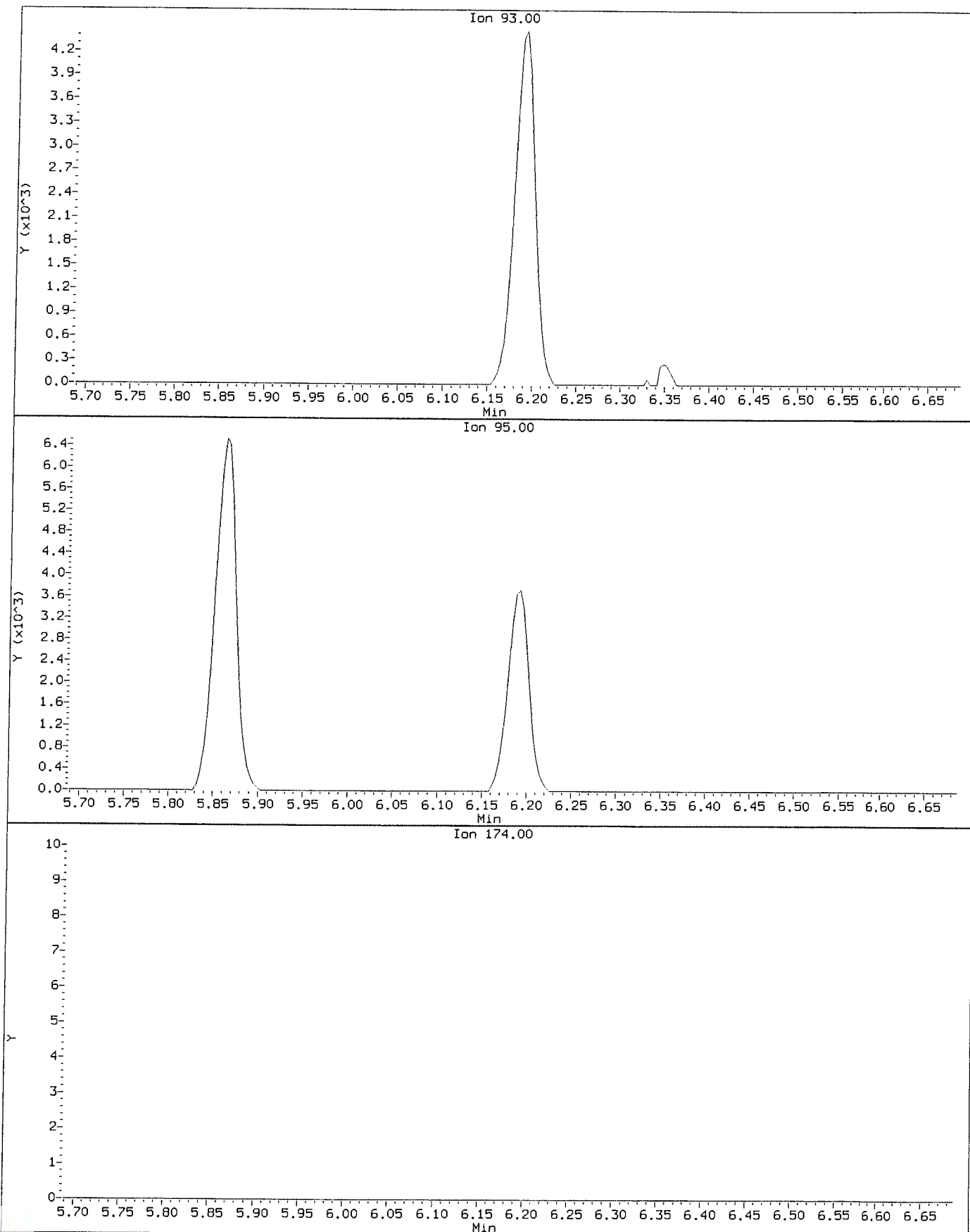
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Injection Date: 13-NOV-2018 13:01
Instrument: VOA9.1
Client Sample ID: VSTD002

Compound: Chloroethane
CAS Number: 75-00-3



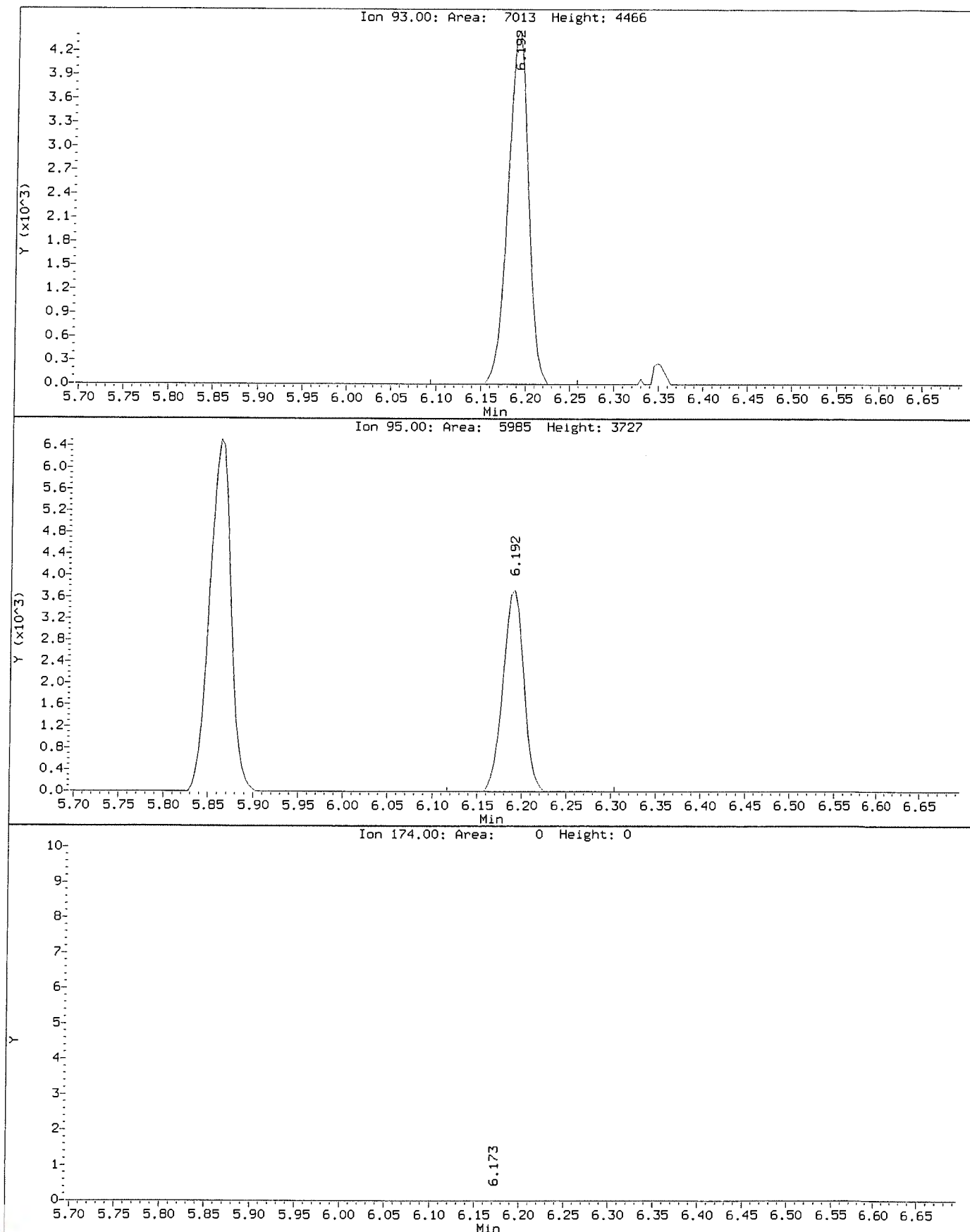
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Injection Date: 13-NOV-2018 13:01
Instrument: VOA9.i
Client Sample ID: VSTD002

Compound: Dibromomethane
CAS Number: 74-95-3



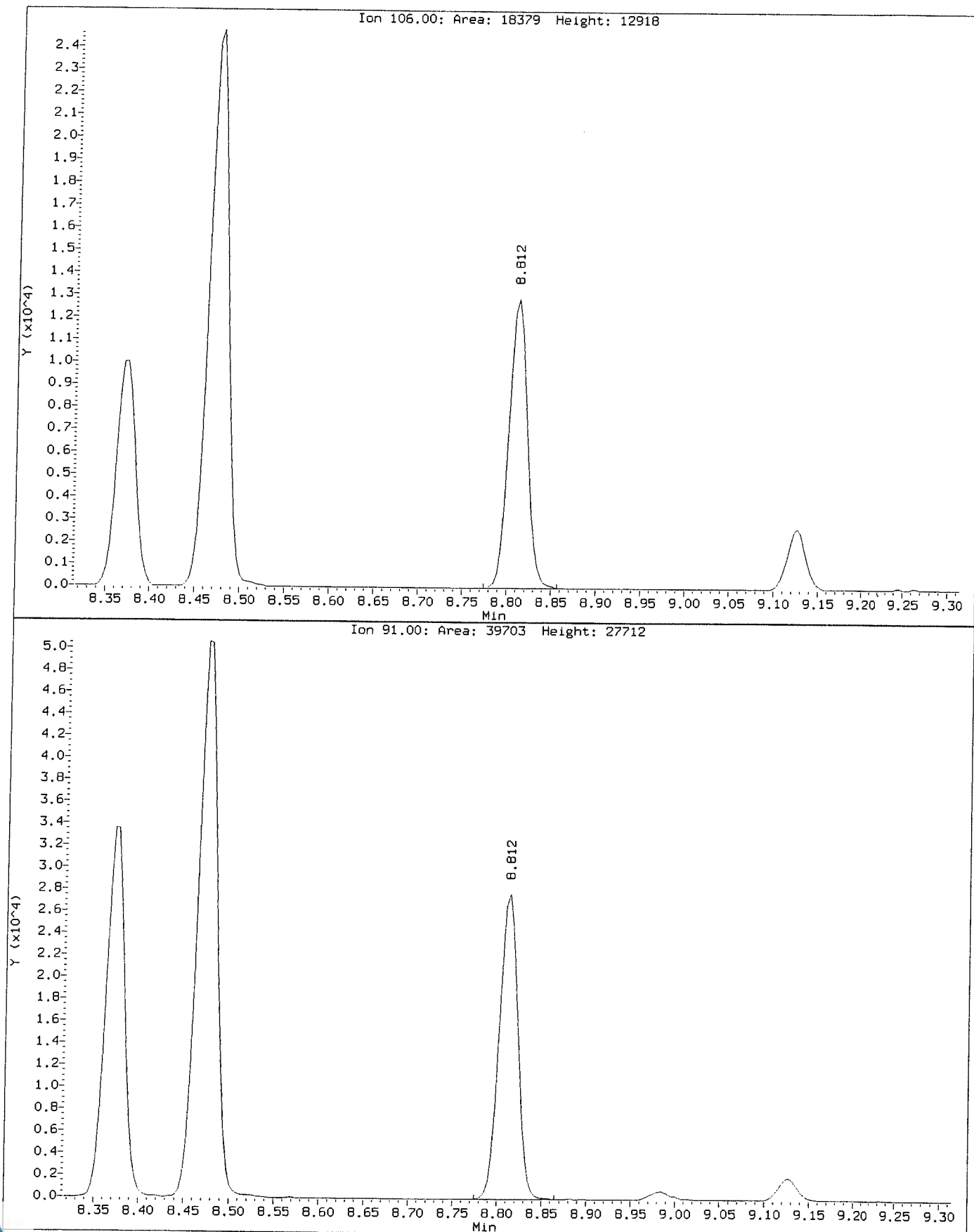
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Instrument: VOA9.i
Client Sample ID: VSTD002

Compound: Dibromomethane
CAS Number: 74-95-3



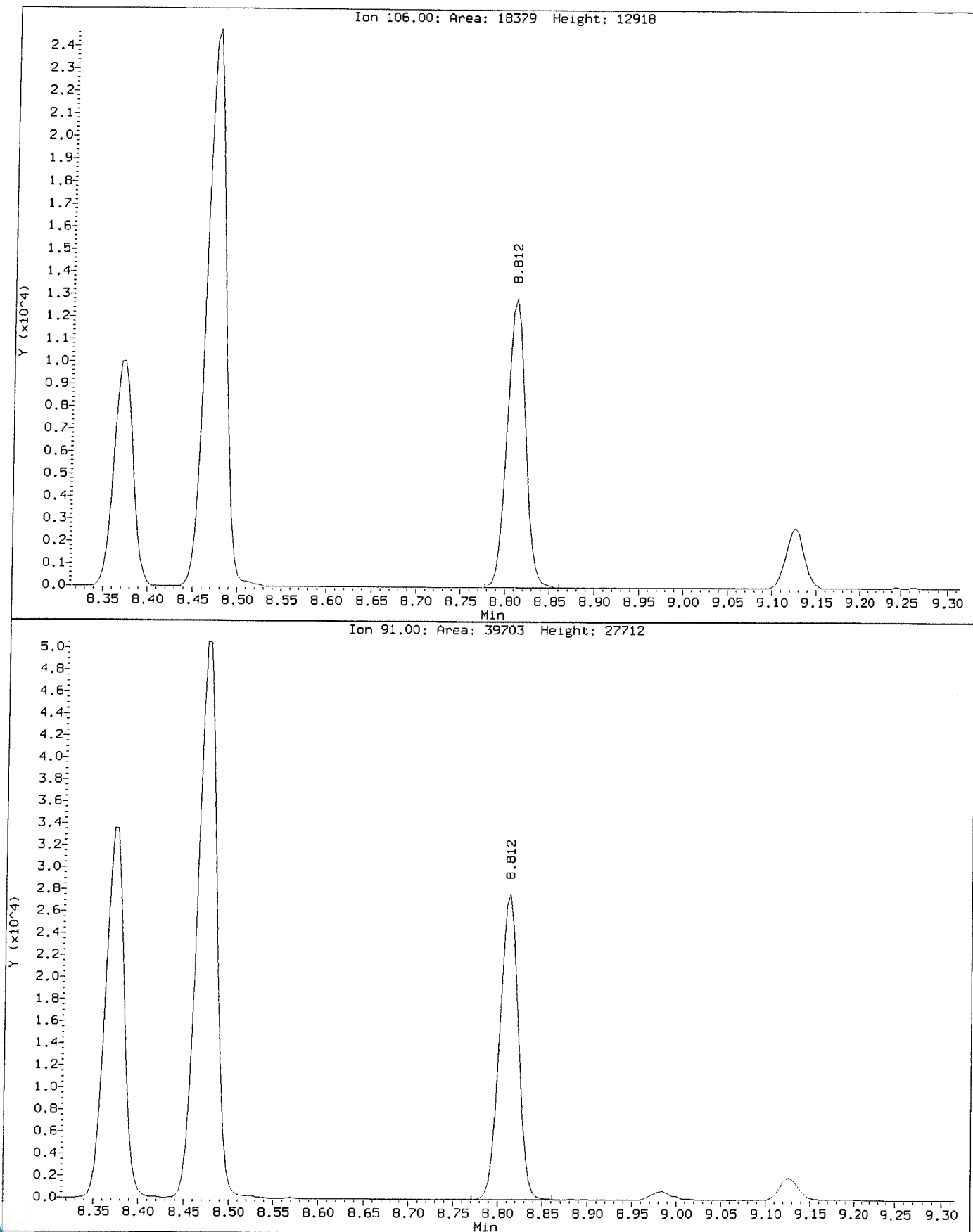
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Injection Date: 13-NOV-2018 13:01
Instrument: VOA9.1
Client Sample ID: VSTD002

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.1\U181113.b\U111305.D
Injection Date: 13-NOV-2018 13:01
Instrument: VOA9.1
Client Sample ID: VSTD002

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111306.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111306.D
 Lab Smp Id: VSTD005 Client Smp ID: VSTD005
 Inj Date : 13-NOV-2018 13:25
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD005;VSTD005;1;5;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 13:01 Cal File: U111305.D
 Als bottle: 7 Calibration Sample, Level: 5
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	AMOUNTS					CAL-AMT (ug/l)	ON-COL (ug/l)
		MASS	RT	EXP RT	REL RT	RESPONSE		
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	422530	50.0000		
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	813180	50.0000		
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	744948	50.0000		
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	348698	50.0000		
\$ 30 Dibromofluoromethane	113	4.830	4.830	(0.987)	27004	5.00000	4.44(a)	
\$ 35 1,2-Dichloroethane-d4	65	5.175	5.175	(1.057)	38984	5.00000	4.54(a)	
\$ 48 Toluene-d8	98	6.990	6.990	(0.847)	112942	5.00000	4.42(a)	
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	41214	5.00000	4.64(a)	
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	21915	5.00000	4.66(a)	
31 1,1,1-Trichloroethane	97	4.827	4.827	(0.986)	31327	5.00000	4.22(a)	
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	42360	5.00000	5.12	
138 Freon TF	101	2.405	2.405	(0.491)	17279	5.00000	4.08(a)	
53 1,1,2-Trichloroethane	83	7.421	7.421	(0.900)	23401	5.00000	4.94(a)	
22 1,1-Dichloroethane	63	3.604	3.604	(0.737)	45506	5.00000	4.69(a)	
11 1,1-Dichloroethene	96	2.401	2.401	(0.491)	20013	5.00000	4.19(a)	
32 1,1-Dichloropropene	75	5.006	5.006	(0.890)	33087	5.00000	4.40(a)	
93 1,2,3-Trichlorobenzene	180	12.335	12.335	(1.205)	31782	5.00000	4.82(a)	
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	42556	5.00000	4.93(a)	
90 1,2,4-Trichlorobenzene	180	11.927	11.927	(1.165)	31441	5.00000	4.58(a)	
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	93787	5.00000	4.84(a)	
89 1,2-Dibromo-3-Chloropropane	155	11.233	11.233	(1.097)	5351	5.00000	5.17	
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	27191	5.00000	4.95(a)	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111306.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	55248	5.00000	4.96(a)
33 1,2-Dichloroethane	62	5.254	5.254	(0.934)	41764	5.00000	4.90(a)
42 1,2-Dichloropropane	63	6.082	6.082	(1.081)	27675	5.00000	4.78(a)
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	88227	5.00000	4.70(a)
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	52374	5.00000	4.77(a)
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	51002	5.00000	5.01
84 1,4-Dichlorobenzene	146	10.255	10.255	(1.002)	55298	5.00000	5.11
26 2,2-Dichloropropane	77	4.275	4.275	(0.874)	25227	5.00000	4.20(a)
24 2-Butanone	43	4.343	4.343	(0.887)	30016	10.00000	10.10
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	82685	5.00000	4.84(a)
52 2-Hexanone	43	7.653	7.653	(0.928)	44101	10.00000	10.18
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	97151	5.00000	4.93(a)
82 p-Isopropyltoluene	119	10.210	10.210	(0.997)	85956	5.00000	4.50(a)
45 4-Methyl-2-Pentanone	43	6.915	6.915	(0.838)	62316	10.00000	10.02
10 Acetone	43	2.484	2.484	(0.508)	24249	10.00000	8.62
37 Benzene	78	5.220	5.220	(0.928)	105397	5.00000	4.68(a)
74 Bromobenzene	156	9.381	9.381	(0.917)	27842	5.00000	4.86(a)
29 Bromochloromethane	128	4.557	4.557	(0.931)	12948	5.00000	5.19
39 Bromodichloromethane	83	6.349	6.349	(1.129)	30775	5.00000	4.55(a)
66 Bromoform	173	8.984	8.984	(1.089)	13391	5.00000	5.07(T)
6 Bromomethane	94	1.674	1.674	(0.342)	17760	5.00000	6.77
19 Carbon Disulfide	76	2.592	2.592	(0.530)	124164	10.00000	8.43
34 Carbon Tetrachloride	117	4.995	4.995	(0.888)	23569	5.00000	4.01(a)
59 Chlorobenzene	112	8.275	8.275	(1.003)	72938	5.00000	4.74(a)
7 Chloroethane	64	1.756	1.756	(0.359)	19233	5.00000	4.36(aMH)
28 Chloroform	83	4.658	4.658	(0.952)	45918	5.00000	4.72(a)
3 Chloromethane	50	1.340	1.340	(0.274)	26431	5.00000	4.89(a)
27 cis-1,2-Dichloroethene	96	4.290	4.290	(0.877)	27696	5.00000	4.62(a)
46 cis-1,3-Dichloropropene	75	6.761	6.761	(1.202)	37232	5.00000	4.37(a)
55 Dibromochloromethane	129	7.758	7.758	(0.940)	21374	5.00000	4.53(a)
44 Dibromomethane	93	6.191	6.191	(1.101)	17227	5.00000	4.74(aM)
2 Dichlorodifluoromethane	85	1.205	1.205	(0.246)	24157	5.00000	4.79(a)
61 Ethylbenzene	106	8.373	8.373	(1.015)	35416	5.00000	4.55(a)
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	8782	5.00000	4.49(a)
67 Isopropylbenzene	105	9.126	9.126	(1.106)	106616	5.00000	4.70(a)
62 m,p-Xylenes	106	8.474	8.474	(1.027)	89708	10.00000	9.36
17 Methylene Chloride	84	2.873	2.873	(0.587)	29126	5.00000	4.77(a)
87 n-Butylbenzene	91	10.558	10.558	(1.031)	79206	5.00000	4.76(a)
73 n-Propylbenzene	91	9.475	9.475	(0.926)	128990	5.00000	4.68(a)
92 Naphthalene	128	12.133	12.133	(1.185)	100680	5.00000	4.87(a)
63 o-Xylene	106	8.811	8.811	(1.068)	46119	5.00000	4.81(aH)
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	99437	5.00000	4.34(a)
64 Styrene	104	8.826	8.826	(1.070)	78010	5.00000	4.81(a)
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	73015	5.00000	4.57(a)
56 Tetrachloroethene	164	7.526	7.526	(0.912)	17447	5.00000	4.47(a)
50 Toluene	91	7.050	7.050	(0.855)	113936	5.00000	4.81(a)
20 trans-1,2-Dichloroethene	96	3.143	3.143	(0.642)	23739	5.00000	4.54(a)
51 trans-1,3-Dichloropropene	75	7.259	7.259	(1.291)	30108	5.00000	4.94(a)
38 Trichloroethene	130	5.861	5.861	(1.042)	24849	5.00000	4.48(a)
8 Trichlorofluoromethane	101	1.955	1.955	(0.400)	34539	5.00000	4.08(a)
5 Vinyl Chloride	62	1.419	1.419	(0.290)	29718	5.00000	4.77(a)



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111306.D Page 3
Report Date: 24-Jan-2019 18:55

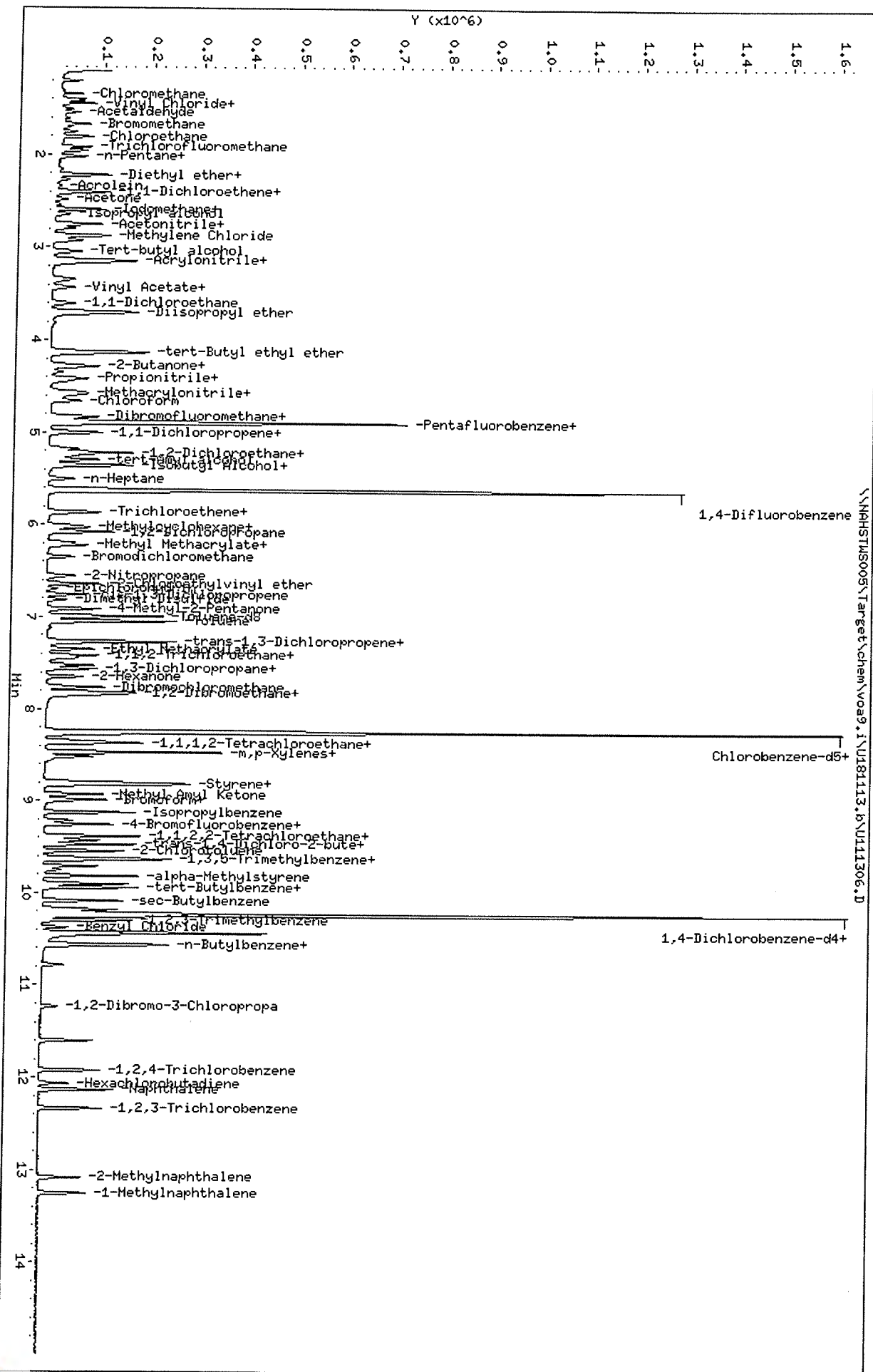
QC Flag Legend

- T - Target compound detected outside RT window.
- a - Target compound detected but, quantitated amount
Below Limit Of Quantitation(BLOQ).
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



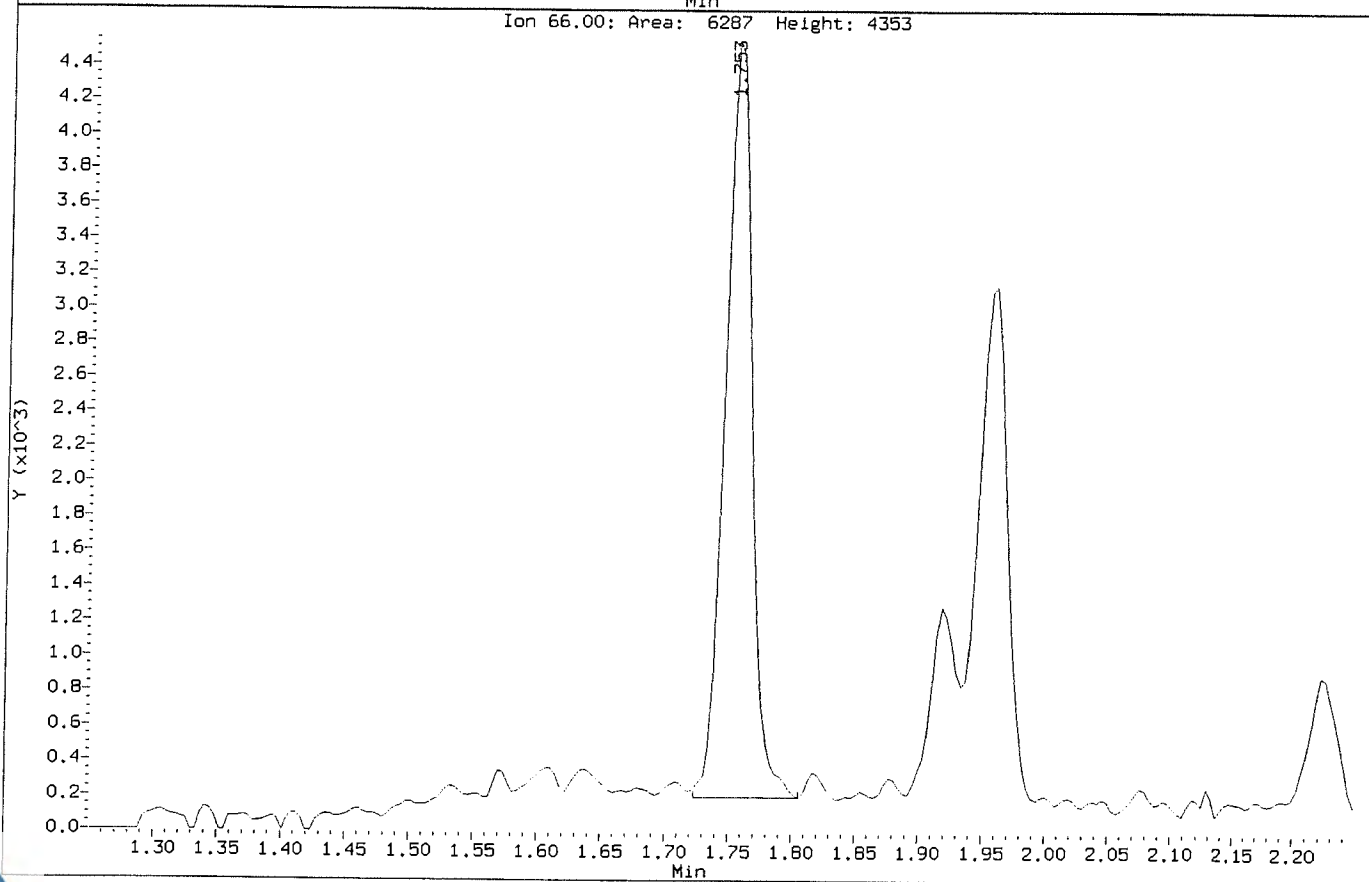
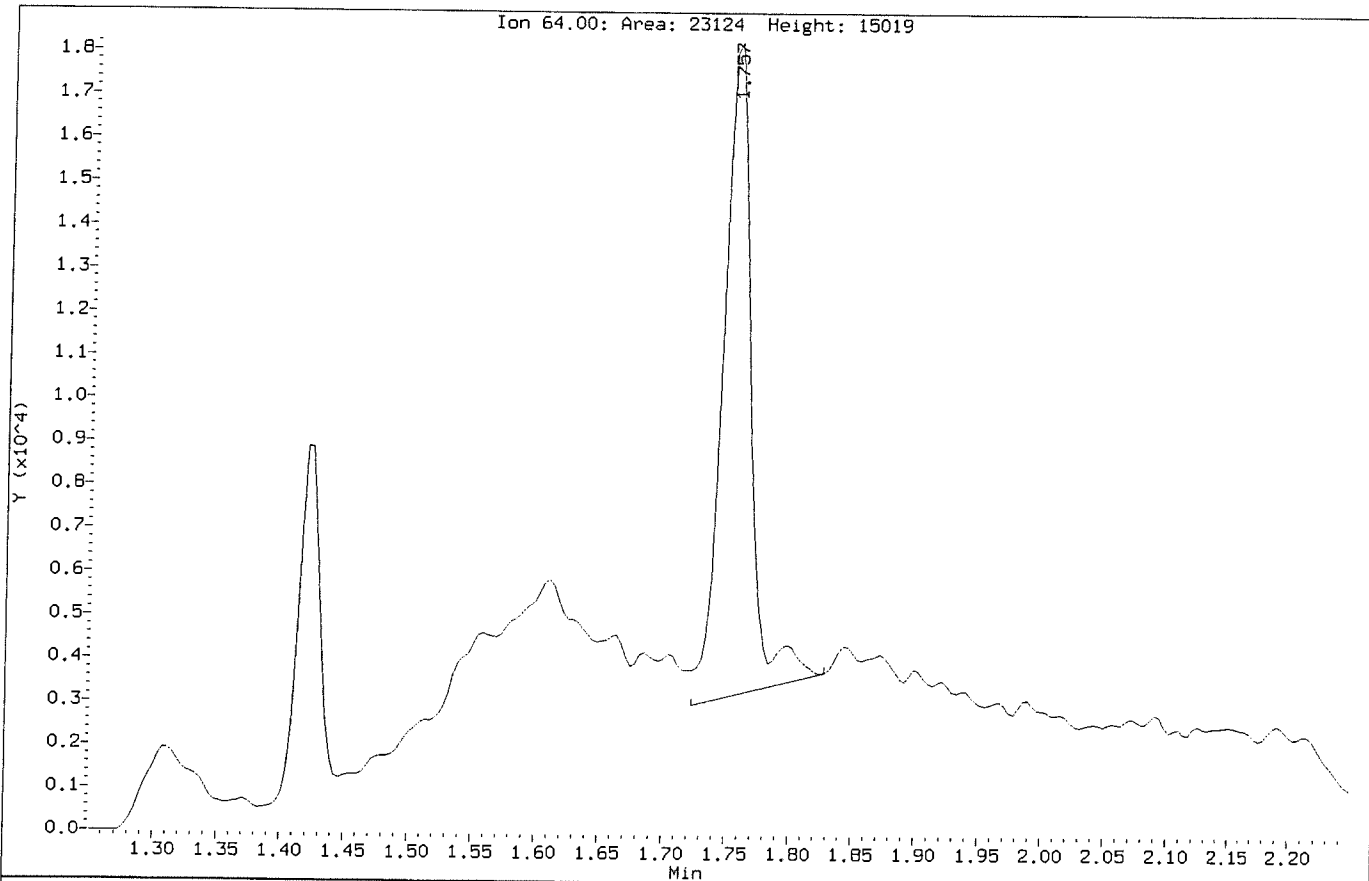
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 Date : 13-NOV-2018 13:25
 Client ID: VSTD005
 Sample Info: VSTD005;VSTD005;1;5;
 Purge Volume: 5.0
 Column phase: DB624

Instrument: VDA9.i
 Operator: PC
 Column diameter: 0.18



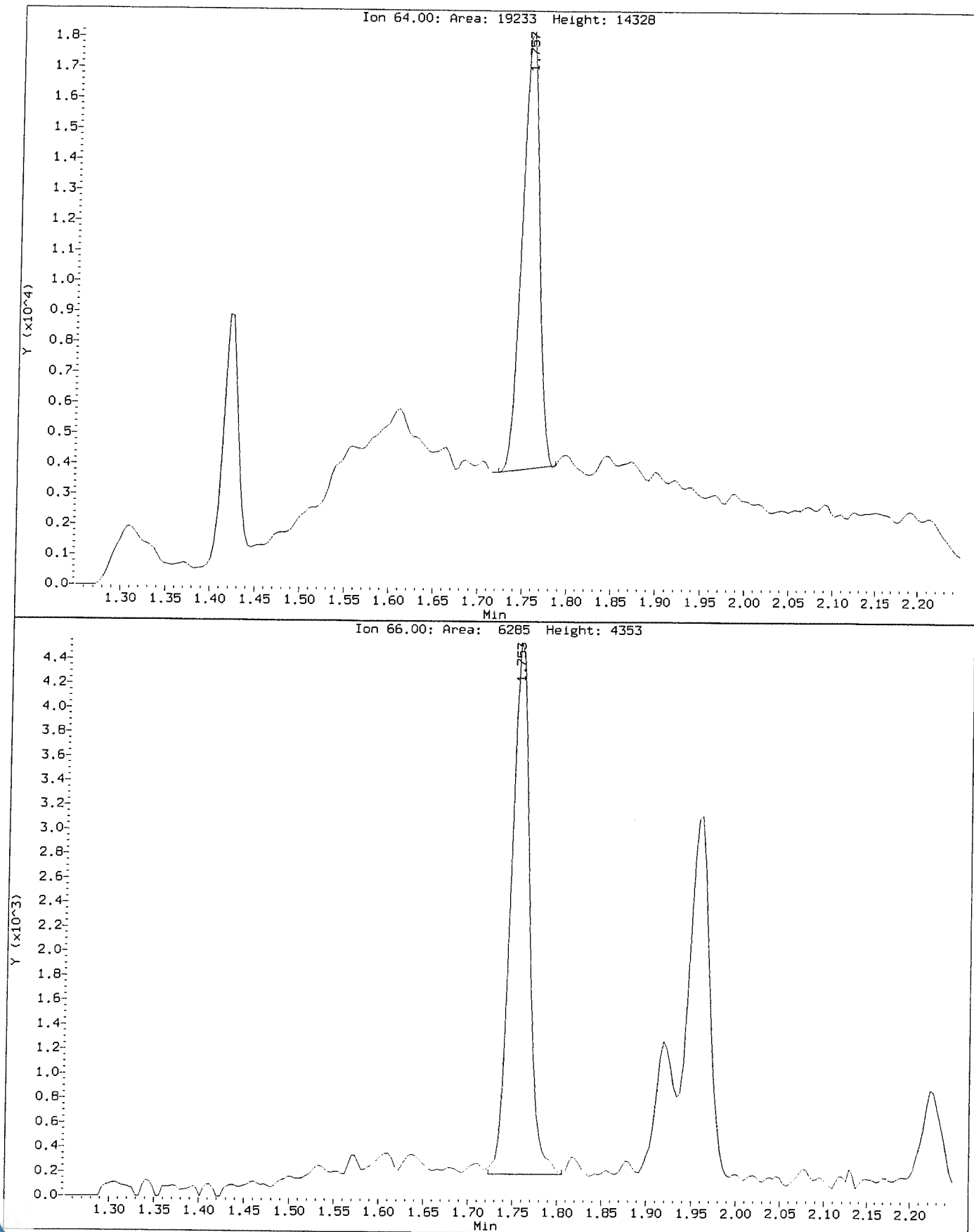
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Injection Date: 13-NOV-2016 13:25
Instrument: VOA9.i
Client Sample ID: VSTD005

Compound: Chloroethane
CAS Number: 75-00-3



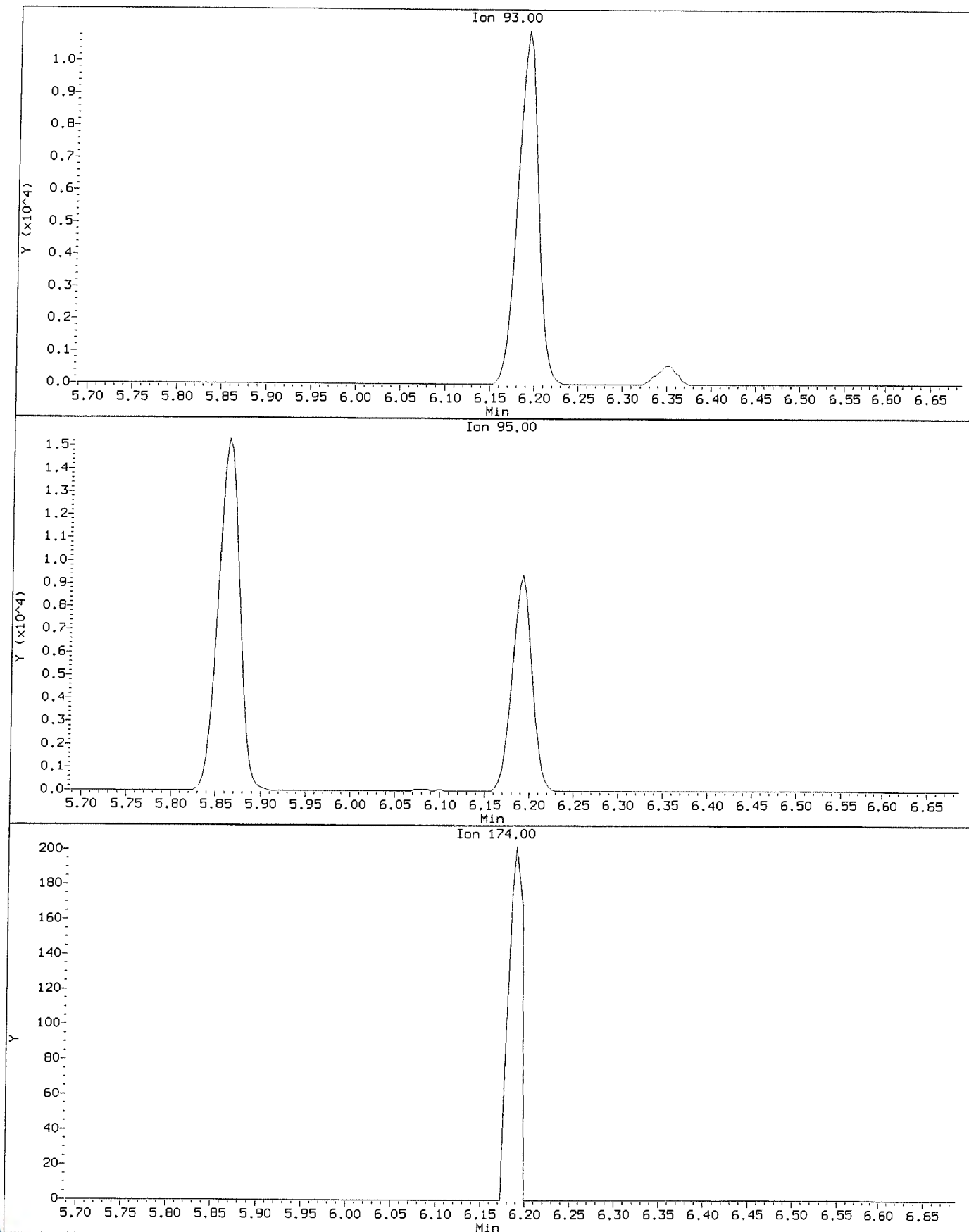
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Instrument: VOA9.1
Client Sample ID: VSTD005

Compound: Chloroethane
CAS Number: 75-00-3



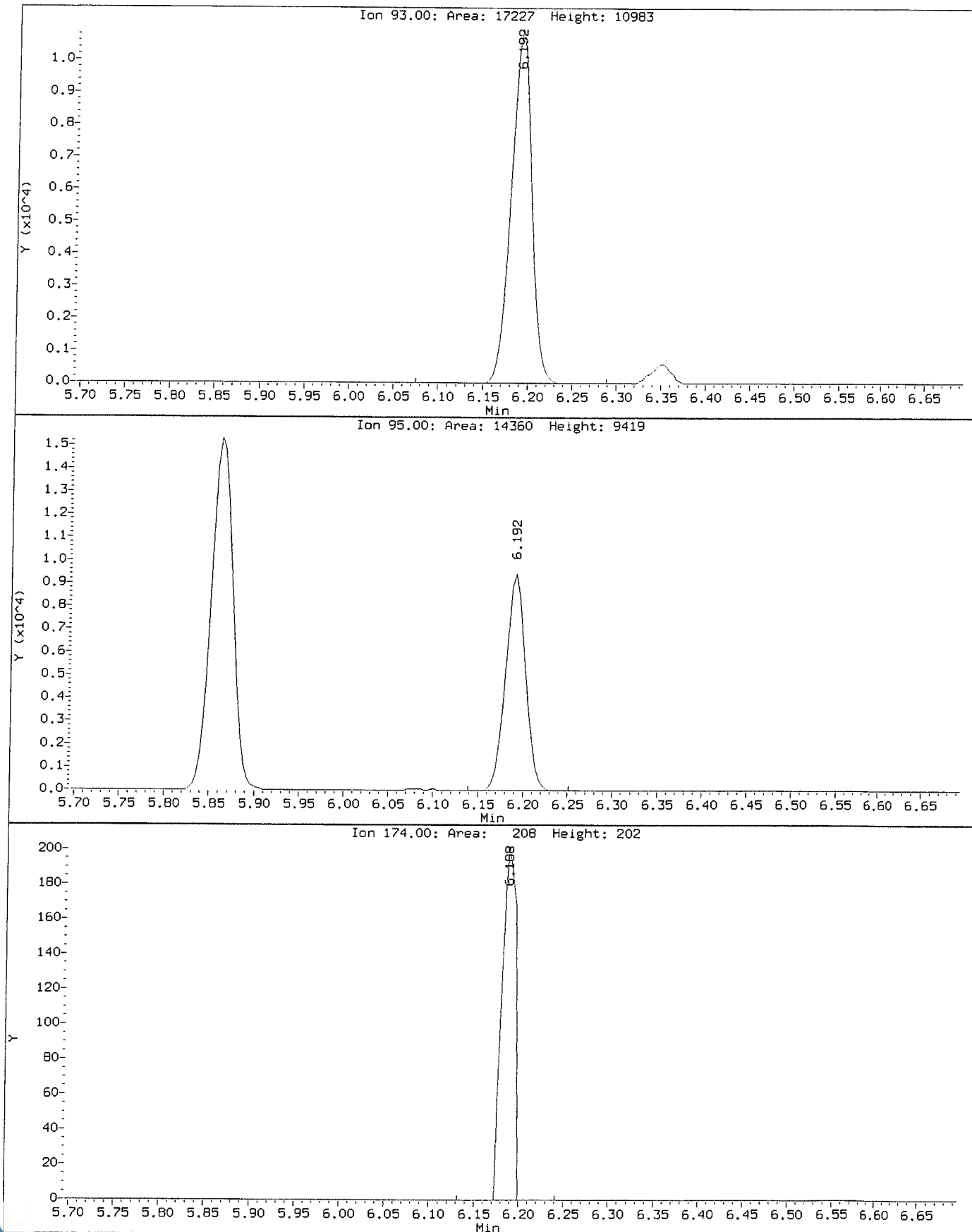
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Instrument: VOA9.1
Client Sample ID: VSTD005

Compound: Dibromomethane
CAS Number: 74-95-3



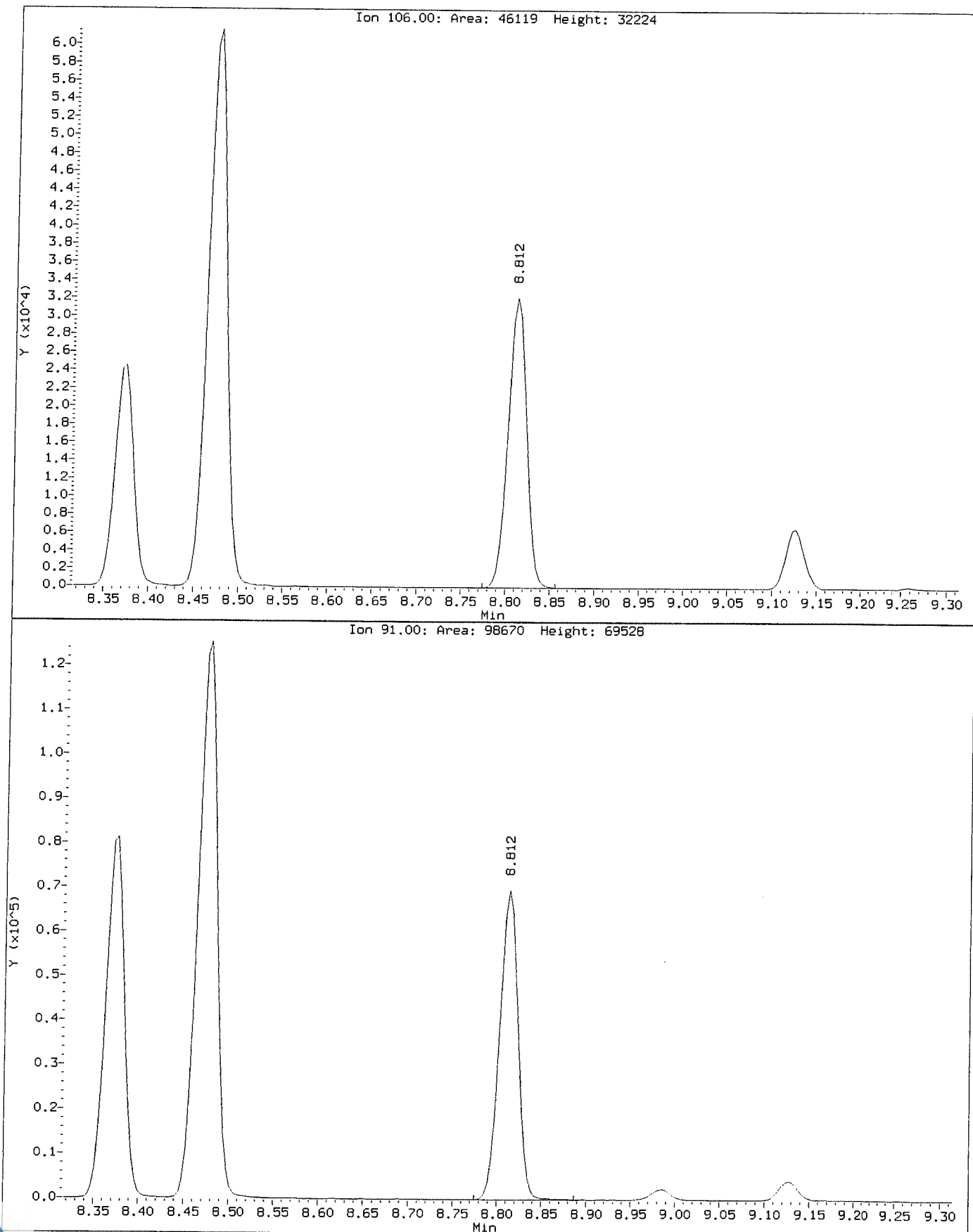
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Injection Date: 13-NOV-2018 13:25
Instrument: VOA9.i
Client Sample ID: VSTD005

Compound: Dibromomethane
CAS Number: 74-95-3



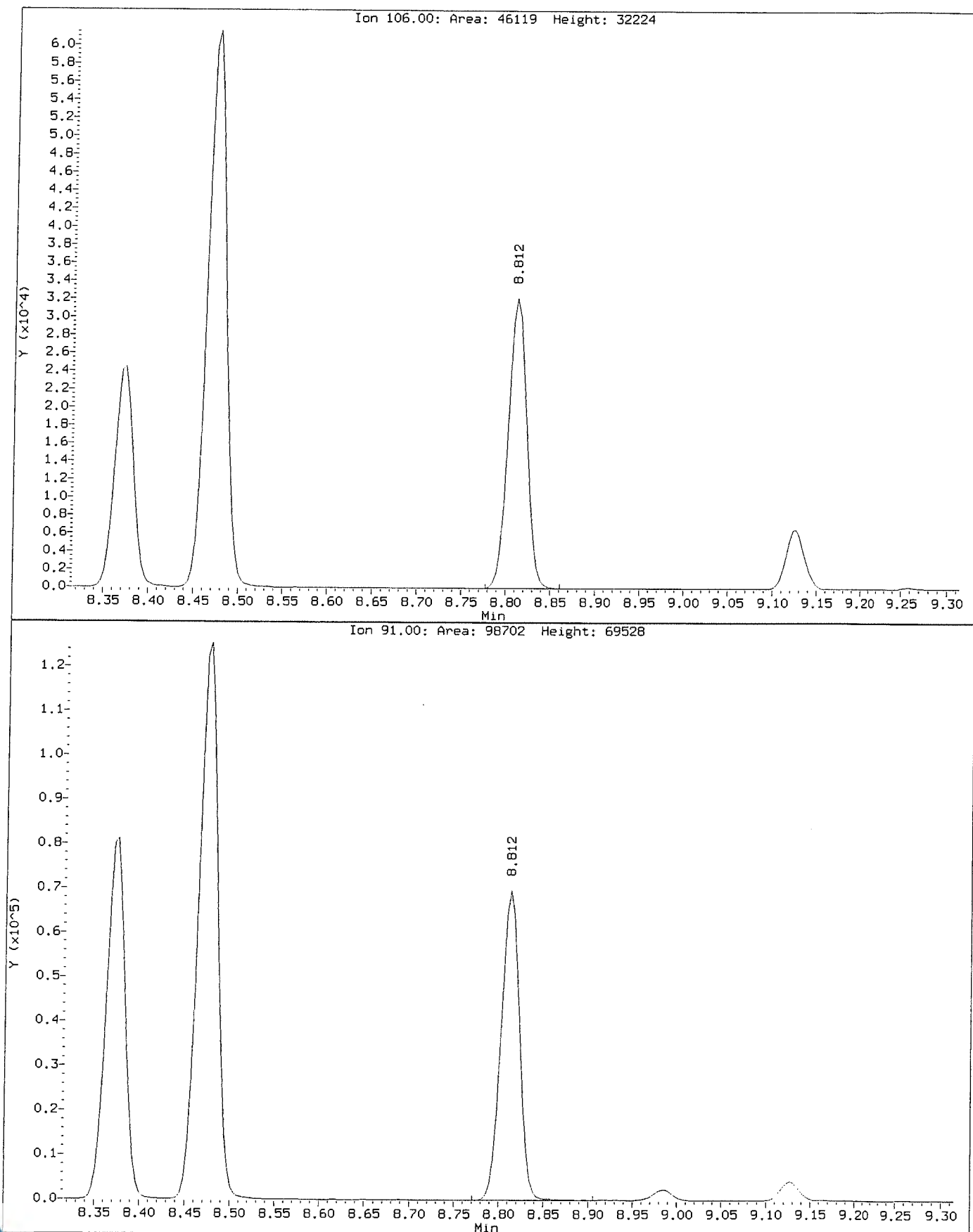
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Injection Date: 13-NOV-2018 13:25
Instrument: VOA9.i
Client Sample ID: VSTD005

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111306.D
Injection Date: 13-NOV-2018 13:25
Instrument: VDA9.1
Client Sample ID: VSTD005

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111307.D Page 1
 Report Date: 24-Jan-2019 18:55

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Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111307.D
 Lab Smp Id: VSTD020 Client Smp ID: VSTD020
 Inj Date : 13-NOV-2018 13:50
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD020;VSTD020;1;6;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 13:25 Cal File: U111306.D
 Als bottle: 8 Calibration Sample, Level: 6
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	AMOUNTS				CAL-AMT (ug/l)	ON-COL (ug/l)
			MASS	RT	EXP RT	REL RT		
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	441430	50.0000		
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	825527	50.0000		
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	767763	50.0000		
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	370767	50.0000		
\$ 30 Dibromofluoromethane	113	4.826	4.826	(0.986)	102630	20.0000	19.47	
\$ 35 1,2-Dichloroethane-d4	65	5.175	5.175	(1.057)	138453	20.0000	19.70	
\$ 48 Toluene-d8	98	6.989	6.989	(0.847)	412445	20.0000	20.02	
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	154498	20.0000	19.71	
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	94836	20.0000	19.58	
31 1,1,1-Trichloroethane	97	4.826	4.826	(0.986)	144369	20.0000	18.61	
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	170546	20.0000	19.41	
138 Freon TF	101	2.397	2.397	(0.490)	81210	20.0000	18.36	
53 1,1,2-Trichloroethane	83	7.420	7.420	(0.900)	94236	20.0000	19.30	
22 1,1-Dichloroethane	63	3.601	3.601	(0.736)	188042	20.0000	18.58	
11 1,1-Dichloroethene	96	2.397	2.397	(0.490)	92859	20.0000	18.63	
32 1,1-Dichloropropene	75	5.003	5.003	(0.889)	142695	20.0000	18.71	
93 1,2,3-Trichlorobenzene	180	12.335	12.335	(1.205)	136668	20.0000	19.52	
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	178640	20.0000	19.49	
90 1,2,4-Trichlorobenzene	180	11.926	11.926	(1.165)	137085	20.0000	18.80	
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	410829	20.0000	19.94	
89 1,2-Dibromo-3-Chloropropane	155	11.233	11.233	(1.097)	24969	20.0000	19.13	
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	112223	20.0000	19.85	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111307.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	222900	20.0000	18.82
33 1,2-Dichloroethane	62	5.250	5.250	(0.933)	165626	20.0000	19.17
42 1,2-Dichloropropane	63	6.082	6.082	(1.081)	114197	20.0000	19.42
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	396843	20.0000	19.92
83 1,3-Dichlorobenzene	146	10.179	10.179	(0.995)	220218	20.0000	18.86
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	202833	20.0000	19.35
84 1,4-Dichlorobenzene	146	10.254	10.254	(1.002)	228900	20.0000	20.02
26 2,2-Dichloropropane	77	4.272	4.272	(0.873)	113941	20.0000	18.18
24 2-Butanone	43	4.335	4.335	(0.886)	118404	40.0000	38.15
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	350860	20.0000	19.32
52 2-Hexanone	43	7.649	7.649	(0.927)	181765	40.0000	40.72
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	410588	20.0000	19.59
82 p-Isopropyltoluene	119	10.209	10.209	(0.997)	404381	20.0000	19.91
45 4-Methyl-2-Pentanone	43	6.914	6.914	(0.838)	257102	40.0000	40.14
10 Acetone	43	2.476	2.476	(0.506)	87797	40.0000	40.81
37 Benzene	78	5.216	5.216	(0.927)	436595	20.0000	19.13
74 Bromobenzene	156	9.381	9.381	(0.917)	115775	20.0000	19.03
29 Bromochloromethane	128	4.553	4.553	(0.930)	53724	20.0000	20.64
39 Bromodichloromethane	83	6.348	6.348	(1.129)	132026	20.0000	19.24
66 Bromoform	173	8.984	8.984	(1.089)	60993	20.0000	18.33(T)
6 Bromomethane	94	1.666	1.666	(0.341)	65394	20.0000	19.07
19 Carbon Disulfide	76	2.585	2.585	(0.528)	579343	40.0000	37.64
34 Carbon Tetrachloride	117	4.995	4.995	(0.888)	113726	20.0000	19.09
59 Chlorobenzene	112	8.275	8.275	(1.003)	302097	20.0000	19.06
7 Chloroethane	64	1.749	1.749	(0.357)	85496	20.0000	18.56(M)
28 Chloroform	83	4.654	4.654	(0.951)	190532	20.0000	18.74
3 Chloromethane	50	1.336	1.336	(0.273)	114171	20.0000	19.51
27 cis-1,2-Dichloroethene	96	4.283	4.283	(0.875)	119044	20.0000	19.03
46 cis-1,3-Dichloropropene	75	6.757	6.757	(1.201)	172818	20.0000	20.00
55 Dibromochloromethane	129	7.758	7.758	(0.940)	96570	20.0000	19.87
44 Dibromomethane	93	6.191	6.191	(1.101)	71419	20.0000	19.38
2 Dichlorodifluoromethane	85	1.201	1.201	(0.246)	119108	20.0000	18.89
61 Ethylbenzene	106	8.369	8.369	(1.015)	154131	20.0000	19.23
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	45235	20.0000	20.48
67 Isopropylbenzene	105	9.126	9.126	(1.106)	456275	20.0000	19.55
62 m,p-Xylenes	106	8.474	8.474	(1.027)	389423	40.0000	39.44
17 Methylene Chloride	84	2.866	2.866	(0.586)	113894	20.0000	19.80
87 n-Butylbenzene	91	10.558	10.558	(1.031)	381141	20.0000	20.10
73 n-Propylbenzene	91	9.475	9.475	(0.926)	574080	20.0000	19.62
92 Naphthalene	128	12.133	12.133	(1.185)	458615	20.0000	20.88
63 o-Xylene	106	8.811	8.811	(1.068)	197393	20.0000	19.97(H)
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	477299	20.0000	19.59
64 Styrene	104	8.826	8.826	(1.070)	337037	20.0000	20.20
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	334318	20.0000	19.69
56 Tetrachloroethene	164	7.525	7.525	(0.912)	75404	20.0000	18.74
50 Toluene	91	7.046	7.046	(0.854)	470397	20.0000	19.29
20 trans-1,2-Dichloroethene	96	3.139	3.139	(0.642)	101477	20.0000	18.58
51 trans-1,3-Dichloropropene	75	7.259	7.259	(1.291)	147063	20.0000	18.17
38 Trichloroethene	130	5.861	5.861	(1.042)	107204	20.0000	19.05
8 Trichlorofluoromethane	101	1.951	1.951	(0.399)	161290	20.0000	18.27
5 Vinyl Chloride	62	1.415	1.415	(0.289)	136290	20.0000	19.25



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111307.D Page 3
Report Date: 24-Jan-2019 18:55

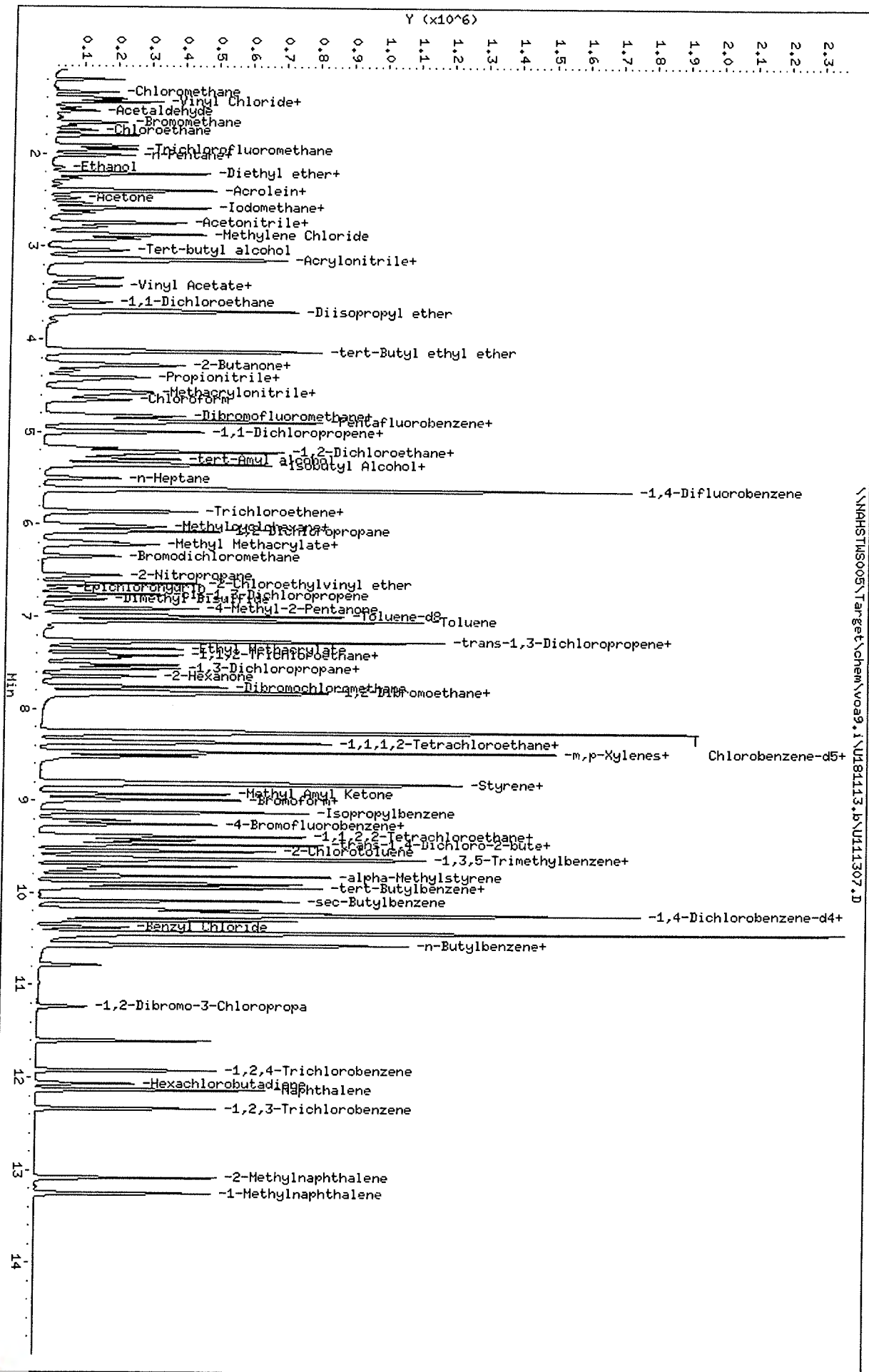
QC Flag Legend

T - Target compound detected outside RT window.
M - Compound response manually integrated.
H - Operator selected an alternate compound hit.



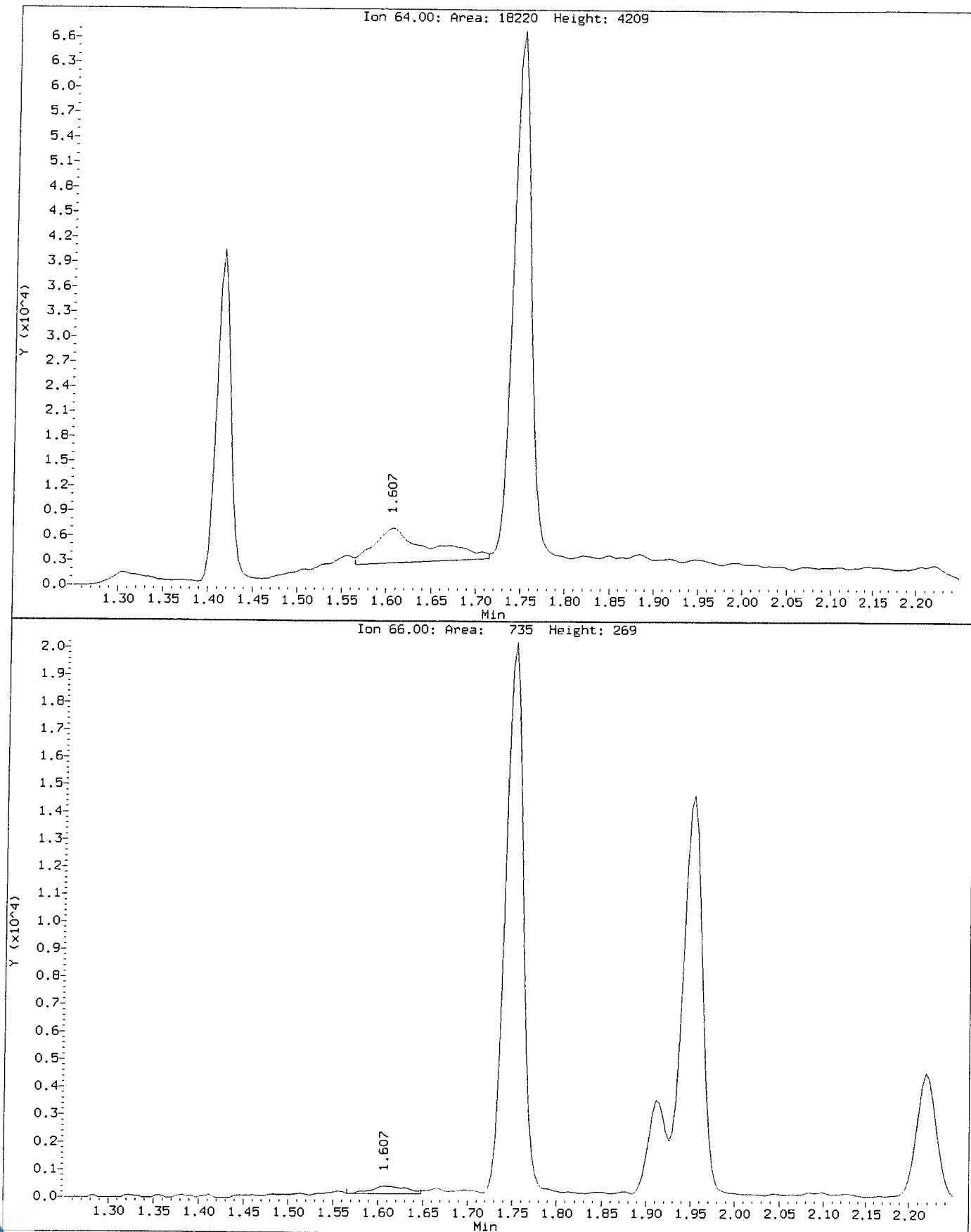
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Sample Info: VSTD020;VSTD020;1;6;
Purge Volume: 5.0
Column phase: DB624

Instrument: V099.i
Operator: PC
Column diameter: 0.18



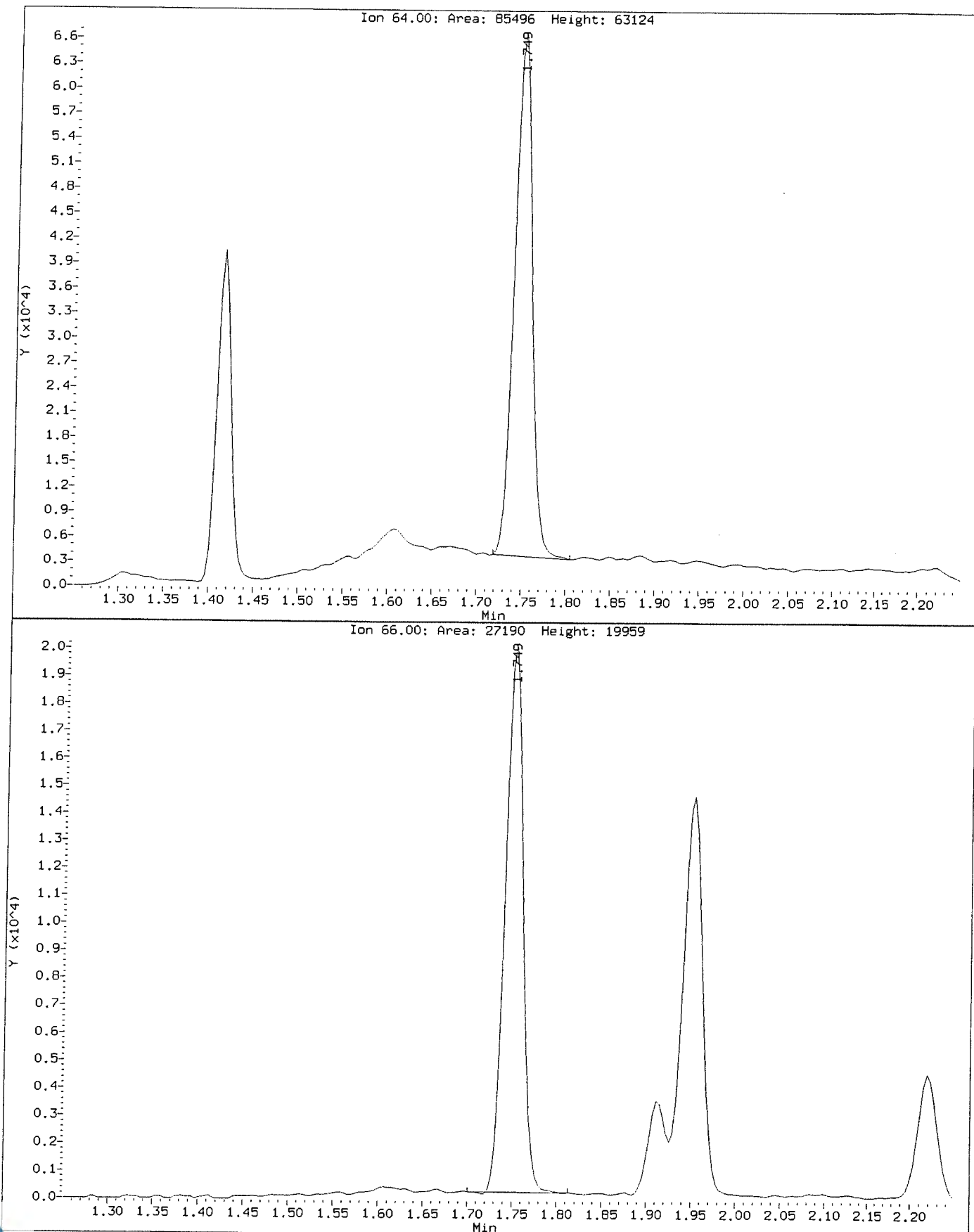
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Instrument: VOA9.i
Client Sample ID: VSTD020

Compound: Chloroethane
CAS Number: 75-00-3



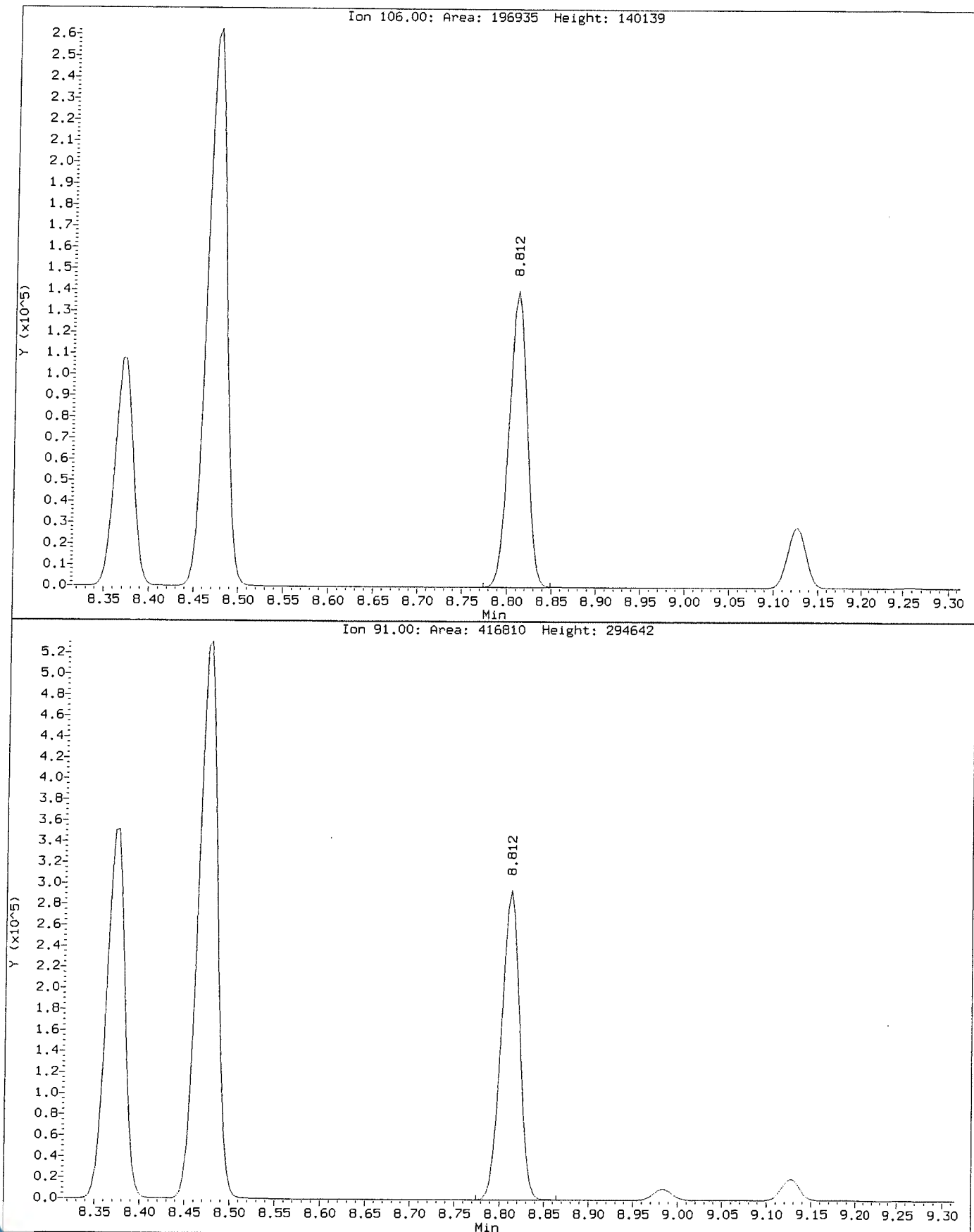
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Instrument: VOA9.1
Client Sample ID: VSTD020

Compound: Chloroethane
CAS Number: 75-00-3



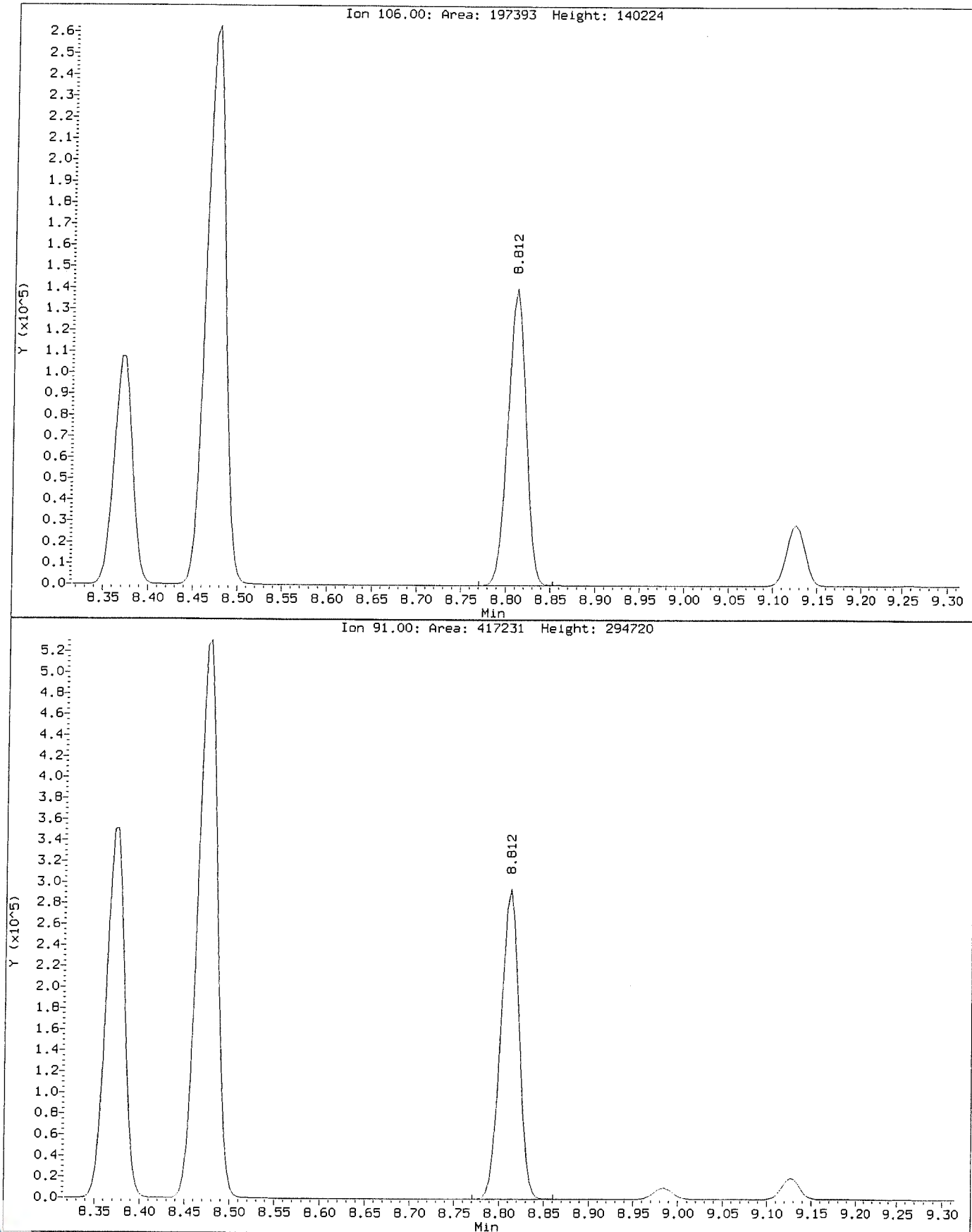
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Injection Date: 13-NOV-2018 13:50
Instrument: VOA9.i
Client Sample ID: VSTD020

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTW5005\Target\chem\voa9.1\U181113.b\U11307.D
Injection Date: 13-NOV-2018 13:50
Instrument: VOA9.1
Client Sample ID: VSTD020

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111308.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111308.D
 Lab Smp Id: VSTD050 Client Smp ID: VSTD050
 Inj Date : 13-NOV-2018 14:15
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD050;VSTD050;1;7;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
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 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 13:50 Cal File: U111307.D
 Als bottle: 9 Calibration Sample, Level: 7
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	AMOUNTS					
			CAL-AMT	ON-COL	RT	EXP RT	REL RT	RESPONSE
	MASS		(ug/l)	(ug/l)				
* 1 Pentafluorobenzene	168		50.0000		4.894	4.894	(1.000)	442076
* 36 1,4-Difluorobenzene	114		50.0000		5.625	5.625	(1.000)	849587
* 47 Chlorobenzene-d5	117		50.0000		8.249	8.249	(1.000)	790877
* 70 1,4-Dichlorobenzene-d4	152		50.0000		10.236	10.236	(1.000)	387587
\$ 30 Dibromofluoromethane	113		50.0000	50.35	4.830	4.830	(0.987)	255826
\$ 35 1,2-Dichloroethane-d4	65		50.0000	50.27	5.175	5.175	(1.057)	335901
\$ 48 Toluene-d8	98		50.0000	51.75	6.989	6.989	(0.847)	1045096
\$ 69 4-Bromofluorobenzene	95		50.0000	50.72	9.257	9.257	(1.122)	396604
60 1,1,1,2-Tetrachloroethane	131		50.0000	50.77	8.350	8.350	(1.012)	253286
31 1,1,1-Trichloroethane	97		50.0000	47.64	4.826	4.826	(0.986)	369959
68 1,1,2,2-Tetrachloroethane	83		50.0000	47.34	9.392	9.392	(0.918)	434682
138 Freon TF	101		50.0000	43.24	2.401	2.401	(0.491)	191531
53 1,1,2-Trichloroethane	83		50.0000	46.54	7.420	7.420	(0.900)	234031
22 1,1-Dichloroethane	63		50.0000	46.99	3.604	3.604	(0.737)	476032
11 1,1-Dichloroethene	96		50.0000	44.91	2.397	2.397	(0.490)	224107
32 1,1-Dichloropropene	75		50.0000	44.78	5.003	5.003	(0.889)	351415
93 1,2,3-Trichlorobenzene	180		50.0000	47.39	12.335	12.335	(1.205)	346843
71 1,2,3-Trichloropropane	75		50.0000	50.17	9.430	9.430	(0.921)	480640
90 1,2,4-Trichlorobenzene	180		50.0000	46.69	11.926	11.926	(1.165)	355891
79 1,2,4-Trimethylbenzene	105		50.0000	47.63	9.943	9.943	(0.971)	1025451
89 1,2-Dibromo-3-Chloropropane	155		50.0000	47.02	11.233	11.233	(1.097)	66581
57 1,2-Dibromoethane	107		50.0000	49.34	7.852	7.852	(0.952)	287318



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111308.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG				AMOUNTS		
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	569575	50.0000	46.01
33 1,2-Dichloroethane	62	5.254	5.254	(0.934)	409387	50.0000	46.04
42 1,2-Dichloropropane	63	6.082	6.082	(1.081)	288083	50.0000	47.62
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	989153	50.0000	47.49
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	558678	50.0000	45.78
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	507685	50.0000	47.02
84 1,4-Dichlorobenzene	146	10.258	10.258	(1.002)	576011	50.0000	48.25
26 2,2-Dichloropropane	77	4.275	4.275	(0.874)	295338	50.0000	47.07
24 2-Butanone	43	4.339	4.339	(0.887)	300842	100.000	96.81
76 2-Chlorotoluene	91	9.550	9.550	(0.933)	883069	50.0000	46.52
52 2-Hexanone	43	7.649	7.649	(0.927)	469848	100.000	102.19
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	1023060	50.0000	46.71
82 p-Isopropyltoluene	119	10.210	10.210	(0.997)	1018309	50.0000	47.97
45 4-Methyl-2-Pentanone	43	6.914	6.914	(0.838)	657858	100.000	99.71
10 Acetone	43	2.480	2.480	(0.507)	201617	100.000	99.32
37 Benzene	78	5.216	5.216	(0.927)	1096105	50.0000	46.67
74 Bromobenzene	156	9.385	9.385	(0.917)	293372	50.0000	46.14
29 Bromochloromethane	128	4.553	4.553	(0.930)	136269	50.0000	52.28
39 Bromodichloromethane	83	6.348	6.348	(1.129)	344674	50.0000	48.82
66 Bromoform	173	8.984	8.984	(1.089)	170886	50.0000	47.61(T)
6 Bromomethane	94	1.666	1.666	(0.341)	154362	50.0000	42.38
19 Carbon Disulfide	76	2.588	2.588	(0.529)	1468777	100.000	95.31
34 Carbon Tetrachloride	117	4.995	4.995	(0.888)	286167	50.0000	46.68
59 Chlorobenzene	112	8.275	8.275	(1.003)	750956	50.0000	46.00
7 Chloroethane	64	1.749	1.749	(0.357)	213028	50.0000	46.18(M)
28 Chloroform	83	4.658	4.658	(0.952)	478555	50.0000	47.01
3 Chloromethane	50	1.340	1.340	(0.274)	285521	50.0000	48.38
27 cis-1,2-Dichloroethene	96	4.287	4.287	(0.876)	293496	50.0000	46.85
46 cis-1,3-Dichloropropene	75	6.757	6.757	(1.201)	464679	50.0000	52.27
55 Dibromochloromethane	129	7.758	7.758	(0.940)	263139	50.0000	52.58
44 Dibromomethane	93	6.191	6.191	(1.101)	181664	50.0000	47.92
2 Dichlorodifluoromethane	85	1.205	1.205	(0.246)	293949	50.0000	45.08
61 Ethylbenzene	106	8.373	8.373	(1.015)	388724	50.0000	47.09
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	110558	50.0000	46.37
67 Isopropylbenzene	105	9.126	9.126	(1.106)	1128288	50.0000	46.94
62 m,p-Xylenes	106	8.474	8.474	(1.027)	962791	100.000	94.66
17 Methylene Chloride	84	2.870	2.870	(0.586)	276248	50.0000	48.96
87 n-Butylbenzene	91	10.558	10.558	(1.031)	954951	50.0000	47.12
73 n-Propylbenzene	91	9.475	9.475	(0.926)	1421540	50.0000	46.47
92 Naphthalene	128	12.133	12.133	(1.185)	1186850	50.0000	51.71
63 o-Xylene	106	8.811	8.811	(1.068)	497517	50.0000	48.87(H)
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	1186653	50.0000	46.59
64 Styrene	104	8.826	8.826	(1.070)	870320	50.0000	50.64
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	827990	50.0000	46.65
56 Tetrachloroethene	164	7.525	7.525	(0.912)	182495	50.0000	44.04
50 Toluene	91	7.049	7.049	(0.855)	1170933	50.0000	46.63
20 trans-1,2-Dichloroethene	96	3.139	3.139	(0.642)	255498	50.0000	46.71
51 trans-1,3-Dichloropropene	75	7.263	7.263	(1.291)	407060	50.0000	46.38
38 Trichloroethene	130	5.861	5.861	(1.042)	263822	50.0000	45.57
8 Trichlorofluoromethane	101	1.951	1.951	(0.399)	392628	50.0000	44.42
5 Vinyl Chloride	62	1.419	1.419	(0.290)	335638	50.0000	46.60



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111308.D Page 3
Report Date: 24-Jan-2019 18:55

QC Flag Legend

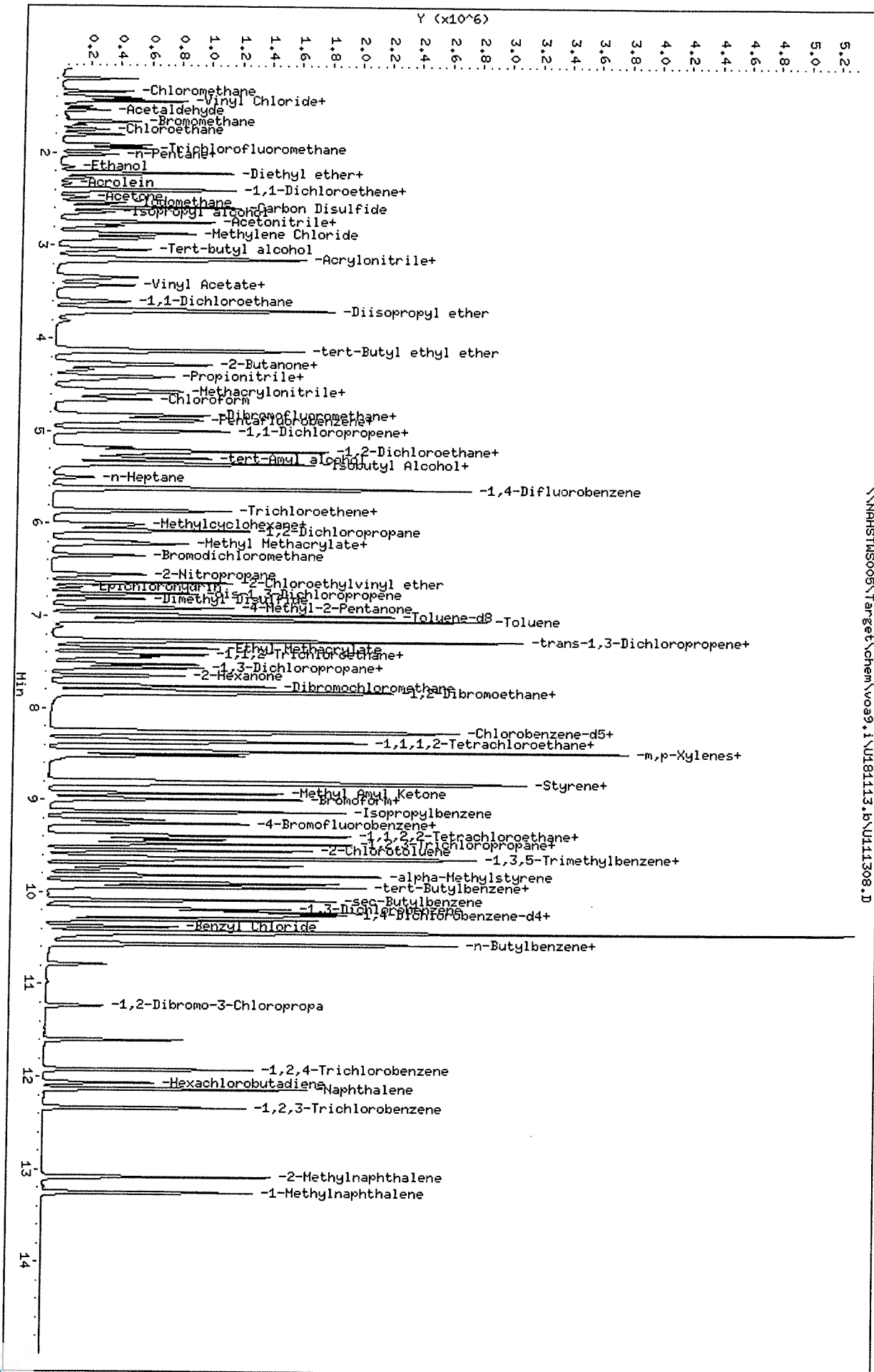
- T - Target compound detected outside RT window.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



Data File: \\NAHSTMS005\Target\chem\voa9.i\N18113.b\N11308.D
Date : 13-NOV-2018 14:15
Client ID: VSTD050
Sample Info: VSTD050;VSTD050;1;7;
Purge Volume: 5.0
Column phase: DB624

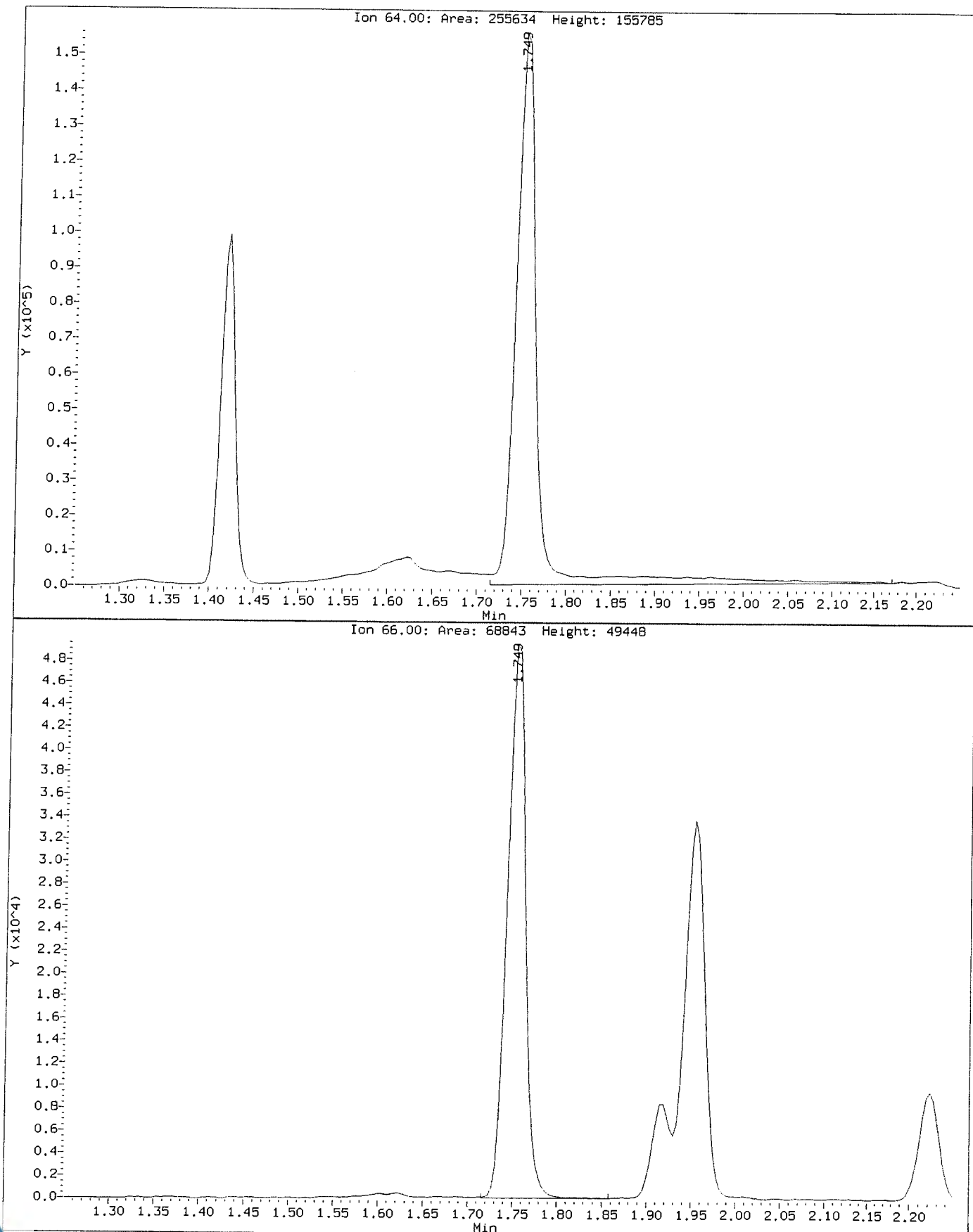
Instrument: W0A9.i
Operator: PC
Column diameter: 0.18

\\NAHSTMS005\Target\chem\voa9.i\N18113.b\N11308.D



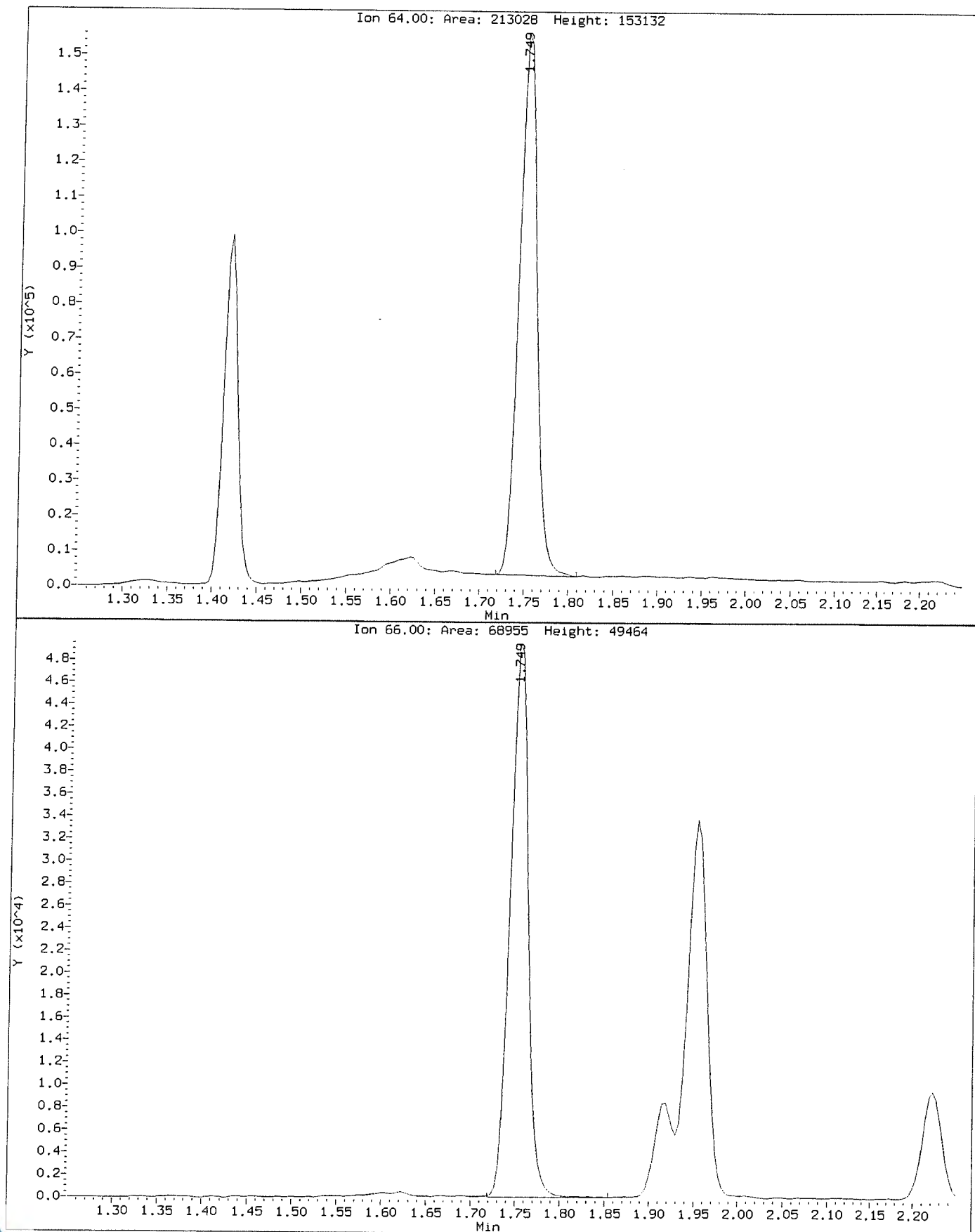
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Injection Date: 13-NOV-2018 14:15
Instrument: VDA9.i
Client Sample ID: VSTD050

Compound: Chloroethane
CAS Number: 75-00-3



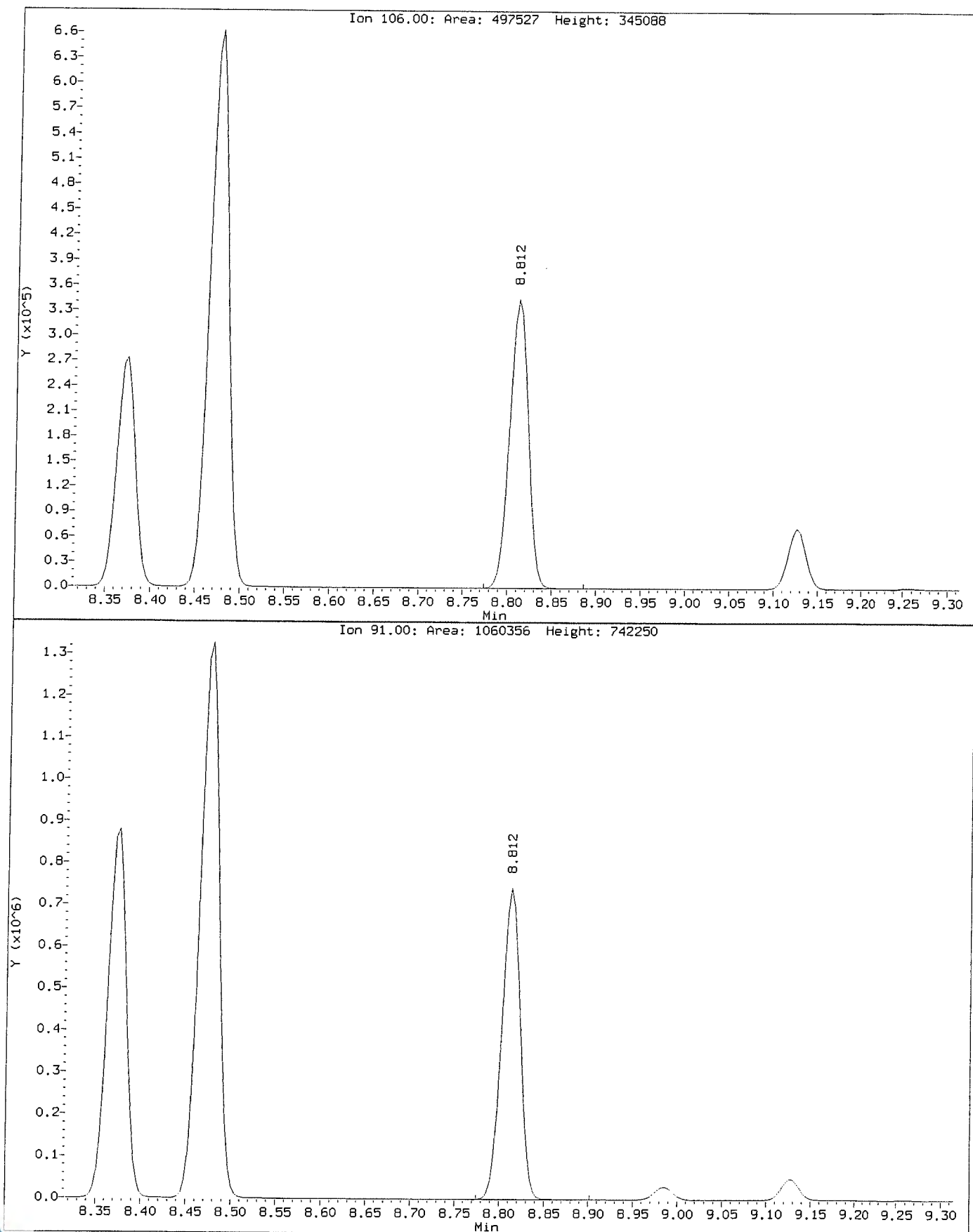
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Injection Date: 13-NOV-2018 14:15
Instrument: VOA9.i
Client Sample ID: VSTD050

Compound: Chloroethane
CAS Number: 75-00-3



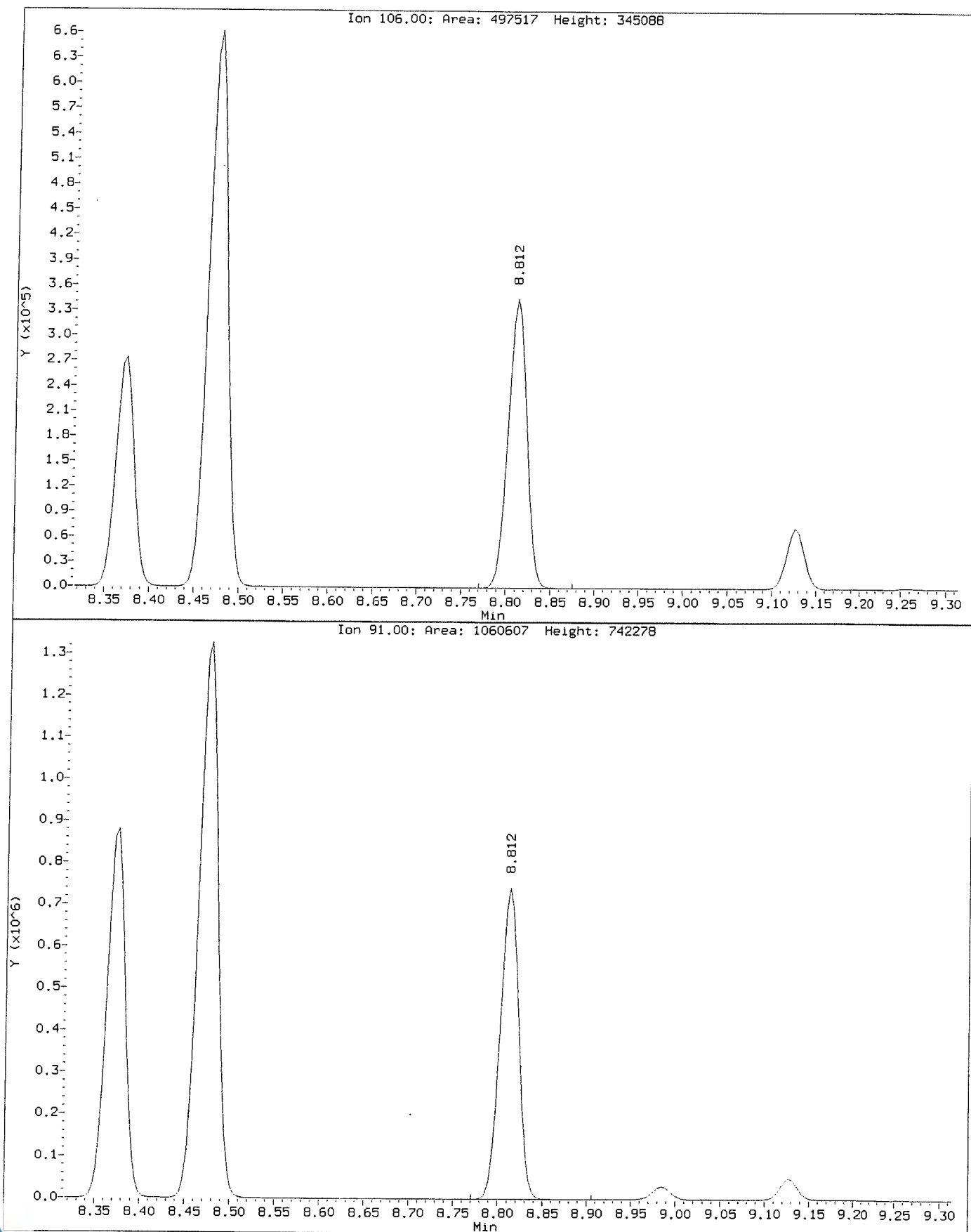
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Injection Date: 13-NOV-2018 14:15
Instrument: VOA9.i
Client Sample ID: VSTD050

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.1\U181113.b\U111308.D
Injection Date: 13-NOV-2018 14:15
Instrument: VOA9.1
Client Sample ID: VSTD050

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111309.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111309.D
 Lab Smp Id: VSTD100 Client Smp ID: VSTD100
 Inj Date : 13-NOV-2018 14:39
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD100;VSTD100;1;8;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 10 Calibration Sample, Level: 8
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG					AMOUNTS	
			MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)
* 1 Pentafluorobenzene	168	====	4.890	4.890	(1.000)	450542	50.0000	
* 36 1,4-Difluorobenzene	114		5.625	5.625	(1.000)	856582	50.0000	
* 47 Chlorobenzene-d5	117		8.249	8.249	(1.000)	821336	50.0000	
* 70 1,4-Dichlorobenzene-d4	152		10.236	10.236	(1.000)	399019	50.0000	
\$ 30 Dibromofluoromethane	113		4.827	4.827	(0.987)	526814	100.000	103.01
\$ 35 1,2-Dichloroethane-d4	65		5.171	5.171	(1.057)	696595	100.000	104.13
\$ 48 Toluene-d8	98		6.990	6.990	(0.847)	2142731	100.000	103.85
\$ 69 4-Bromofluorobenzene	95		9.258	9.258	(1.122)	823996	100.000	102.55
60 1,1,1,2-Tetrachloroethane	131		8.350	8.350	(1.012)	547424	100.000	105.66
31 1,1,1-Trichloroethane	97		4.823	4.823	(0.986)	826811	100.000	104.47
68 1,1,2,2-Tetrachloroethane	83		9.392	9.392	(0.918)	905540	100.000	95.80
138 Freon TF	101		2.394	2.394	(0.490)	456374	100.000	101.11
53 1,1,2-Trichloroethane	83		7.421	7.421	(0.900)	496464	100.000	95.07
22 1,1-Dichloroethane	63		3.597	3.597	(0.736)	991209	100.000	96.00
11 1,1-Dichloroethene	96		2.394	2.394	(0.490)	489401	100.000	96.23
32 1,1-Dichloropropene	75		4.999	4.999	(0.889)	782616	100.000	98.92
93 1,2,3-Trichlorobenzene	180		12.339	12.339	(1.205)	769767	100.000	102.18
71 1,2,3-Trichloropropane	75		9.430	9.430	(0.921)	1033869	100.000	104.82
90 1,2,4-Trichlorobenzene	180		11.927	11.927	(1.165)	795857	100.000	101.42
79 1,2,4-Trimethylbenzene	105		9.944	9.944	(0.971)	2240667	100.000	101.09
89 1,2-Dibromo-3-Chloropropane	155		11.237	11.237	(1.098)	149236	100.000	100.38
57 1,2-Dibromoethane	107		7.852	7.852	(0.952)	601967	100.000	99.55



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111309.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG		AMOUNTS				
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	1221135	100.000	95.83
33 1,2-Dichloroethane	62	5.250	5.250	(0.933)	845716	100.000	94.35
42 1,2-Dichloropropane	63	6.079	6.079	(1.081)	607568	100.000	99.62
75 1,3,5-Trimethylbenzene	105	9.629	9.629	(0.941)	2179527	100.000	101.66
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	1206946	100.000	96.07
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	1063077	100.000	94.81
84 1,4-Dichlorobenzene	146	10.258	10.258	(1.002)	1240019	100.000	100.95
26 2,2-Dichloropropane	77	4.268	4.268	(0.873)	666488	100.000	104.24
24 2-Butanone	43	4.332	4.332	(0.886)	626648	200.000	197.86
76 2-Chlorotoluene	91	9.550	9.550	(0.933)	1897282	100.000	97.10
52 2-Hexanone	43	7.649	7.649	(0.927)	1009723	200.000	211.48 (A)
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	2209785	100.000	98.01
82 p-Isopropyltoluene	119	10.213	10.213	(0.998)	2303162	100.000	105.40
45 4-Methyl-2-Pentanone	43	6.915	6.915	(0.838)	1416197	200.000	206.69 (A)
10 Acetone	43	2.476	2.476	(0.506)	430350	200.000	212.87 (A)
37 Benzene	78	5.216	5.216	(0.927)	2279526	100.000	96.27
74 Bromobenzene	156	9.385	9.385	(0.917)	640019	100.000	97.78
29 Bromochloromethane	128	4.549	4.549	(0.930)	269321	100.000	101.39
39 Bromodichloromethane	83	6.345	6.345	(1.128)	740758	100.000	104.07
66 Bromoform	173	8.984	8.984	(1.089)	383125	100.000	100.58
6 Bromomethane	94	1.655	1.655	(0.339)	367005	100.000	96.33
19 Carbon Disulfide	76	2.581	2.581	(0.528)	3169516	200.000	201.81 (A)
34 Carbon Tetrachloride	117	4.991	4.991	(0.887)	664499	100.000	107.51
59 Chlorobenzene	112	8.275	8.275	(1.003)	1606902	100.000	94.79
7 Chloroethane	64	1.741	1.741	(0.356)	444513	100.000	94.55 (M)
28 Chloroform	83	4.654	4.654	(0.952)	994934	100.000	95.91
3 Chloromethane	50	1.333	1.333	(0.273)	595590	100.000	98.79
27 cis-1,2-Dichloroethene	96	4.283	4.283	(0.876)	613605	100.000	96.11
46 cis-1,3-Dichloropropene	75	6.757	6.757	(1.201)	1019257	100.000	113.72
55 Dibromochloromethane	129	7.758	7.758	(0.940)	576245	100.000	110.87
44 Dibromomethane	93	6.187	6.187	(1.100)	378115	100.000	98.92
2 Dichlorodifluoromethane	85	1.198	1.198	(0.245)	680233	100.000	101.10
61 Ethylbenzene	106	8.373	8.373	(1.015)	844002	100.000	98.46
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	270039	100.000	102.92
67 Isopropylbenzene	105	9.126	9.126	(1.106)	2535018	100.000	101.55
62 m,p-Xylenes	106	8.474	8.474	(1.027)	2096282	200.000	198.47
17 Methylene Chloride	84	2.866	2.866	(0.586)	577262	100.000	101.12
87 n-Butylbenzene	91	10.558	10.558	(1.031)	2203969	100.000	102.68
73 n-Propylbenzene	91	9.479	9.479	(0.926)	3164922	100.000	100.51
92 Naphthalene	128	12.133	12.133	(1.185)	2576616	100.000	109.04
63 o-Xylene	106	8.811	8.811	(1.068)	1070218	100.000	101.23 (H)
81 sec-Butylbenzene	105	10.090	10.090	(0.986)	2704732	100.000	103.16
64 Styrene	104	8.826	8.826	(1.070)	1856392	100.000	104.02
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	1862977	100.000	101.96
56 Tetrachloroethene	164	7.526	7.526	(0.912)	415217	100.000	96.50
50 Toluene	91	7.046	7.046	(0.854)	2482746	100.000	95.21
20 trans-1,2-Dichloroethene	96	3.136	3.136	(0.641)	538852	100.000	96.67
51 trans-1,3-Dichloropropene	75	7.259	7.259	(1.291)	912116	100.000	101.29
38 Trichloroethene	130	5.861	5.861	(1.042)	566649	100.000	97.08
8 Trichlorofluoromethane	101	1.944	1.944	(0.398)	897271	100.000	99.61
5 Vinyl Chloride	62	1.411	1.411	(0.289)	744825	100.000	100.88



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111309.D Page 3
Report Date: 24-Jan-2019 18:55

QC Flag Legend

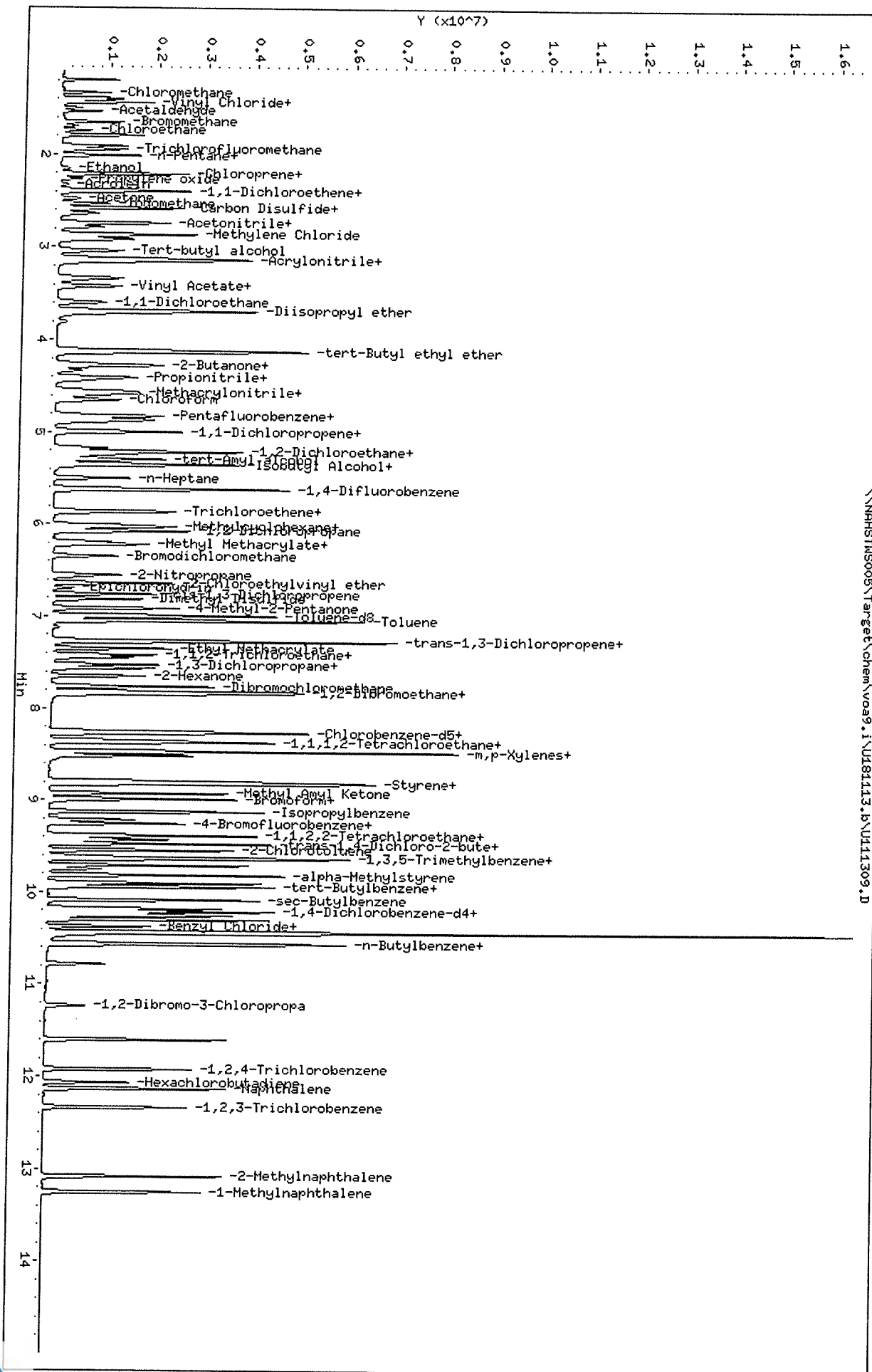
- A - Target compound detected but, quantitated amount exceeded maximum amount.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



Data File: \\NAHSTMS005\Target\chem\voa9.i\1181113.b\1111309.D
 Date : 13-NOV-2018 14:39
 Client ID: VSTD100
 Sample Info: VSTD100;VSTD100;1:8;
 Purge Volume: 5.0
 Column phase: DB624

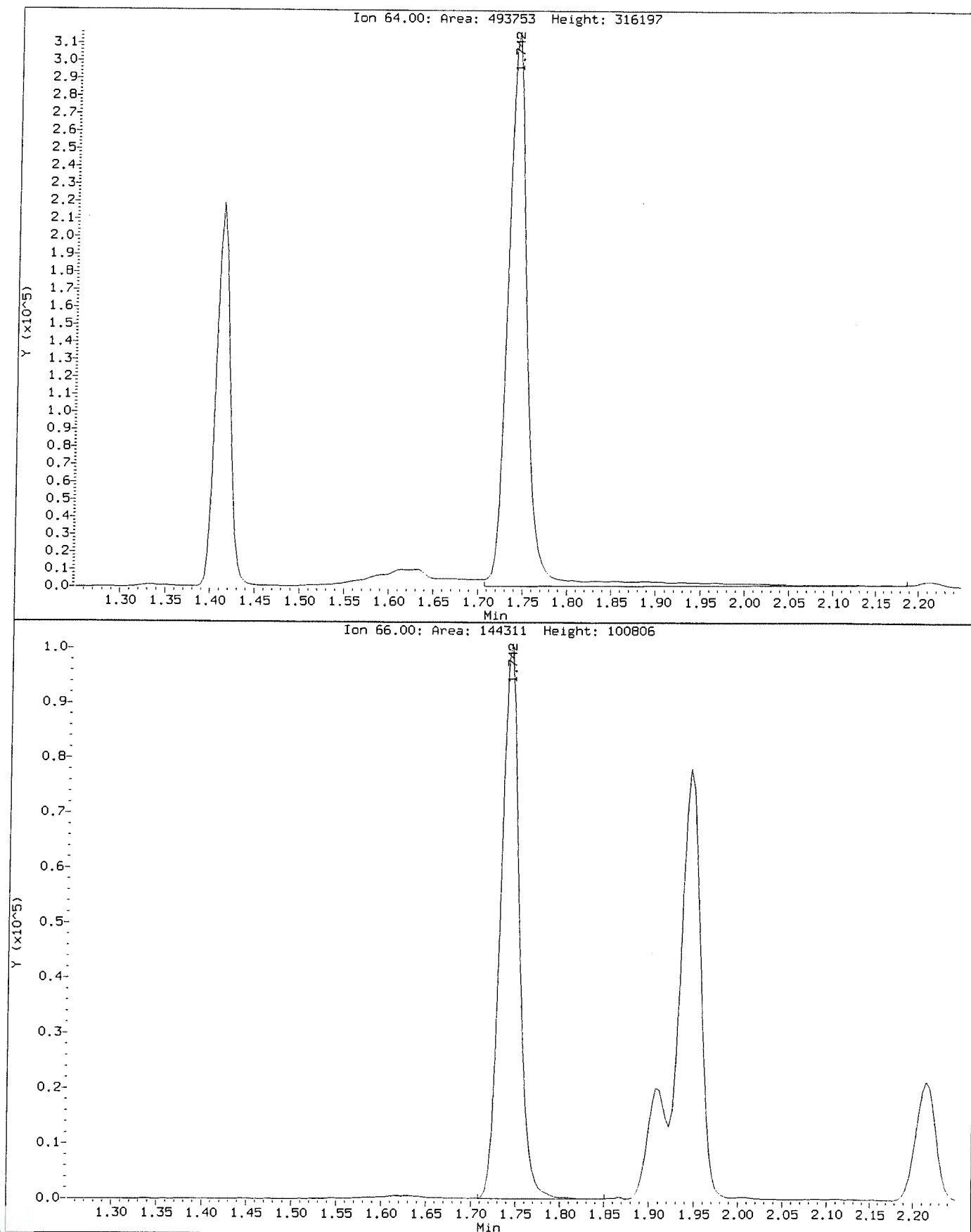
Instrument: VOA9.i
 Operator: PC
 Column diameter: 0.18

\\NAHSTMS005\Target\chem\voa9.i\1181113.b\1111309.D



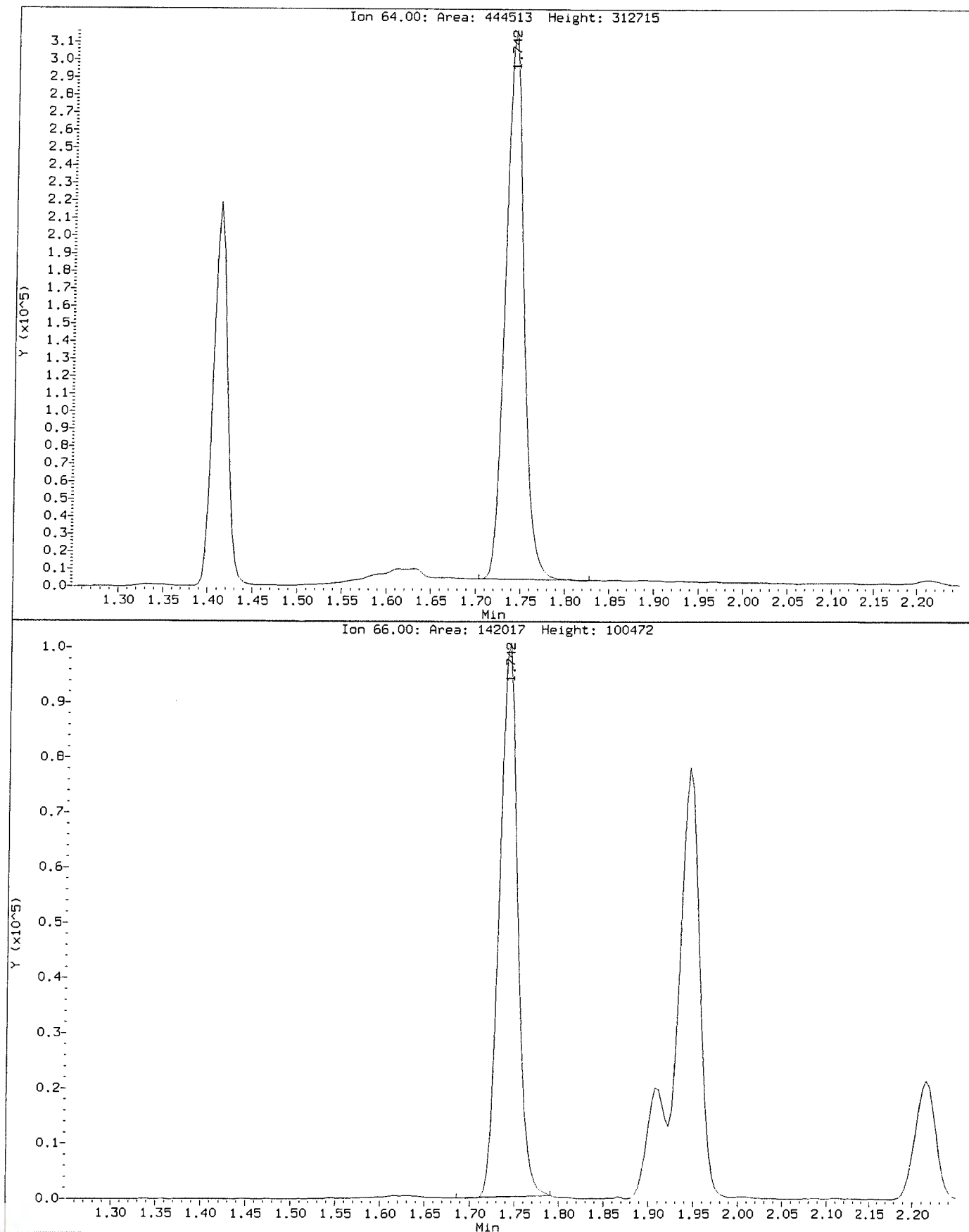
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Injection Date: 13-NOV-2018 14:39
Instrument: VOA9.i
Client Sample ID: VSTD100

Compound: Chloroethane
CAS Number: 75-00-3



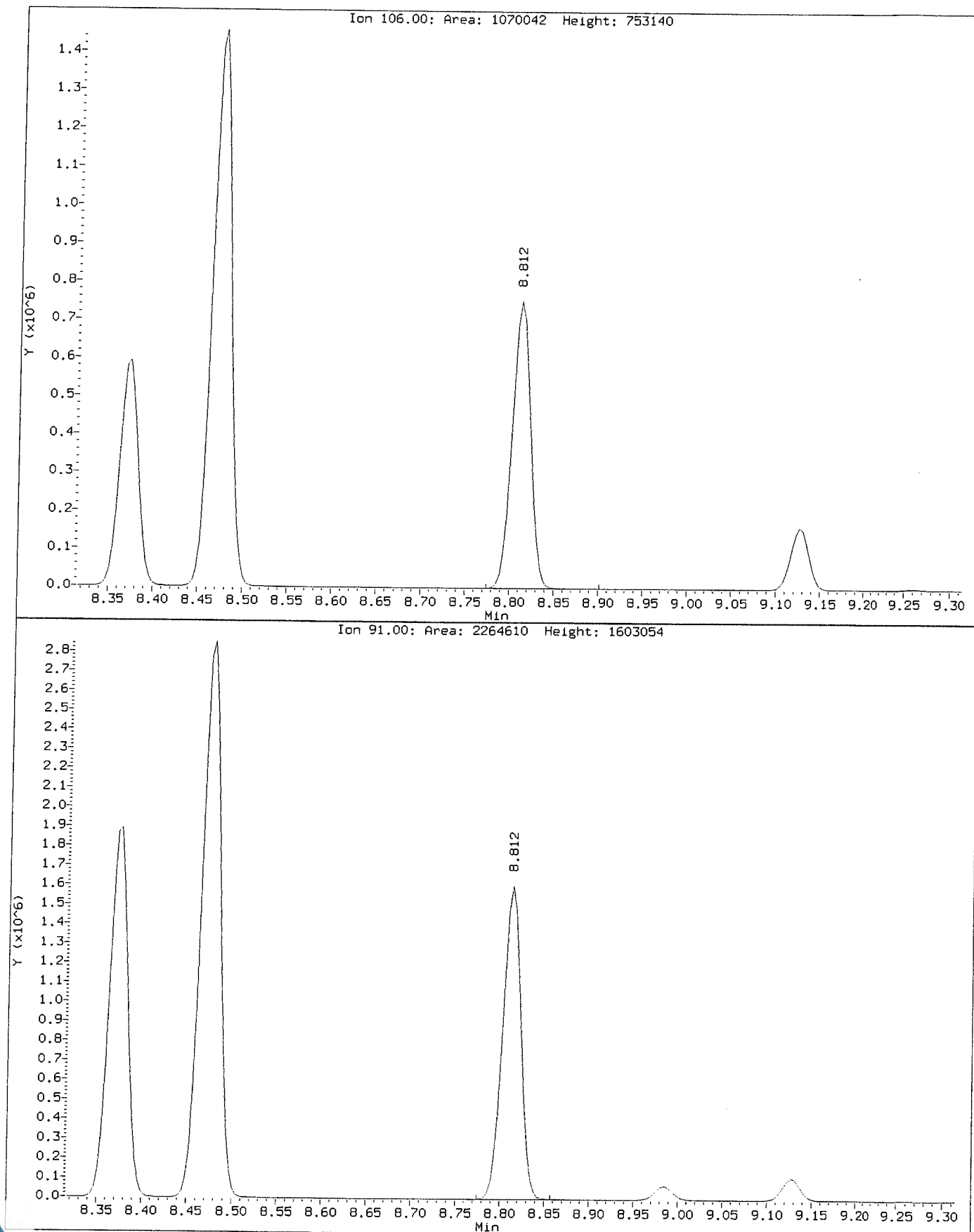
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Instrument: VOA9.1
Client Sample ID: VSTD100

Compound: Chloroethane
CAS Number: 75-00-3



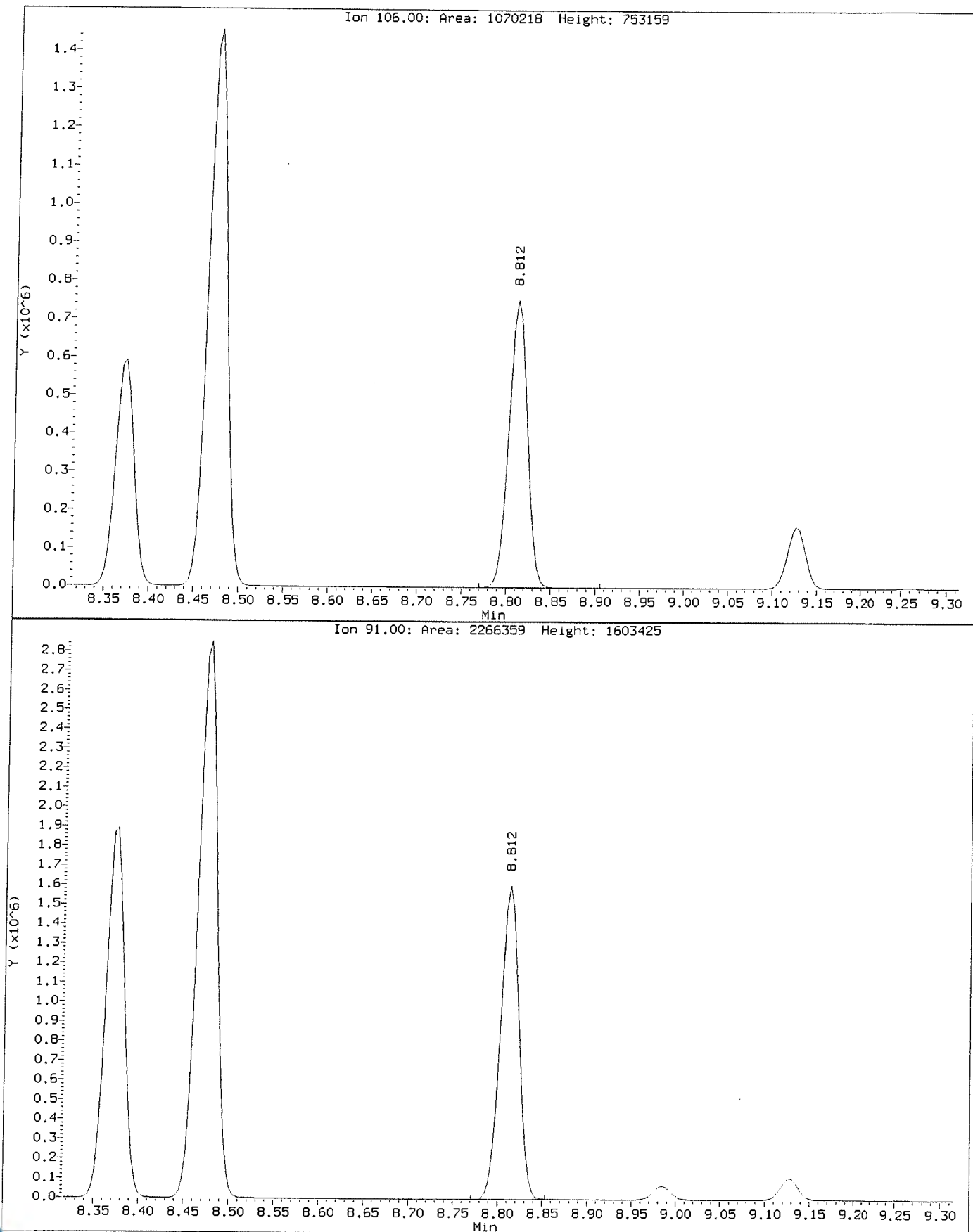
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Injection Date: 13-NOV-2018 14:39
Instrument: VOA9.i
Client Sample ID: VSTD100

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTW5005\Target\chem\voa9.1\U181113.b\U111309.D
Injection Date: 13-NOV-2018 14:39
Instrument: VOA9.i
Client Sample ID: VSTD100

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111310.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111310.D
 Lab Smp Id: VSTD150 Client Smp ID: VSTD150
 Inj Date : 13-NOV-2018 15:04
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD150;VSTD150;1;9;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:39 Cal File: U111309.D
 Als bottle: 11 Calibration Sample, Level: 9
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT	SIG	AMOUNTS				CAL-AMT (ug/l)	ON-COL (ug/l)
			MASS	RT	EXP RT	REL RT		
* 1 Pentafluorobenzene	168		4.894	4.894	(1.000)	463456	50.0000	
* 36 1,4-Difluorobenzene	114		5.625	5.625	(1.000)	890799	50.0000	
* 47 Chlorobenzene-d5	117		8.249	8.249	(1.000)	854800	50.0000	
* 70 1,4-Dichlorobenzene-d4	152		10.236	10.236	(1.000)	408174	50.0000	
\$ 30 Dibromofluoromethane	113		4.827	4.827	(0.986)	809097	150.000	154.42
\$ 35 1,2-Dichloroethane-d4	65		5.175	5.175	(1.057)	1062364	150.000	155.24
\$ 48 Toluene-d8	98		6.990	6.990	(0.847)	3238162	150.000	151.57
\$ 69 4-Bromofluorobenzene	95		9.258	9.258	(1.122)	1261578	150.000	151.36
60 1,1,1,2-Tetrachloroethane	131		8.350	8.350	(1.012)	858465	150.000	159.21
31 1,1,1-Trichloroethane	97		4.827	4.827	(0.986)	1329534	150.000	163.31
68 1,1,2,2-Tetrachloroethane	83		9.392	9.392	(0.918)	1394090	150.000	144.19
138 Freon TF	101		2.397	2.397	(0.490)	735007	150.000	158.31
53 1,1,2-Trichloroethane	83		7.421	7.421	(0.900)	767365	150.000	141.20
22 1,1-Dichloroethane	63		3.601	3.601	(0.736)	1566973	150.000	147.54
11 1,1-Dichloroethene	96		2.394	2.394	(0.489)	789917	150.000	150.99
32 1,1-Dichloropropene	75		5.003	5.003	(0.889)	1250083	150.000	151.94
93 1,2,3-Trichlorobenzene	180		12.335	12.335	(1.205)	1201294	150.000	155.88
71 1,2,3-Trichloropropane	75		9.430	9.430	(0.921)	1603390	150.000	158.92
90 1,2,4-Trichlorobenzene	180		11.927	11.927	(1.165)	1246599	150.000	155.30
79 1,2,4-Trimethylbenzene	105		9.944	9.944	(0.971)	3425922	150.000	151.11
89 1,2-Dibromo-3-Chloropropane	155		11.233	11.233	(1.097)	234820	150.000	152.68
57 1,2-Dibromoethane	107		7.852	7.852	(0.952)	923726	150.000	146.78



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111310.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG				AMOUNTS		
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	1872904	150.000	143.68
33 1,2-Dichloroethane	62	5.250	5.250	(0.933)	1302354	150.000	139.71
42 1,2-Dichloropropane	63	6.082	6.082	(1.081)	950163	150.000	149.81
75 1,3,5-Trimethylbenzene	105	9.629	9.629	(0.941)	3367895	150.000	153.56
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	1864119	150.000	145.05
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	1651259	150.000	141.50
84 1,4-Dichlorobenzene	146	10.258	10.258	(1.002)	1907496	150.000	151.83
26 2,2-Dichloropropane	77	4.272	4.272	(0.873)	1089497	150.000	165.65
24 2-Butanone	43	4.336	4.336	(0.886)	955107	300.000	293.17(A)
76 2-Chlorotoluene	91	9.550	9.550	(0.933)	2905161	150.000	145.35
52 2-Hexanone	43	7.653	7.653	(0.928)	1563288	300.000	314.60(A)
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	3405553	150.000	147.66
82 p-Isopropyltoluene	119	10.213	10.213	(0.998)	3559913	150.000	159.26
45 4-Methyl-2-Pentanone	43	6.915	6.915	(0.838)	2161270	300.000	303.08(A)
10 Acetone	43	2.480	2.480	(0.507)	636098	300.000	307.81(A)
37 Benzene	78	5.216	5.216	(0.927)	3563563	150.000	144.72
74 Bromobenzene	156	9.385	9.385	(0.917)	991935	150.000	148.14
29 Bromochloromethane	128	4.553	4.553	(0.930)	396590	150.000	145.15
39 Bromodichloromethane	83	6.349	6.349	(1.129)	1169536	150.000	158.01
66 Bromoform	173	8.984	8.984	(1.089)	610635	150.000	152.14
6 Bromomethane	94	1.655	1.655	(0.338)	609177	150.000	154.28
19 Carbon Disulfide	76	2.585	2.585	(0.528)	5076235	300.000	314.21(A)
34 Carbon Tetrachloride	117	4.995	4.995	(0.888)	1081499	150.000	168.27
59 Chlorobenzene	112	8.275	8.275	(1.003)	2477986	150.000	140.45
7 Chloroethane	64	1.741	1.741	(0.356)	708077	150.000	146.42(M)
28 Chloroform	83	4.654	4.654	(0.951)	1558200	150.000	146.02
3 Chloromethane	50	1.337	1.337	(0.273)	963160	150.000	155.17
27 cis-1,2-Dichloroethene	96	4.283	4.283	(0.875)	965405	150.000	147.00
46 cis-1,3-Dichloropropene	75	6.757	6.757	(1.201)	1623779	150.000	174.21
55 Dibromochloromethane	129	7.758	7.758	(0.940)	908238	150.000	167.91
44 Dibromomethane	93	6.191	6.191	(1.101)	587501	150.000	147.80
2 Dichlorodifluoromethane	85	1.202	1.202	(0.246)	1086406	150.000	156.42
61 Ethylbenzene	106	8.373	8.373	(1.015)	1325471	150.000	148.58
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	426795	150.000	149.03
67 Isopropylbenzene	105	9.126	9.126	(1.106)	3937489	150.000	151.57
62 m,p-Xylenes	106	8.474	8.474	(1.027)	3241074	300.000	294.85(A)
17 Methylene Chloride	84	2.866	2.866	(0.586)	911224	150.000	155.55
87 n-Butylbenzene	91	10.558	10.558	(1.031)	3347969	150.000	149.04
73 n-Propylbenzene	91	9.479	9.479	(0.926)	4851775	150.000	150.63
92 Naphthalene	128	12.133	12.133	(1.185)	3933835	150.000	162.75
63 o-Xylene	106	8.811	8.811	(1.068)	1663910	150.000	151.23(H)
81 sec-Butylbenzene	105	10.090	10.090	(0.986)	4188144	150.000	156.16
64 Styrene	104	8.826	8.826	(1.070)	2865534	150.000	154.28
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	2893337	150.000	154.80
56 Tetrachloroethene	164	7.526	7.526	(0.912)	658457	150.000	147.04
50 Toluene	91	7.050	7.050	(0.855)	3865080	150.000	142.41
20 trans-1,2-Dichloroethene	96	3.140	3.140	(0.642)	855722	150.000	149.25
51 trans-1,3-Dichloropropene	75	7.259	7.259	(1.291)	1439550	150.000	152.95
38 Trichloroethene	130	5.861	5.861	(1.042)	898620	150.000	148.04
8 Trichlorofluoromethane	101	1.948	1.948	(0.398)	1438631	150.000	155.26
5 Vinyl Chloride	62	1.415	1.415	(0.289)	1189386	150.000	156.33



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111310.D Page 3
Report Date: 24-Jan-2019 18:55

QC Flag Legend

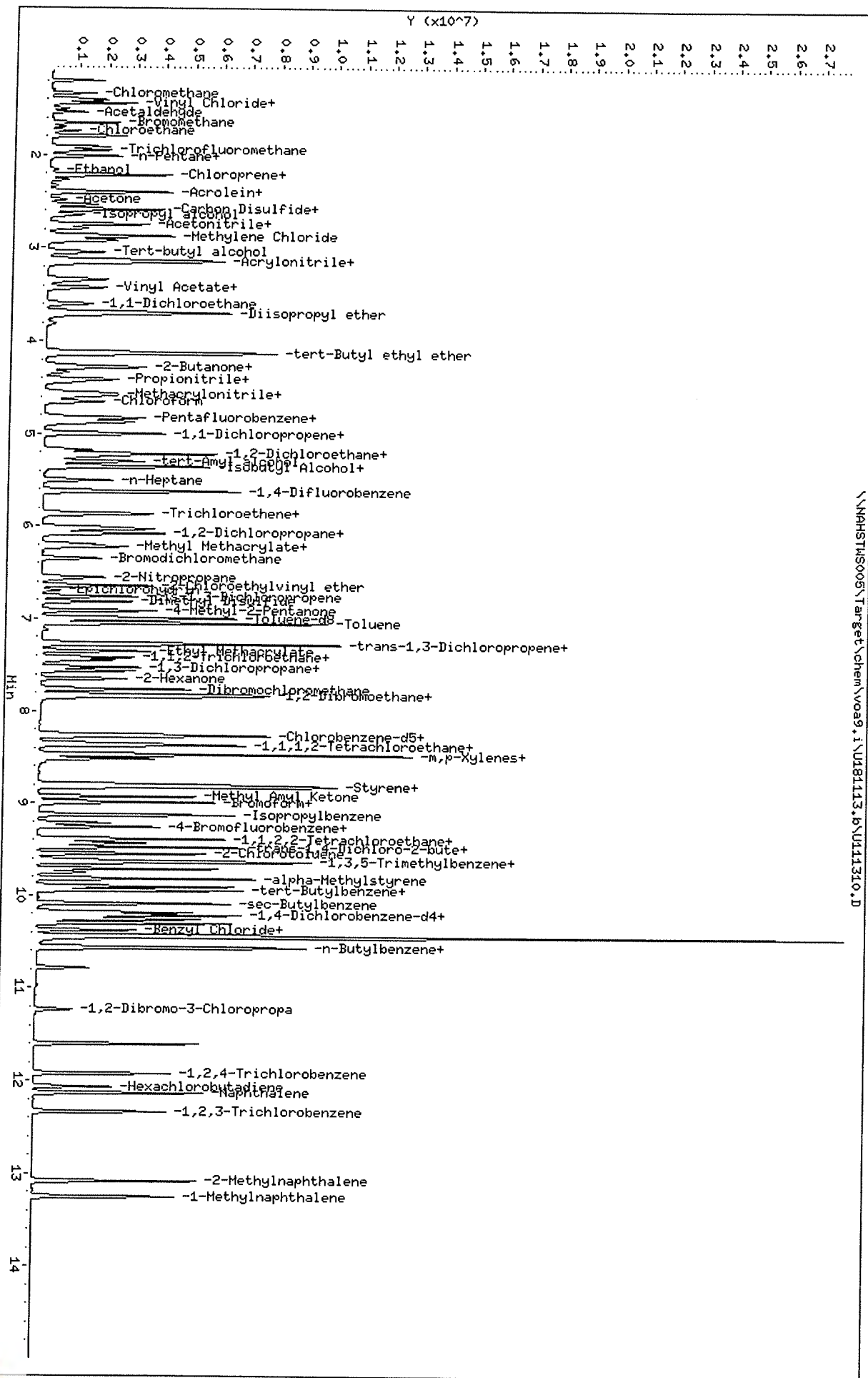
- A - Target compound detected but, quantitated amount exceeded maximum amount.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



Data File: \\NAHSTMS005\Target\chem\voa9.1\U181113.B\U111310.D
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Sample Info: VSTD150;VSTD150;119;
Purge Volume: 5.0
Column phase: DB624

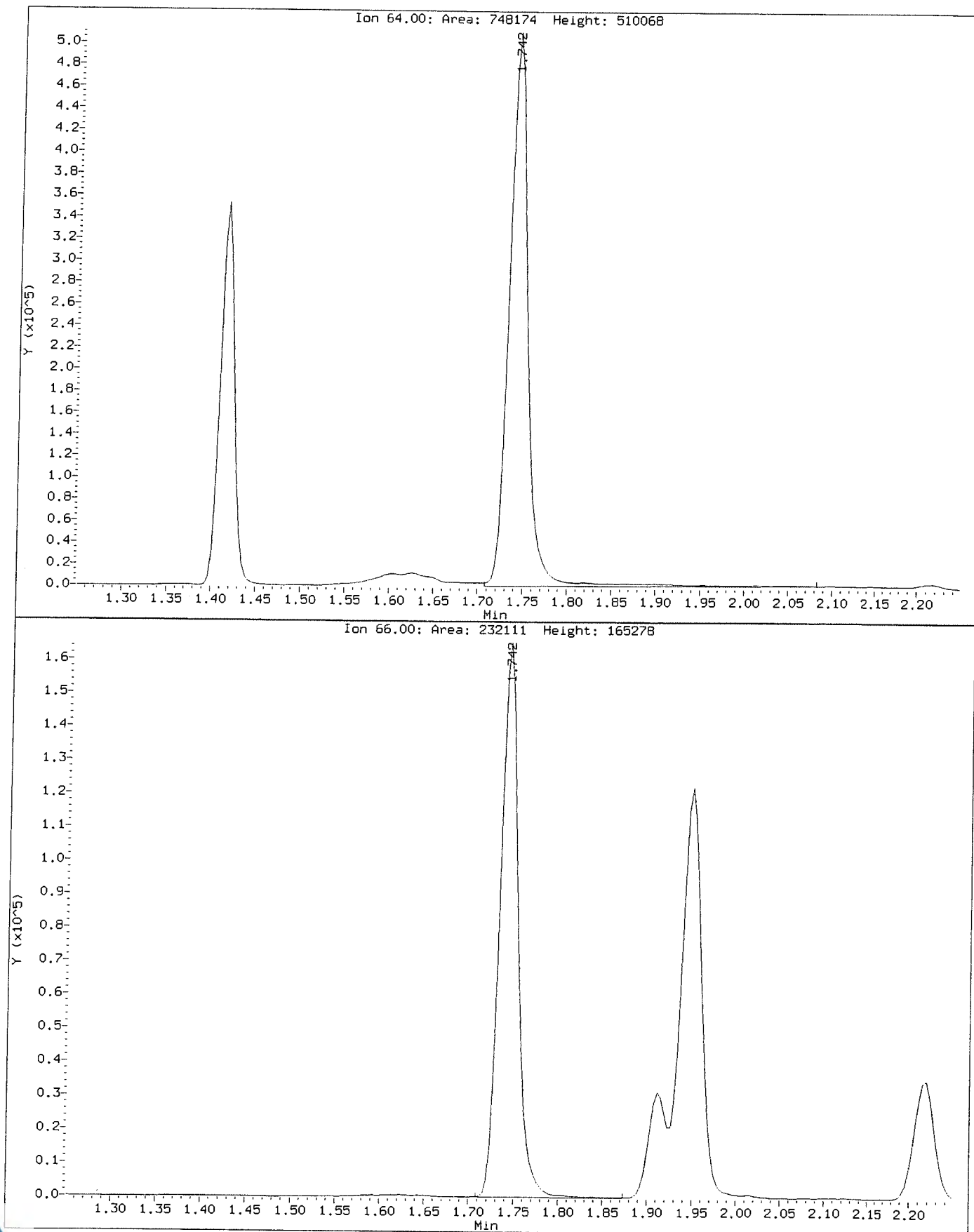
Instrument: V099.1
Operator: PC
Column diameter: 0.18

\\NAHSTMS005\Target\chem\voa9.1\U181113.B\U111310.D



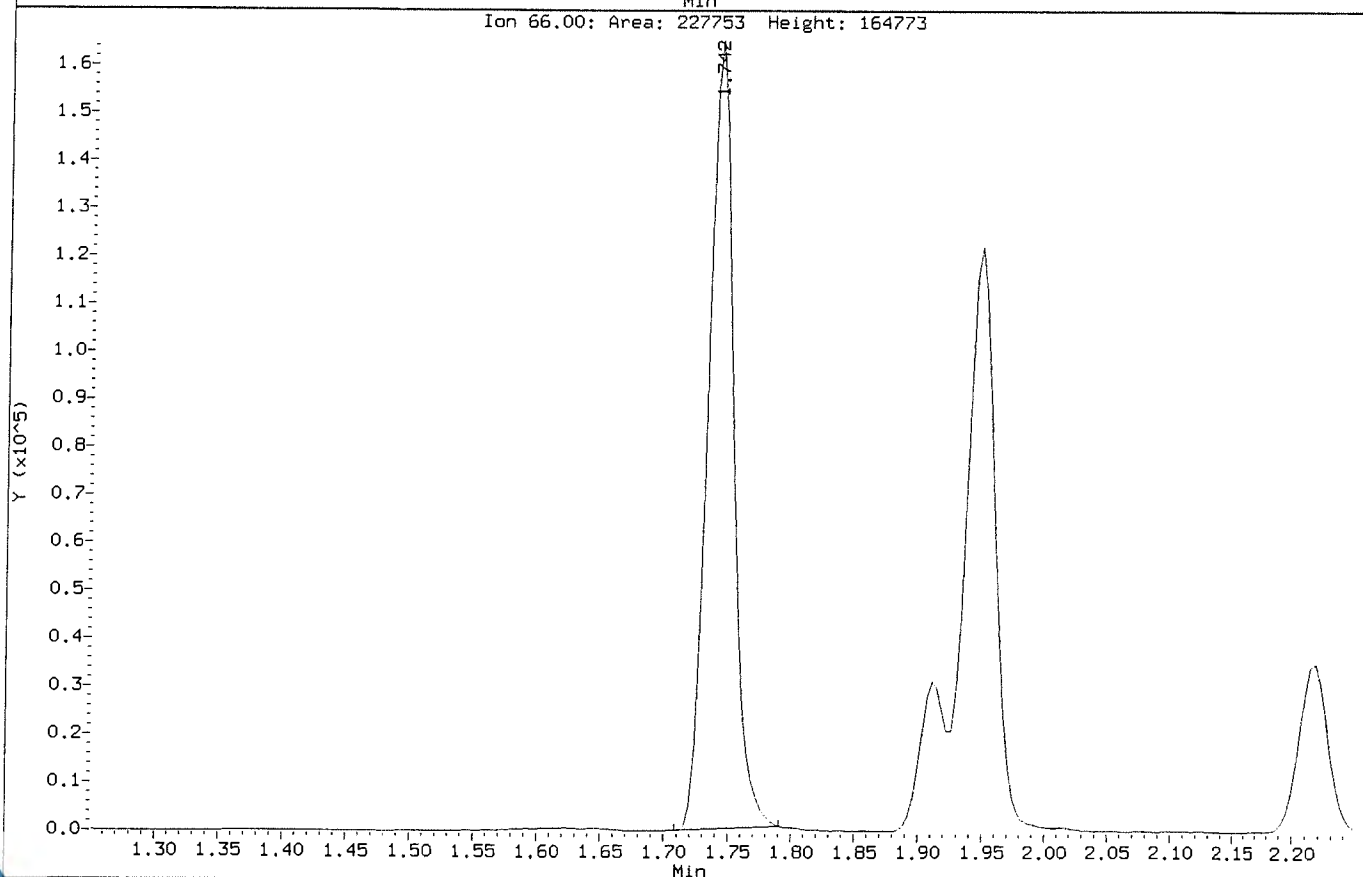
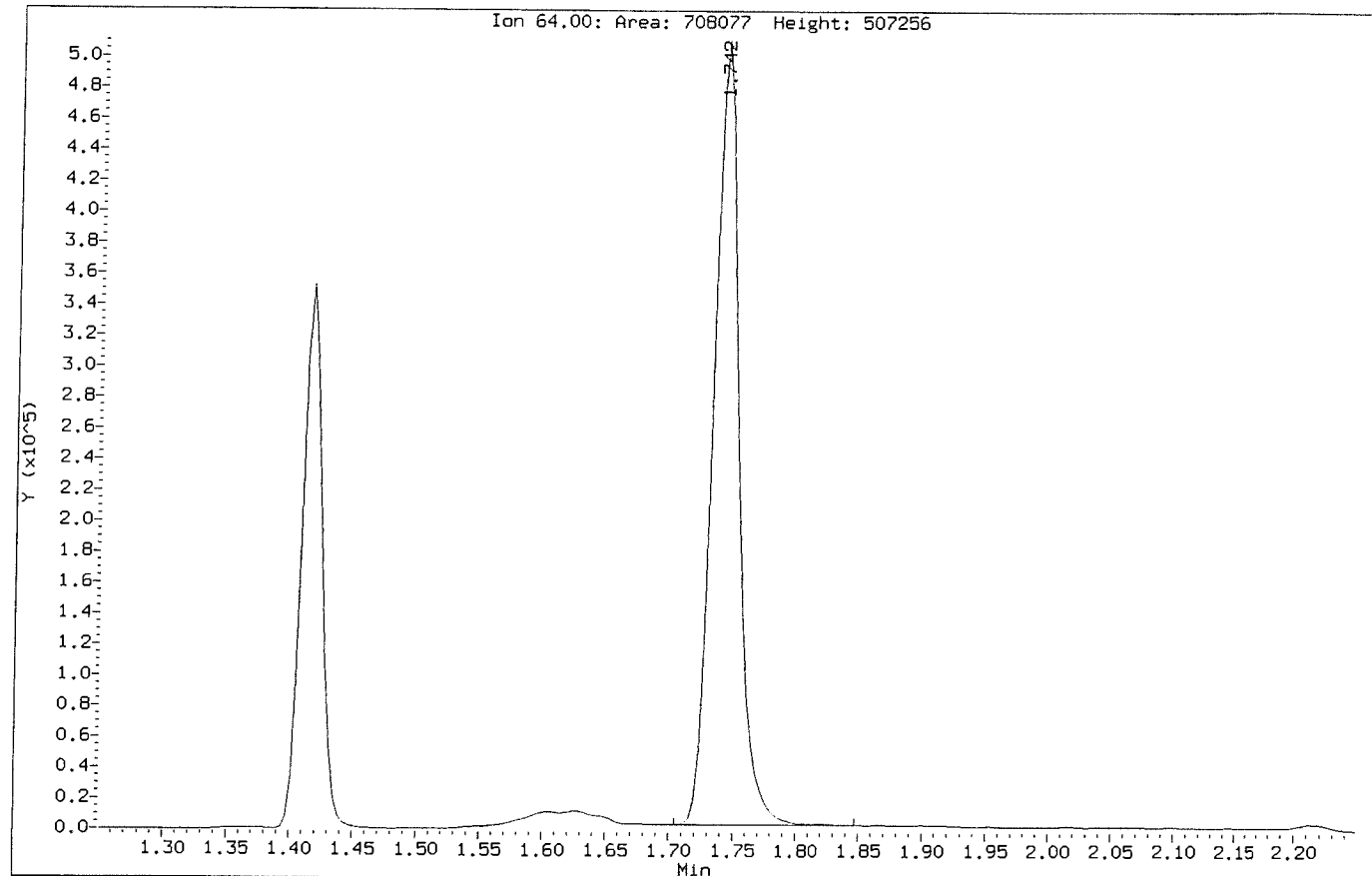
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Instrument: VOA9.1
Client Sample ID: VSTD150

Compound: Chloroethane
CAS Number: 75-00-3



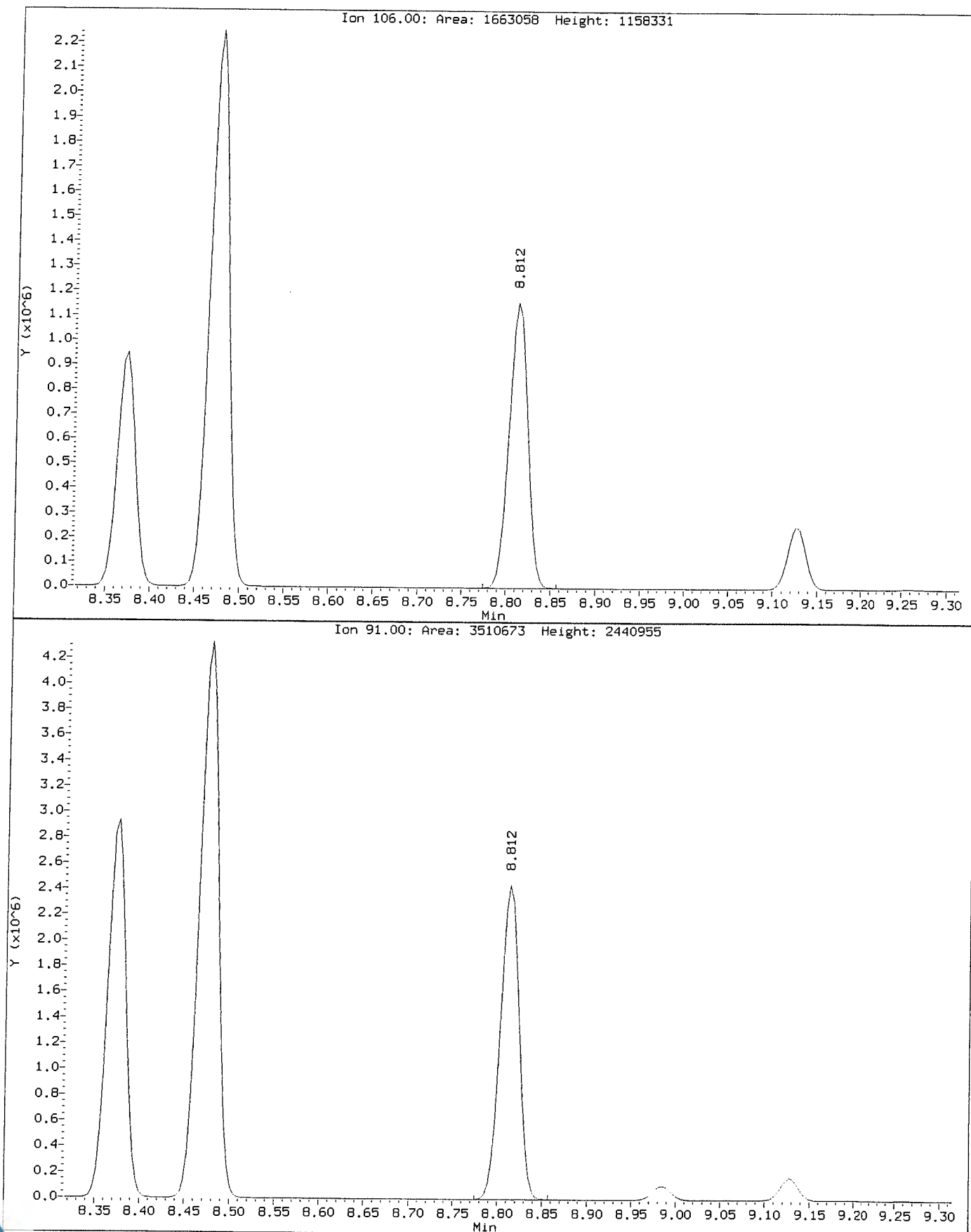
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Instrument: VOA9.i
Client Sample ID: VSTD150

Compound: Chloroethane
CAS Number: 75-00-3



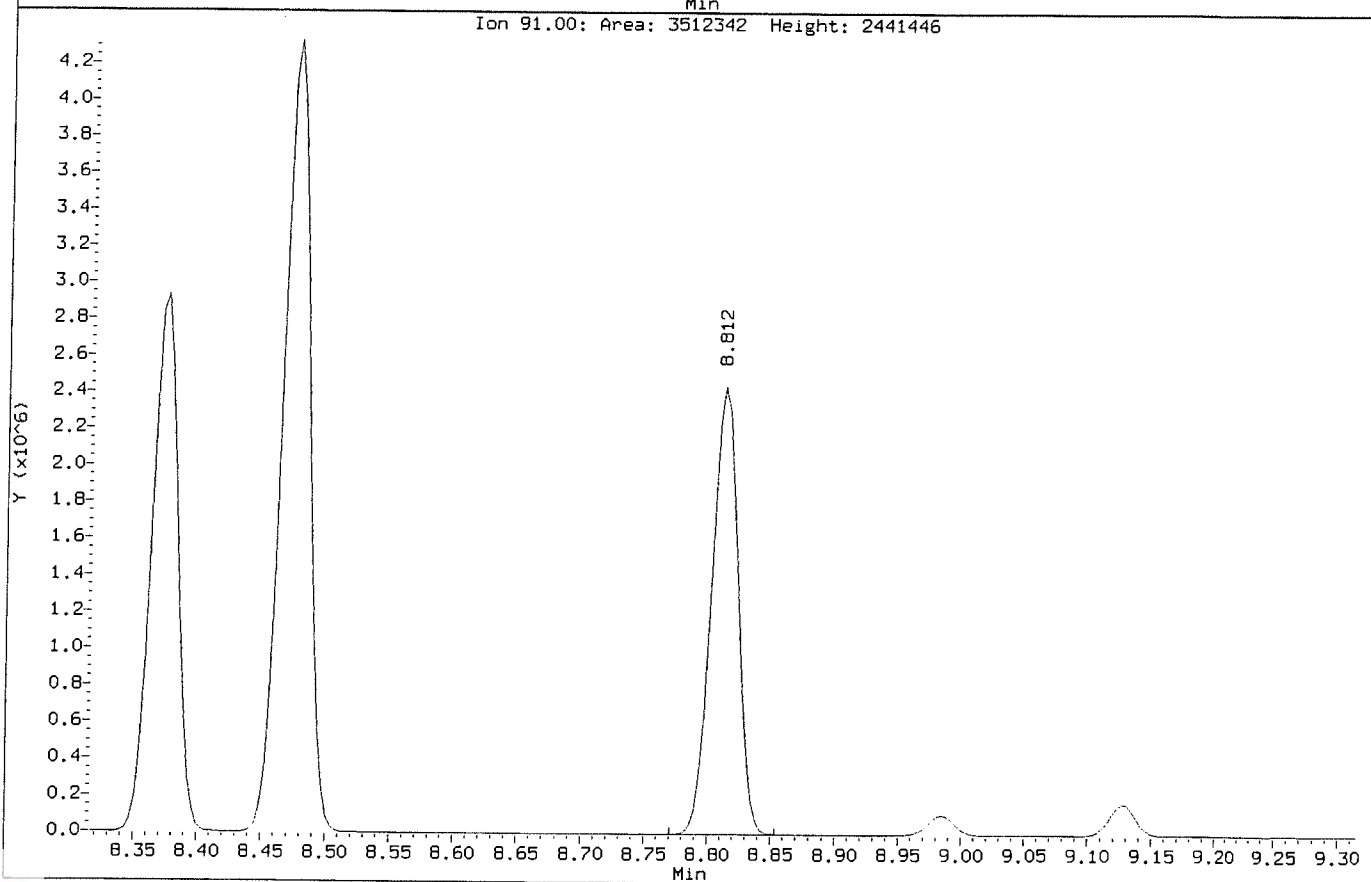
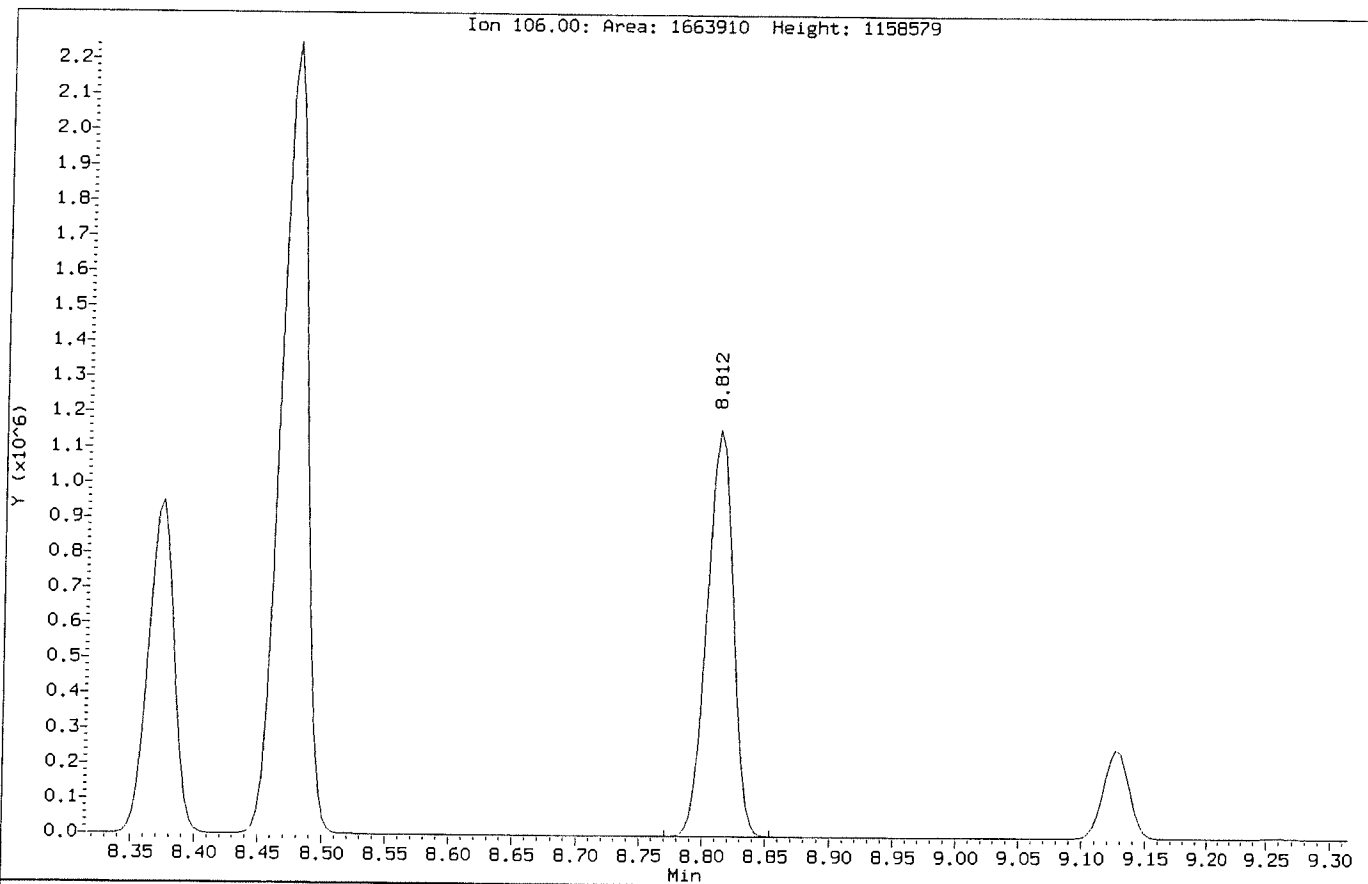
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Injection Date: 13-NOV-2018 15:04
Instrument: VOA9.i
Client Sample ID: VSTD150

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTW005\Target\chem\voa9.i\U181113.b\U111310.D
Injection Date: 13-NOV-2018 15:04
Instrument: VOA9.i
Client Sample ID: VSTD150

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111311.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111311.D
 Lab Smp Id: VSTD200 Client Smp ID: VSTD200
 Inj Date : 13-NOV-2018 15:28
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD200;VSTD200;1;10;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 15:04 Cal File: U111310.D
 Als bottle: 12 Calibration Sample, Level: 10
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG				AMOUNTS		
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.898	(1.000)	505644	50.0000	
* 36 1,4-Difluorobenzene	114	5.629	5.629	(1.000)	946062	50.0000	
* 47 Chlorobenzene-d5	117	8.253	8.253	(1.000)	896349	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.239	10.239	(1.000)	426550	50.0000	
\$ 30 Dibromofluoromethane	113	4.834	4.834	(0.987)	1112100	200.000	194.87
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.179	(1.057)	1441844	200.000	193.54
\$ 48 Toluene-d8	98	6.993	6.993	(0.847)	4386438	200.000	196.29
\$ 69 4-Bromofluorobenzene	95	9.261	9.261	(1.122)	1723103	200.000	197.48
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	1191494	200.000	210.73(A)
31 1,1,1-Trichloroethane	97	4.830	4.830	(0.986)	1824622	200.000	205.42(A)
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.917)	1884161	200.000	186.48
138 Freon TF	101	2.405	2.405	(0.491)	1007022	200.000	198.80
53 1,1,2-Trichloroethane	83	7.424	7.424	(0.900)	1043779	200.000	183.16
22 1,1-Dichloroethane	63	3.608	3.608	(0.737)	2138732	200.000	184.57
11 1,1-Dichloroethane	96	2.405	2.405	(0.491)	1081090	200.000	189.41
32 1,1-Dichloropropene	75	5.010	5.010	(0.890)	1707700	200.000	195.44
93 1,2,3-Trichlorobenzene	180	12.339	12.339	(1.205)	1636643	200.000	203.23(A)
71 1,2,3-Trichloropropane	75	9.434	9.434	(0.921)	2176006	200.000	206.38(A)
90 1,2,4-Trichlorobenzene	180	11.926	11.926	(1.165)	1712716	200.000	204.18(A)
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	4631542	200.000	195.48
89 1,2-Dibromo-3-Chloropropane	155	11.237	11.237	(1.097)	321770	200.000	198.50
57 1,2-Dibromoethane	107	7.855	7.855	(0.952)	1265930	200.000	191.84



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111311.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT SIG					AMOUNTS	
	MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.573	(1.033)	2535364	200.000	186.12
33 1,2-Dichloroethane	62	5.257	5.257	(0.934)	1770998	200.000	178.89
42 1,2-Dichloropropane	63	6.086	6.086	(1.081)	1294674	200.000	192.20
75 1,3,5-Trimethylbenzene	105	9.628	9.628	(0.940)	4535917	200.000	197.91
83 1,3-Dichlorobenzene	146	10.183	10.183	(0.995)	2548608	200.000	189.77
54 1,3-Dichloropropane	76	7.567	7.567	(0.917)	2235193	200.000	182.67
84 1,4-Dichlorobenzene	146	10.258	10.258	(1.002)	2606334	200.000	198.53
26 2,2-Dichloropropane	77	4.279	4.279	(0.874)	1492829	200.000	208.03(A)
24 2-Butanone	43	4.343	4.343	(0.887)	1287268	400.000	362.16(A)
76 2-Chlorotoluene	91	9.550	9.550	(0.933)	3934986	200.000	188.39
52 2-Hexanone	43	7.653	7.653	(0.927)	2091584	400.000	401.41(A)
77 4-Chlorotoluene	91	9.643	9.643	(0.942)	4615489	200.000	191.50
82 p-Isopropyltoluene	119	10.213	10.213	(0.997)	4810748	200.000	205.94(A)
45 4-Methyl-2-Pentanone	43	6.918	6.918	(0.838)	2944165	400.000	393.73(A)
10 Acetone	43	2.491	2.491	(0.509)	869596	400.000	386.81(A)
37 Benzene	78	5.224	5.224	(0.928)	4810692	200.000	183.96(M)
74 Bromobenzene	156	9.385	9.385	(0.917)	1358088	200.000	194.09
29 Bromochloromethane	128	4.560	4.560	(0.931)	537789	200.000	180.41
39 Bromodichloromethane	83	6.352	6.352	(1.129)	1603888	200.000	204.03(A)
66 Bromoform	173	8.987	8.987	(1.089)	844522	200.000	198.75
6 Bromomethane	94	1.662	1.662	(0.340)	794649	200.000	184.09
19 Carbon Disulfide	76	2.596	2.596	(0.530)	6896612	400.000	391.27(A)
34 Carbon Tetrachloride	117	4.999	4.999	(0.888)	1483776	200.000	217.37(A)
59 Chlorobenzene	112	8.279	8.279	(1.003)	3391468	200.000	183.32
7 Chloroethane	64	1.749	1.749	(0.357)	955282	200.000	181.06(M)
28 Chloroform	83	4.661	4.661	(0.952)	2117243	200.000	181.86
3 Chloromethane	50	1.344	1.344	(0.274)	1334230	200.000	196.96
27 cis-1,2-Dichloroethene	96	4.290	4.290	(0.876)	1322041	200.000	184.51
46 cis-1,3-Dichloropropene	75	6.761	6.761	(1.201)	2224591	200.000	224.72(A)
55 Dibromochloromethane	129	7.762	7.762	(0.940)	1258930	200.000	221.96(A)
44 Dibromomethane	93	6.195	6.195	(1.101)	803081	200.000	190.24
2 Dichlorodifluoromethane	85	1.213	1.213	(0.248)	1483355	200.000	195.50
61 Ethylbenzene	106	8.373	8.373	(1.015)	1806514	200.000	193.12(M)
91 Hexachlorobutadiene	225	12.065	12.065	(1.178)	582613	200.000	183.44
67 Isopropylbenzene	105	9.130	9.130	(1.106)	5274717	200.000	193.63
62 m,p-Xylenes	106	8.474	8.474	(1.027)	4413064	400.000	382.86(AM)
17 Methylene Chloride	84	2.877	2.877	(0.587)	1246871	200.000	195.27
87 n-Butylbenzene	91	10.558	10.558	(1.031)	4531778	200.000	189.12
73 n-Propylbenzene	91	9.478	9.478	(0.926)	6433909	200.000	191.14
92 Naphthalene	128	12.136	12.136	(1.185)	5280797	200.000	209.07(A)
63 o-Xylene	106	8.815	8.815	(1.068)	2277275	200.000	197.39(MH)
81 sec-Butylbenzene	105	10.090	10.090	(0.985)	5603961	200.000	199.95
64 Styrene	104	8.826	8.826	(1.069)	3881020	200.000	199.27
78 tert-Butylbenzene	119	9.906	9.906	(0.967)	3887715	200.000	199.04
56 Tetrachloroethene	164	7.525	7.525	(0.912)	907758	200.000	193.32
50 Toluene	91	7.049	7.049	(0.854)	5182031	200.000	182.09
20 trans-1,2-Dichloroethene	96	3.147	3.147	(0.643)	1173691	200.000	187.63
51 trans-1,3-Dichloropropene	75	7.263	7.263	(1.290)	1983766	200.000	198.03
38 Trichloroethene	130	5.865	5.865	(1.042)	1238549	200.000	192.12
8 Trichlorofluoromethane	101	1.955	1.955	(0.399)	1950000	200.000	192.89
5 Vinyl Chloride	62	1.426	1.426	(0.291)	1622522	200.000	195.34



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Report Date: 24-Jan-2019 18:55

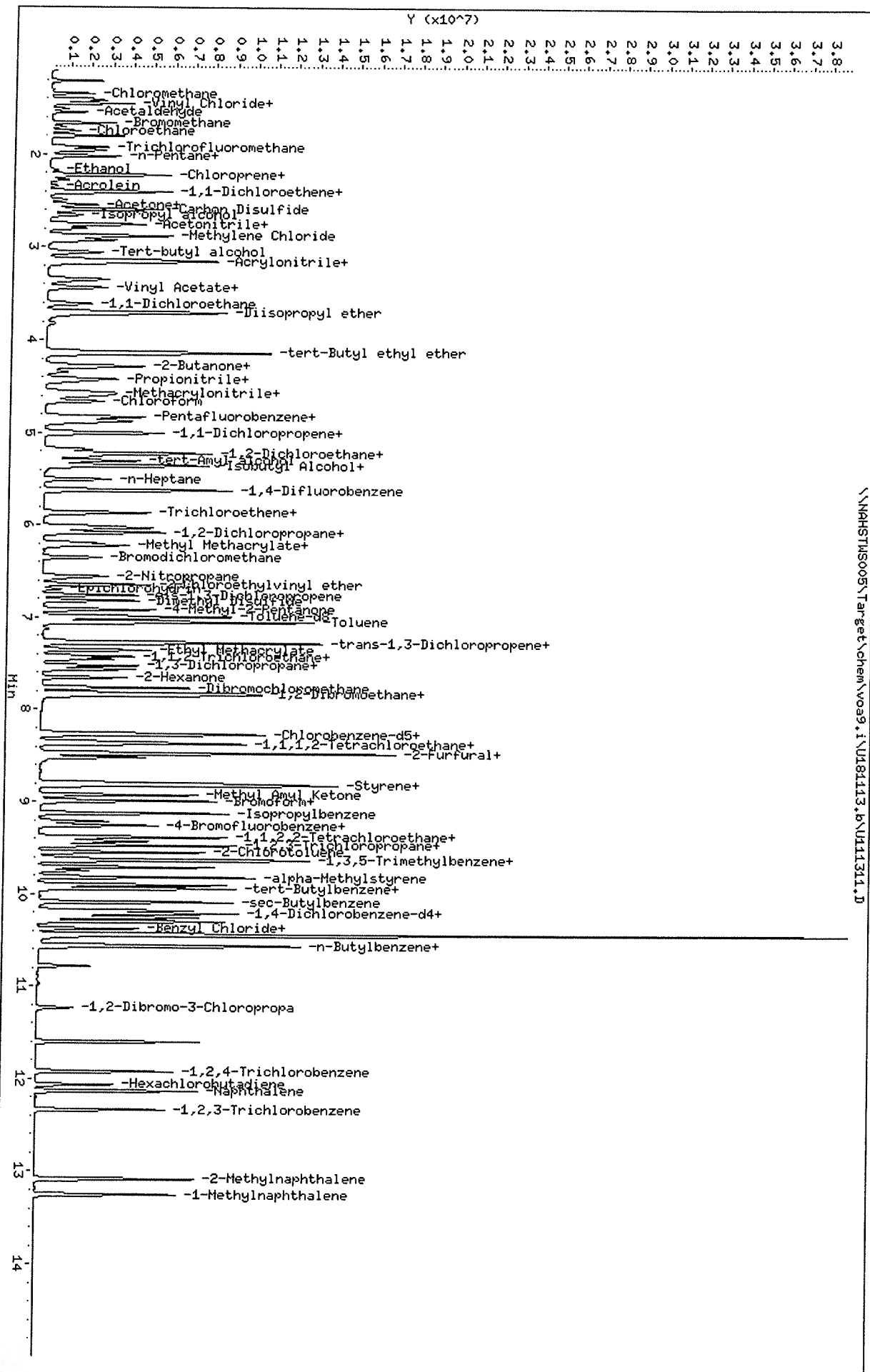
QC Flag Legend

- A - Target compound detected but, quantitated amount exceeded maximum amount.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



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 Purge Volume: 5.0
 Column phase: DB624

Instrument: V099.i
 Operator: PC
 Column diameter: 0.18

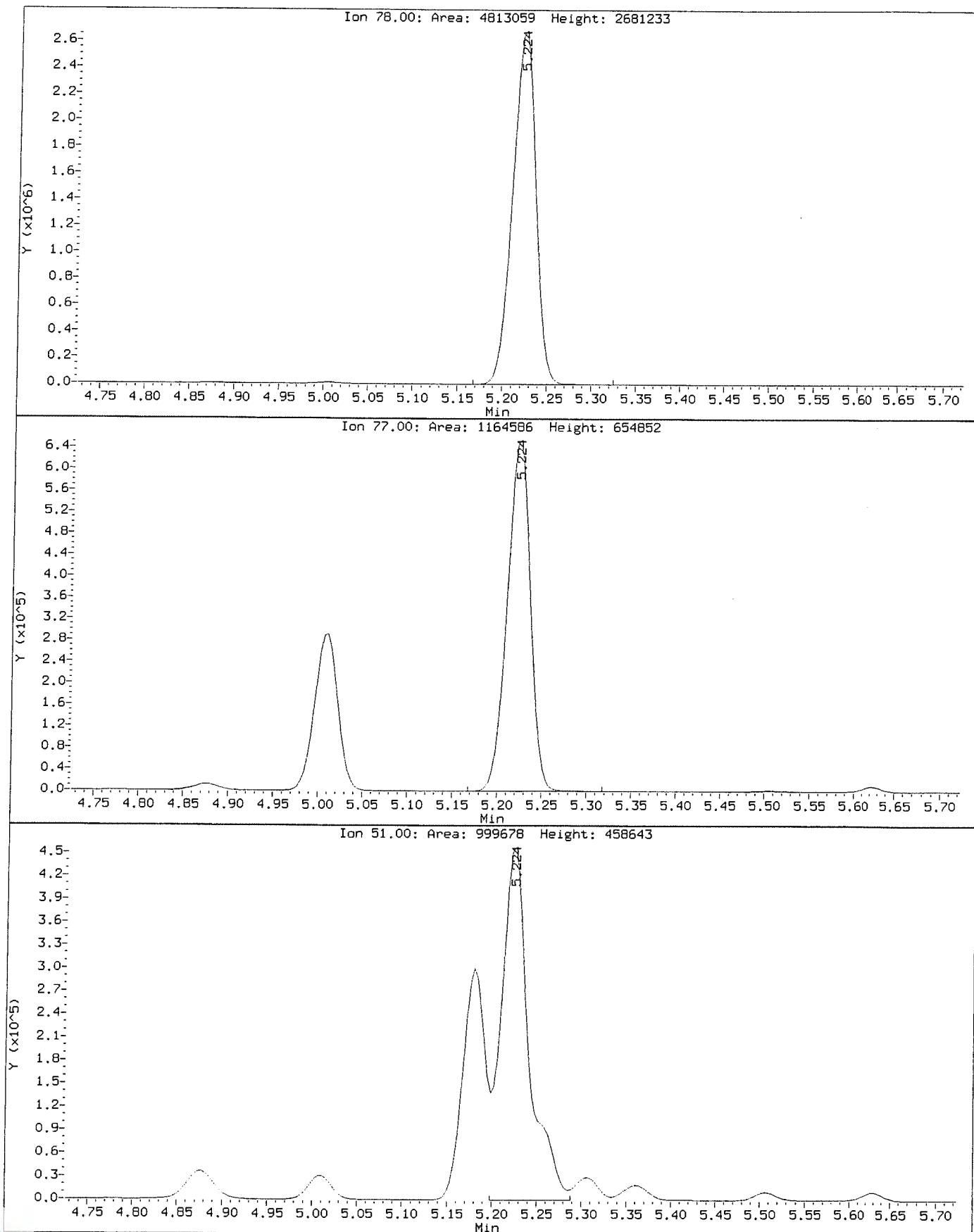


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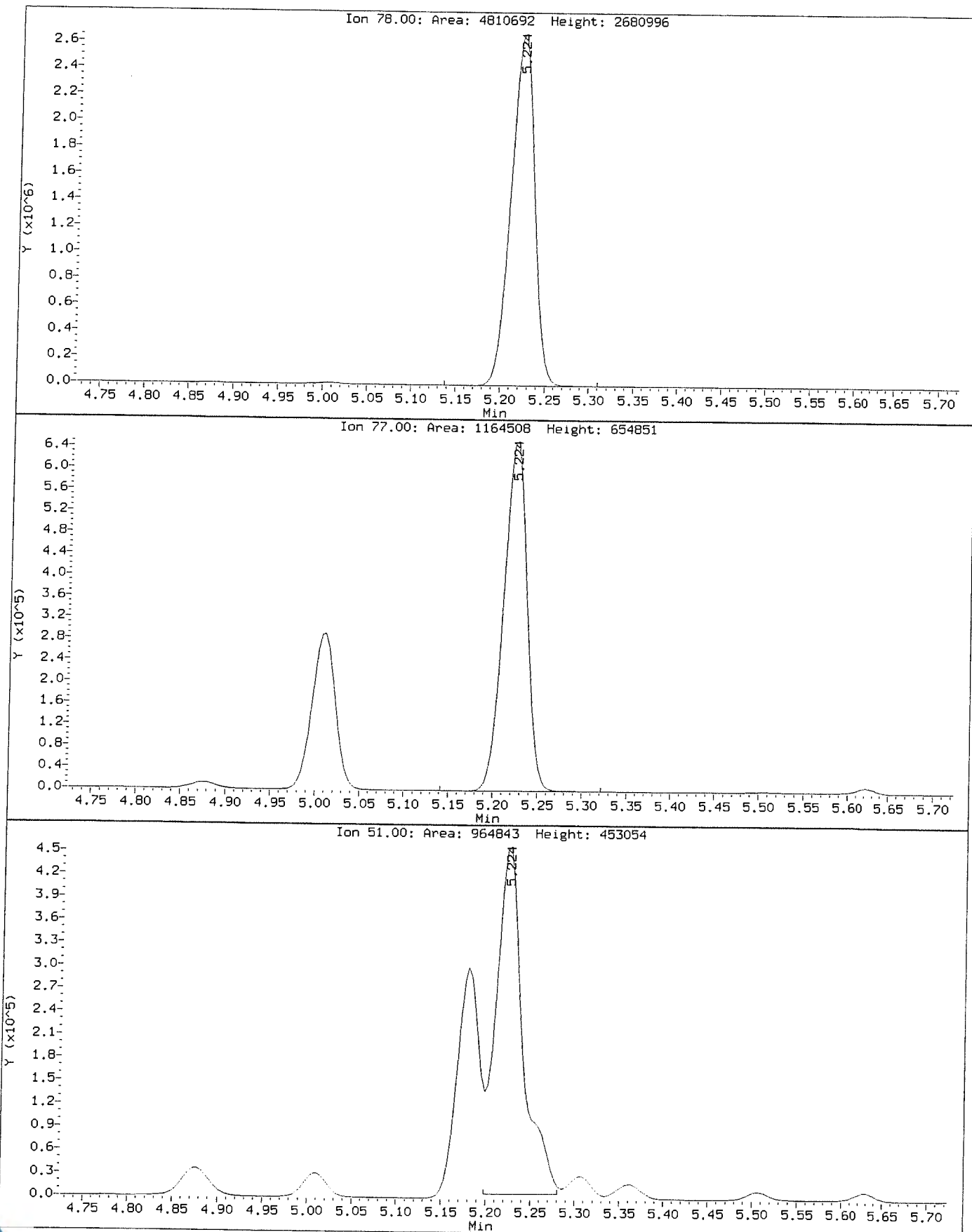
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Instrument: VOA9.1
Client Sample ID: VSTD200

Compound: Benzene
CAS Number: 71-43-2



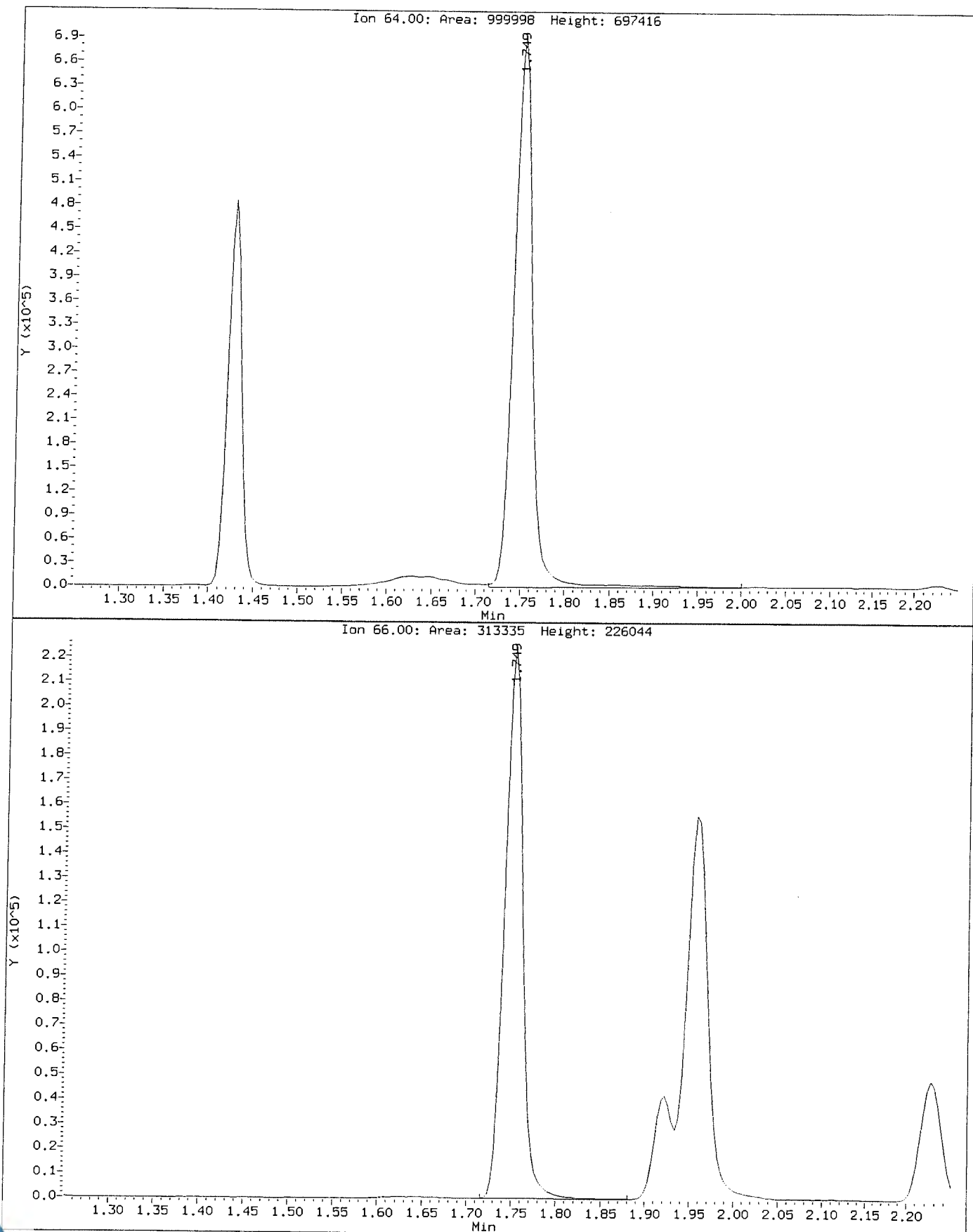
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Compound: Benzene
CAS Number: 71-43-2



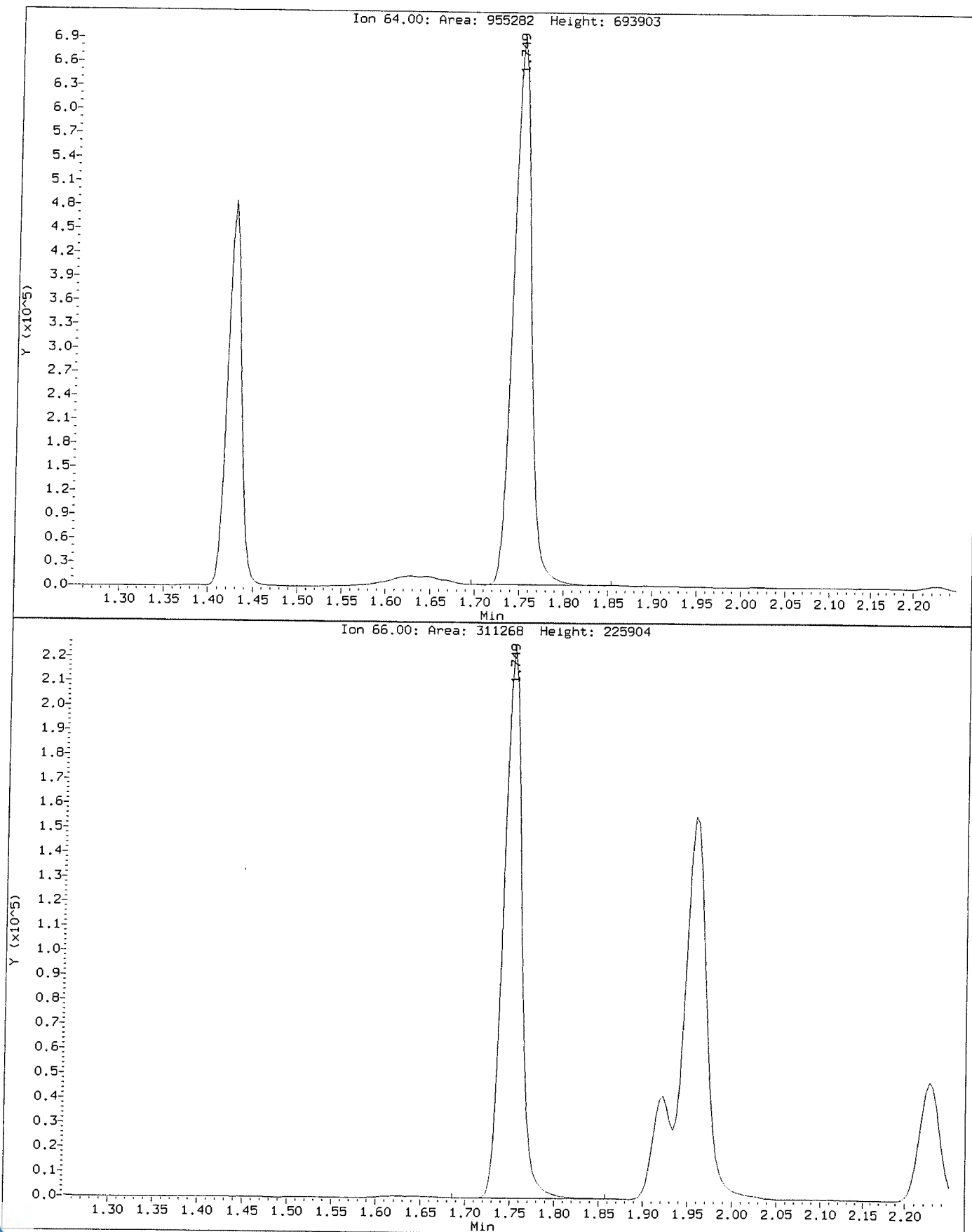
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Injection Date: 13-NOV-2018 15:28
Instrument: VDA9.1
Client Sample ID: VSTD200

Compound: Chloroethane
CAS Number: 75-00-3



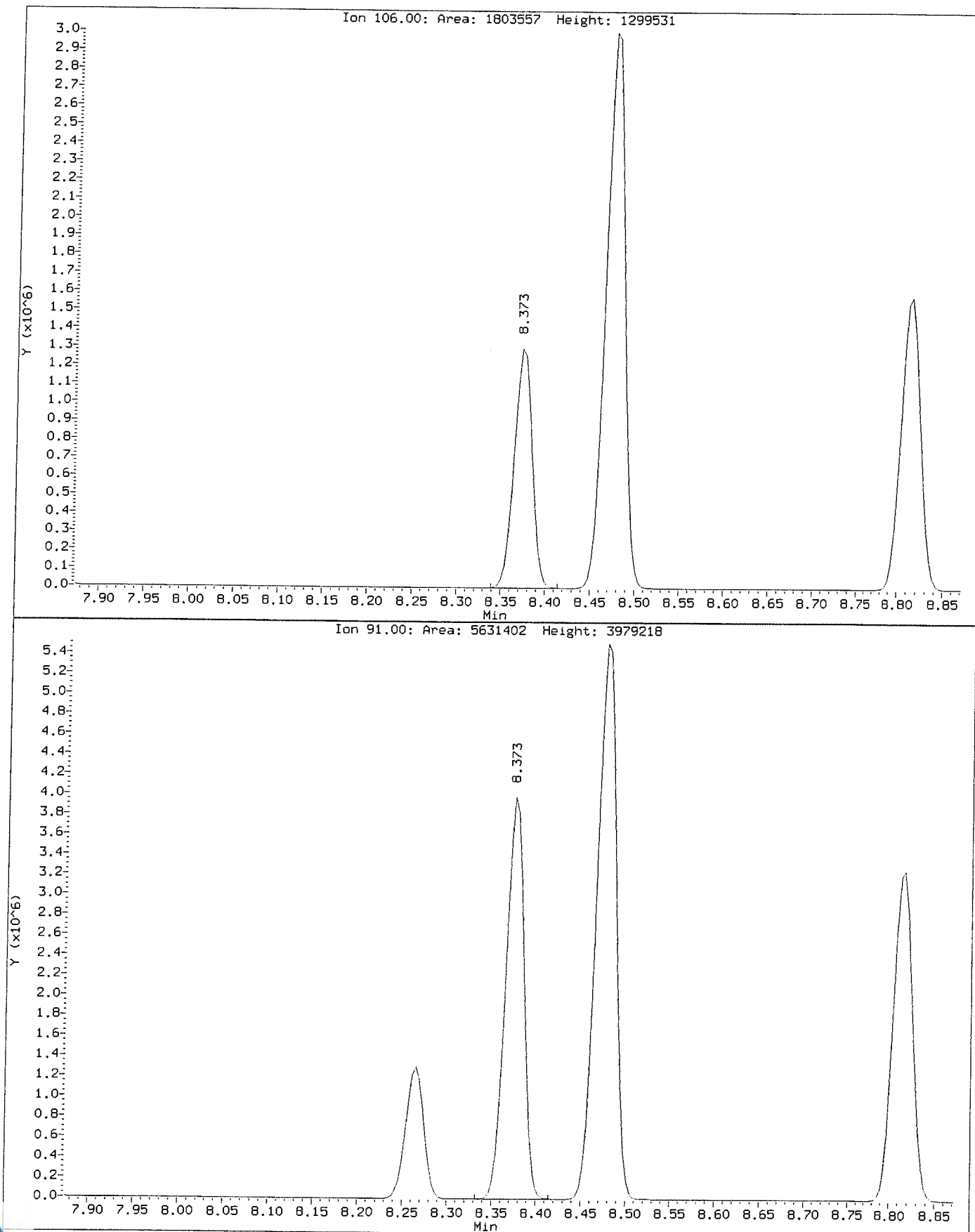
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Instrument: VOA9.i
Client Sample ID: VSTD200

Compound: Chloroethane
CAS Number: 75-00-3



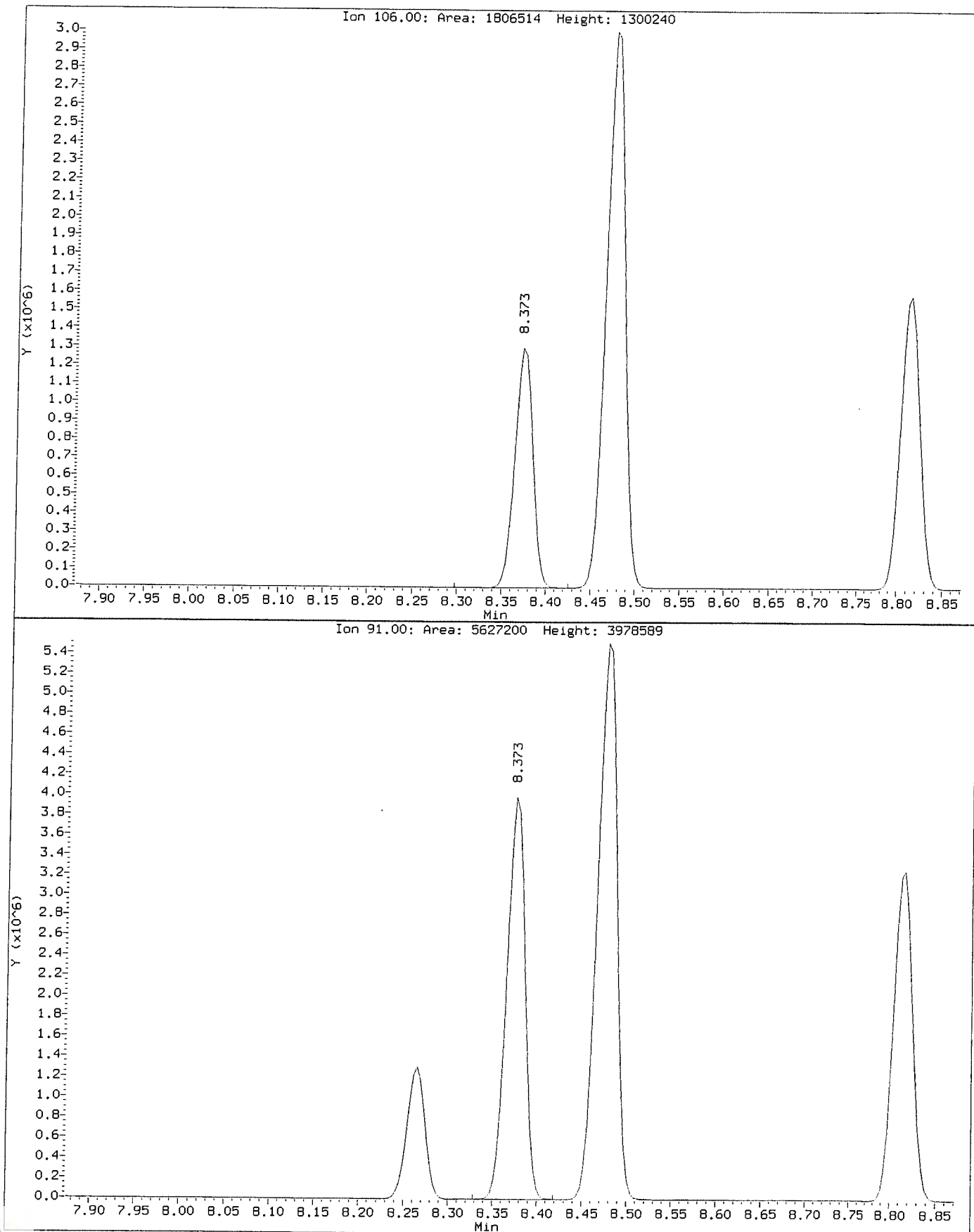
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Instrument: VOA9.i
Client Sample ID: VSTD200

Compound: Ethylbenzene
CAS Number: 100-41-4



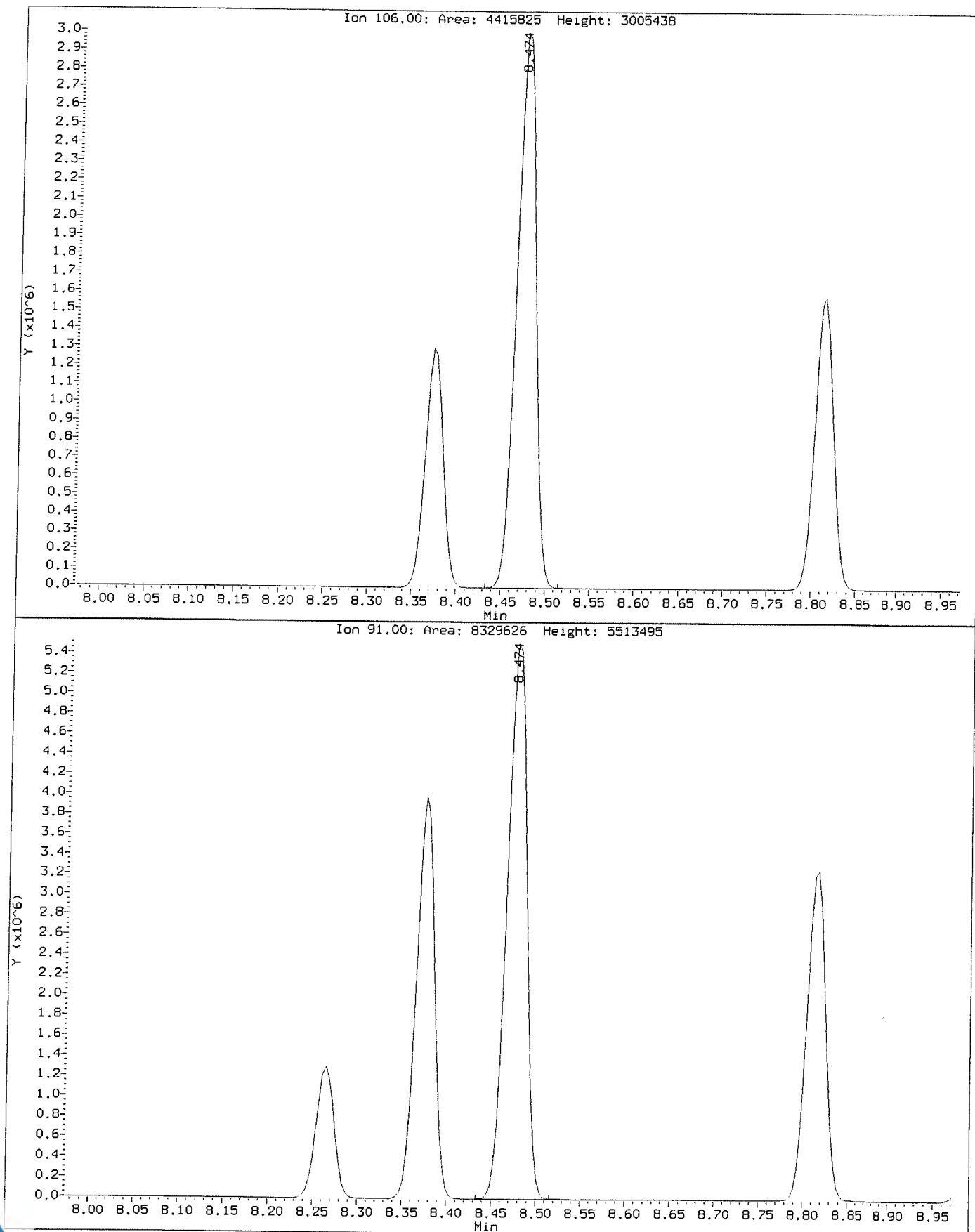
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Instrument: VDA9.i
Client Sample ID: VSTD200

Compound: Ethylbenzene
CAS Number: 100-41-4



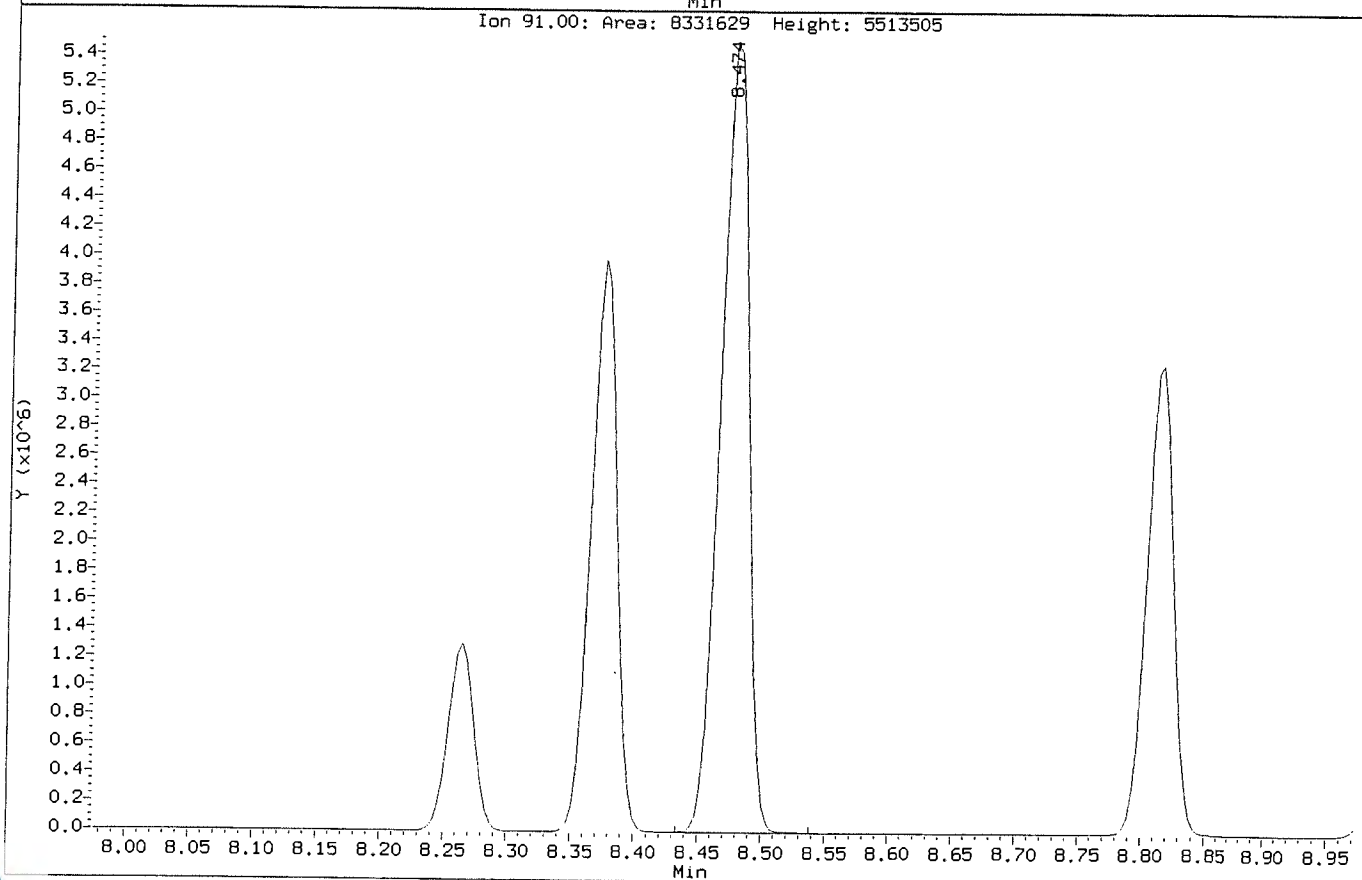
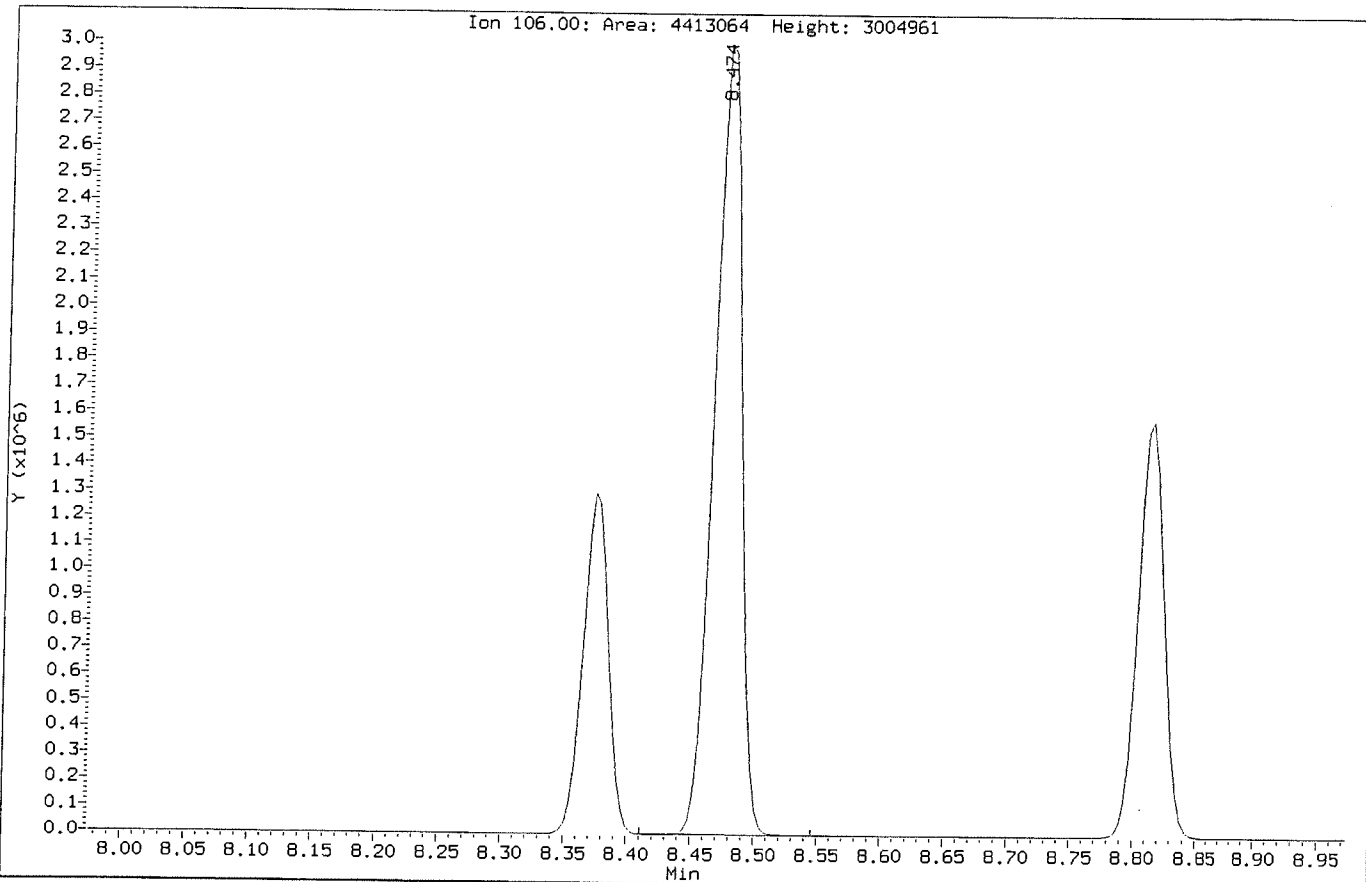
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Instrument: VOA9.i
Client Sample ID: VSTD200

Compound: m,p-Xylenes
CAS Number:



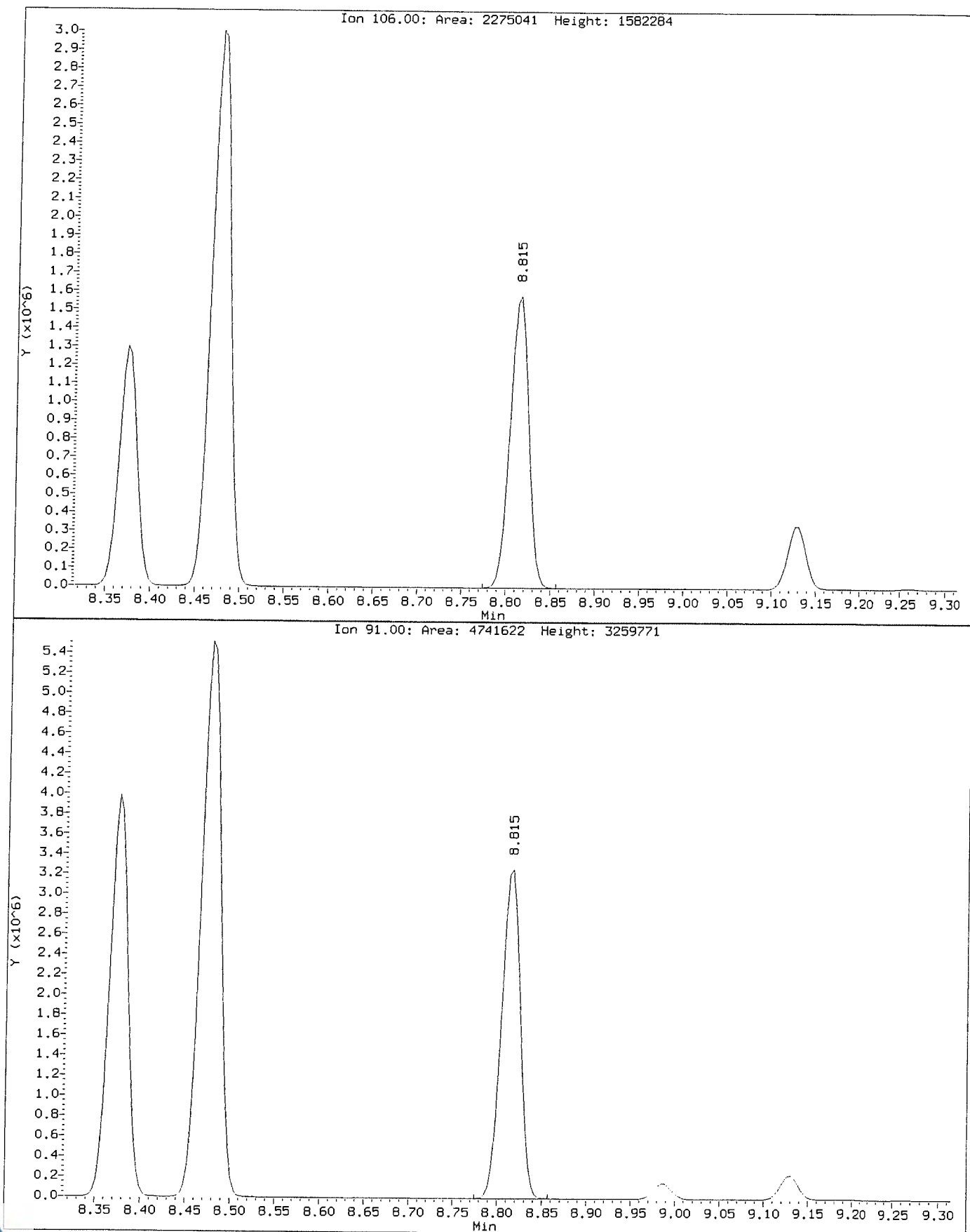
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Instrument: VDA9.1
Client Sample ID: VSTD200

Compound: m,p-Xylenes
CAS Number:



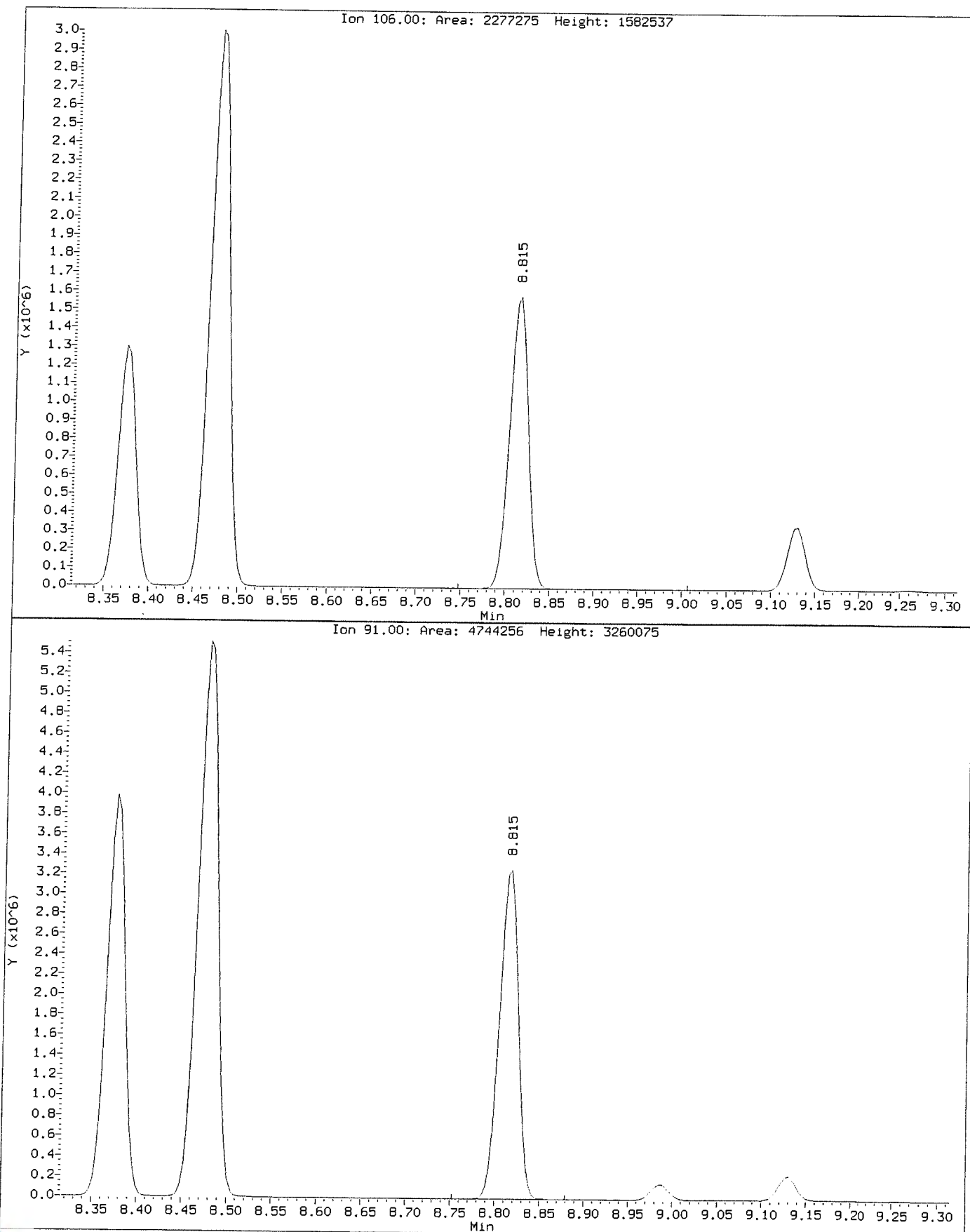
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Injection Date: 13-NOV-2018 15:28
Instrument: VDA9.1
Client Sample ID: VSTD200

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111311.D
Injection Date: 13-NOV-2018 15:28
Instrument: VOA9.i
Client Sample ID: VSTD200

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111313.D Page 1
 Report Date: 24-Jan-2019 18:55

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111313.D
 Lab Smp Id: VICV050 Client Smp ID: VICV050
 Inj Date : 13-NOV-2018 16:18
 Operator : PC Inst ID: VOA9.i
 Smp Info : VICV050;VICV050;2;;ICV
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181113.b\8260C.m
 Meth Date : 24-Jan-2019 18:55 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 15:28 Cal File: U111311.D
 Als bottle: 14 QC Sample: METHSPIKE
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.898	(1.000)	492069	50.0000	
* 36 1,4-Difluorobenzene	114	5.629	5.629	(1.000)	927951	50.0000	
* 47 Chlorobenzene-d5	117	8.253	8.253	(1.000)	869612	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.239	(1.000)	424507	50.0000	
\$ 30 Dibromofluoromethane	113	4.834	4.834	(0.987)	285495	50.4842	50.48
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.179	(1.057)	379384	51.0366	51.03
\$ 48 Toluene-d8	98	6.990	6.993	(0.847)	1145021	51.5678	51.56
\$ 69 4-Bromofluorobenzene	95	9.258	9.261	(1.122)	441801	51.4072	51.40
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	274179	49.9849	49.98
31 1,1,1-Trichloroethane	97	4.830	4.830	(0.986)	416232	48.1554	48.15
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	462139	45.9603	45.96
138 Freon TF	101	2.409	2.405	(0.492)	226905	46.0312	46.03
53 1,1,2-Trichloroethane	83	7.421	7.424	(0.899)	253396	45.8339	45.83
22 1,1-Dichloroethane	63	3.608	3.608	(0.737)	513902	45.5744	45.57
11 1,1-Dichloroethene	96	2.405	2.405	(0.491)	253613	45.6608	45.66
32 1,1-Dichloropropene	75	5.006	5.010	(0.889)	399473	46.6112	46.61
93 1,2,3-Trichlorobenzene	180	12.335	12.339	(1.205)	387107	48.3006	48.30
71 1,2,3-Trichloropropane	75	9.430	9.434	(0.921)	517101	49.2816	49.28
90 1,2,4-Trichlorobenzene	180	11.923	11.926	(1.165)	396308	47.4733	47.47
79 1,2,4-Trimethylbenzene	105	9.944	9.943	(0.971)	1125014	47.7134	47.71
89 1,2-Dibromo-3-Chloropropane	155	11.233	11.237	(1.097)	74206	47.8284	47.82
57 1,2-Dibromoethane	107	7.852	7.855	(0.951)	310006	48.4238	48.42



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111313.D Page 2
 Report Date: 24-Jan-2019 18:55

Compounds	QUANT	SIG						CONCENTRATIONS	
			MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/l)	FINAL (ug/l)
88 1,2-Dichlorobenzene	146		10.573	10.573	(1.033)	609149	44.9344	44.93	
33 1,2-Dichloroethane	62		5.254	5.257	(0.933)	438739	45.1829	45.18	
42 1,2-Dichloropropane	63		6.082	6.086	(1.081)	312363	47.2789	47.27	
75 1,3,5-Trimethylbenzene	105		9.629	9.628	(0.941)	1087962	47.7004	47.70	
83 1,3-Dichlorobenzene	146		10.180	10.183	(0.995)	608880	45.5562	45.55	
54 1,3-Dichloropropane	76		7.567	7.567	(0.917)	548032	46.1653	46.16	
84 1,4-Dichlorobenzene	146		10.258	10.258	(1.002)	631760	48.3226	48.32	
26 2,2-Dichloropropane	77		4.279	4.279	(0.874)	357603	51.2102	51.21	
24 2-Butanone	43		4.343	4.343	(0.887)	319364	92.3297	92.32	
76 2-Chlorotoluene	91		9.550	9.550	(0.933)	950278	45.7148	45.71	
52 2-Hexanone	43		7.653	7.653	(0.927)	501801	99.2665	99.26	
77 4-Chlorotoluene	91		9.640	9.643	(0.942)	1123555	46.8423	46.84	
82 p-Isopropyltoluene	119		10.213	10.213	(0.998)	1142647	49.1524	49.15	
45 4-Methyl-2-Pentanone	43		6.915	6.918	(0.838)	699041	96.3600	96.36	
10 Acetone	43		2.487	2.491	(0.508)	216067	95.4641	95.46	
37 Benzene	78		5.220	5.224	(0.927)	1188078	46.3202	46.32	
74 Bromobenzene	156		9.385	9.385	(0.917)	321699	46.1973	46.19	
29 Bromochloromethane	128		4.557	4.560	(0.930)	147658	50.9008	50.90	
39 Bromodichloromethane	83		6.349	6.352	(1.128)	374186	48.5304	48.53	
66 Bromoform	173		8.984	8.987	(1.089)	188320	47.7183	47.71(T)	
6 Bromomethane	94		1.670	1.662	(0.341)	208101	50.9317	50.93	
19 Carbon Disulfide	76		2.596	2.596	(0.530)	1686616	98.3287	98.32	
34 Carbon Tetrachloride	117		4.999	4.999	(0.888)	330709	49.3949	49.39	
59 Chlorobenzene	112		8.275	8.279	(1.003)	813973	45.3514	45.35	
7 Chloroethane	64		1.756	1.749	(0.359)	230904	44.9734	44.97(M)	
28 Chloroform	83		4.662	4.661	(0.952)	517425	45.6713	45.67	
3 Chloromethane	50		1.344	1.344	(0.274)	306744	46.7074	46.70	
27 cis-1,2-Dichloroethene	96		4.290	4.290	(0.876)	320390	45.9503	45.95	
46 cis-1,3-Dichloropropene	75		6.761	6.761	(1.201)	516764	53.2223	53.22	
55 Dibromochloromethane	129		7.758	7.762	(0.940)	286771	52.1150	52.11	
44 Dibromomethane	93		6.191	6.195	(1.100)	195311	47.1700	47.16	
2 Dichlorodifluoromethane	85		1.209	1.213	(0.247)	347869	47.8726	47.87	
61 Ethylbenzene	106		8.373	8.373	(1.015)	424849	46.8143	46.81	
91 Hexachlorobutadiene	225		12.065	12.065	(1.179)	145238	55.0778	55.07	
67 Isopropylbenzene	105		9.126	9.130	(1.106)	1266469	47.9214	47.92	
62 m,p-Xylenes	106		8.474	8.474	(1.027)	1058456	94.6512	94.65	
17 Methylene Chloride	84		2.877	2.877	(0.587)	301983	48.0719	48.07	
87 n-Butylbenzene	91		10.558	10.558	(1.031)	1100279	49.5091	49.50	
73 n-Propylbenzene	91		9.475	9.478	(0.926)	1593592	47.5729	47.57	
92 Naphthalene	128		12.133	12.136	(1.185)	1287047	51.2004	51.20	
63 o-Xylene	106		8.811	8.815	(1.068)	530431	47.3909	47.39(H)	
81 sec-Butylbenzene	105		10.086	10.090	(0.985)	1350043	48.4022	48.40	
64 Styrene	104		8.826	8.826	(1.069)	941191	49.8117	49.81	
78 tert-Butylbenzene	119		9.902	9.906	(0.967)	929608	47.8232	47.82	
56 Tetrachloroethene	164		7.526	7.525	(0.912)	207441	45.5377	45.53	
50 Toluene	91		7.050	7.049	(0.854)	1274589	46.1658	46.16	
20 trans-1,2-Dichloroethene	96		3.147	3.147	(0.643)	282235	46.3646	46.36	
51 trans-1,3-Dichloropropene	75		7.263	7.263	(1.290)	454600	47.3993	47.39	
38 Trichloroethene	130		5.865	5.865	(1.042)	293292	46.3847	46.38	
8 Trichlorofluoromethane	101		1.959	1.955	(0.400)	460052	46.7647	46.76	
5 Vinyl Chloride	62		1.423	1.426	(0.291)	382169	47.6661	47.66	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111313.D Page 3
Report Date: 24-Jan-2019 18:55

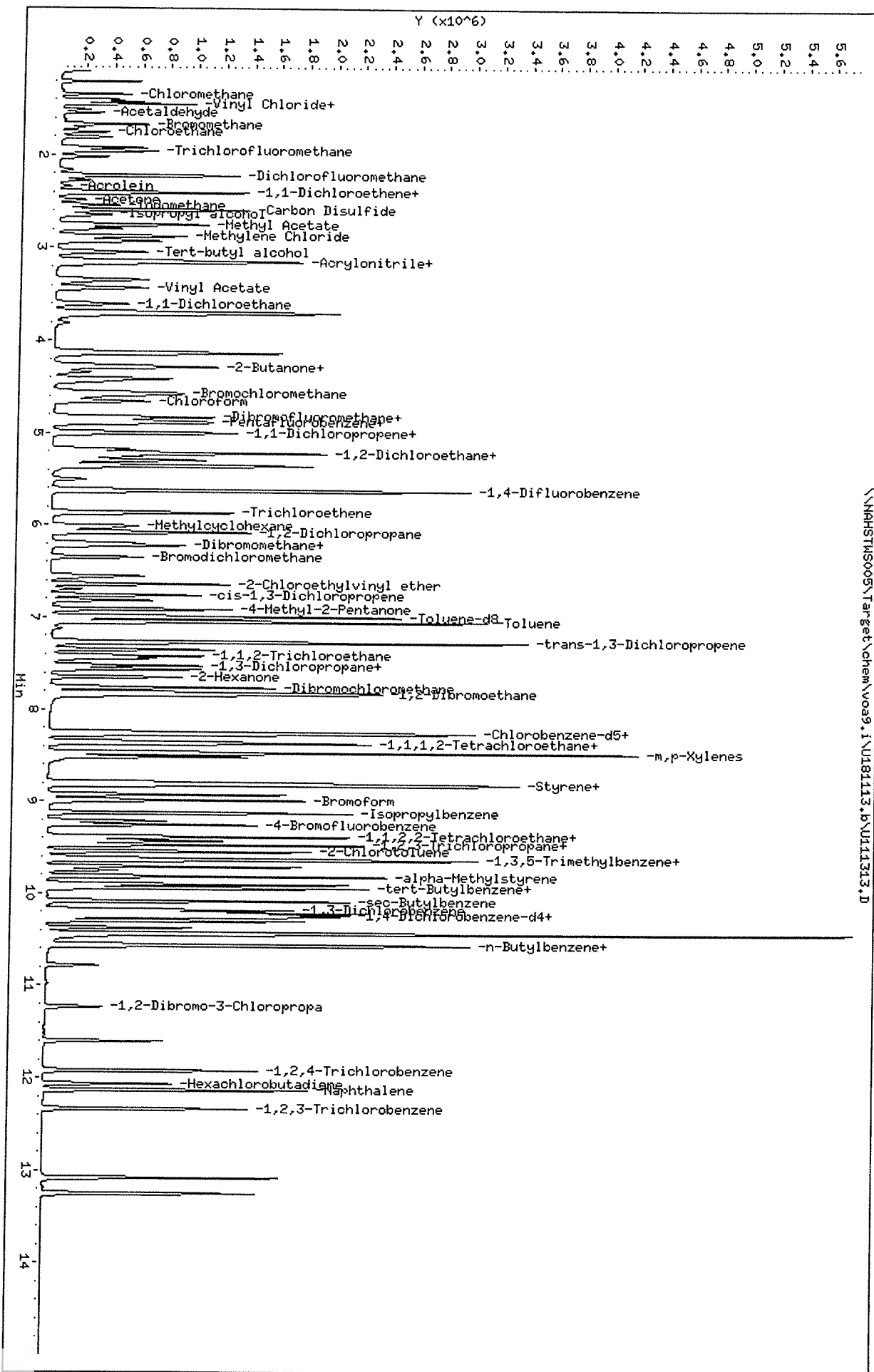
QC Flag Legend

- T - Target compound detected outside RT window.
- M - Compound response manually integrated.
- H - Operator selected an alternate compound hit.



Data File: \\NAHSTMS005\Target\chem\voa9.1\U481113.B\U48111313.D
 Date: 13-NOV-2018 16:18
 Client ID: VICV050
 Sample Info: VICV050;VICV050;2;ICV
 Purge Volume: 5.0
 Column phase: DB624

Instrument: V099.1
 Operator: PC
 Column diameter: 0.18

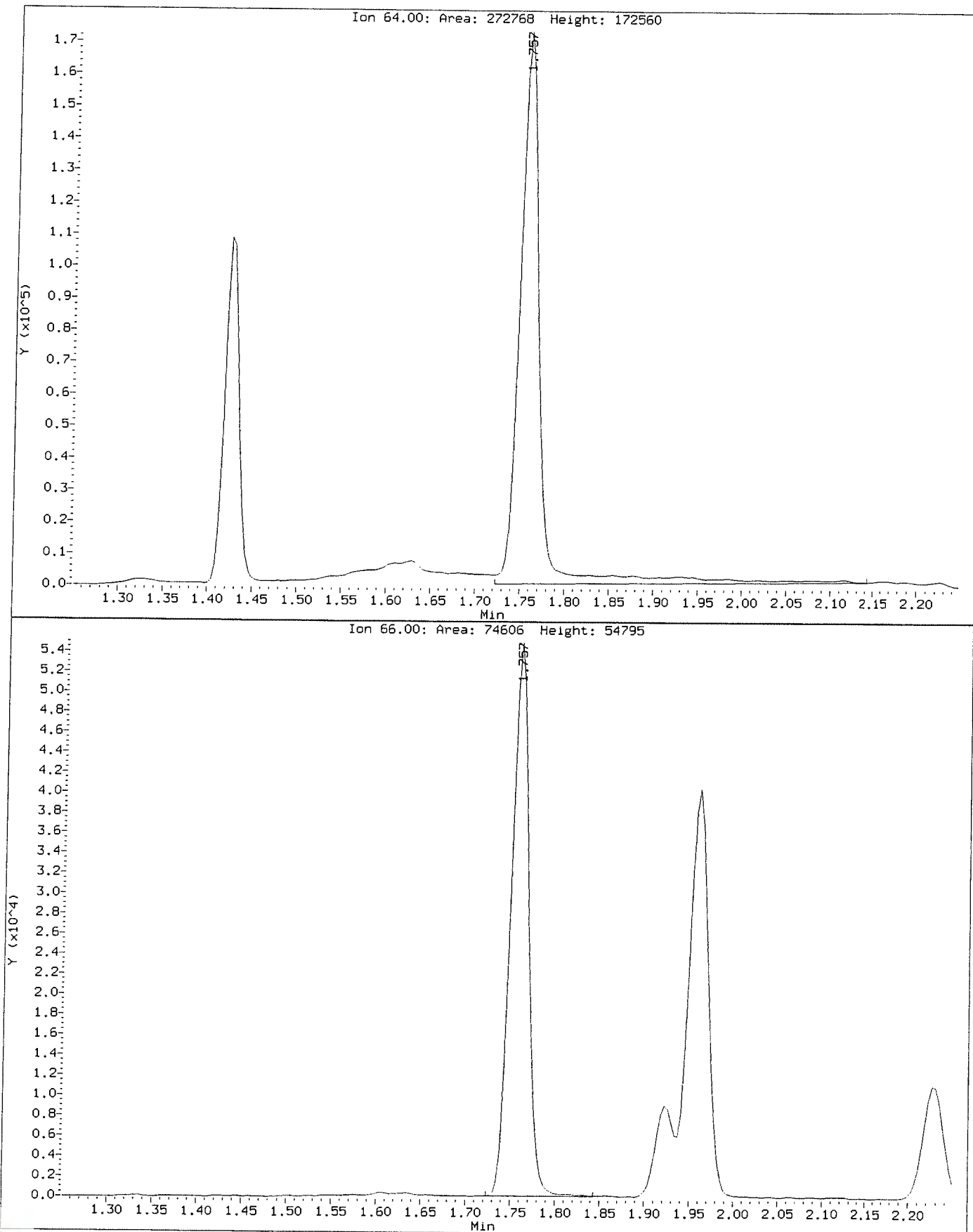


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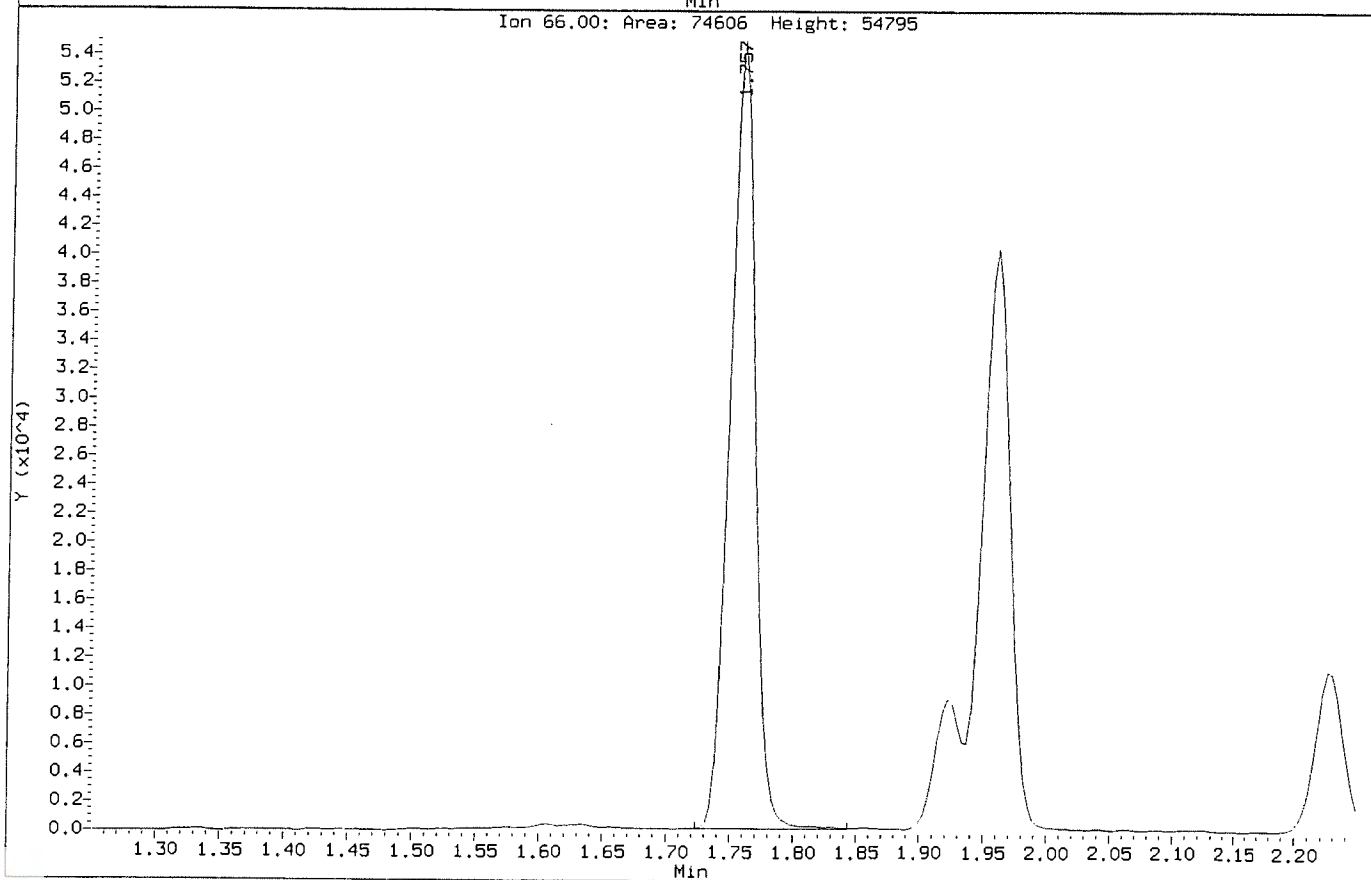
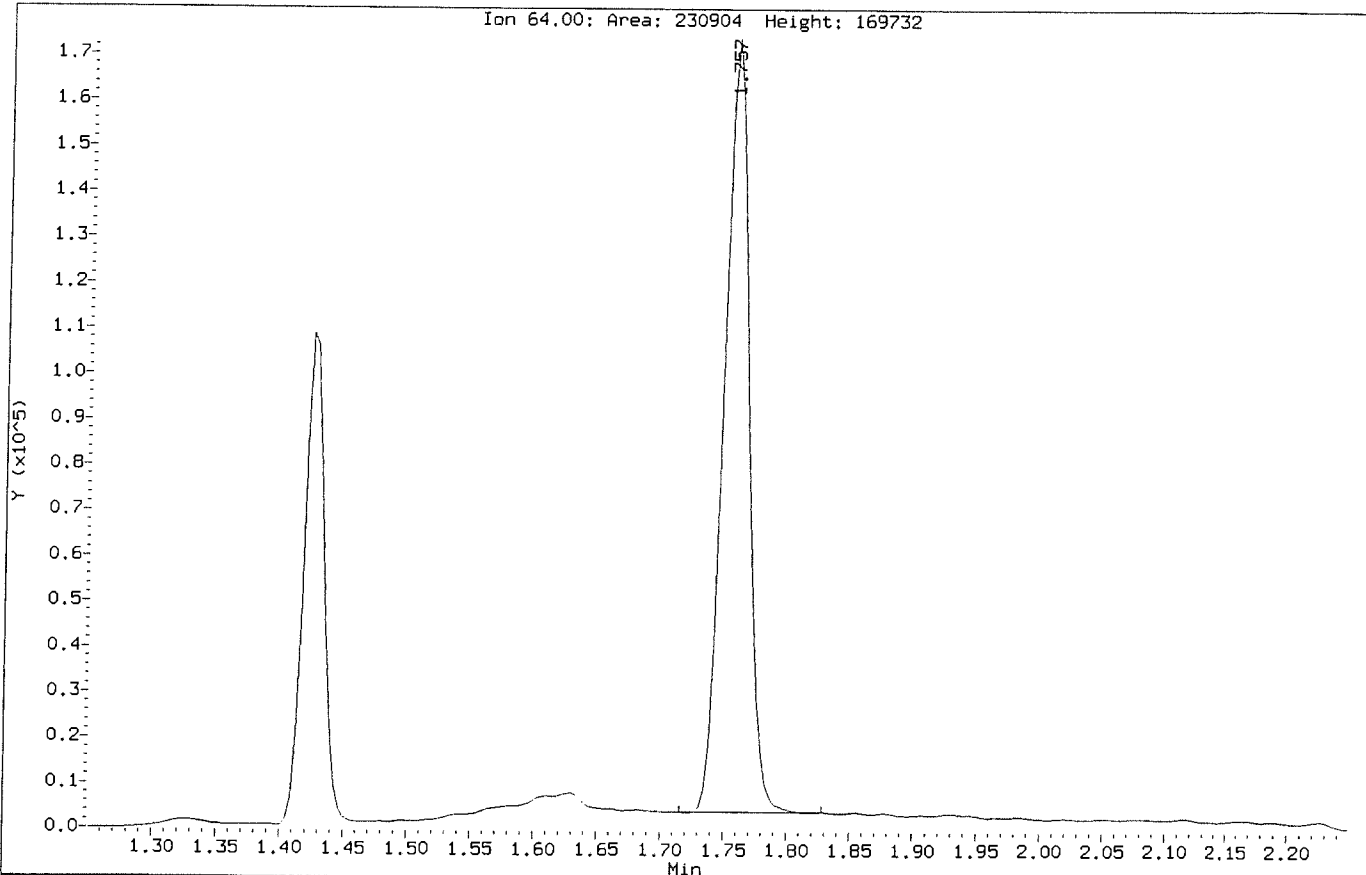
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Injection Date: 13-NOV-2018 16:18
Instrument: VOA9.1
Client Sample ID: VICV050

Compound: Chloroethane
CAS Number: 75-00-3



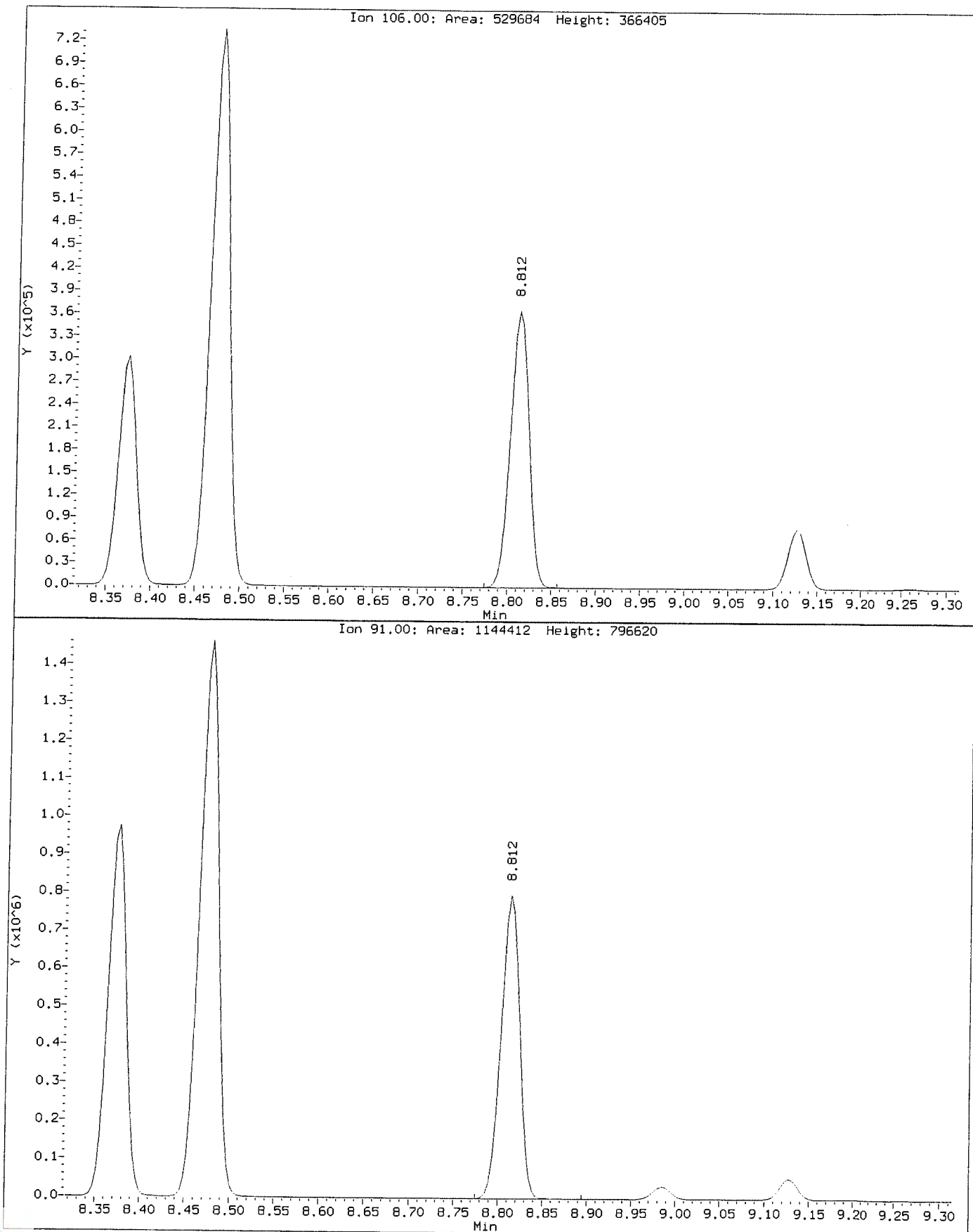
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Instrument: VDA9.i
Client Sample ID: VICV050

Compound: Chloroethane
CAS Number: 75-00-3



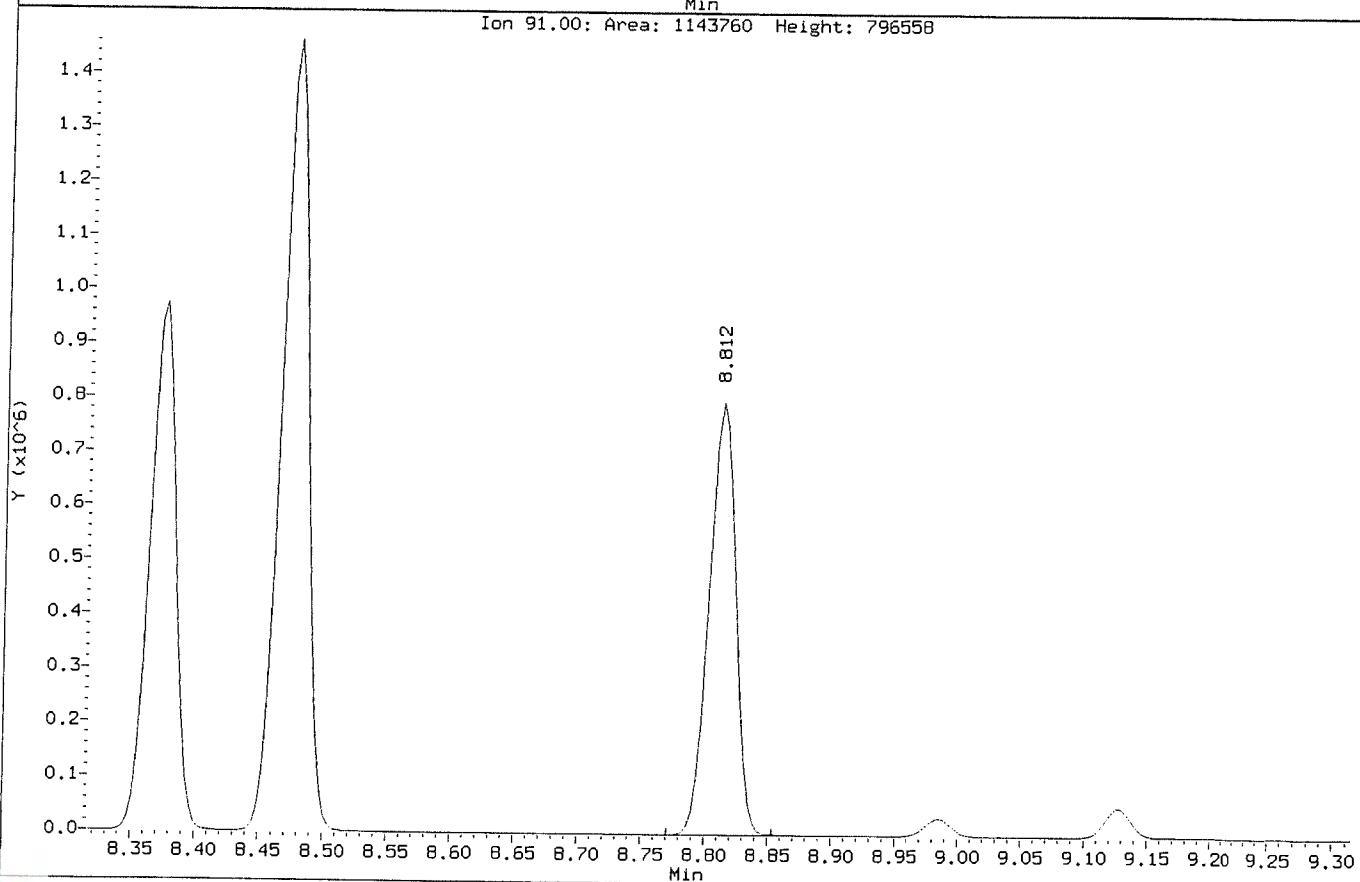
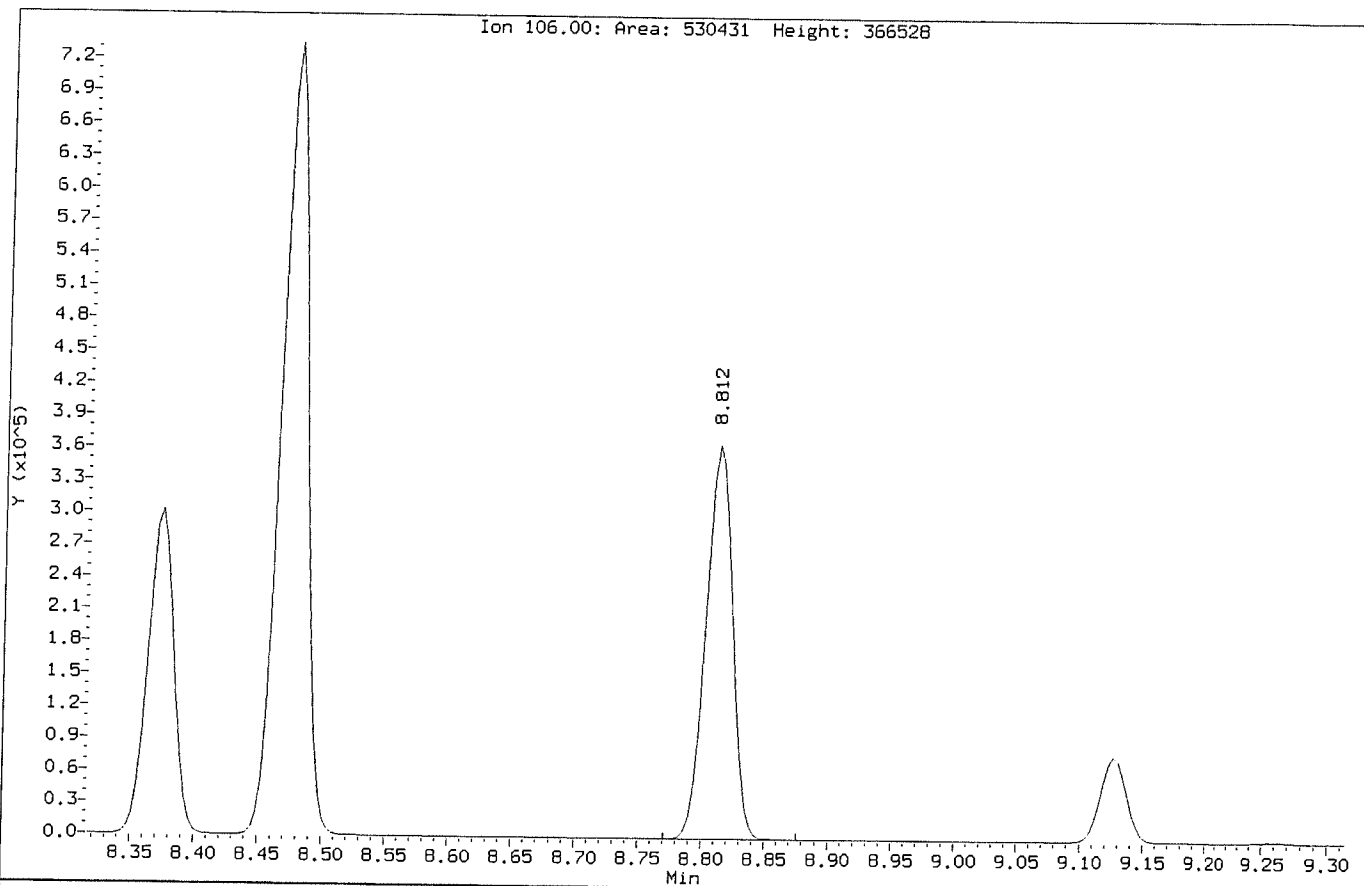
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Injection Date: 13-NOV-2018 16:18
Instrument: VOA9.1
Client Sample ID: VICV050

Compound: o-Xylene
CAS Number: 95-47-6



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181113.b\U111313.D
Injection Date: 13-NOV-2018 16:18
Instrument: VOA9.i
Client Sample ID: VICV050

Compound: o-Xylene
CAS Number: 95-47-6



MSVOA09 -Logbook

Batch: 34000
 Date: 12-18-2018
 Method: 8260
 Comments:

Analyst: Presenta Cabascango
 Reviewer:
 Laboratory: Houston

#	Samp ID	Type	Analyzed	DF	Init Wt/Vol	Final Vol	File ID	Matrix	Status	pH
1	BFB	TUNE	12-18-2018 11:30 am	1.00	50 mL	50 mL	U121801.D	Liquid	Y	NA
	<i>Auto find/purged</i>									
2	VSTD050	CCV	12-18-2018 11:54 am	1.00	50 mL	50 mL	U121802.D	Liquid	Y	NA
	<i>10 uL cal std/50 mL DI</i>									
3	BLANK	MBLK	12-18-2018 12:19 pm	1.00	50 mL	50 mL	U121803.D	Liquid	Y	NA
4	BLANK	SAMP	12-18-2018 12:44 pm	1.00	50 mL	50 mL	U121804.D	Liquid	Y	NA
5	BLANK	SAMP	12-18-2018 01:09 pm	1.00	50 mL	50 mL	U121805.D	Liquid	Y	NA
6	VBLKW-181218	MBLK	12-18-2018 01:34 pm	1.00	50 mL	50 mL	U121806.D	Liquid	Y	NA
7	VLCSW-1812018	LCS	12-18-2018 01:58 pm	1.00	50 mL	50 mL	U121807.D	Liquid	Y	NA
	<i>4 uL ICV std/50 mL DI</i>									
8	HS18120442-01	SAMP	12-18-2018 02:23 pm	1.00	50 mL	50 mL	U121808.D	Liquid	Y	<2
9	HS18120278-05	SAMP	12-18-2018 02:48 pm	1.00	50 mL	50 mL	U121809.D	Liquid	Y	<2
10	HS18120278-02	SAMP	12-18-2018 03:13 pm	1.00	50 mL	50 mL	U121810.D	Liquid	Y	<2
11	HS18120278-04	SAMP	12-18-2018 03:37 pm	1.00	50 mL	50 mL	U121811.D	Liquid	Y	<2
12	HS18120278-03	SAMP	12-18-2018 04:02 pm	1.00	50 mL	50 mL	U121812.D	Liquid	Y	<2
13	HS18120278-01	SAMP	12-18-2018 04:27 pm	1.00	50 mL	50 mL	U121813.D	Liquid	Y	<2
14	HS18120375-01	SAMP	12-18-2018 04:51 pm	1.00	50 mL	50 mL	U121814.D	Liquid	Y	<2
15	HS18120375-03	SAMP	12-18-2018 05:16 pm	1.00	50 mL	50 mL	U121815.D	Liquid	Y	<2
16	HS18120278-03MS	MS	12-18-2018 05:41 pm	1.00	50 mL	50 mL	U121816.D	Liquid	Y	<2
	<i>3.5 uL cal std/43 mL Sample</i>									
17	HS18120278-03MSD	MSD	12-18-2018 06:06 pm	1.00	50 mL	50 mL	U121817.D	Liquid	Y	<2
	<i>3.5 uL cal std/43 mL Sample</i>									
18	HS18120360-01	SAMP	12-18-2018 06:31 pm	1.00	50 mL	50 mL	U121818.D	Liquid	Y	<2
19	HS18120360-02	SAMP	12-18-2018 06:55 pm	1.00	50 mL	50 mL	U121819.D	Liquid	Y	<2
20	HS18120412-02	SAMP	12-18-2018 07:20 pm	1.00	50 mL	50 mL	U121820.D	Liquid	Y	<2
21	HS18120412-04	SAMP	12-18-2018 07:45 pm	1.00	50 mL	50 mL	U121821.D	Liquid	Y	<2
22	HS18120412-05	SAMP	12-18-2018 08:09 pm	1.00	50 mL	50 mL	U121822.D	Liquid	Y	<2
23	HS18120412-08	SAMP	12-18-2018 08:34 pm	1.00	50 mL	50 mL	U121823.D	Liquid	Y	<2
24	HS18120412-09	SAMP	12-18-2018 08:59 pm	1.00	50 mL	50 mL	U121824.D	Liquid	Y	<2
25	HS18120412-10	SAMP	12-18-2018 09:24 pm	1.00	50 mL	50 mL	U121825.D	Liquid	Y	<2
26	HS18120412-11	SAMP	12-18-2018 09:48 pm	1.00	50 mL	50 mL	U121826.D	Liquid	Y	<2
27	HS18120412-12	SAMP	12-18-2018 10:13 pm	1.00	50 mL	50 mL	U121827.D	Liquid	Y	<2
28	VSTD050-END	CCV	12-18-2018 10:38 pm	1.00	50 mL	50 mL	U121828.D	Liquid	Y	NA
	<i>10 uL cal std/50 mL DI</i>									
29	BFB	TUNE	12-18-2018 11:03 pm	1.00	50 mL	50 mL	V121801.D	Liquid	Y	NA
30	VSTD050	CCV	12-18-2018 11:27 pm	1.00	50 mL	50 mL	V121802.D	Liquid	Y	NA
	<i>10 uL cal std/50 mL DI</i>									
31	CCB	SAMP	12-18-2018 11:52 pm	1.00	50 mL	50 mL	V121803.D	Liquid	Y	NA
	CCB									
32	VLCSW-1812018	LCS	12-19-2018 12:17 am	1.00	50 mL	50 mL	V121804.D	Liquid	Y	NA
	<i>4 uL ICV std/50 mL DI</i>									
33	BLANK	SAMP	12-19-2018 12:42 am	1.00	50 mL	50 mL	V121805.D	Liquid	Y	NA
34	VBLKW-181218	MBLK	12-19-2018 01:06 am	1.00	50 mL	50 mL	V121806.D	Liquid	Y	NA
35	HS18120336-11	SAMP	12-19-2018 01:31 am	1.00	50 mL	50 mL	V121807.D	Liquid	Y	<2
36	HS18120336-09	SAMP	12-19-2018 01:56 am	1.00	50 mL	50 mL	V121808.D	Liquid	Y	<2



MSVOA09 -Logbook

#	<u>Samp ID</u>	<u>Type</u>	<u>Analyzed</u>	<u>DF</u>	<u>Init Wt/Vol</u>	<u>Final Vol</u>	<u>File ID</u>	<u>Matrix</u>	<u>Status</u>	<u>pH</u>	
37	HS18120336-10	SAMP	12-19-2018 02:21 am	1.00	50 mL	50 mL	V121809.D	Liquid	Y	<2	
38	HS18120336-13	SAMP	12-19-2018 02:45 am	1.00	50 mL	50 mL	V121810.D	Liquid	Y	<2	
39	HS18120336-14	SAMP	12-19-2018 03:10 am	1.00	50 mL	50 mL	V121811.D	Liquid	Y	<2	
40	HS18120336-07	SAMP	12-19-2018 03:35 am	1.00	50 mL	50 mL	V121812.D	Liquid	Y	<2	
41	HS18120336-14MS	MS	12-19-2018 04:00 am	1.00	50 mL	50 mL	V121813.D	Liquid	Y	<2	
	<i>3.5 uL cal std/43 mL Sample</i>										
42	HS18120336-14MSD	MSD	12-19-2018 04:25 am	1.00	50 mL	50 mL	V121814.D	Liquid	Y	<2	
	<i>3.5 uL cal std/43 mL Sample</i>										
43	BLANK	SAMP	12-19-2018 04:49 am	1.00	50 mL	50 mL	V121815.D	Liquid	Y	NA	
	<i>Cleanup blk</i>										
44	HS18120336-15	SAMP	12-19-2018 05:14 am	1.00	50 mL	50 mL	V121816.D	Liquid	Y	<2	
45	HS18120336-16	SAMP	12-19-2018 05:39 am	1.00	50 mL	50 mL	V121817.D	Liquid	Y	<2	
46	HS18120336-17	SAMP	12-19-2018 06:04 am	1.00	50 mL	50 mL	V121818.D	Liquid	Y	<2	
47	HS18120336-18	SAMP	12-19-2018 06:28 am	1.00	50 mL	50 mL	V121819.D	Liquid	Y	<2	
48	HS18120336-19	SAMP	12-19-2018 06:53 am	1.00	50 mL	50 mL	V121820.D	Liquid	Y	<2	
49	HS18120336-20	SAMP	12-19-2018 07:18 am	1.00	50 mL	50 mL	V121821.D	Liquid	Y	<2	
50	HS18120336-03	SAMP	12-19-2018 07:43 am	10.00	5 mL	50 mL	V121822.D	Liquid	Y	<2	
51	HS18120336-21	SAMP	12-19-2018 08:07 am	10.00	5 mL	50 mL	V121823.D	Liquid	Y	<2	
52	HS18120336-01	SAMP	12-19-2018 08:32 am	1.00	50 mL	50 mL	V121824.D	Liquid	Y	<2	
53	HS18120336-02	SAMP	12-19-2018 08:57 am	1.00	50 mL	50 mL	V121825.D	Liquid	Y	<2	
54	HS18120336-02	SAMP	12-19-2018 09:22 am	10.00	5 mL	50 mL	V121826.D	Liquid	Y	<2	
55	HS18120336-04	SAMP	12-19-2018 09:46 am	25.00	2 mL	50 mL	V121827.D	Liquid	Y	<2	
56	HS18120336-05	SAMP	12-19-2018 10:11 am	100.00	500 µL	50 mL	V121828.D	Liquid	Y	<2	
57	HS18120336-08	SAMP	12-19-2018 10:36 am	10.00	5 mL	50 mL	V121829.D	Liquid	Y	<2	

Chemical	Value
SURR SPK ID	30502-52-03
IS ID	30502-52-04
ICV STD ID	30603-45-01
LCS/MS ID	30603-45-01
CAL STD ID	30502-55-01/02
BFB Ion 95 Response	30502-52-03



FORM 2
WATER VOLATILE SYSTEM MONITORING COMPOUND RECOVERY

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

	CLIENT SAMPLE NO.	SMC1 (DCE) #	SMC2 #	SMC3 (TOL) #	OTHER #	TOT OUT
01	VBLKW-181218	90	87	104	98	0
02	VLCSW-181218	88	89	105	102	0
03	HS18120278-0	89	88	106	98	0
04	HS18120278-0	90	88	105	98	0
05	HS18120278-0	91	88	105	97	0
06	HS18120278-0	91	89	105	98	0
07	HS18120278-0	91	87	106	99	0
08	HS18120278-0	88	90	106	103	0
09	HS18120278-0	87	90	105	102	0
10						
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28						

QC LIMITS

SMC1 (DCE) = 1,2-Dichloroethane-d4 (0-130)
 SMC2 = Dibromofluoromethane (0-130)
 SMC3 (TOL) = Toluene-d8 (0-130)
 OTHER = 4-Bromofluorobenzene (0-130)

Column to be used to flag recovery values

* Values outside of contract required QC limits

D System Monitoring Compound diluted out



FORM 4
VOLATILE METHOD BLANK SUMMARY

CLIENT SAMPLE NO.

VBLKW-181218

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Lab File ID: U121806

Lab Sample ID: VBLKW-181218

Date Analyzed: 12/18/18

Time Analyzed: 1334

GC Column: DB624 ID: 0.18 (mm)

Heated Purge: (Y/N) N

Instrument ID: VOA9

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES, MS and MSD:

	SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	TIME ANALYZED
01	VLCSW-181218	VLCSW-1812018	U121807	1358
02	HS18120278-0	HS18120278-05	U121809	1448
03	HS18120278-0	HS18120278-02	U121810	1513
04	HS18120278-0	HS18120278-04	U121811	1537
05	HS18120278-0	HS18120278-03	U121812	1602
06	HS18120278-0	HS18120278-01	U121813	1627
07	HS18120278-0	HS18120278-03M	U121816	1741
08	HS18120278-0	HS18120278-03M	U121817	1806
09				
10				
11				
12				
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29				
30				

COMMENTS:



FORM 5
VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name: ALS LABORATORY GROUP Contract:
 Lab Code: ALS-HS Case No.: SAS No.: SDG No.: HS18120278
 Lab File ID: U121801 BFB Injection Date: 12/18/18
 Instrument ID: VOA9 BFB Injection Time: 1130
 GC Column: DB624 ID: 0.25 (mm) Heated Purge: (Y/N) N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	19.0
75	30.0 - 60.0% of mass 95	49.8
95	Base Peak, 100% relative abundance	100.0
96	5.0 - 9.0% of mass 95	6.8
173	Less than 2.0% of mass 174	0.6 (0.9)1
174	Greater than 50.0% of mass 95	67.5
175	5.0 - 9.0% of mass 174	5.0 (7.5)1
176	95.0 - 101.0% of mass 174	64.3 (95.2)1
177	5.0 - 9.0% of mass 176	4.4 (6.8)2

1-Value is % mass 174

2-Value is % mass 176

THIS CHECK APPLIES TO THE FOLLOWING SAMPLES, MS, MSD, BLANKS, AND STANDARDS:

	EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
01	VSTD050	VSTD050	U121802	12/18/18	1154
02	VBLKW-181218	VBLKW-181218	U121806	12/18/18	1334
03	VLCSW-181218	VLCSW-1812018	U121807	12/18/18	1358
04	HS18120278-0	HS18120278-05	U121809	12/18/18	1448
05	HS18120278-0	HS18120278-02	U121810	12/18/18	1513
06	HS18120278-0	HS18120278-04	U121811	12/18/18	1537
07	HS18120278-0	HS18120278-03	U121812	12/18/18	1602
08	HS18120278-0	HS18120278-01	U121813	12/18/18	1627
09	HS18120278-0	HS18120278-03M	U121816	12/18/18	1741
10	HS18120278-0	HS18120278-03M	U121817	12/18/18	1806
11	VSTD050-END	VSTD050-END	U121828	12/18/18	2238
12					
13					
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18					
19					
20					
21					
22					

page 1 of 1

FORM V VOA



FORM 7B
VOLATILE CALIBRATION VERIFICATION SUMMARY

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Instrument ID: VOA9

Calibration Date: 12/18/18

Time: 1154

Lab File ID: U121802

Init. Calib. Date(s): 11/13/18

11/13/18

Init. Calib. Times: 1123

1528

GC Column: DB624

ID: 0.18 (mm)

COMPOUND	RRF or AMOUNT	RRF50.000 or AMOUNT	CCAL RRF50.000	MIN RRF	%D or %DRIFT	MAX %D or %DRIFT	CURV TYPE
cis-1,3-Dichloropropene	0.5230000	0.6465575	0.6465575	0.2	-23.62	20.00	AVRG
trans-1,3-Dichloropropene	52.675131	50.000000	0.5461798	0.1	-5.35	20.00	LINR
1,3-Dichlorobenzene	1.5740000	1.5484671	1.5484671	0.6	1.62	20.00	AVRG
2,2-Dichloropropane	0.7100000	0.7232802	0.7232802	0.1	-1.87	20.00	AVRG
1,1-Dichloropropene	0.4620000	0.4695681	0.4695681	0.1	-1.64	20.00	AVRG
Dibromomethane	0.2230000	0.2383912	0.2383912	0.1	-6.90	20.00	AVRG
1,2-Dibromoethane	0.3680000	0.4105056	0.4105056	0.1	-11.55	20.00	AVRG
trans-1,2-Dichloroethene	0.6180000	0.6145443	0.6145443	0.1	0.56	20.00	AVRG
1,1,1,2-Tetrachloroethane	0.3150000	0.3490986	0.3490986	0.1	-10.82	20.00	AVRG
1,1,1-Trichloroethane	0.8780000	0.8174927	0.8174927	0.1	6.89	20.00	AVRG
1,1,2,2-Tetrachloroethane	1.1840000	1.2619648	1.2619648	0.3	-6.58	20.00	AVRG
Freon TF	0.5010000	0.4411715	0.4411715	0.1	11.94	20.00	AVRG
1,1,2-Trichloroethane	0.3180000	0.3468275	0.3468275	0.1	-9.06	20.00	AVRG
1,1-Dichloroethane	1.1460000	1.1593607	1.1593607	0.2	-1.16	20.00	AVRG
1,1-Dichloroethene	0.5640000	0.5229133	0.5229133	0.1	7.28	20.00	AVRG
Trichlorofluoromethane	1.0000000	0.8793023	0.8793023	0.1	12.07	20.00	AVRG
1,2,3-Trichlorobenzene	0.9440000	0.9415381	0.9415381	0.1	0.26	20.00	AVRG
Toluene	1.5870000	1.6824304	1.6824304	0.4	-6.01	20.00	AVRG
1,2,4-Trichlorobenzene	0.9830000	0.9593526	0.9593526	0.2	2.40	20.00	AVRG
1,2,4-Trimethylbenzene	2.7770000	2.8110755	2.8110755	0.1	-1.23	20.00	AVRG
Tetrachloroethene	0.2620000	0.2582452	0.2582452	0.2	1.43	20.00	AVRG
Trichloroethene	0.3410000	0.3482725	0.3482725	0.2	-2.13	20.00	AVRG
1,2-Dichlorobenzene	1.5970000	1.5610963	1.5610963	0.4	2.25	20.00	AVRG
1,2-Dichloroethane	0.5230000	0.5358117	0.5358117	0.1	-2.45	20.00	AVRG
1,2-Dichloropropane	0.3560000	0.4081818	0.4081818	0.1	-14.66	20.00	AVRG
1,3,5-Trimethylbenzene	2.6860000	2.6747190	2.6747190	0.1	0.42	20.00	AVRG
1,3-Dichloropropane	0.6820000	0.7530752	0.7530752	0.1	-10.42	20.00	AVRG
1,4-Dichlorobenzene	51.442257	50.000000	1.5842138	0.4	-2.88	20.00	LINR
2-Butanone	0.3510000	0.3780984	0.3780984	0.1	-7.72	20.00	AVRG
2-Chlorotoluene	2.4480000	2.4612724	2.4612724	0.1	-0.54	20.00	AVRG
2-Hexanone	0.2910000	0.3474431	0.3474431	0.1	-19.40	20.00	AVRG
4-Chlorotoluene	2.8250000	2.8609952	2.8609952	0.1	-1.27	20.00	AVRG
tert-Butylbenzene	2.2890000	2.1959507	2.1959507	0.1	4.06	20.00	AVRG
4-Methyl-2-Pentanone	0.4170000	0.4939195	0.4939195	0.1	-18.44	20.00	AVRG
Acetone	116.62060	100.00000	0.2660475	0.1	-16.62	20.00	LINR
Benzene	1.3820000	1.5172987	1.5172987	0.5	-9.79	20.00	AVRG
Bromobenzene	0.8200000	0.8421597	0.8421597	0.1	-2.70	20.00	AVRG
Bromochloromethane	0.2950000	0.3212433	0.3212433	0.1	-8.90	20.00	AVRG
Bromodichloromethane	0.4160000	0.4489122	0.4489122	0.2	-7.91	20.00	AVRG
Bromoform	51.101037	50.000000	0.2324213	0.1	-2.20	20.00	2RDR
Bromomethane	49.310917	50.000000	0.4089300	0.1	1.38	20.00	LINR
Carbon Disulfide	1.7430000	1.7437932	1.7437932	0.1	-0.04	20.00	AVRG

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FORM VII VOA



FORM 7B
VOLATILE CALIBRATION VERIFICATION SUMMARY

Lab Name: ALS LABORATORY GROUP Contract:

Lab Code: ALS-HS Case No.: SAS No.: SDG No.: HS1812027

Instrument ID: VOA9 Calibration Date: 12/18/18 Time: 2238

Lab File ID: U121828 Init. Calib. Date(s): 11/13/18 11/13/18

Init. Calib. Times: 1123 1528

GC Column: DB624 ID: 0.18 (mm)

COMPOUND	RRF or AMOUNT	RRF50.000 or AMOUNT	CCAL RRF50.000	MIN RRF	%D or %DRIFT	MAX %D or %DRIFT	CURV TYPE
Carbon Tetrachloride	0.3610000	0.3940237	0.3940237	0.1	-9.15	50.00	AVRG
Chlorobenzene	1.0320000	1.1394195	1.1394195	0.5	-10.41	50.00	AVRG
Chloroethane	0.5220000	0.6632847	0.6632847	0.1	-27.07	50.00	AVRG
Chloroform	1.1510000	1.2121159	1.2121159	0.2	-5.31	50.00	AVRG
Chloromethane	53.566027	50.000000	0.7153657	0.1	-7.13	50.00	LINR
cis-1,2-Dichloroethene	0.7080000	0.7735681	0.7735681	0.1	-9.26	50.00	AVRG
Dibromochloromethane	0.3160000	0.3813545	0.3813545	0.1	-20.68	50.00	AVRG
Dichlorodifluoromethane	52.388430	50.000000	0.7750621	0.1	-4.78	50.00	LINR
Ethylbenzene	0.5220000	0.5882191	0.5882191	0.1	-12.68	50.00	AVRG
Hexachlorobutadiene	55.694377	50.000000	0.3462056	0.1	-11.39	50.00	2RDR
Isopropylbenzene	1.5190000	1.7162602	1.7162602	0.1	-12.99	50.00	AVRG
m,p-Xylenes	0.6430000	0.7307393	0.7307393	0.1	-13.64	50.00	AVRG
Methylene Chloride	59.054019	50.000000	0.7518827	0.1	-18.11	50.00	LINR
n-Butylbenzene	55.289556	50.000000	2.9039165	0.5	-10.58	50.00	2RDR
n-Propylbenzene	3.9460000	4.4409755	4.4409755	0.1	-12.54	50.00	AVRG
Naphthalene	2.9610000	3.6557846	3.6557846	0.2	-23.46	50.00	AVRG
o-Xylene	0.6430000	0.7445051	0.7445051	0.3	-15.79	50.00	AVRG
sec-Butylbenzene	3.2850000	3.6733768	3.6733768	0.1	-11.82	50.00	AVRG
Styrene	1.0860000	1.3008233	1.3008233	0.3	-19.78	50.00	AVRG
Vinyl Chloride	59.125421	50.000000	0.9653842	0.1	-18.25	50.00	LINR
1,2,3-Trichloropropane	1.2360000	1.4416158	1.4416158	0.1	-16.64	50.00	AVRG
p-Isopropyltoluene	2.7380000	3.1397602	3.1397602	0.1	-14.67	50.00	AVRG
1,2-Dibromo-3-Chloropropane	53.275827	50.000000	0.1953092	0.05	-6.55	50.00	2RDR
1,2-Dichloroethane-d4	43.667004	50.000000	0.6634090	0.1	12.66	50.00	LINR
Dibromofluoromethane	45.002139	50.000000	0.5187186	0.1	10.0	50.00	LINR
Toluene-d8	52.704142	50.000000	1.3447872	0.1	-5.41	50.00	LINR
4-Bromofluorobenzene	51.164093	50.000000	0.5056900	0.1	-2.33	50.00	LINR



FORM 8
VOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Lab File ID (Standard): U121802

Date Analyzed: 12/18/18

Instrument ID: VOA9

Time Analyzed: 1154

GC Column: DB624

ID: 0.18 (mm)

Heated Purge: (Y/N) N

	IS1 (DCB) AREA #	RT #	IS2 (CBZ) AREA #	RT #	IS3 (DFB) AREA #	RT #
=====	=====	=====	=====	=====	=====	=====
12 HOUR STD	323219	10.24	657840	8.25	710257	5.63
UPPER LIMIT	646438	10.74	1315680	8.75	1420514	6.13
LOWER LIMIT	161610	9.74	328920	7.75	355129	5.13
=====	=====	=====	=====	=====	=====	=====
CLIENT SAMPLE NO.						
=====	=====	=====	=====	=====	=====	=====
01 VBLKW-181218	292418	10.24	636133	8.25	694572	5.63
02 VLCSW-181218	316393	10.24	655559	8.25	714075	5.63
03 HS18120278-0	290869	10.24	640292	8.25	702137	5.63
04 HS18120278-0	291140	10.24	631599	8.25	697785	5.63
05 HS18120278-0	294161	10.24	641402	8.25	700337	5.63
06 HS18120278-0	290407	10.24	635273	8.25	700993	5.63
07 HS18120278-0	293976	10.24	634287	8.25	701381	5.63
08 HS18120278-0	308103	10.24	641586	8.25	699438	5.63
09 HS18120278-0	315859	10.24	656441	8.25	706692	5.63
10						
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						

IS1 (DCB) = 1,4-Dichlorobenzene-d4

IS2 (CBZ) = Chlorobenzene-d5

IS3 (DFB) = 1,4-Difluorobenzene

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = - 50% of internal standard area

RT UPPER LIMIT = + 0.50 minutes of internal standard RT

RT LOWER LIMIT = - 0.50 minutes of internal standard RT

Column used to flag values outside QC limits with an asterisk.

* Values outside of QC limits.

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FORM VIII VOA



FORM 8
VOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: ALS LABORATORY GROUP

Contract:

Lab Code: ALS-HS

Case No.:

SAS No.:

SDG No.: HS18120278

Lab File ID (Standard): U121802

Date Analyzed: 12/18/18

Instrument ID: VOA9

Time Analyzed: 1154

GC Column: DB624

ID: 0.18 (mm)

Heated Purge: (Y/N) N

	IS4 AREA #	RT #	AREA #	RT #	AREA #	RT #
=====	=====	=====	=====	=====	=====	=====
12 HOUR STD	402866	4.89				
UPPER LIMIT	805732	5.39				
LOWER LIMIT	201433	4.39				
=====	=====	=====	=====	=====	=====	=====
CLIENT SAMPLE NO.						
=====	=====	=====	=====	=====	=====	=====
01 VBLKW-181218	385775	4.90				
02 VLCSW-181218	400551	4.90				
03 HS18120278-0	395597	4.90				
04 HS18120278-0	388196	4.90				
05 HS18120278-0	392367	4.89				
06 HS18120278-0	386598	4.89				
07 HS18120278-0	389639	4.89				
08 HS18120278-0	392642	4.90				
09 HS18120278-0	400954	4.90				
10						
11						
12						
13						
14						
15						
16						
17						
18						
19						
20						

IS4 = Pentafluorobenzene

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = - 50% of internal standard area

RT UPPER LIMIT = + 0.50 minutes of internal standard RT

RT LOWER LIMIT = - 0.50 minutes of internal standard RT

Column used to flag values outside QC limits with an asterisk.

* Values outside of QC limits.



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121801.D

Page 1

Date : 18-DEC-2018 11:30

Client ID: BFB

Instrument: VOA9.i

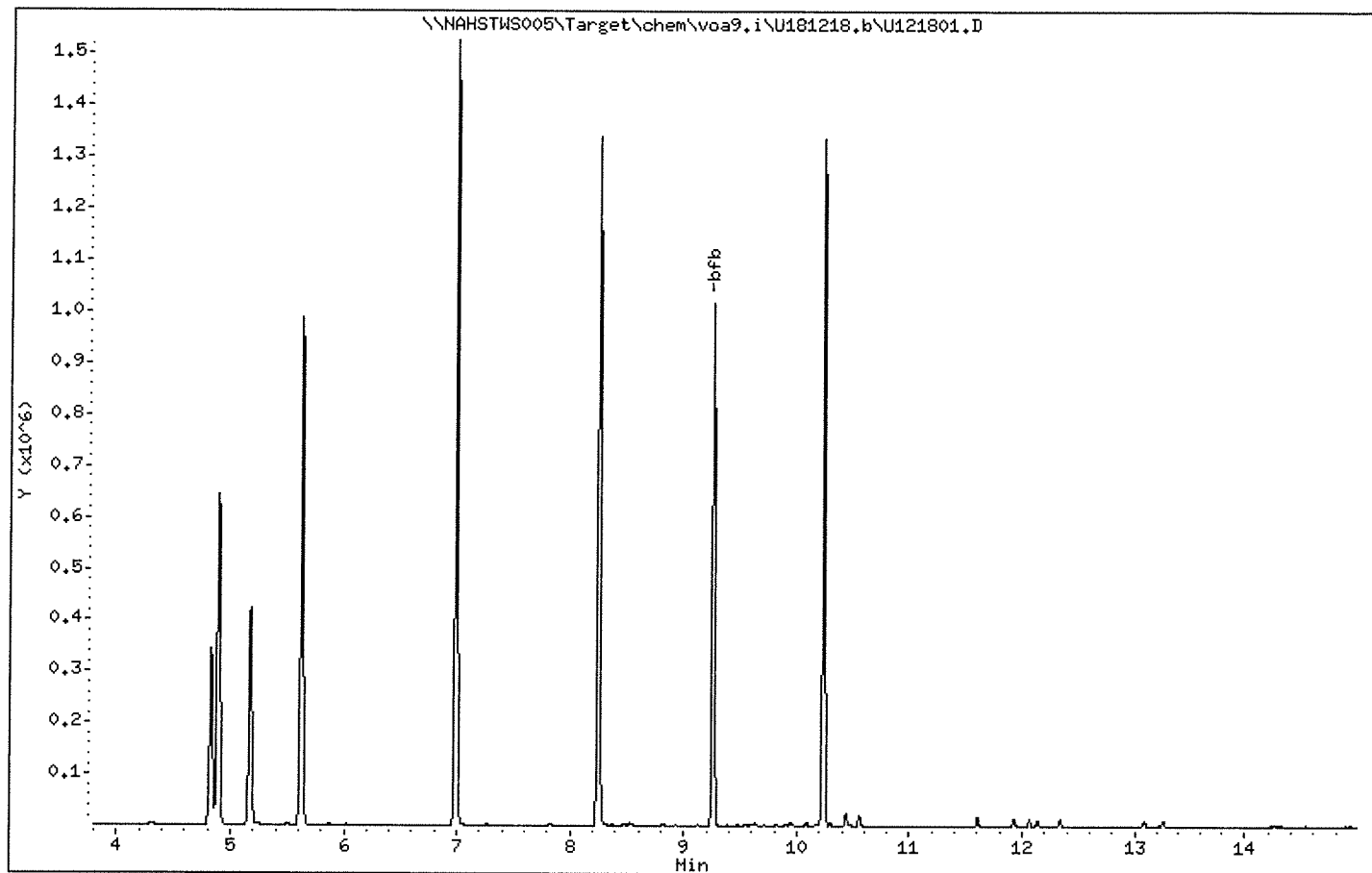
Sample Info: BFB;BFB;3;;BFB

Volume Injected (uL): 2.0

Operator: PC

Column phase: DB624

Column diameter: 0,25



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121801.D

Page 2

Date : 18-DEC-2018 11:30

Client ID: BFB

Instrument: VOA9.i

Sample Info: BFB;BFB;3;;BFB

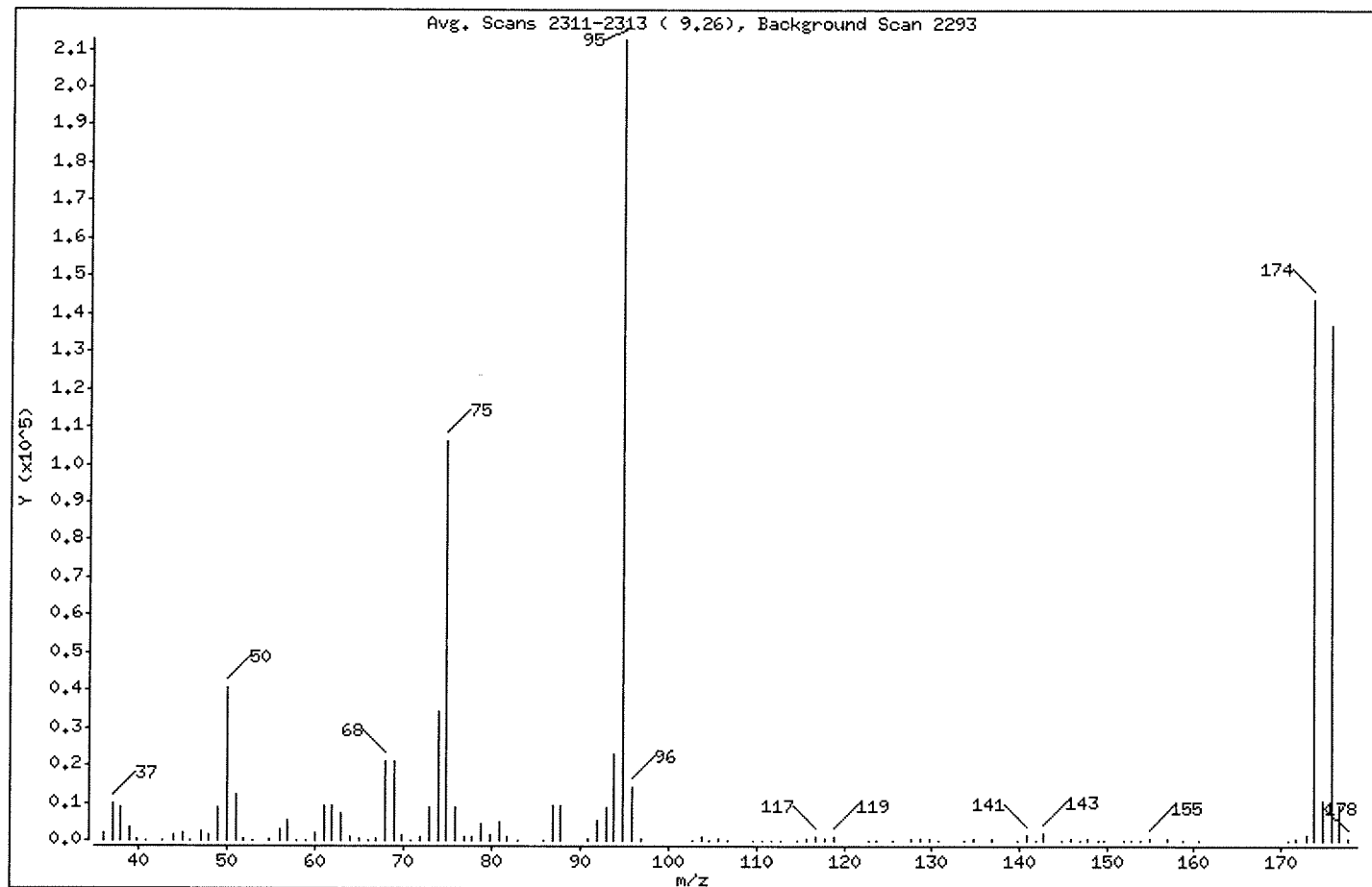
Volume Injected (uL): 2.0

Operator: PC

Column phase: DB624

Column diameter: 0.25

1 bfb



m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
95	Base Peak, 100% relative abundance	100.00
50	15.00 - 40.00% of mass 95	18.97
75	30.00 - 60.00% of mass 95	49.82
96	5.00 - 9.00% of mass 95	6.77
173	Less than 2.00% of mass 174	0.58 (0.86)
174	Greater than 50.00% of mass 95	67.51
175	5.00 - 9.00% of mass 174	5.03 (7.45)
176	95.00 - 101.00% of mass 174	64.26 (95.19)
177	5.00 - 9.00% of mass 176	4.37 (6.81)



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121801.D

Page 3

Date : 18-DEC-2018 11:30

Client ID: BFB

Instrument: VOA9.i

Sample Info: BFB;BFB;3;;BFB

Volume Injected (uL): 2.0

Operator: PC

Column phase: DB624

Column diameter: 0.25

Data File: U121801.D

Spectrum: Avg. Scans 2311-2313 (9.26), Background Scan 2293

Location of Maximum: 94.95

Number of points: 105

m/z	Y	m/z	Y	m/z	Y	m/z	Y
36.00	1745	65.00	498	95.90	14398	140.80	1565
37.00	9863	66.00	107	96.90	464	141.80	168
38.00	8582	67.00	526	102.70	99	142.70	1760
39.00	3513	67.90	21248	103.80	776	144.70	199
39.90	474	68.90	20816	104.70	243	145.70	256
40.90	63	69.90	1621	105.80	661	146.70	84
42.80	125	70.90	95	106.80	195	147.70	442
43.90	1399	71.90	1053	109.70	103	148.80	106
45.00	1717	72.90	8643	110.80	121	149.60	172
45.90	235	73.90	34248	111.80	100	151.80	97
47.00	2542	74.90	105976	112.70	110	152.60	124
47.90	1329	75.90	8769	114.70	104	153.70	83
49.00	8644	76.90	1149	115.70	565	154.70	418
50.00	40360	77.80	942	116.70	942	156.70	306
51.00	12336	78.80	4640	117.80	537	158.70	199
51.90	520	79.90	1305	118.80	798	160.60	120
53.00	58	80.80	4993	122.80	55	170.80	70
54.90	574	81.80	1016	123.70	105	171.70	550
56.00	2742	82.90	170	125.60	74	172.80	1236
57.00	5368	85.80	210	127.70	539	173.70	143616
57.90	243	86.90	9507	128.70	330	174.70	10700
58.90	56	87.80	9344	129.70	649	175.70	136704
60.00	1763	90.80	654	130.70	234	176.70	9304
61.00	9399	91.90	5372	133.60	54	177.70	263
61.90	9161	92.90	8580	134.70	303		
63.00	7143	93.90	22776	136.70	247		
63.90	856	94.90	212736	139.70	98		



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121802.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121802.D
 Lab Smp Id: VSTD050 Client Smp ID: VSTD050
 Inj Date : 18-DEC-2018 11:54
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD050;VSTD050;2;;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 2 Continuing Calibration Sample
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	AMOUNTS					CAL-AMT (ug/l)	ON-COL (ug/l)
		MASS	RT	EXP RT	REL RT	RESPONSE		
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	402866	50.0000		
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	710257	50.0000		
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	657840	50.0000		
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	323219	50.0000		
\$ 30 Dibromofluoromethane	113	4.827	4.827	(0.986)	207074	50.0000	44.58	
\$ 35 1,2-Dichloroethane-d4	65	5.171	5.171	(1.057)	264783	50.0000	43.24	
\$ 48 Toluene-d8	98	6.990	6.990	(0.847)	883203	50.0000	52.61	
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	334527	50.0000	51.45	
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	229651	50.0000	55.34	
31 1,1,1-Trichloroethane	97	4.827	4.827	(0.986)	329340	50.0000	46.53	
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	407891	50.0000	53.27	
138 Freon TF	101	2.397	2.397	(0.490)	177733	50.0000	44.03	
53 1,1,2-Trichloroethane	83	7.421	7.421	(0.900)	228157	50.0000	54.55	
22 1,1-Dichloroethane	63	3.601	3.601	(0.736)	467067	50.0000	50.59	
11 1,1-Dichloroethene	96	2.397	2.397	(0.490)	210664	50.0000	46.32	
32 1,1-Dichloropropene	75	5.003	5.003	(0.889)	333514	50.0000	50.84	
93 1,2,3-Trichlorobenzene	180	12.335	12.335	(1.205)	304323	50.0000	49.87	
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	455503	50.0000	57.01	
90 1,2,4-Trichlorobenzene	180	11.923	11.923	(1.165)	310081	50.0000	48.78	
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	908593	50.0000	50.61	
89 1,2-Dibromo-3-Chloropropane	155	11.233	11.233	(1.097)	59352	50.0000	50.17	
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	270047	50.0000	55.76	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121802.D Page 2
 Report Date: 25-Jan-2019 20:37

Compounds	QUANT	SIG						AMOUNTS	
			MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)	ON-COL (ug/l)
88 1,2-Dichlorobenzene	146		10.570	10.570	(1.033)	504576	50.0000	48.88	
33 1,2-Dichloroethane	62		5.250	5.250	(0.933)	380564	50.0000	51.20	
42 1,2-Dichloropropane	63		6.079	6.079	(1.081)	289914	50.0000	57.33	
75 1,3,5-Trimethylbenzene	105		9.625	9.625	(0.940)	864520	50.0000	49.78	
83 1,3-Dichlorobenzene	146		10.180	10.180	(0.995)	500494	50.0000	49.18	
54 1,3-Dichloropropane	76		7.563	7.563	(0.917)	495403	50.0000	55.16	
84 1,4-Dichlorobenzene	146		10.255	10.255	(1.002)	512048	50.0000	51.44	
26 2,2-Dichloropropane	77		4.272	4.272	(0.873)	291385	50.0000	50.96	
24 2-Butanone	43		4.335	4.335	(0.886)	304646	100.000	107.57	
76 2-Chlorotoluene	91		9.546	9.546	(0.933)	795530	50.0000	50.26	
52 2-Hexanone	43		7.649	7.649	(0.927)	457124	100.000	119.53	
77 4-Chlorotoluene	91		9.640	9.640	(0.942)	924728	50.0000	50.63	
82 p-Isopropyltoluene	119		10.210	10.210	(0.997)	876034	50.0000	49.49	
45 4-Methyl-2-Pentanone	43		6.915	6.915	(0.838)	649840	100.000	118.41	
10 Acetone	43		2.476	2.476	(0.506)	214363	100.000	116.62	
37 Benzene	78		5.216	5.216	(0.927)	1077672	50.0000	54.89	
74 Bromobenzene	156		9.381	9.381	(0.917)	272202	50.0000	51.33	
29 Bromochloromethane	128		4.553	4.553	(0.930)	129418	50.0000	54.49	
39 Bromodichloromethane	83		6.348	6.348	(1.129)	318843	50.0000	54.02	
66 Bromoform	173		8.984	8.984	(1.089)	152896	50.0000	51.10 (T)	
6 Bromomethane	94		1.663	1.663	(0.340)	164744	50.0000	49.31	
19 Carbon Disulfide	76		2.585	2.585	(0.528)	1405030	100.000	100.04	
34 Carbon Tetrachloride	117		4.991	4.991	(0.887)	250572	50.0000	48.89	
59 Chlorobenzene	112		8.275	8.275	(1.003)	703155	50.0000	51.78	
7 Chloroethane	64		1.745	1.745	(0.357)	237474	50.0000	56.49	
28 Chloroform	83		4.654	4.654	(0.951)	453514	50.0000	48.89	
3 Chloromethane	50		1.336	1.336	(0.273)	293918	50.0000	54.62	
27 cis-1,2-Dichloroethene	96		4.283	4.283	(0.875)	288356	50.0000	50.51	
46 cis-1,3-Dichloropropene	75		6.757	6.757	(1.201)	459222	50.0000	61.79	
55 Dibromochloromethane	129		7.758	7.758	(0.940)	241039	50.0000	57.90	
44 Dibromomethane	93		6.187	6.187	(1.100)	169319	50.0000	53.42	
2 Dichlorodifluoromethane	85		1.202	1.202	(0.246)	271170	50.0000	45.62	
61 Ethylbenzene	106		8.369	8.369	(1.015)	353954	50.0000	51.56	
91 Hexachlorobutadiene	225		12.065	12.065	(1.179)	105243	50.0000	52.56	
67 Isopropylbenzene	105		9.126	9.126	(1.106)	1002208	50.0000	50.13	
62 m,p-Xylenes	106		8.474	8.474	(1.027)	880711	100.000	104.10	
17 Methylene Chloride	84		2.866	2.866	(0.586)	281642	50.0000	54.85	
87 n-Butylbenzene	91		10.558	10.558	(1.031)	812665	50.0000	48.06	
73 n-Propylbenzene	91		9.475	9.475	(0.926)	1254341	50.0000	49.17	
92 Naphthalene	128		12.133	12.133	(1.185)	1050116	50.0000	54.86	
63 o-Xylene	106		8.811	8.811	(1.068)	452543	50.0000	53.45	
81 sec-Butylbenzene	105		10.086	10.086	(0.985)	1011602	50.0000	47.63	
64 Styrene	104		8.826	8.826	(1.070)	802660	50.0000	56.15	
78 tert-Butylbenzene	119		9.902	9.902	(0.967)	709773	50.0000	47.95	
56 Tetrachloroethene	164		7.522	7.522	(0.912)	169884	50.0000	49.29	
50 Toluene	91		7.046	7.046	(0.854)	1106770	50.0000	52.99	
20 trans-1,2-Dichloroethene	96		3.136	3.136	(0.641)	247579	50.0000	49.67	
51 trans-1,3-Dichloropropene	75		7.259	7.259	(1.291)	387928	50.0000	52.67	
38 Trichloroethene	130		5.861	5.861	(1.042)	247363	50.0000	51.11	
8 Trichlorofluoromethane	101		1.948	1.948	(0.398)	354241	50.0000	43.98	
5 Vinyl Chloride	62		1.415	1.415	(0.289)	340847	50.0000	51.87	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121802.D Page 3
Report Date: 25-Jan-2019 20:37

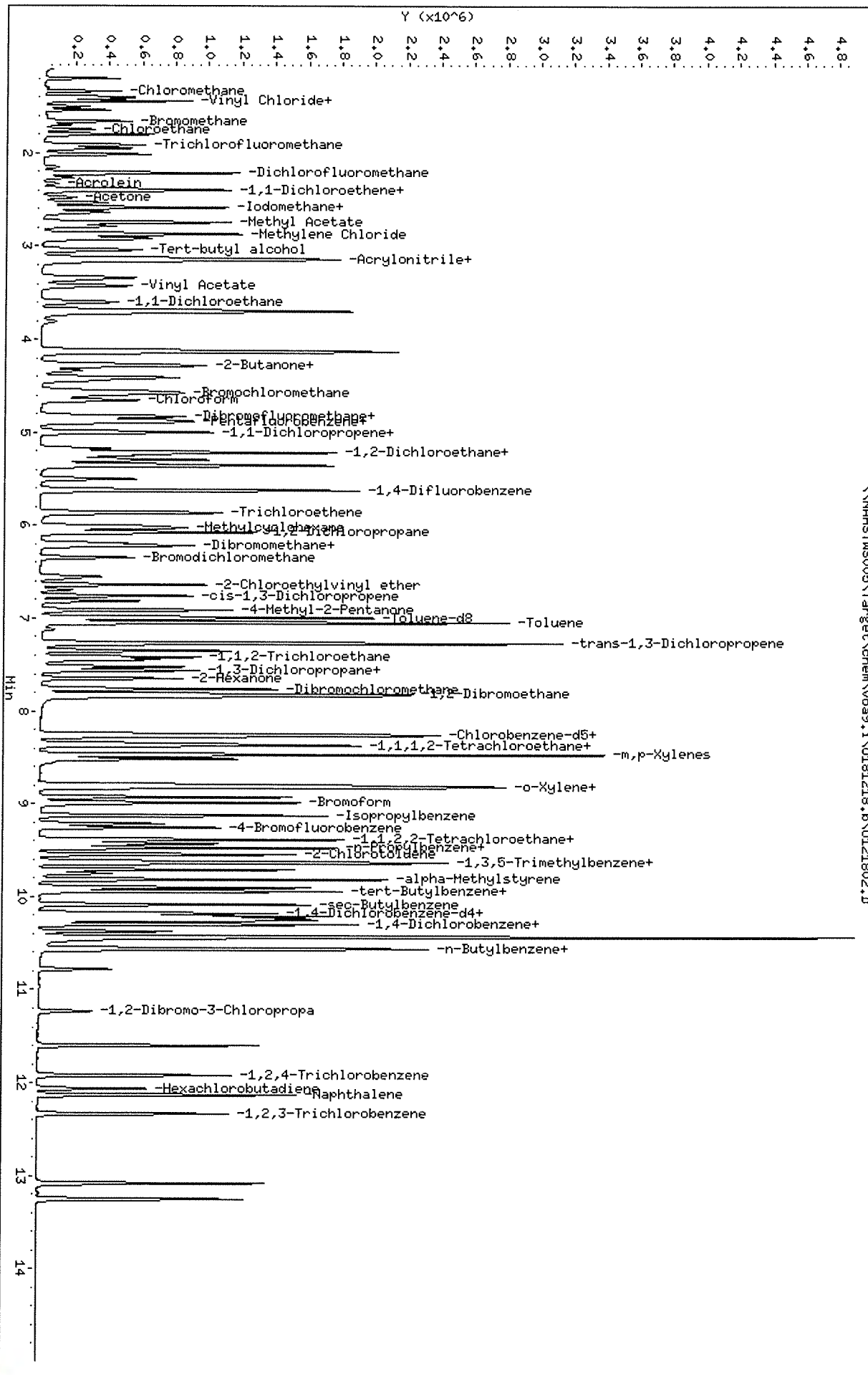
QC Flag Legend

T - Target compound detected outside RT window.



Data File: \\NAHSTMS005\Target\chem\voa9.i\UJ81218.b\UJ21802.D
 Date : 18-DEC-2018 11:54
 Client ID: VSTD050
 Sample Info: VSTD050;VSTD050;2;;
 Purge Volume: 5.0
 Column phase: DB624

Instrument: VOA9.i
 Operator: PC
 Column diameter: 0.18



\\NAHSTMS005\Target\chem\voa9.i\UJ81218.b\UJ21802.D



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121806.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121806.D
 Lab Smp Id: VBLKW-181218 Client Smp ID: VBLKW-181218
 Inj Date : 18-DEC-2018 13:34
 Operator : PC Inst ID: VOA9.i
 Smp Info : VBLKW-181218;VBLKW-181218;3;;BLANK
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 6 QC Sample: BLANK
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

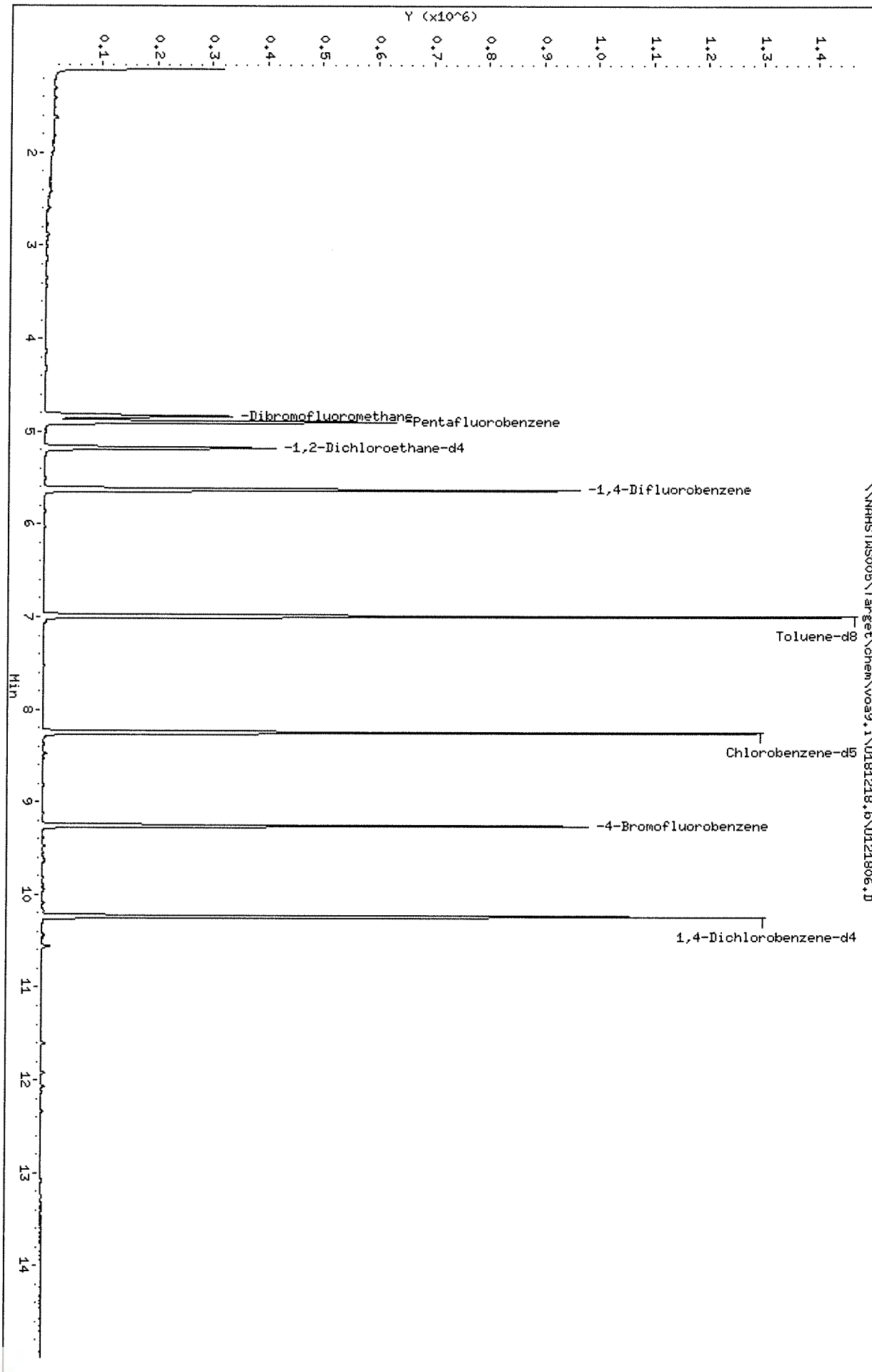
Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.894	(1.000)	385775	50.0000	
* 36 1,4-Difluorobenzene	114	5.629	5.625	(1.000)	694572	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	636133	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	292418	50.0000	
\$ 30 Dibromofluoromethane	113	4.834	4.827	(0.987)	194447	43.6934	43.69
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.171	(1.057)	264506	45.1904	45.19
\$ 48 Toluene-d8	98	6.989	6.990	(0.847)	845972	52.1004	52.10
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	307890	48.9242	48.92



Data File: \\NAHSTMS005\Target\chem\voa9.i\U181218.b\U121806.D
Date : 18-DEC-2018 13:34
Client ID: VBULK-181218
Sample Info: VBULK-181218;VBULK-181218;3;:BLANK
Purge Volume: 5.0
Column phase: DB624

Instrument: VOA9.1
Operator: PC
Column diameter: 0.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121807.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121807.D
 Lab Smp Id: VLCSW-1812018 Client Smp ID: VLCSW-181218
 Inj Date : 18-DEC-2018 13:58
 Operator : PC Inst ID: VOA9.i
 Smp Info : VLCSW-1812018;VLCSW-181218;3;;LCS
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 7 QC Sample: LCS
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG						CONCENTRATIONS	
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168		4.897	4.894	(1.000)	400551	50.0000	
* 36 1,4-Difluorobenzene	114		5.625	5.625	(1.000)	714075	50.0000	
* 47 Chlorobenzene-d5	117		8.249	8.249	(1.000)	655559	50.0000	
* 70 1,4-Dichlorobenzene-d4	152		10.235	10.236	(1.000)	316393	50.0000	
\$ 30 Dibromofluoromethane	113		4.830	4.827	(0.986)	205791	44.5608	44.56
\$ 35 1,2-Dichloroethane-d4	65		5.175	5.171	(1.057)	268992	44.2250	44.22
\$ 48 Toluene-d8	98		6.989	6.990	(0.847)	881545	52.7015	52.70
\$ 69 4-Bromofluorobenzene	95		9.257	9.257	(1.122)	331590	51.1768	51.17
60 1,1,1,2-Tetrachloroethane	131		8.350	8.350	(1.012)	89740	21.7022	21.70
31 1,1,1-Trichloroethane	97		4.830	4.827	(0.986)	141124	20.0576	20.05
68 1,1,2,2-Tetrachloroethane	83		9.392	9.392	(0.918)	161937	21.6080	21.60
138 Freon TF	101		2.404	2.397	(0.491)	86910	21.6594	21.65
53 1,1,2-Trichloroethane	83		7.420	7.421	(0.900)	91272	21.8997	21.89
22 1,1-Dichloroethane	63		3.604	3.601	(0.736)	192391	20.9601	20.96
11 1,1-Dichloroethene	96		2.401	2.397	(0.490)	93258	20.6265	20.62
32 1,1-Dichloropropene	75		5.006	5.003	(0.890)	149174	22.6192	22.61
93 1,2,3-Trichlorobenzene	180		12.335	12.335	(1.205)	132556	22.1911	22.19
71 1,2,3-Trichloropropane	75		9.426	9.426	(0.921)	171993	21.9927	21.99
90 1,2,4-Trichlorobenzene	180		11.926	11.923	(1.165)	136376	21.9186	21.91
79 1,2,4-Trimethylbenzene	105		9.943	9.943	(0.971)	405685	23.0850	23.08
89 1,2-Dibromo-3-Chloropropane	155		11.233	11.233	(1.097)	21955	19.6877	19.68
57 1,2-Dibromoethane	107		7.851	7.852	(0.952)	107069	22.1853	22.18



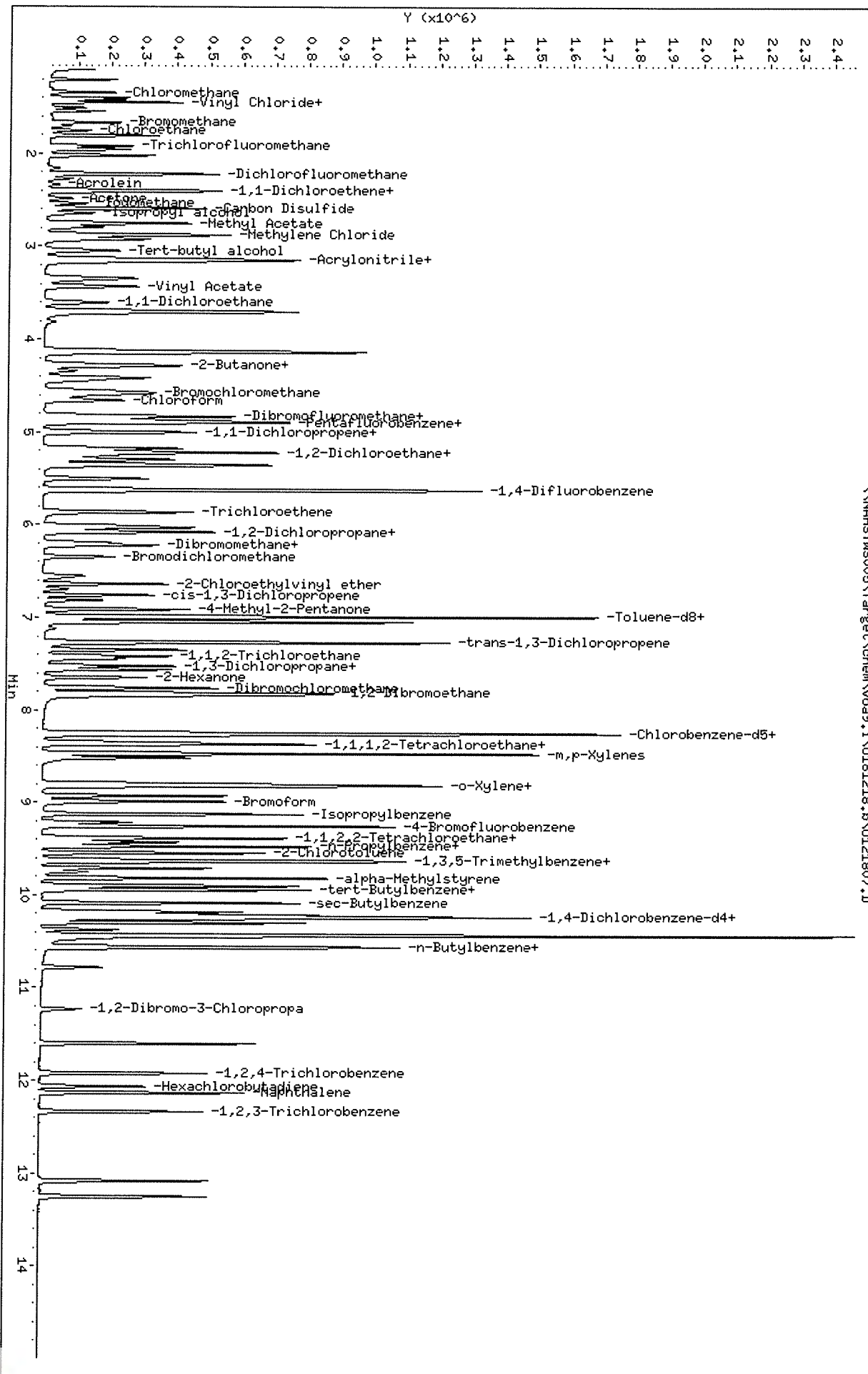
Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121807.D Page 2
 Report Date: 25-Jan-2019 20:37

Compounds	QUANT SIG MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
88 1,2-Dichlorobenzene	146	10.573	10.570	(1.033)	214077	21.1877	21.18
33 1,2-Dichloroethane	62	5.253	5.250	(0.934)	155334	20.7882	20.78
42 1,2-Dichloropropane	63	6.082	6.079	(1.081)	117753	23.1612	23.16
75 1,3,5-Trimethylbenzene	105	9.624	9.625	(0.940)	394362	23.1986	23.19
83 1,3-Dichlorobenzene	146	10.179	10.180	(0.995)	216187	21.7022	21.70
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	199341	22.2751	22.27
84 1,4-Dichlorobenzene	146	10.254	10.255	(1.002)	223543	22.9189	22.91
26 2,2-Dichloropropane	77	4.275	4.272	(0.873)	120949	21.2777	21.27
24 2-Butanone	43	4.339	4.335	(0.886)	119439	42.4199	42.41
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	348424	22.4891	22.48
52 2-Hexanone	43	7.649	7.649	(0.927)	177505	46.5796	46.57
77 4-Chlorotoluene	91	9.639	9.640	(0.942)	405202	22.6659	22.66
82 p-Isopropyltoluene	119	10.209	10.210	(0.997)	413164	23.8459	23.84
45 4-Methyl-2-Pentanone	43	6.914	6.915	(0.838)	254420	46.5221	46.52
10 Acetone	43	2.483	2.476	(0.507)	83187	42.8169	42.81
37 Benzene	78	5.220	5.216	(0.928)	446907	22.6413	22.64
74 Bromobenzene	156	9.381	9.381	(0.917)	112076	21.5942	21.59
29 Bromochloromethane	128	4.556	4.553	(0.930)	52445	22.2096	22.20
39 Bromodichloromethane	83	6.348	6.348	(1.129)	123749	20.8569	20.85
66 Bromoform	173	8.983	8.984	(1.089)	55015	19.2965	19.29
6 Bromomethane	94	1.670	1.663	(0.341)	73853	23.2782	23.27
19 Carbon Disulfide	76	2.592	2.585	(0.529)	604634	43.3037	43.30
34 Carbon Tetrachloride	117	4.999	4.991	(0.889)	111990	21.7369	21.73
59 Chlorobenzene	112	8.275	8.275	(1.003)	294755	21.7849	21.78
7 Chloroethane	64	1.752	1.745	(0.358)	96365	23.0575	23.05
28 Chloroform	83	4.657	4.654	(0.951)	184450	20.0006	20.00
3 Chloromethane	50	1.344	1.336	(0.274)	127839	24.0254	24.02
27 cis-1,2-Dichloroethene	96	4.286	4.283	(0.875)	118161	20.8186	20.81
46 cis-1,3-Dichloropropene	75	6.757	6.757	(1.201)	170301	22.7929	22.79
55 Dibromochloromethane	129	7.758	7.758	(0.940)	90639	21.8502	21.85
44 Dibromomethane	93	6.191	6.187	(1.101)	68292	21.4333	21.43
2 Dichlorodifluoromethane	85	1.209	1.202	(0.247)	125172	21.7200	21.71
61 Ethylbenzene	106	8.372	8.369	(1.015)	153825	22.4885	22.48
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	50567	26.6066	26.60
67 Isopropylbenzene	105	9.126	9.126	(1.106)	460914	23.1350	23.13
62 m,p-Xylenes	106	8.474	8.474	(1.027)	384675	45.6281	45.62
17 Methylene Chloride	84	2.873	2.866	(0.587)	117281	22.5682	22.56
87 n-Butylbenzene	91	10.558	10.558	(1.031)	393164	24.1807	24.18
73 n-Propylbenzene	91	9.475	9.475	(0.926)	584534	23.4126	23.41
92 Naphthalene	128	12.132	12.133	(1.185)	427693	22.8281	22.82
63 o-Xylene	106	8.811	8.811	(1.068)	193177	22.8984	22.89
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	490713	23.6050	23.60
64 Styrene	104	8.826	8.826	(1.070)	333195	23.3919	23.39
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	335135	23.1322	23.13
56 Tetrachloroethene	164	7.525	7.522	(0.912)	76788	22.3606	22.36
50 Toluene	91	7.049	7.046	(0.855)	469480	22.5570	22.55
20 trans-1,2-Dichloroethene	96	3.143	3.136	(0.642)	102720	20.7300	20.72
51 trans-1,3-Dichloropropene	75	7.263	7.259	(1.291)	140882	19.9713	19.97
38 Trichloroethene	130	5.865	5.861	(1.043)	106178	21.8218	21.82
8 Trichlorofluoromethane	101	1.955	1.948	(0.399)	163417	20.4069	20.40
5 Vinyl Chloride	62	1.422	1.415	(0.290)	148850	23.0721	23.07



Data File: \\NAHSTMS005\Target\chem\voa9.i\U181218.b\U121807.D
Date: 18-DEC-2018 13:58
Client ID: VLC5M-181218
Sample Info: VLC5M-1812018;VLC5M-181218;3;LCS
Purge Volume: 5.0
Column Phase: DB624

Instrument: V0A9.i
Operator: PC
Column diameter: 0.18



\\NAHSTMS005\Target\chem\voa9.i\U181218.b\U121807.D



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121809.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121809.D
 Lab Smp Id: HS18120278-05 Client Smp ID: HS18120278-05
 Inj Date : 18-DEC-2018 14:48
 Operator : PC Inst ID: VOA9.i
 Smp Info : HS18120278-05;HS18120278-05;;;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 9
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.894	(1.000)	395597	50.0000	
* 36 1,4-Difluorobenzene	114	5.629	5.625	(1.000)	702137	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	640292	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	290869	50.0000	
\$ 30 Dibromofluoromethane	113	4.834	4.827	(0.987)	200474	43.9360	43.93
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.171	(1.057)	268570	44.7280	44.72
\$ 48 Toluene-d8	98	6.990	6.990	(0.847)	864694	52.9340	52.93
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	310925	49.0890	49.08
10 Acetone	43	2.487	2.476	(0.508)	12388	2.69362	2.69(a)

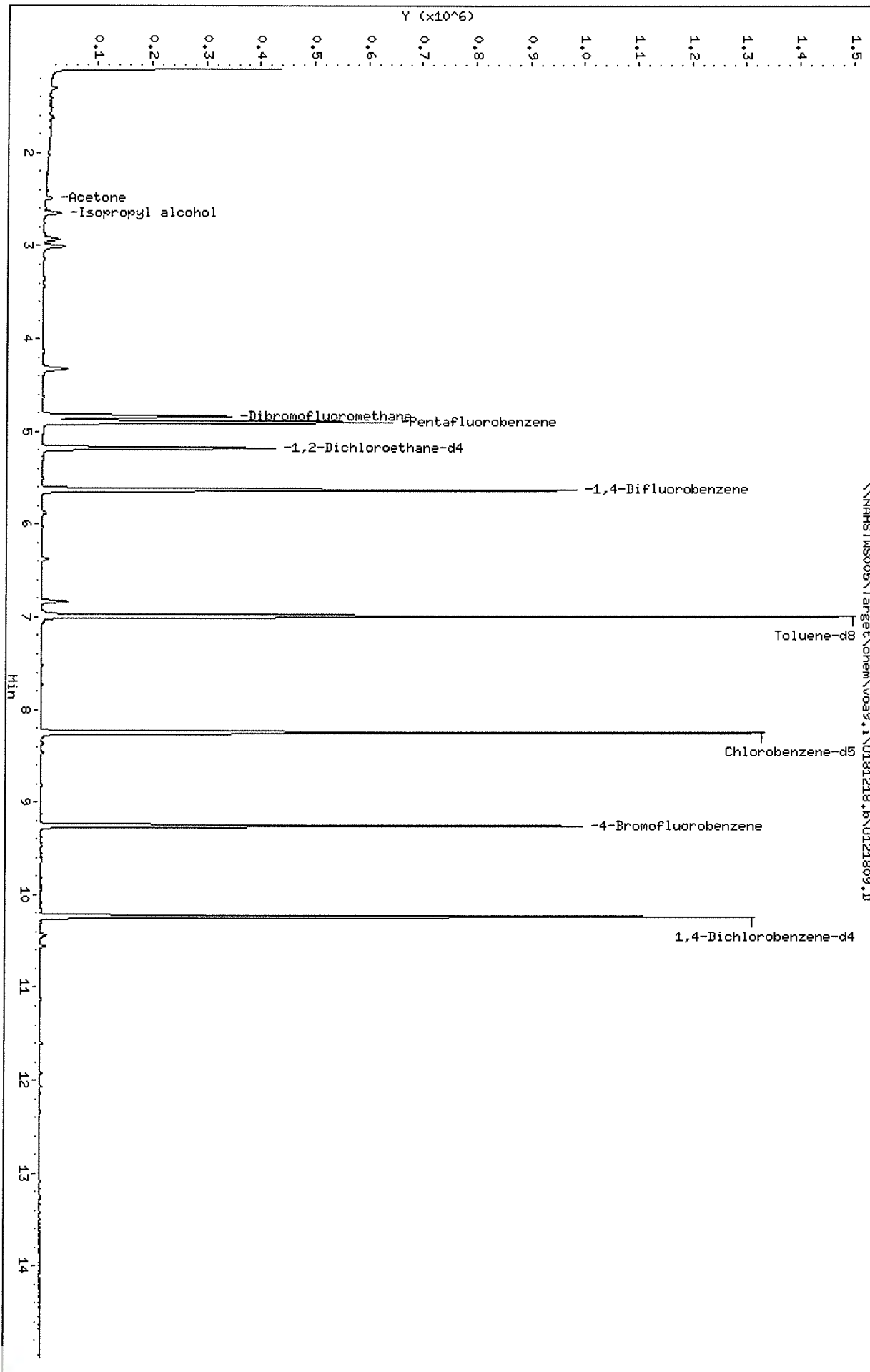
QC Flag Legend

a - Target compound detected but, quantitated amount
 Below Limit Of Quantitation(BLOQ).



Data File: \\NAHSTMS005\Target\chem\voa9.i\U181218.b\U121809.D
Date : 18-DEC-2018 14:48
Client ID: HS18120278-05
Sample Info: HS18120278-05;HS18120278-05;;
Purge Volume: 5.0
Column phase: DB624

Instrument: VDA9.i
Operator: PC
Column diameter: 0.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121809.D

Page 3

Date : 18-DEC-2018 14:48

Client ID: HS18120278-05

Instrument: VOA9.i

Sample Info: HS18120278-05;HS18120278-05;;;

Purge Volume: 5.0

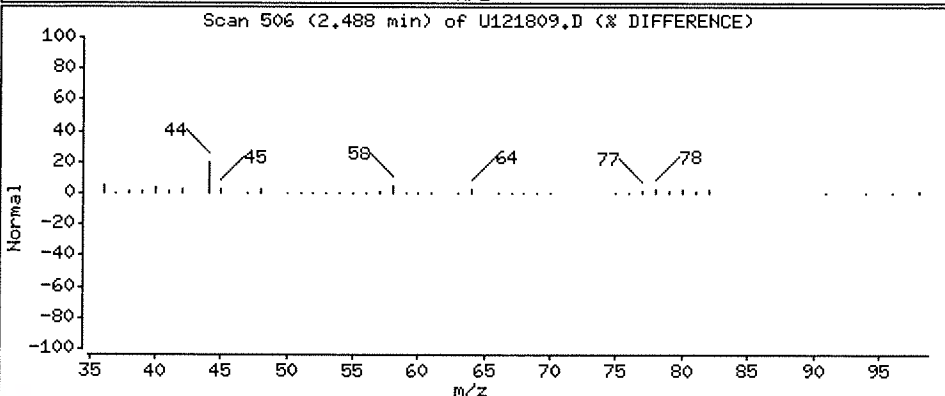
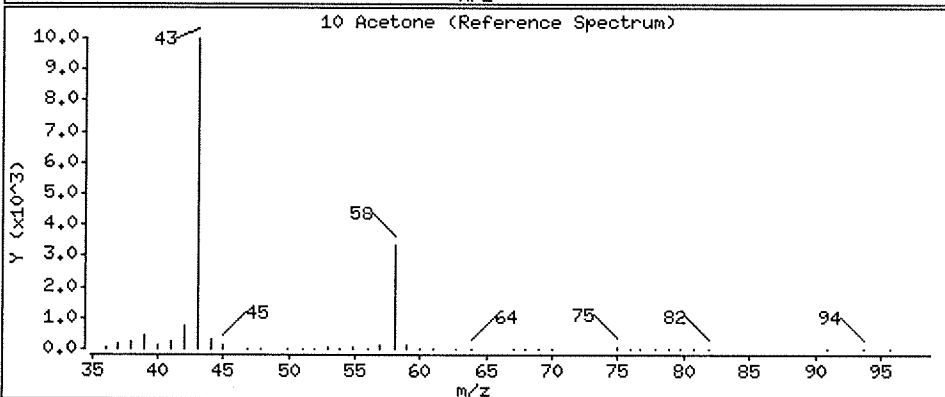
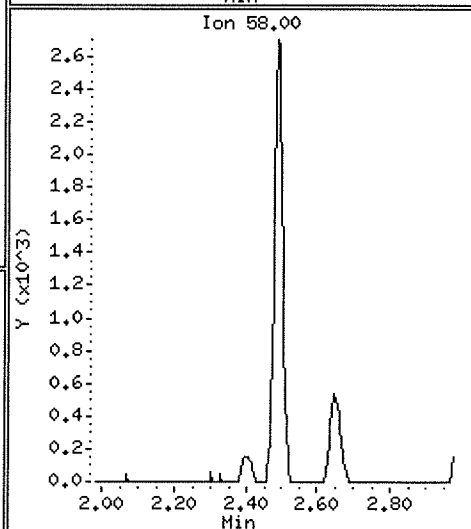
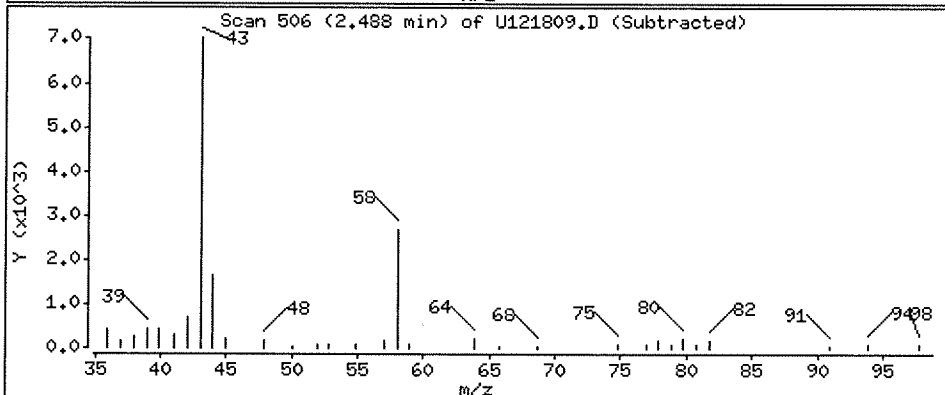
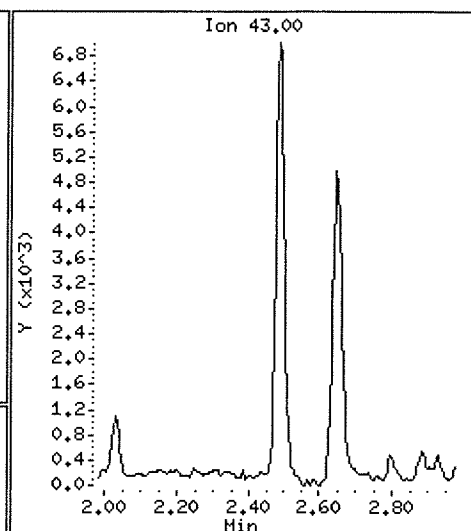
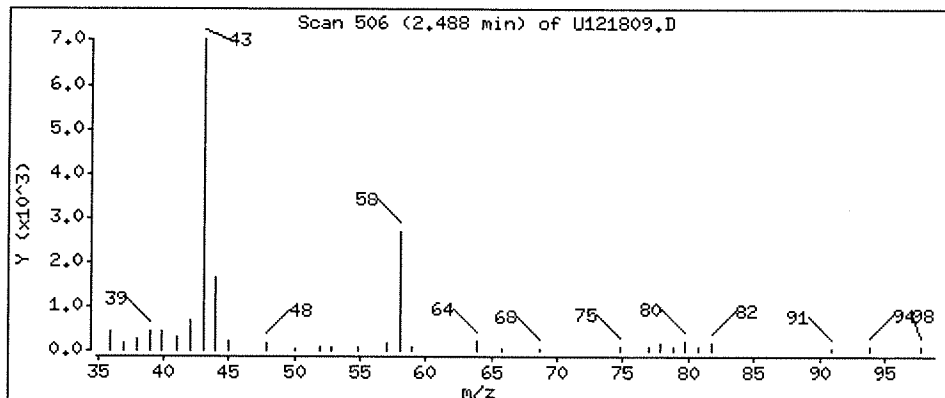
Operator: PC

Column phase: DB624

Column diameter: 0.18

10 Acetone

Concentration: 2.69 ug/l



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121810.D Page 1
 Report Date: 25-Jan-2019 20:37

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Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121810.D
 Lab Smp Id: HS18120278-02 Client Smp ID: HS18120278-02
 Inj Date : 18-DEC-2018 15:13
 Operator : PC Inst ID: VOA9.i
 Smp Info : HS18120278-02;HS18120278-02;;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 10
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

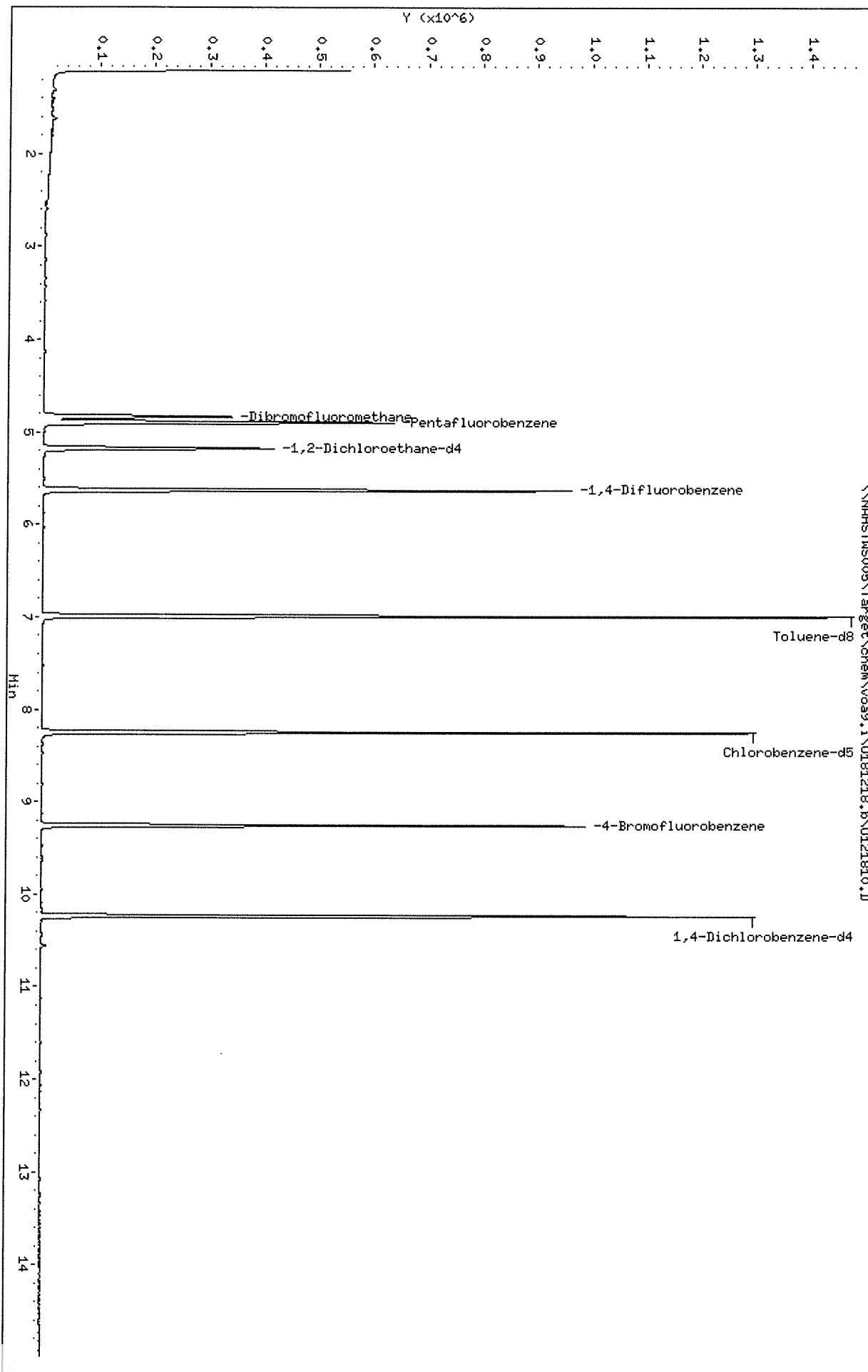
Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG				CONCENTRATIONS		
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.894	(1.000)	388196	50.0000	
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	697785	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	631599	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	291140	50.0000	
\$ 30 Dibromofluoromethane	113	4.834	4.827	(0.987)	197885	44.2028	44.20
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.171	(1.057)	266481	45.2460	45.24
\$ 48 Toluene-d8	98	6.990	6.990	(0.847)	849387	52.7055	52.70
\$ 69 4-Bromofluorobenzene	95	9.258	9.257	(1.122)	305442	48.8827	48.88



Data File: \\NHSTMS005\Target\chem\voa9.i\U181218.b\U121810.D
Date : 18-DEC-2018 15:13
Client ID: HS18120278-02
Sample Info: HS18120278-02;HS18120278-02;;
Purge Volume: 5.0
Column phase: DB624

Instrument: WDA9.i
Operator: PC
Column diameter: 0.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121811.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121811.D
 Lab Smp Id: HS18120278-04 Client Smp ID: HS18120278-04
 Inj Date : 18-DEC-2018 15:37
 Operator : PC Inst ID: VOA9.i
 Smp Info : HS18120278-04;HS18120278-04;;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 11
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

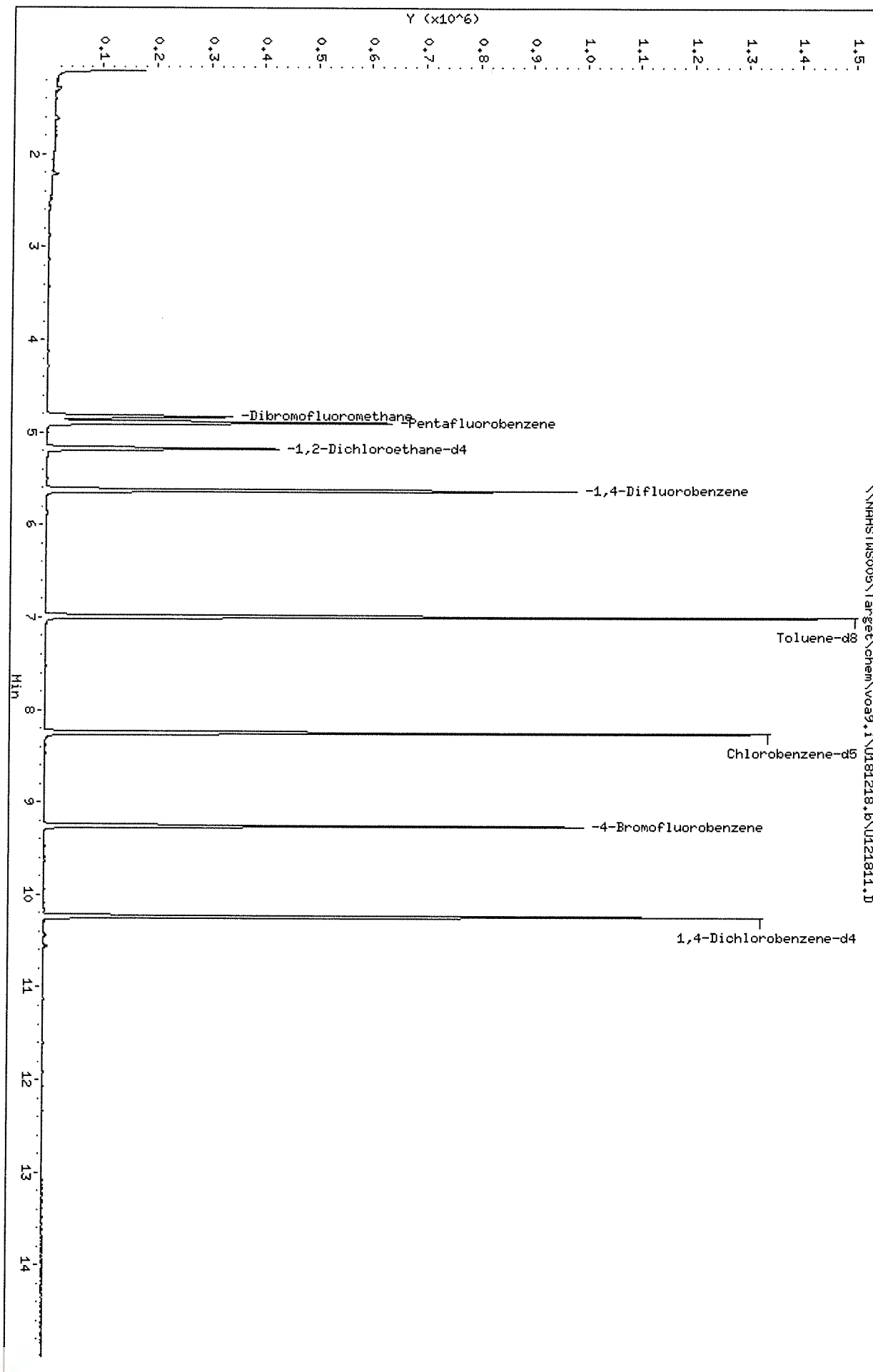
Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG					CONCENTRATIONS	
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	392367	50.0000	
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	700337	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	641402	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	294161	50.0000	
\$ 30 Dibromofluoromethane	113	4.830	4.827	(0.987)	199144	44.0057	44.00
\$ 35 1,2-Dichloroethane-d4	65	5.175	5.171	(1.057)	271864	45.6858	45.68
\$ 48 Toluene-d8	98	6.990	6.990	(0.847)	859102	52.4867	52.48
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	307071	48.3816	48.38



Data File: \\NAHSTMS005\Target\chem\voa9.i\U181218.b\U121811.D
Date: 18-DEC-2018 15:37
Client ID: HS18120278-04
Sample Info: HS18120278-04;HS18120278-04;;
Purge Volume: 5.0
Column phase: DB624

Instrument: W089.i
Operator: PC
Column diameter: 0.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121812.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121812.D
 Lab Smp Id: HS18120278-03 Client Smp ID: HS18120278-03
 Inj Date : 18-DEC-2018 16:02
 Operator : PC Inst ID: VOA9.i
 Smp Info : HS18120278-03;HS18120278-03;;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 12
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG				CONCENTRATIONS		
	MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	386598	50.0000	
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	700993	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	635273	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	290407	50.0000	
\$ 30 Dibromofluoromethane	113	4.830	4.827	(0.987)	198561	44.5467	44.54
\$ 35 1,2-Dichloroethane-d4	65	5.175	5.171	(1.057)	268050	45.7183	45.71
\$ 48 Toluene-d8	98	6.990	6.990	(0.847)	853104	52.6275	52.62
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	306625	48.7862	48.78
11 1,1-Dichloroethene	96	2.401	2.397	(0.491)	3101	0.71062	0.71 (a)
59 Chlorobenzene	112	8.275	8.275	(1.003)	20224	1.54246	1.54 (a)
27 cis-1,2-Dichloroethene	96	4.287	4.283	(0.876)	118604	21.6508	21.65
38 Trichloroethene	130	5.861	5.861	(1.042)	227152	47.5556	47.55

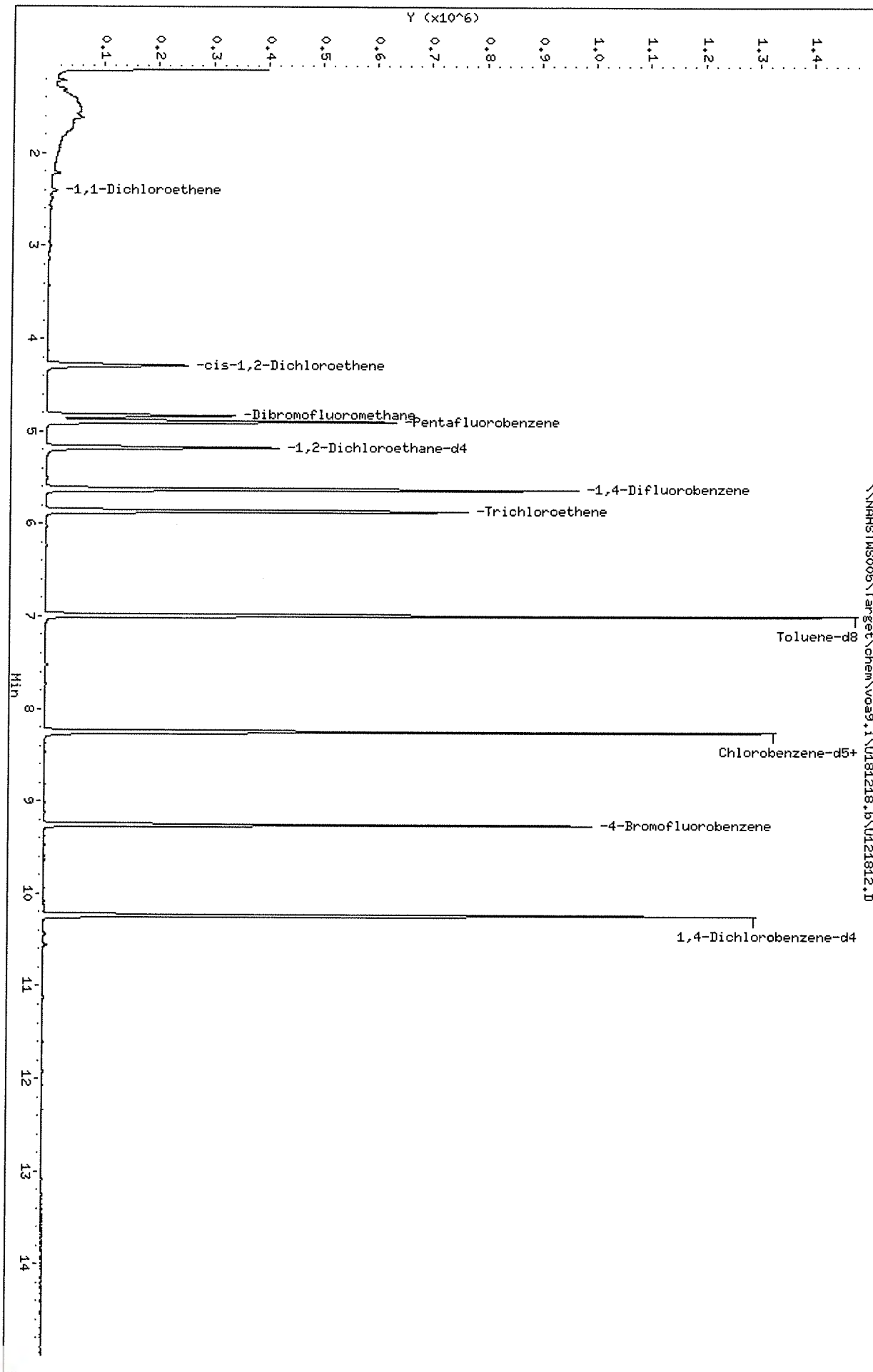
QC Flag Legend

a - Target compound detected but, quantitated amount
 Below Limit Of Quantitation(BLOQ).



Data File: \\NAHSTMS005\Target\chem\voa9.i\U181218.b\U121812.D
Date: 18-DEC-2018 16:02
Client ID: HS18120278-03
Sample Info: HS18120278-03;HS18120278-03;;
Purge Volume: 5.0
Column Phase: DB624

Instrument: V089.i
Operator: PC
Column diameter: 0.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U121812.D

Page 3

Date : 18-DEC-2018 16:02

Client ID: HS18120278-03

Instrument: VOA9.i

Sample Info: HS18120278-03;HS18120278-03;;;

Purge Volume: 5.0

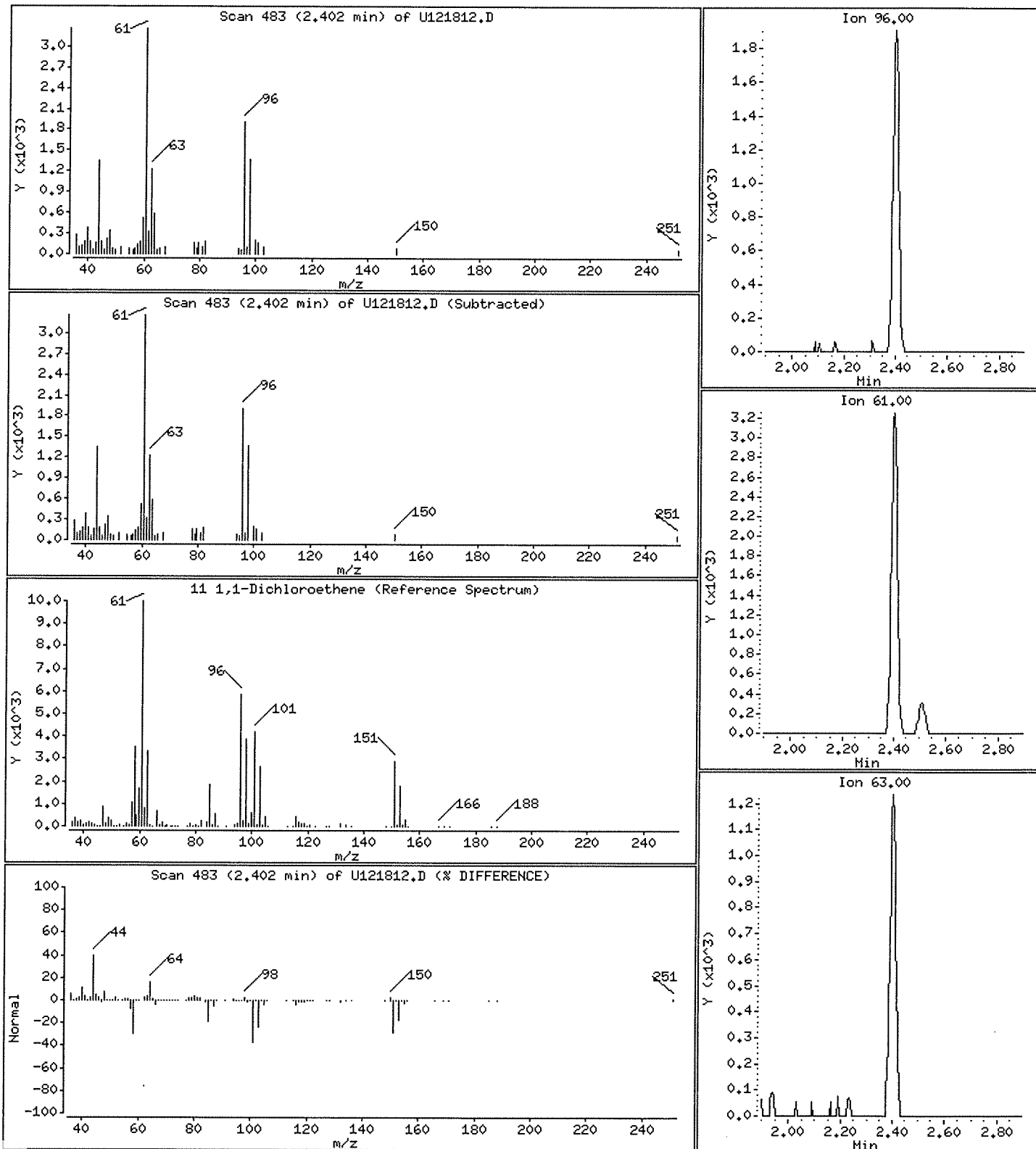
Operator: PC

Column phase: DB624

Column diameter: 0.18

11 1,1-Dichloroethene

Concentration: 0.71 ug/l



Data File: \\NAHSTWS005\Target\chem\voa9.i\U121812.D

Page 4

Date : 18-DEC-2018 16:02

Client ID: HS18120278-03

Instrument: VOA9.i

Sample Info: HS18120278-03;HS18120278-03;;;

Purge Volume: 5.0

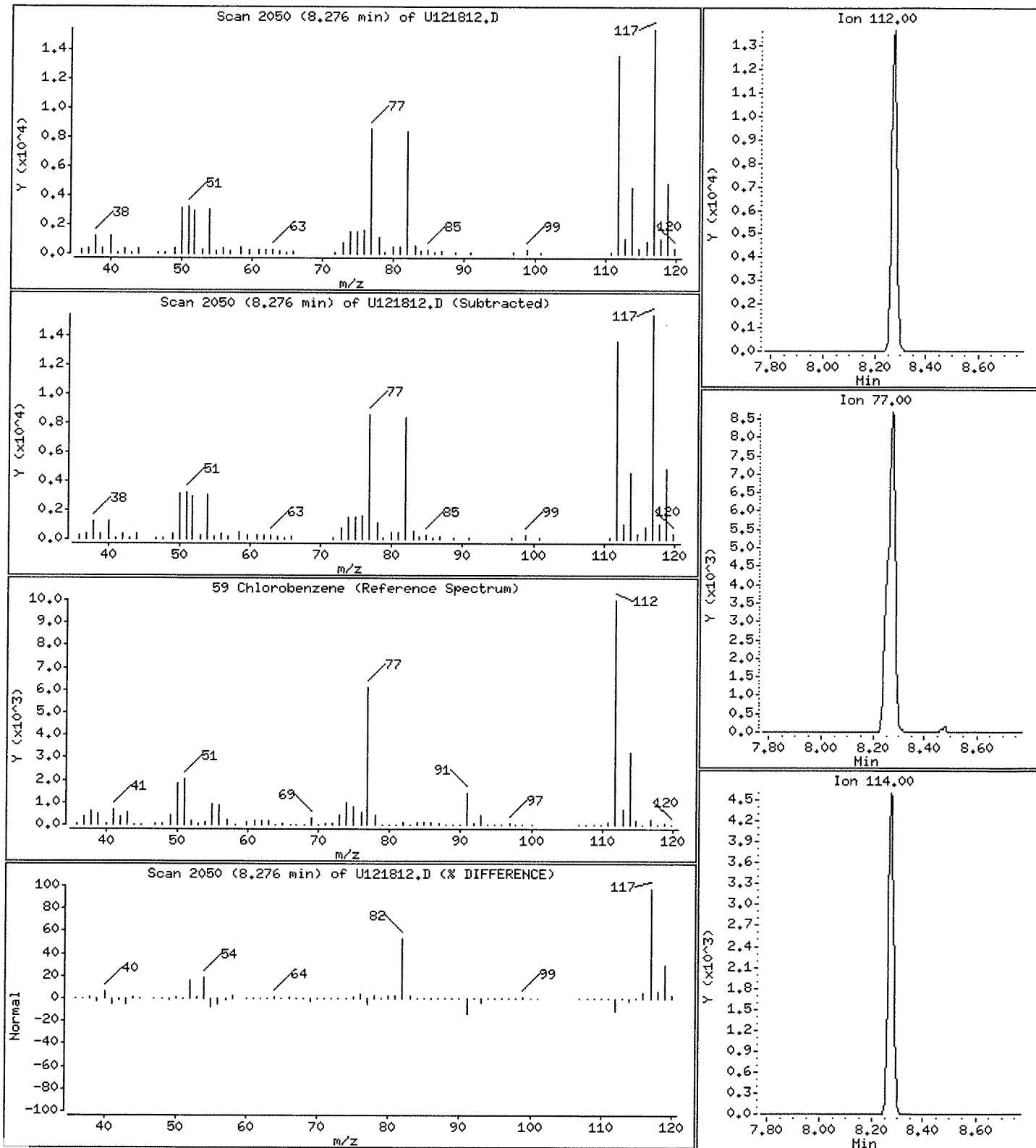
Operator: PC

Column phase: DB624

Column diameter: 0.18

59 Chlorobenzene

Concentration: 1.54 ug/l



Data File: \\NAHSTWS005\Target\chem\voa9.i\U121812.D

Page 5

Date : 18-DEC-2018 16:02

Client ID: HS18120278-03

Instrument: VOA9.i

Sample Info: HS18120278-03;HS18120278-03;;;

Purge Volume: 5.0

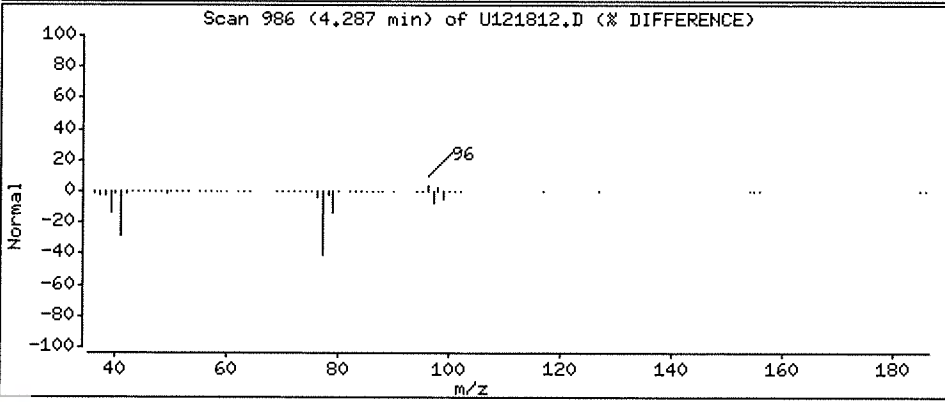
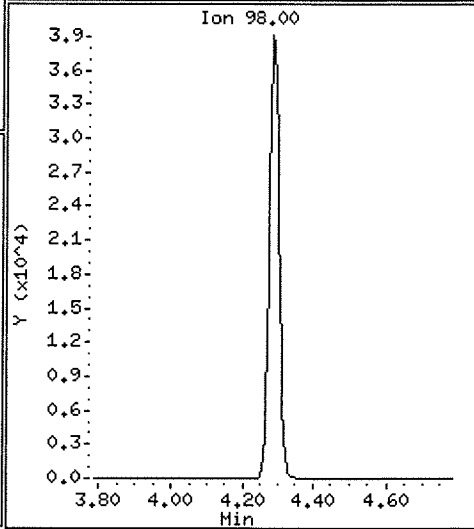
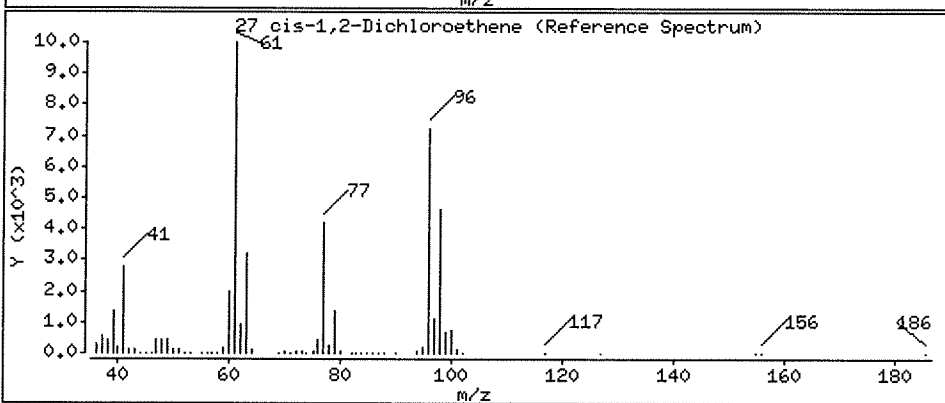
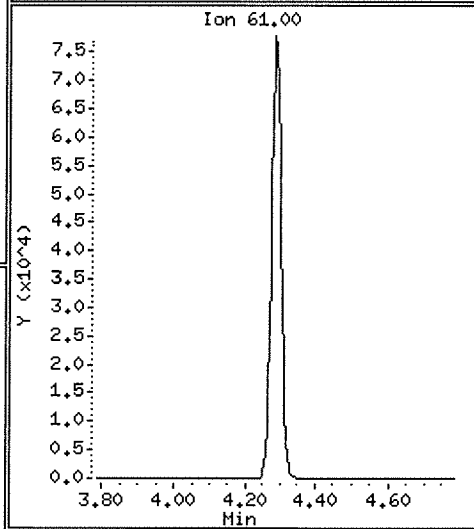
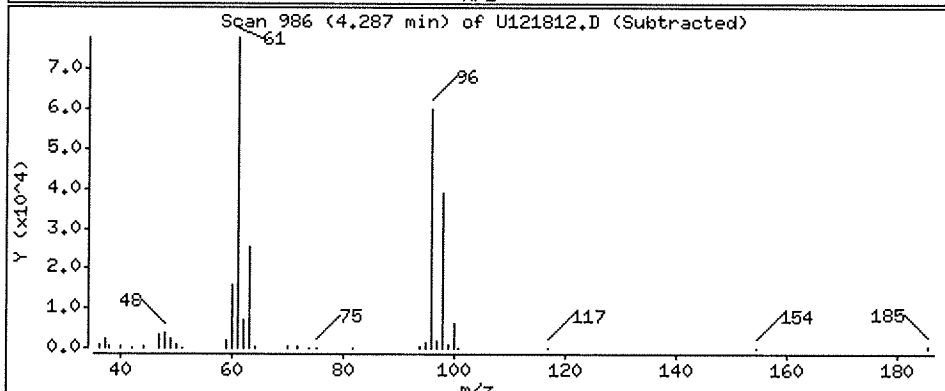
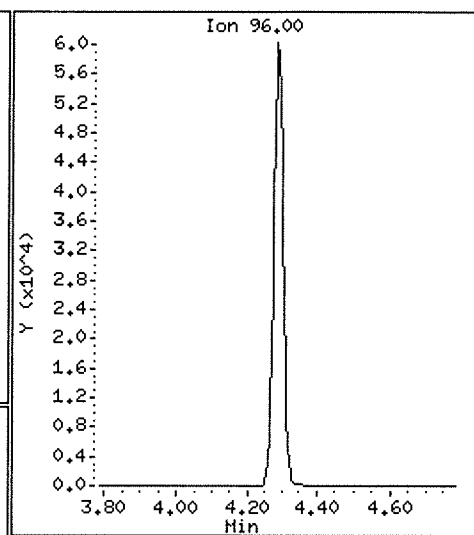
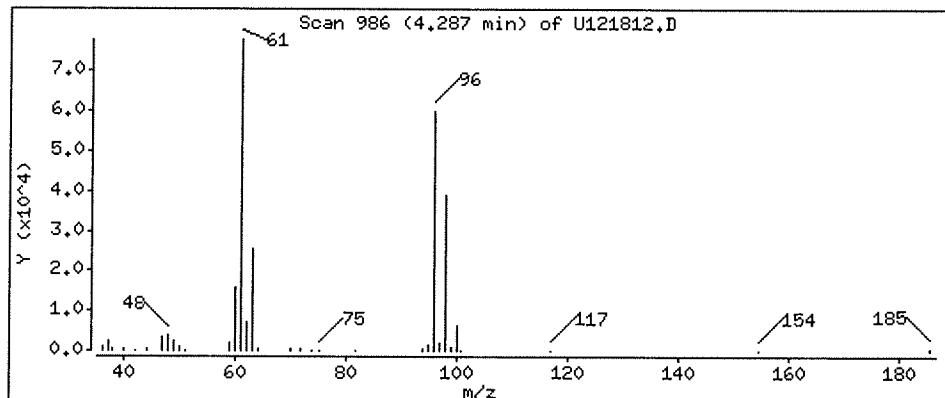
Operator: PC

Column phase: DB624

Column diameter: 0.18

27 cis-1,2-Dichloroethene

Concentration: 21.65 ug/l



Data File: \\NAHSTWS005\Target\chem\voa9.i\U121812.D

Page 6

Date : 18-DEC-2018 16:02

Client ID: HS18120278-03

Instrument: VOA9.i

Sample Info: HS18120278-03;HS18120278-03;;

Purge Volume: 5.0

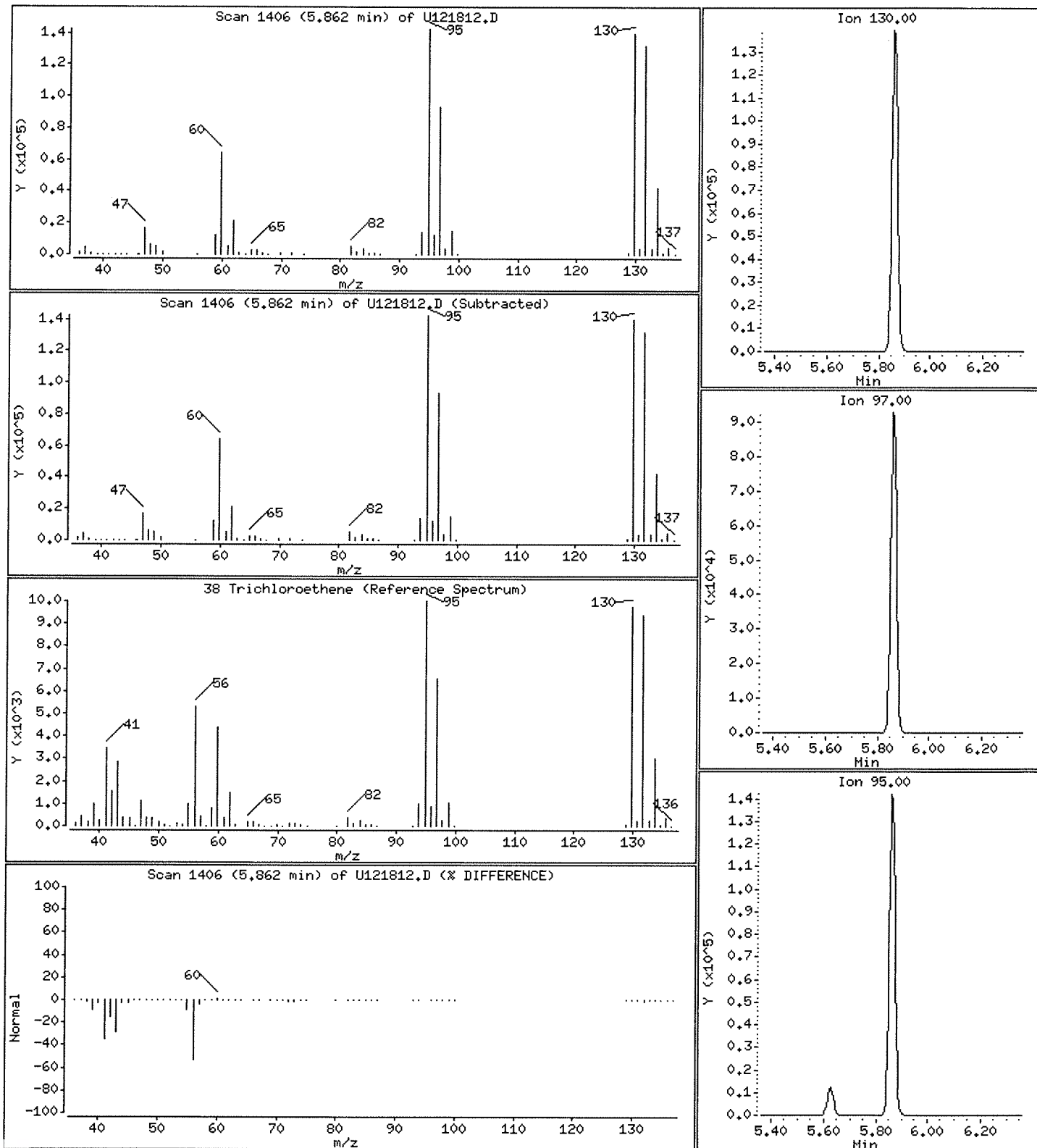
Operator: PC

Column phase: DB624

Column diameter: 0.18

38 Trichloroethene

Concentration: 47.55 ug/l



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121813.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121813.D
 Lab Smp Id: HS18120278-01 Client Smp ID: HS18120278-01
 Inj Date : 18-DEC-2018 16:27
 Operator : PC Inst ID: VOA9.i
 Smp Info : HS18120278-01;HS18120278-01;;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 13
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

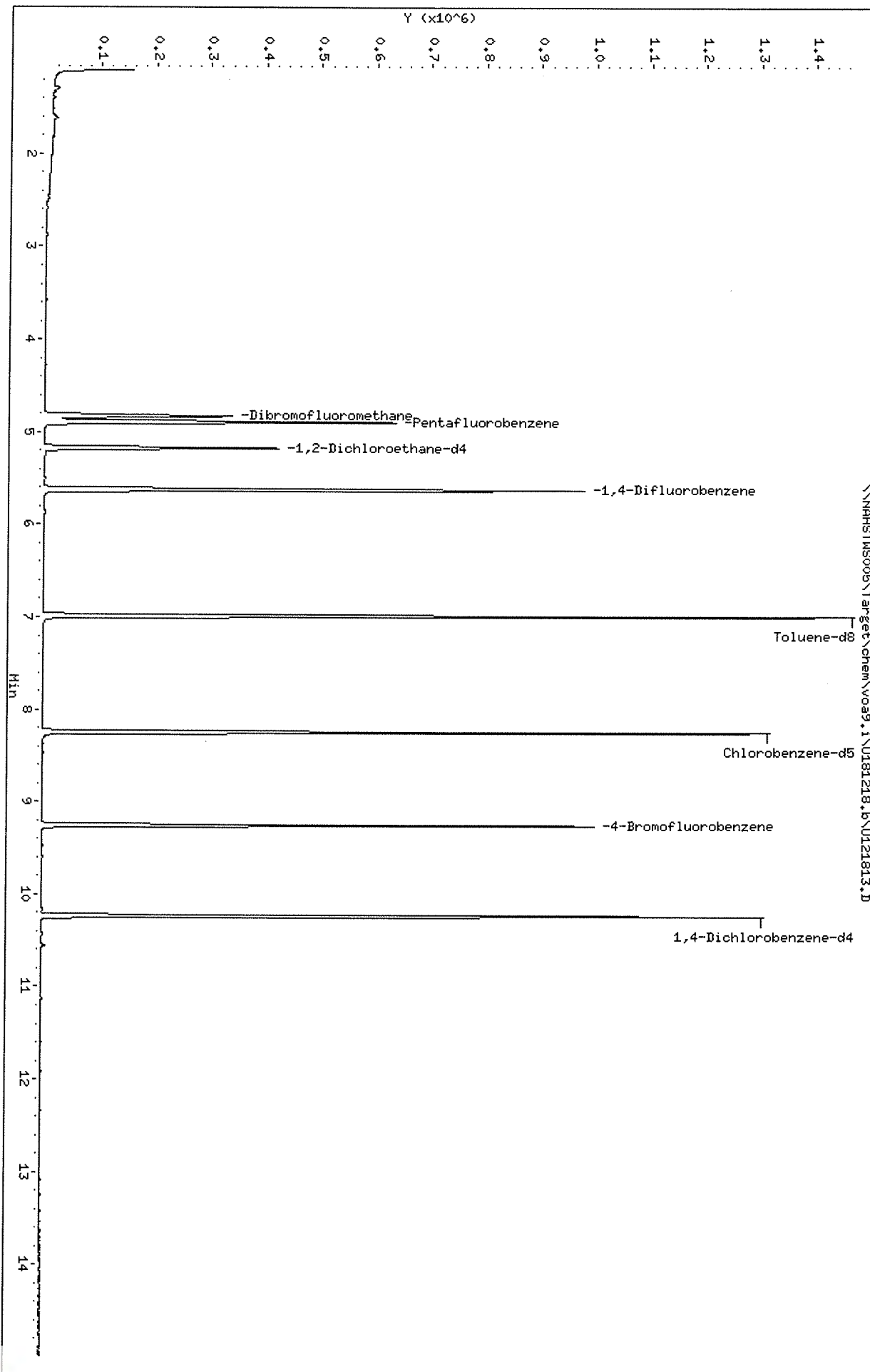
Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	389639	50.0000	
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	701381	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	634287	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	293976	50.0000	
\$ 30 Dibromofluoromethane	113	4.826	4.827	(0.986)	196405	43.6958	43.69
\$ 35 1,2-Dichloroethane-d4	65	5.175	5.171	(1.057)	270169	45.7202	45.72
\$ 48 Toluene-d8	98	6.989	6.990	(0.847)	859210	53.1015	53.10
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	311170	49.6037	49.60



Data File: \\NAHSTMS005\Target\chem\voa9.i\U181218.b\U121813.D
Date: 18-DEC-2018 16:27
Client ID: HS18120278-01
Sample Info: HS18120278-01;HS18120278-01;;
Purge Volume: 5.0
Column Phase: DB624

Instrument: VOA9.i
Operator: PC
Column diameter: 0.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121816.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121816.D
 Lab Smp Id: HS18120278-03MS Client Smp ID: HS18120278-03MS
 Inj Date : 18-DEC-2018 17:41
 Operator : PC Inst ID: VOA9.i
 Smp Info : HS18120278-03MS;HS18120278-03MS;3;;MS
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 16 QC Sample: MS
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG						CONCENTRATIONS	
		MASS	RT	EXP RT	REL RT	RESPONSE	ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.894	(1.000)	392642	50.0000		
* 36 1,4-Difluorobenzene	114	5.629	5.625	(1.000)	699438	50.0000		
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	641586	50.0000		
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	308103	50.0000		
\$ 30 Dibromofluoromethane	113	4.834	4.827	(0.987)	203783	45.0276	45.02	
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.171	(1.057)	263347	44.1668	44.16	
\$ 48 Toluene-d8	98	6.989	6.990	(0.847)	867952	53.0293	53.02	
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	326216	51.4494	51.44	
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	96623	23.8757	23.87	
31 1,1,1-Trichloroethane	97	4.830	4.827	(0.986)	156457	22.6848	22.68	
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	174691	23.9370	23.93	
138 Freon TF	101	2.409	2.397	(0.492)	87323	22.2007	22.20	
53 1,1,2-Trichloroethane	83	7.421	7.421	(0.900)	98833	24.2304	24.23	
22 1,1-Dichloroethane	63	3.608	3.601	(0.737)	209461	23.2795	23.27	
11 1,1-Dichloroethene	96	2.405	2.397	(0.491)	102742	23.1819	23.18	
32 1,1-Dichloropropene	75	5.006	5.003	(0.889)	166686	25.8035	25.80	
93 1,2,3-Trichlorobenzene	180	12.335	12.335	(1.205)	142561	24.5082	24.50	
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	177052	23.2487	23.24	
90 1,2,4-Trichlorobenzene	180	11.923	11.923	(1.165)	144320	23.8195	23.81	
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	439382	25.6752	25.67	
89 1,2-Dibromo-3-Chloropropane	155	11.233	11.233	(1.097)	23803	21.7951	21.79	
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	114145	24.1666	24.16	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121816.D Page 2
 Report Date: 25-Jan-2019 20:37

Compounds	QUANT SIG MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
88 1,2-Dichlorobenzene	146	10.569	10.570	(1.033)	228735	23.2475	23.24
33 1,2-Dichloroethane	62	5.254	5.250	(0.933)	165741	22.6451	22.64
42 1,2-Dichloropropane	63	6.082	6.079	(1.081)	126016	25.3052	25.30
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	426833	25.7843	25.78
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	228183	23.5228	23.52
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	213709	24.4008	24.40
84 1,4-Dichlorobenzene	146	10.255	10.255	(1.002)	244511	25.7484	25.74
26 2,2-Dichloropropane	77	4.279	4.272	(0.874)	122187	21.9285	21.92
24 2-Butanone	43	4.343	4.335	(0.887)	125547	45.4874	45.48
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	373050	24.7265	24.72
52 2-Hexanone	43	7.653	7.649	(0.928)	190474	51.0714	51.07
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	437841	25.1507	25.15
82 p-Isopropyltoluene	119	10.210	10.210	(0.997)	453828	26.8976	26.89
45 4-Methyl-2-Pentanone	43	6.915	6.915	(0.838)	269990	50.4444	50.44
10 Acetone	43	2.487	2.476	(0.508)	87042	46.0022	46.00
37 Benzene	78	5.220	5.216	(0.927)	489084	25.2966	25.29
74 Bromobenzene	156	9.381	9.381	(0.917)	120200	23.7826	23.78
29 Bromochloromethane	128	4.557	4.553	(0.930)	56796	24.5366	24.53
39 Bromodichloromethane	83	6.348	6.348	(1.128)	133158	22.9123	22.91
66 Bromoform	173	8.984	8.984	(1.089)	58291	20.7885	20.78
6 Bromomethane	94	1.674	1.663	(0.342)	34511	12.0926	12.09
19 Carbon Disulfide	76	2.596	2.585	(0.530)	613362	44.8136	44.81
34 Carbon Tetrachloride	117	4.999	4.991	(0.888)	121349	24.0463	24.04
59 Chlorobenzene	112	8.275	8.275	(1.003)	339216	25.6170	25.61
7 Chloroethane	64	1.756	1.745	(0.359)	101272	24.7197	24.71 (M)
28 Chloroform	83	4.662	4.654	(0.952)	199507	22.0691	22.06
3 Chloromethane	50	1.344	1.336	(0.274)	81391	15.6848	15.68
27 cis-1,2-Dichloroethene	96	4.290	4.283	(0.876)	249819	44.9018	44.90
46 cis-1,3-Dichloropropene	75	6.761	6.757	(1.201)	177823	24.2977	24.29
55 Dibromochloromethane	129	7.758	7.758	(0.940)	96375	23.7390	23.73
44 Dibromomethane	93	6.191	6.187	(1.100)	71390	22.8745	22.87
2 Dichlorodifluoromethane	85	1.209	1.202	(0.247)	55770	10.4181	10.41
61 Ethylbenzene	106	8.373	8.369	(1.015)	168348	25.1477	25.14
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	52202	28.1493	28.14
67 Isopropylbenzene	105	9.126	9.126	(1.106)	507707	26.0387	26.03
62 m,p-Xylenes	106	8.474	8.474	(1.027)	420565	50.9716	50.97
17 Methylene Chloride	84	2.877	2.866	(0.587)	123918	24.3803	24.38
87 n-Butylbenzene	91	10.558	10.558	(1.031)	426638	26.8733	26.87
73 n-Propylbenzene	91	9.475	9.475	(0.926)	640382	26.3397	26.33
92 Naphthalene	128	12.133	12.133	(1.185)	462810	25.3671	25.36
63 o-Xylene	106	8.811	8.811	(1.068)	210076	25.4439	25.44
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	541706	26.7590	26.75
64 Styrene	104	8.826	8.826	(1.070)	358679	25.7295	25.72
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	367701	26.0629	26.06
56 Tetrachloroethene	164	7.526	7.522	(0.912)	85796	25.5278	25.52
50 Toluene	91	7.049	7.046	(0.855)	512572	25.1638	25.16
20 trans-1,2-Dichloroethene	96	3.147	3.136	(0.643)	114024	23.4748	23.47
51 trans-1,3-Dichloropropene	75	7.263	7.259	(1.290)	148464	21.3745	21.37
38 Trichloroethene	130	5.865	5.861	(1.042)	345285	72.4482	72.44
8 Trichlorofluoromethane	101	1.959	1.948	(0.400)	164233	20.9219	20.92
5 Vinyl Chloride	62	1.423	1.415	(0.291)	127229	20.1831	20.18



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121816.D Page 3
Report Date: 25-Jan-2019 20:37

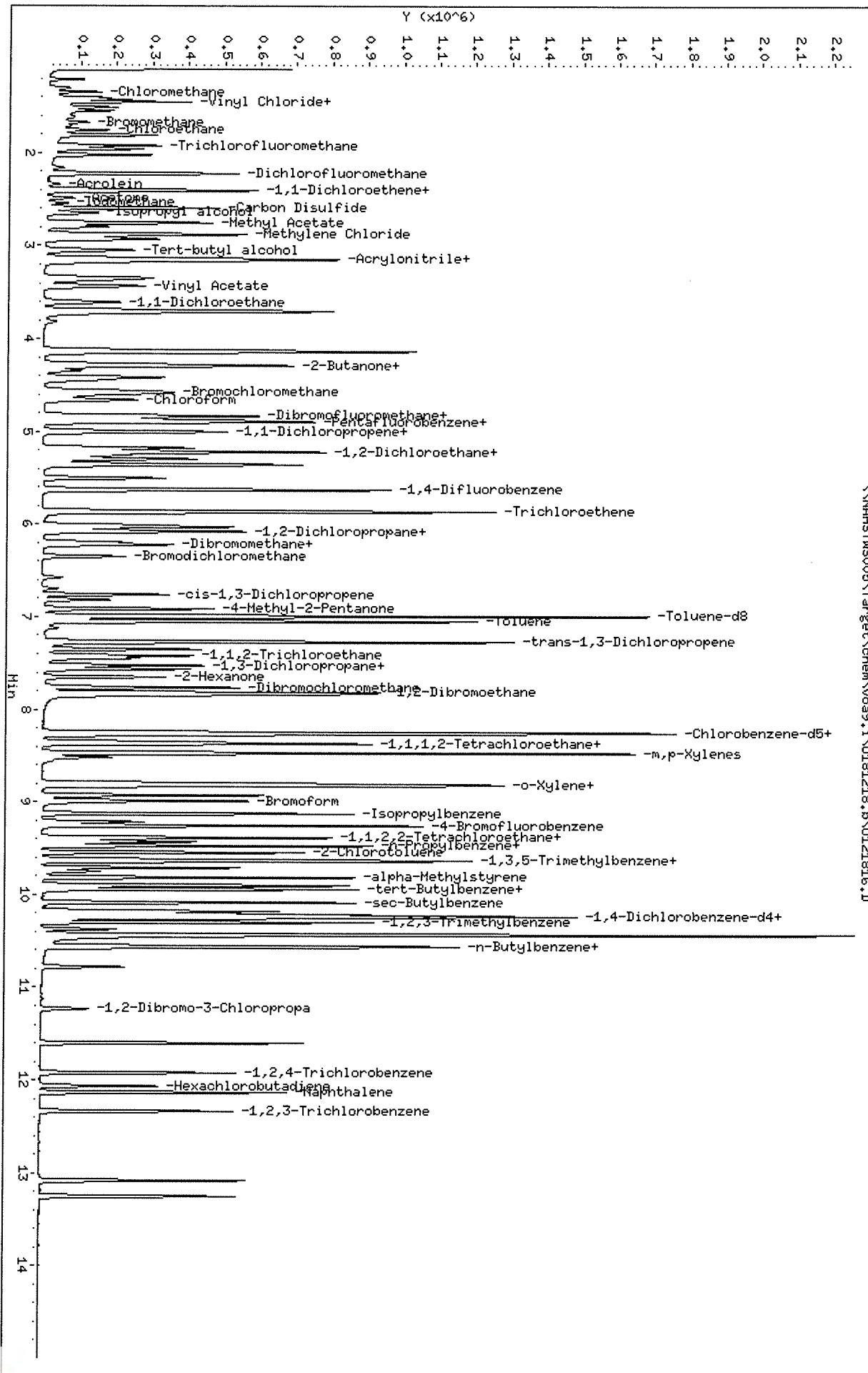
QC Flag Legend

M - Compound response manually integrated.



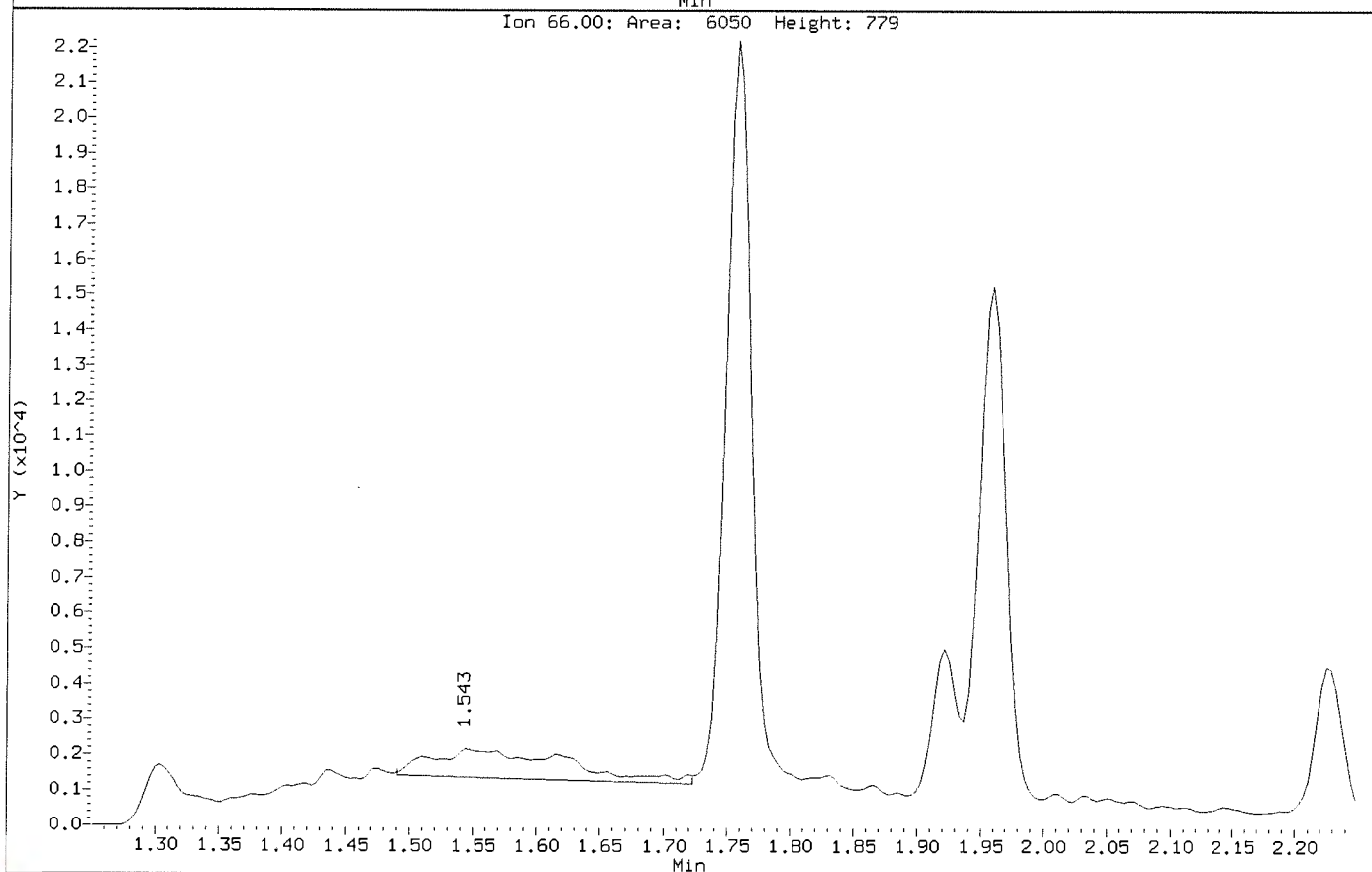
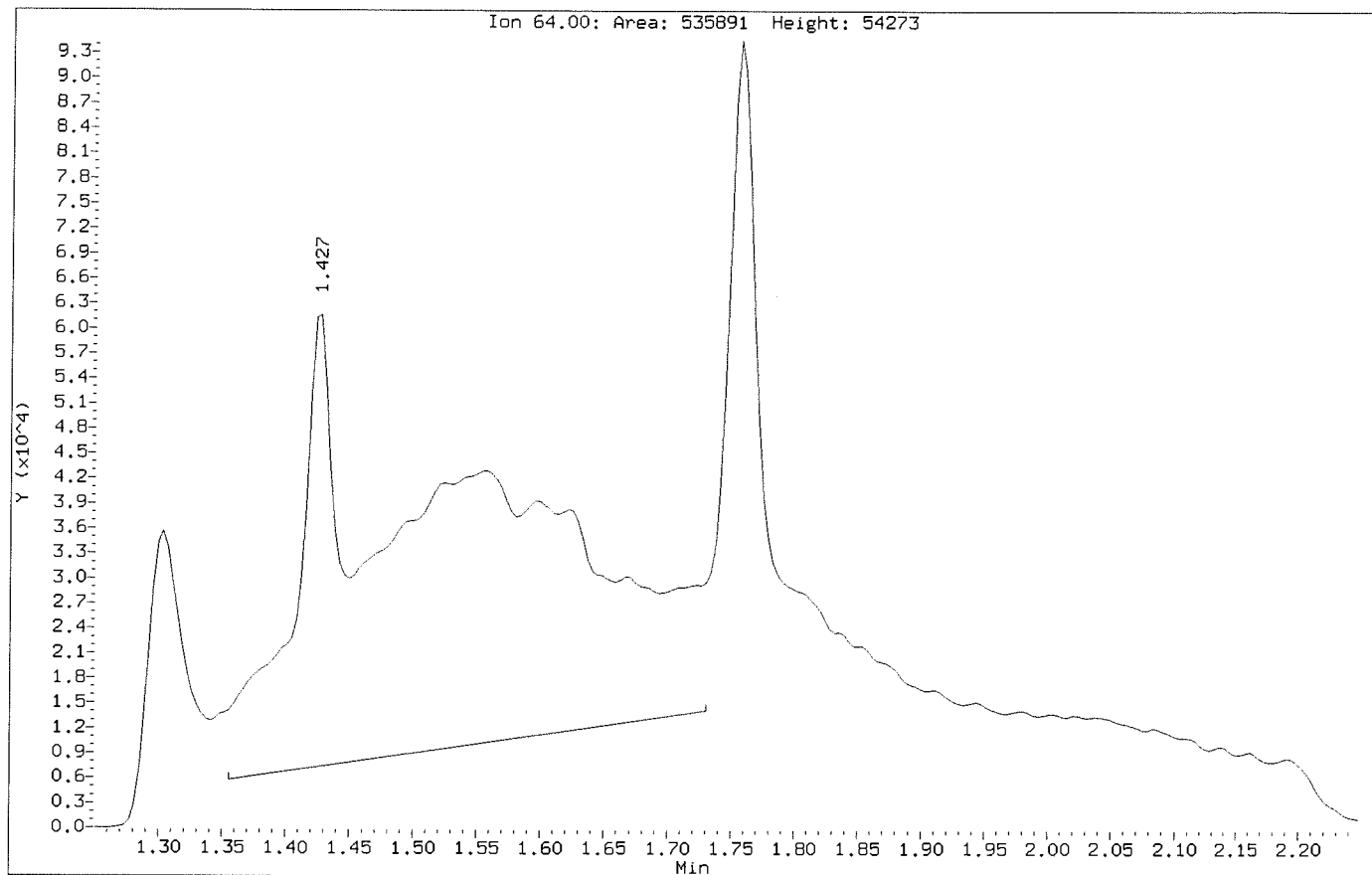
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Sample Info: HS18120278-03HS;HS18120278-03HS;3;HS
Purge Volume: 5.0
Column phase: DB624

Instrument: V089.i
Operator: PC
Column diameter: 0.18



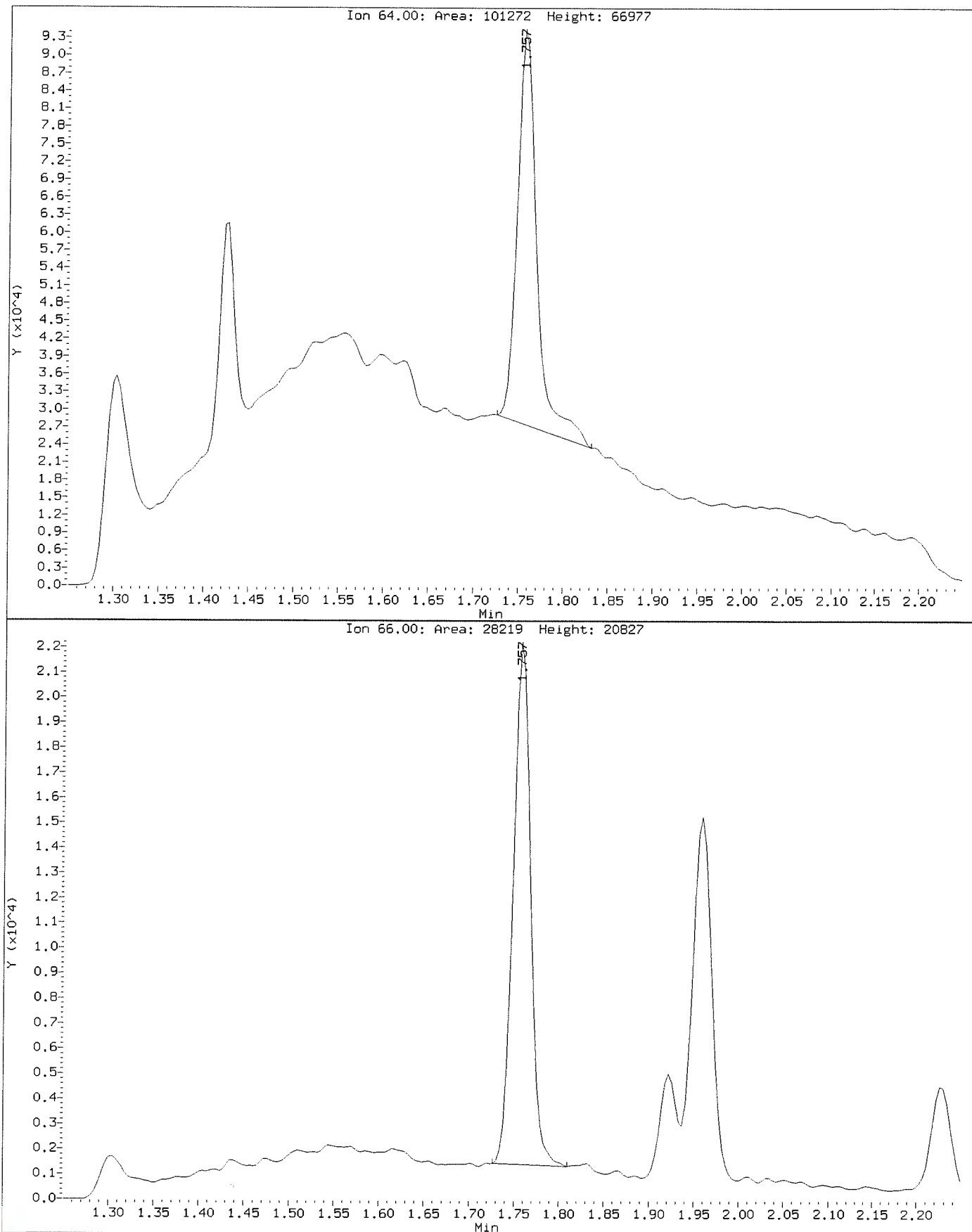
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Instrument: VDA9.i
Client Sample ID: HS18120278-03MS

Compound: Chloroethane
CAS Number: 75-00-3



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121816.D
Injection Date: 18-DEC-2018 17:41
Instrument: VOA9.i
Client Sample ID: HS1812027B-03MS

Compound: Chloroethane
CAS Number: 75-00-3



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121817.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121817.D
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 Inj Date : 18-DEC-2018 18:06
 Operator : PC Inst ID: VOA9.i
 Smp Info : HS18120278-03MSD;HS18120278-03MSD;3;;MSD
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 16 QC Sample: MSD
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
* 1 Pentafluorobenzene	168	4.898	4.894	(1.000)	400954	50.0000	
* 36 1,4-Difluorobenzene	114	5.629	5.625	(1.000)	706692	50.0000	
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	656441	50.0000	
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	315859	50.0000	
\$ 30 Dibromofluoromethane	113	4.834	4.827	(0.987)	207657	44.9298	44.92
\$ 35 1,2-Dichloroethane-d4	65	5.179	5.171	(1.057)	265380	43.5617	43.56
\$ 48 Toluene-d8	98	6.989	6.990	(0.847)	881002	52.5949	52.59
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	331476	51.0886	51.08
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	94992	22.9415	22.94
31 1,1,1-Trichloroethane	97	4.834	4.827	(0.987)	151124	21.4573	21.45
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	172184	23.0141	23.01
138 Freon TF	101	2.412	2.397	(0.493)	83980	20.9082	20.90
53 1,1,2-Trichloroethane	83	7.421	7.421	(0.900)	96652	23.1594	23.15
22 1,1-Dichloroethane	63	3.612	3.601	(0.737)	203872	22.1886	22.18
11 1,1-Dichloroethene	96	2.409	2.397	(0.492)	97491	21.5411	21.54
32 1,1-Dichloropropene	75	5.010	5.003	(0.890)	158542	24.2908	24.29
93 1,2,3-Trichlorobenzene	180	12.335	12.335	(1.205)	137085	22.9881	22.98
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	171588	21.9780	21.97
90 1,2,4-Trichlorobenzene	180	11.923	11.923	(1.165)	142107	22.8783	22.87
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	427765	24.3825	24.38
89 1,2-Dibromo-3-Chloropropane	155	11.233	11.233	(1.097)	22869	20.4946	20.49
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	112372	23.2529	23.25



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121817.D Page 2
 Report Date: 25-Jan-2019 20:37

Compounds	QUANT SIG MASS	RT	EXP RT	REL RT	RESPONSE	CONCENTRATIONS	
						ON-COLUMN (ug/l)	FINAL (ug/l)
88 1,2-Dichlorobenzene	146	10.569	10.570	(1.033)	222695	22.0779	22.07
33 1,2-Dichloroethane	62	5.258	5.250	(0.934)	162518	21.9768	21.97
42 1,2-Dichloropropane	63	6.082	6.079	(1.081)	124559	24.7558	24.75
75 1,3,5-Trimethylbenzene	105	9.625	9.625	(0.940)	414054	24.3981	24.39
83 1,3-Dichlorobenzene	146	10.180	10.180	(0.995)	223718	22.4962	22.49
54 1,3-Dichloropropane	76	7.563	7.563	(0.917)	209267	23.3529	23.35
84 1,4-Dichlorobenzene	146	10.255	10.255	(1.002)	239174	24.5660	24.56
26 2,2-Dichloropropane	77	4.279	4.272	(0.874)	117836	20.7093	20.70
24 2-Butanone	43	4.343	4.335	(0.887)	127361	45.1880	45.18
76 2-Chlorotoluene	91	9.546	9.546	(0.933)	365721	23.6455	23.64
52 2-Hexanone	43	7.649	7.649	(0.927)	188514	49.4020	49.40
77 4-Chlorotoluene	91	9.640	9.640	(0.942)	427325	23.9439	23.94
82 p-Isopropyltoluene	119	10.210	10.210	(0.997)	436533	25.2372	25.23
45 4-Methyl-2-Pentanone	43	6.914	6.915	(0.838)	266360	48.6400	48.63
10 Acetone	43	2.487	2.476	(0.508)	84551	43.5433	43.54
37 Benzene	78	5.224	5.216	(0.928)	477730	24.4557	24.45
74 Bromobenzene	156	9.381	9.381	(0.917)	115891	22.3670	22.36
29 Bromochloromethane	128	4.560	4.553	(0.931)	55899	23.6485	23.64
39 Bromodichloromethane	83	6.348	6.348	(1.128)	131031	22.3149	22.31
66 Bromoform	173	8.984	8.984	(1.089)	57698	20.1518	20.15
6 Bromomethane	94	1.678	1.663	(0.343)	39508	13.3262	13.32
19 Carbon Disulfide	76	2.600	2.585	(0.531)	590001	42.2132	42.21
34 Carbon Tetrachloride	117	4.999	4.991	(0.888)	118878	23.3149	23.31
59 Chlorobenzene	112	8.275	8.275	(1.003)	329663	24.3322	24.33
7 Chloroethane	64	1.760	1.745	(0.359)	86486	20.6729	20.67 (M)
28 Chloroform	83	4.662	4.654	(0.952)	195384	21.1649	21.16
3 Chloromethane	50	1.348	1.336	(0.275)	79127	14.9434	14.94
27 cis-1,2-Dichloroethene	96	4.290	4.283	(0.876)	243227	42.8107	42.81
46 cis-1,3-Dichloropropene	75	6.761	6.757	(1.201)	176358	23.8502	23.85
55 Dibromochloromethane	129	7.758	7.758	(0.940)	95950	23.0995	23.09
44 Dibromomethane	93	6.191	6.187	(1.100)	72191	22.8937	22.89
2 Dichlorodifluoromethane	85	1.213	1.202	(0.248)	53634	9.86969	9.86
61 Ethylbenzene	106	8.373	8.369	(1.015)	163448	23.8632	23.86
91 Hexachlorobutadiene	225	12.065	12.065	(1.179)	50484	26.6078	26.60
67 Isopropylbenzene	105	9.126	9.126	(1.106)	491242	24.6241	24.62
62 m,p-Xylenes	106	8.474	8.474	(1.027)	409050	48.4541	48.45
17 Methylene Chloride	84	2.881	2.866	(0.588)	119110	22.9073	22.90
87 n-Butylbenzene	91	10.558	10.558	(1.031)	415629	25.5691	25.56
73 n-Propylbenzene	91	9.475	9.475	(0.926)	617141	24.7604	24.76
92 Naphthalene	128	12.133	12.133	(1.185)	452641	24.2005	24.20
63 o-Xylene	106	8.811	8.811	(1.068)	202778	24.0042	24.00
81 sec-Butylbenzene	105	10.086	10.086	(0.985)	520853	25.0972	25.09
64 Styrene	104	8.826	8.826	(1.070)	348574	24.4387	24.43
78 tert-Butylbenzene	119	9.902	9.902	(0.967)	357619	24.7259	24.72
56 Tetrachloroethene	164	7.526	7.522	(0.912)	82749	24.0641	24.06
50 Toluene	91	7.049	7.046	(0.855)	495483	23.7744	23.77
20 trans-1,2-Dichloroethene	96	3.151	3.136	(0.643)	109083	21.9920	21.99
51 trans-1,3-Dichloropropene	75	7.263	7.259	(1.290)	146343	20.8889	20.88
38 Trichloroethene	130	5.865	5.861	(1.042)	332180	68.9831	68.98
8 Trichlorofluoromethane	101	1.962	1.948	(0.401)	154789	19.3100	19.31
5 Vinyl Chloride	62	1.426	1.415	(0.291)	120522	18.7596	18.75



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121817.D Page 3
Report Date: 25-Jan-2019 20:37

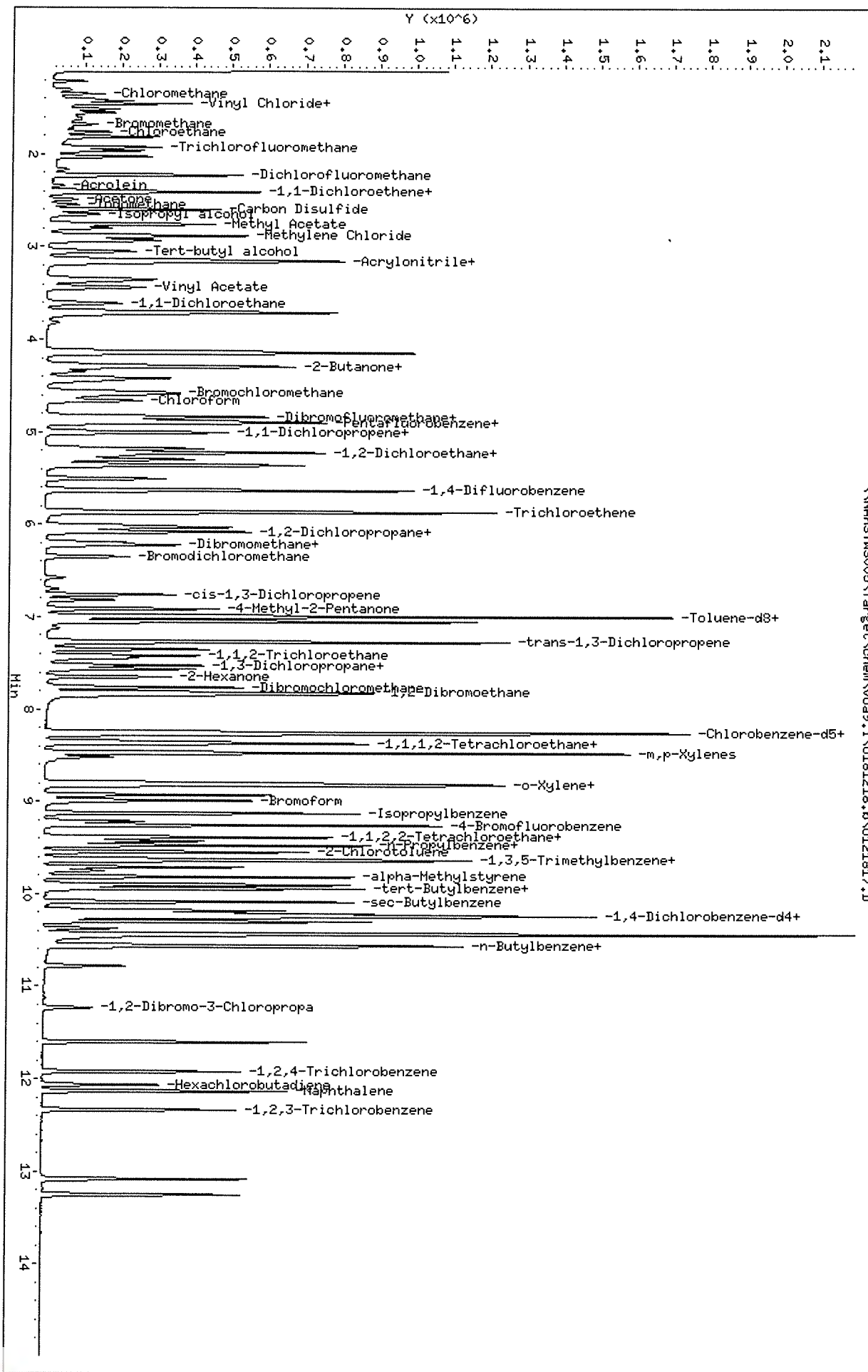
QC Flag Legend

M - Compound response manually integrated.



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 Column phase: DB624

Instrument: VOA9.1
 Operator: PC
 Column diameter: 0.18

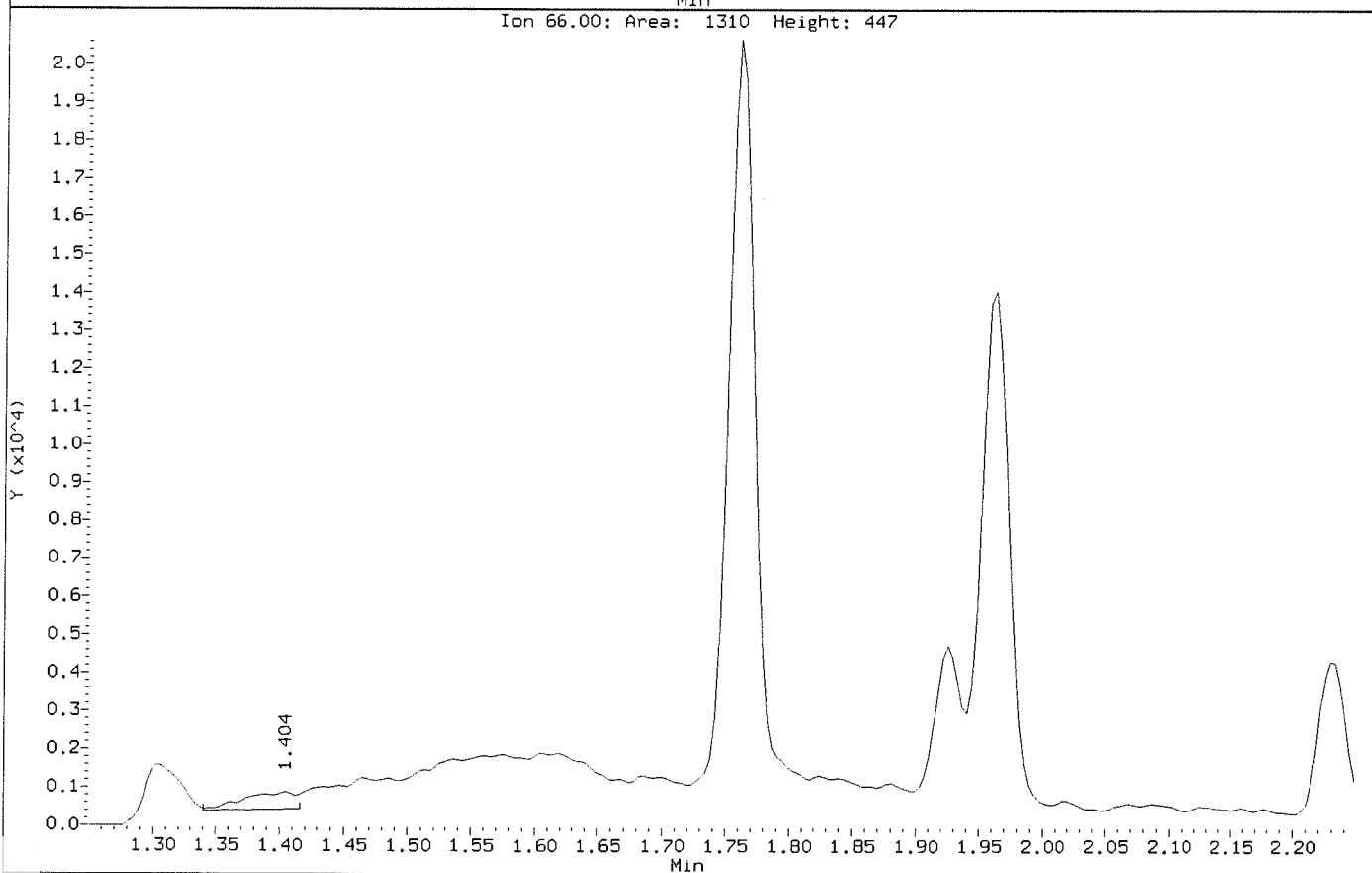
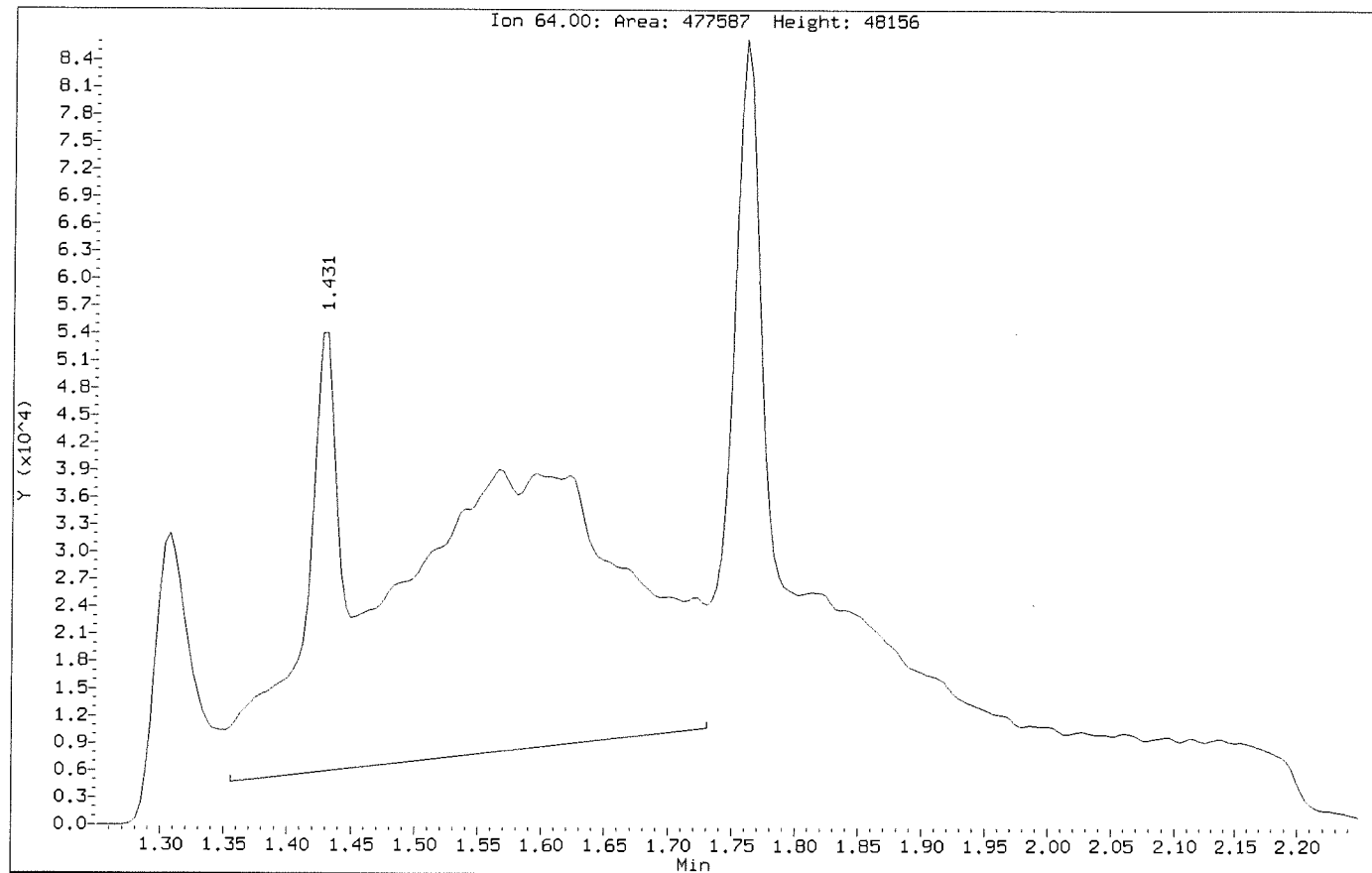


\\NAHSTMS006\Target\chem\voa9.i\U181218.b\U121817.D



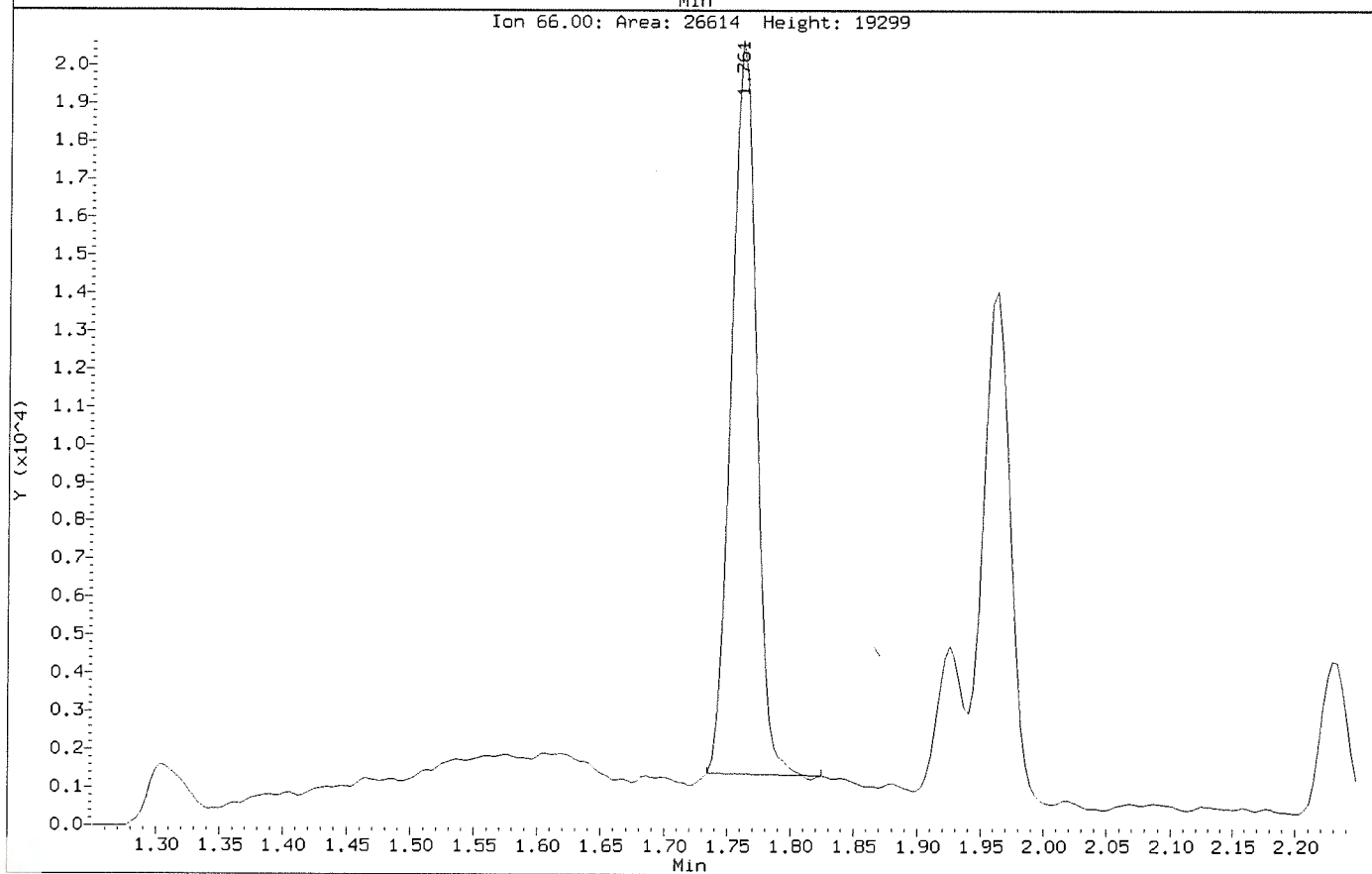
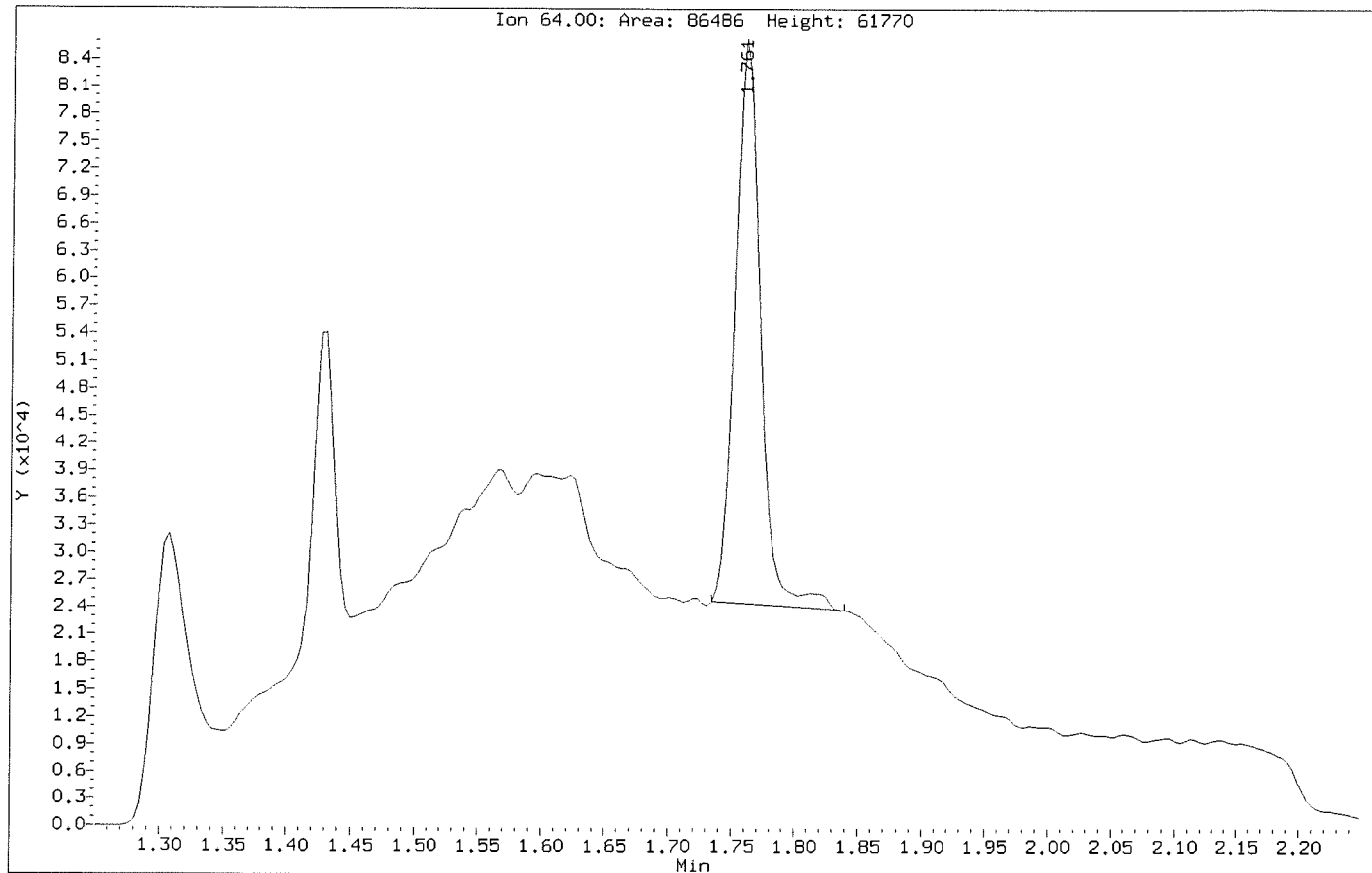
Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\Before\U121817.D
Injection Date: 18-DEC-2018 18:06
Instrument: VDA9.i
Client Sample ID: HS18120278-03MSD

Compound: Chloroethane
CAS Number: 75-00-3



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121817.D
Injection Date: 18-DEC-2018 18:06
Instrument: VDA9.1
Client Sample ID: HS1812027B-03MSD

Compound: Chloroethane
CAS Number: 75-00-3



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121828.D Page 1
 Report Date: 25-Jan-2019 20:37

ALS Laboratory Group

Data file : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121828.D
 Lab Smp Id: VSTD050-END Client Smp ID: VSTD050-END
 Inj Date : 18-DEC-2018 22:38
 Operator : PC Inst ID: VOA9.i
 Smp Info : VSTD050-END;VSTD050-END;2;;
 Misc Info : 180315V9;WATER;0;1;
 Comment :
 Method : \\NAHSTWS005\Target\chem\voa9.i\U181218.b\8260C.m
 Meth Date : 25-Jan-2019 20:37 VOA9.i Quant Type: ISTD
 Cal Date : 13-NOV-2018 14:15 Cal File: U111308.D
 Als bottle: 27 Continuing Calibration Sample
 Dil Factor: 1.00000
 Integrator: HP RTE Compound Sublist: LHAAP.sub
 Target Version: 4.14
 Processing Host: NAHSTW7087

Concentration Formula: Amt * DF * (Uf/Vo)*1 * CpndVariable

Name	Value	Description
DF	1.000	Dilution Factor
Uf	5.000	ng unit correction factor
Vo	5.000	sample purged
Cpnd Variable		Local Compound Variable

Compounds	QUANT SIG	AMOUNTS					CAL-AMT (ug/l)	ON-COL (ug/l)
		MASS	RT	EXP RT	REL RT	RESPONSE		
* 1 Pentafluorobenzene	168	4.894	4.894	(1.000)	378180	50.0000		
* 36 1,4-Difluorobenzene	114	5.625	5.625	(1.000)	671815	50.0000		
* 47 Chlorobenzene-d5	117	8.249	8.249	(1.000)	626627	50.0000		
* 70 1,4-Dichlorobenzene-d4	152	10.236	10.236	(1.000)	302976	50.0000		
\$ 30 Dibromofluoromethane	113	4.830	4.830	(0.987)	196169	50.0000	45.00	
\$ 35 1,2-Dichloroethane-d4	65	5.175	5.175	(1.057)	250888	50.0000	43.66	
\$ 48 Toluene-d8	98	6.989	6.989	(0.847)	842680	50.0000	52.70	
\$ 69 4-Bromofluorobenzene	95	9.257	9.257	(1.122)	316879	50.0000	51.16	
60 1,1,1,2-Tetrachloroethane	131	8.350	8.350	(1.012)	229533	50.0000	58.07	
31 1,1,1-Trichloroethane	97	4.826	4.826	(0.986)	344273	50.0000	51.82	
68 1,1,2,2-Tetrachloroethane	83	9.392	9.392	(0.918)	410071	50.0000	57.14	
138 Freon TF	101	2.401	2.401	(0.491)	189722	50.0000	50.07	
53 1,1,2-Trichloroethane	83	7.420	7.420	(0.900)	228744	50.0000	57.41	
22 1,1-Dichloroethane	63	3.601	3.601	(0.736)	478009	50.0000	55.15	
11 1,1-Dichloroethene	96	2.397	2.397	(0.490)	225962	50.0000	52.93	
32 1,1-Dichloropropene	75	5.003	5.003	(0.889)	355153	50.0000	57.23	
93 1,2,3-Trichlorobenzene	180	12.335	12.335	(1.205)	327297	50.0000	57.21	
71 1,2,3-Trichloropropane	75	9.426	9.426	(0.921)	436775	50.0000	58.32	
90 1,2,4-Trichlorobenzene	180	11.923	11.923	(1.165)	331419	50.0000	55.62	
79 1,2,4-Trimethylbenzene	105	9.943	9.943	(0.971)	961099	50.0000	57.11	
89 1,2-Dibromo-3-Chloropropane	155	11.233	11.233	(1.097)	59174	50.0000	53.27	
57 1,2-Dibromoethane	107	7.852	7.852	(0.952)	266717	50.0000	57.81	



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121828.D Page 2
 Report Date: 25-Jan-2019 20:37

Compounds	QUANT	SIG	AMOUNTS					
			MASS	RT	EXP RT	REL RT	RESPONSE	CAL-AMT (ug/l)
88 1,2-Dichlorobenzene	146		10.569	10.569	(1.033)	520553	50.0000	53.80
33 1,2-Dichloroethane	62		5.254	5.254	(0.934)	379556	50.0000	53.99
42 1,2-Dichloropropane	63		6.078	6.078	(1.081)	293890	50.0000	61.44
75 1,3,5-Trimethylbenzene	105		9.625	9.625	(0.940)	925420	50.0000	56.84
83 1,3-Dichlorobenzene	146		10.180	10.180	(0.995)	518297	50.0000	54.33
54 1,3-Dichloropropane	76		7.563	7.563	(0.917)	492889	50.0000	57.62
84 1,4-Dichlorobenzene	146		10.255	10.255	(1.002)	531293	50.0000	56.94
26 2,2-Dichloropropane	77		4.272	4.272	(0.873)	242214	50.0000	45.13
24 2-Butanone	43		4.335	4.335	(0.886)	303521	100.000	114.17
76 2-Chlorotoluene	91		9.546	9.546	(0.933)	827082	50.0000	55.74
52 2-Hexanone	43		7.649	7.649	(0.927)	458733	100.000	125.93
77 4-Chlorotoluene	91		9.640	9.640	(0.942)	967226	50.0000	56.50
82 p-Isopropyltoluene	119		10.210	10.210	(0.997)	951272	50.0000	57.33
45 4-Methyl-2-Pentanone	43		6.914	6.914	(0.838)	653968	100.000	125.10
10 Acetone	43		2.480	2.480	(0.507)	210321	100.000	122.09
37 Benzene	78		5.216	5.216	(0.927)	1101084	50.0000	59.29
74 Bromobenzene	156		9.381	9.381	(0.917)	275484	50.0000	55.42
29 Bromochloromethane	128		4.553	4.553	(0.930)	129015	50.0000	57.86
39 Bromodichloromethane	83		6.348	6.348	(1.129)	320311	50.0000	57.38
66 Bromoform	173		8.984	8.984	(1.089)	150873	50.0000	52.87(T)
6 Bromomethane	94		1.663	1.663	(0.340)	139467	50.0000	44.65
19 Carbon Disulfide	76		2.588	2.588	(0.529)	1482971	100.000	112.49
34 Carbon Tetrachloride	117		4.995	4.995	(0.888)	264711	50.0000	54.61
59 Chlorobenzene	112		8.275	8.275	(1.003)	713991	50.0000	55.20
7 Chloroethane	64		1.749	1.749	(0.357)	250841	50.0000	63.56
28 Chloroform	83		4.658	4.658	(0.952)	458398	50.0000	52.64
3 Chloromethane	50		1.336	1.336	(0.273)	270537	50.0000	53.56
27 cis-1,2-Dichloroethene	96		4.287	4.287	(0.876)	292548	50.0000	54.59
46 cis-1,3-Dichloropropene	75		6.757	6.757	(1.201)	434455	50.0000	61.80
55 Dibromochloromethane	129		7.758	7.758	(0.940)	238967	50.0000	60.26
44 Dibromomethane	93		6.191	6.191	(1.101)	170582	50.0000	56.90
2 Dichlorodifluoromethane	85		1.201	1.201	(0.246)	293113	50.0000	52.38
61 Ethylbenzene	106		8.373	8.373	(1.015)	368594	50.0000	56.37
91 Hexachlorobutadiene	225		12.065	12.065	(1.179)	104892	50.0000	55.69
67 Isopropylbenzene	105		9.126	9.126	(1.106)	1075455	50.0000	56.47
62 m,p-Xylenes	106		8.474	8.474	(1.027)	915802	100.000	113.64
17 Methylene Chloride	84		2.870	2.870	(0.586)	284347	50.0000	59.05
87 n-Butylbenzene	91		10.558	10.558	(1.031)	879817	50.0000	55.28
73 n-Propylbenzene	91		9.475	9.475	(0.926)	1345509	50.0000	56.27
92 Naphthalene	128		12.133	12.133	(1.185)	1107615	50.0000	61.73
63 o-Xylene	106		8.811	8.811	(1.068)	466527	50.0000	57.85
81 sec-Butylbenzene	105		10.086	10.086	(0.985)	1112945	50.0000	55.90
64 Styrene	104		8.823	8.823	(1.070)	815131	50.0000	59.86
78 tert-Butylbenzene	119		9.902	9.902	(0.967)	771287	50.0000	55.59
56 Tetrachloroethene	164		7.525	7.525	(0.912)	177630	50.0000	54.11
50 Toluene	91		7.046	7.046	(0.854)	1133358	50.0000	56.96
20 trans-1,2-Dichloroethene	96		3.139	3.139	(0.642)	252130	50.0000	53.89
51 trans-1,3-Dichloropropene	75		7.259	7.259	(1.291)	370216	50.0000	53.13
38 Trichloroethene	130		5.861	5.861	(1.042)	251920	50.0000	55.03
8 Trichlorofluoromethane	101		1.951	1.951	(0.399)	382135	50.0000	50.54
5 Vinyl Chloride	62		1.419	1.419	(0.290)	365089	50.0000	59.12



Data File: \\NAHSTWS005\Target\chem\voa9.i\U181218.b\U121828.D Page 3
Report Date: 25-Jan-2019 20:37

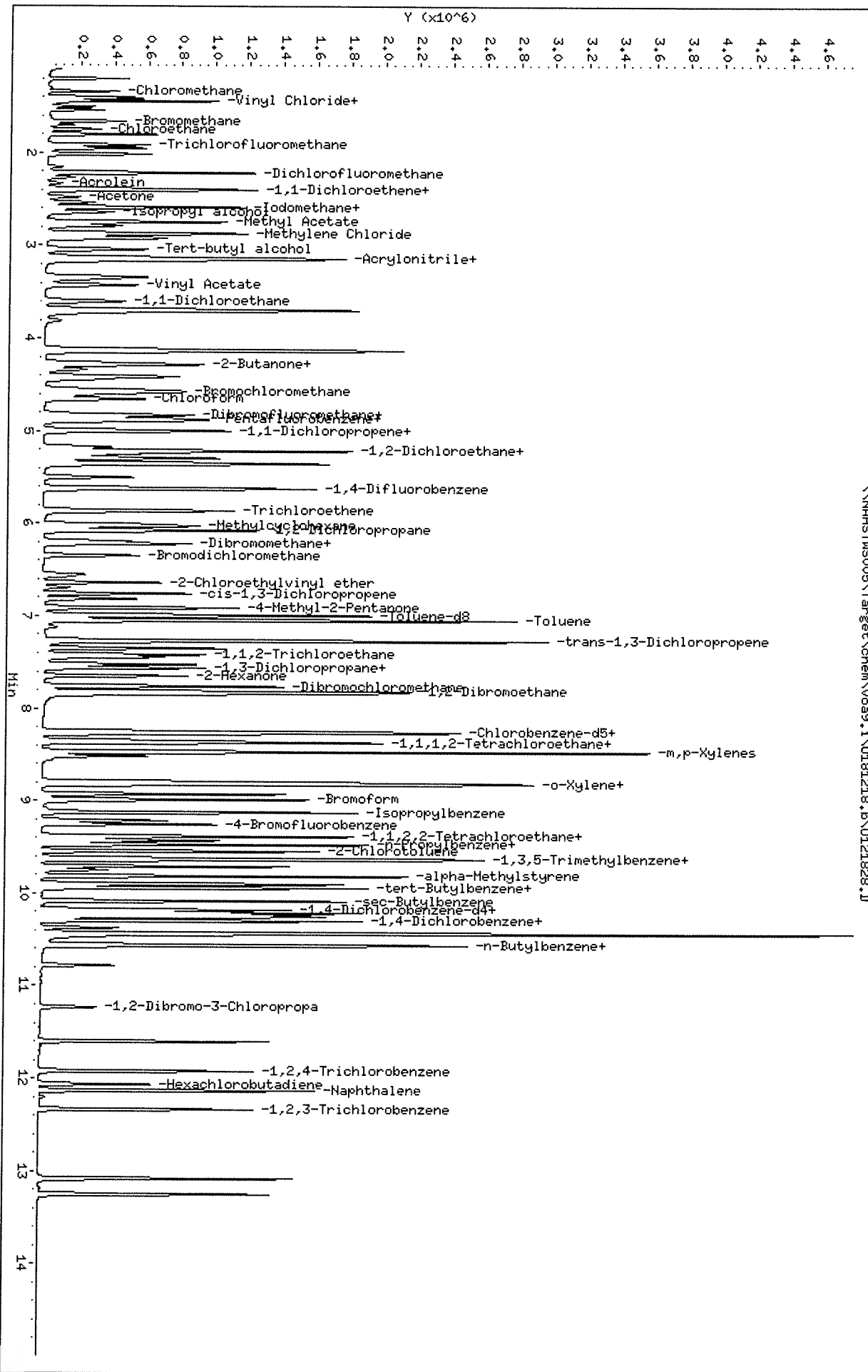
QC Flag Legend

T - Target compound detected outside RT window.



Data File: \\NAHSTMS005\Target\chem\voa9.1\U181218.b\U121828.D
 Date : 18-DEC-2018 22:38
 Client ID: VSTD050-END
 Sample Info: VSTD050-END;VSTD050-END;2;
 Purge Volume: 5.0
 Column phase: DB624

Instrument: WDA9.1
 Operator: PC
 Column diameter: 0.18



\\NAHSTMS005\Target\chem\voa9.1\U181218.b\U121828.D





10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

December 19, 2018

Susan Huang
Aptim Environmental & Infrastructure, Inc.
2500 City West Blvd., Suite 1700
Houston, TX 77042

Work Order: **HS18120278**

Laboratory Results for: **LHAARP - 12**

Dear Susan,

ALS Environmental received 5 sample(s) on Dec 05, 2018 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
Work Order: HS18120278

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS18120278-01	12WW21-181204	Groundwater		04-Dec-2018 10:05	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-02	12WW21-181204 FD	Groundwater		04-Dec-2018 10:05	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-03	12WW24-181204	Groundwater		04-Dec-2018 13:10	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-04	12WW20-181204	Groundwater		04-Dec-2018 12:05	05-Dec-2018 09:30	<input type="checkbox"/>
HS18120278-05	Trip Blank ALS-071918-78	Water		04-Dec-2018 00:00	05-Dec-2018 09:30	<input type="checkbox"/>

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
Work Order: HS18120278

CASE NARRATIVE

GCMS Volatiles by Method SW8260**Batch ID: R329535****Sample ID: VLCSW-1812018**

- 1,4_Dichlorobenzene and Hexachlorobutadiene exceeded QC limits for LCS. CCV is OK.

Sample ID: VSTD050

- cis-1,3-Dichloropropene exceeded %D limits for CCV. Samples are ND for this compound.

Sample ID: 12WW24-181204 (HS18120278-03MS)

- MS/MSD recovered outside the control limits for multiple compounds
-

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-01
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 16:27	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-01
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:27	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:27	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 16:27	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:27	
Surr: 1,2-Dichloroethane-d4	91.4			0	70-126	%REC	1	18-Dec-2018 16:27	
Surr: 4-Bromofluorobenzene	99.2			0	81-113	%REC	1	18-Dec-2018 16:27	
Surr: Dibromofluoromethane	87.4			0	77-123	%REC	1	18-Dec-2018 16:27	
Surr: Toluene-d8	106			0	82-127	%REC	1	18-Dec-2018 16:27	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204 FD
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT

WorkOrder:HS18120278
 Lab ID:HS18120278-02
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 15:13	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW21-181204 FD
 Collection Date: 04-Dec-2018 10:05

ANALYTICAL REPORT

WorkOrder:HS18120278
 Lab ID:HS18120278-02
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:13	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:13	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 15:13	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:13	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>90.5</i>			0	<i>70-126</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:13</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>97.8</i>			0	<i>81-113</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:13</i>	
<i>Surr: Dibromofluoromethane</i>	<i>88.4</i>			0	<i>77-123</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:13</i>	
<i>Surr: Toluene-d8</i>	<i>105</i>			0	<i>82-127</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:13</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW24-181204
 Collection Date: 04-Dec-2018 13:10

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-03
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1-Dichloroethene	0.71	J	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 16:02	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Chlorobenzene	1.5		0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW24-181204
 Collection Date: 04-Dec-2018 13:10

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-03
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
cis-1,2-Dichloroethene	22		0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 16:02	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 16:02	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 16:02	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Trichloroethene	48		0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 16:02	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>91.4</i>			0	<i>70-126</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>97.6</i>			0	<i>81-113</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	
<i>Surr: Dibromofluoromethane</i>	<i>89.1</i>			0	<i>77-123</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	
<i>Surr: Toluene-d8</i>	<i>105</i>			0	<i>82-127</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 16:02</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW20-181204
 Collection Date: 04-Dec-2018 12:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-04
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
Acetone	2.0	U	2.0	2.0	2.0	ug/L	1	18-Dec-2018 15:37	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: 12WW20-181204
 Collection Date: 04-Dec-2018 12:05

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-04
 Matrix:Groundwater

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 15:37	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 15:37	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 15:37	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 15:37	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>91.4</i>			0	<i>70-126</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:37</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>96.8</i>			0	<i>81-113</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:37</i>	
<i>Surr: Dibromofluoromethane</i>	<i>88.0</i>			0	<i>77-123</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:37</i>	
<i>Surr: Toluene-d8</i>	<i>105</i>			0	<i>82-127</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 15:37</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: Trip Blank ALS-071918-78
 Collection Date: 04-Dec-2018 00:00

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-05
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dibromo-3-chloropropane	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
2-Butanone	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
2-Hexanone	1.0	U	1.0	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
Acetone	2.7		2.0	2.0	2.0	ug/L	1	18-Dec-2018 14:48	
Benzene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromobenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromochloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromoform	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Bromomethane	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Carbon disulfide	1.0	U	0.60	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Chlorobenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Chloroethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 Sample ID: Trip Blank ALS-071918-78
 Collection Date: 04-Dec-2018 00:00

ANALYTICAL REPORT
 WorkOrder:HS18120278
 Lab ID:HS18120278-05
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
LOW LEVEL VOLATILES BY SW8260C		Method:SW8260						Analyst: PC	
Chloroform	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Chloromethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Dibromomethane	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Ethylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	ug/L	1	18-Dec-2018 14:48	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
m,p-Xylene	1.0	U	0.50	1.0	2.0	ug/L	1	18-Dec-2018 14:48	
Methylene chloride	0.50	U	1.0	0.50	2.0	ug/L	1	18-Dec-2018 14:48	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Naphthalene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
o-Xylene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Styrene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Toluene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Trichloroethene	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
Vinyl chloride	0.50	U	0.20	0.50	1.0	ug/L	1	18-Dec-2018 14:48	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>89.5</i>			0	<i>70-126</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 14:48</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>98.2</i>			0	<i>81-113</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 14:48</i>	
<i>Surr: Dibromofluoromethane</i>	<i>87.9</i>			0	<i>77-123</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 14:48</i>	
<i>Surr: Toluene-d8</i>	<i>106</i>			0	<i>82-127</i>	<i>%REC</i>	<i>1</i>	<i>18-Dec-2018 14:48</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R329535	Test Name : LOW LEVEL VOLATILES BY SW8260C			Matrix: Water		
HS18120278-05	Trip Blank ALS-071918-78	04 Dec 2018 00:00			18 Dec 2018 14:48	1
Batch ID R329535	Test Name : LOW LEVEL VOLATILES BY SW8260C			Matrix: Groundwater		
HS18120278-01	12WW21-181204	04 Dec 2018 10:05			18 Dec 2018 16:27	1
HS18120278-02	12WW21-181204 FD	04 Dec 2018 10:05			18 Dec 2018 15:13	1
HS18120278-03	12WW24-181204	04 Dec 2018 13:10			18 Dec 2018 16:02	1
HS18120278-04	12WW20-181204	04 Dec 2018 12:05			18 Dec 2018 15:37	1

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MBLK	Sample ID: VBLKW-181218	Units: ug/L			Analysis Date: 18-Dec-2018 13:34					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872159	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	0.50	1.0								U
1,1,1-Trichloroethane	0.50	1.0								U
1,1,2,2-Tetrachloroethane	0.50	1.0								U
1,1,2-Trichlor-1,2,2-trifluoroethane	0.50	1.0								U
1,1,2-Trichloroethane	0.50	1.0								U
1,1-Dichloroethane	0.50	1.0								U
1,1-Dichloroethene	0.50	1.0								U
1,1-Dichloropropene	0.50	1.0								U
1,2,3-Trichlorobenzene	0.50	1.0								U
1,2,3-Trichloropropane	0.50	1.0								U
1,2,4-Trichlorobenzene	0.50	1.0								U
1,2,4-Trimethylbenzene	0.50	1.0								U
1,2-Dibromo-3-chloropropane	1.0	1.0								U
1,2-Dibromoethane	0.50	1.0								U
1,2-Dichlorobenzene	0.50	1.0								U
1,2-Dichloroethane	0.50	1.0								U
1,2-Dichloropropane	0.50	1.0								U
1,3,5-Trimethylbenzene	0.50	1.0								U
1,3-Dichlorobenzene	0.50	1.0								U
1,3-Dichloropropane	0.50	1.0								U
1,4-Dichlorobenzene	0.50	1.0								U
2,2-Dichloropropane	0.50	1.0								U
2-Butanone	1.0	2.0								U
2-Chlorotoluene	0.50	1.0								U
2-Hexanone	1.0	2.0								U
4-Chlorotoluene	0.50	1.0								U
4-Isopropyltoluene	0.50	1.0								U
4-Methyl-2-pentanone	1.0	2.0								U
Acetone	2.0	2.0								U
Benzene	0.50	1.0								U
Bromobenzene	0.50	1.0								U
Bromochloromethane	0.50	1.0								U
Bromodichloromethane	0.50	1.0								U
Bromoform	0.50	1.0								U

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MBLK	Sample ID: VBLKW-181218	Units: ug/L			Analysis Date: 18-Dec-2018 13:34					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872159	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Bromomethane	0.50	1.0								U
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>45.19</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>90.4</i>	<i>70 - 123</i>				
<i>Surr: 4-Bromofluorobenzene</i>	<i>48.92</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>97.8</i>	<i>82 - 115</i>				
<i>Surr: Dibromofluoromethane</i>	<i>43.69</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>87.4</i>	<i>73 - 126</i>				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MBLK	Sample ID: VBLKW-181218	Units: ug/L		Analysis Date: 18-Dec-2018 13:34						
Client ID:	Run ID: VOA9_329535	SeqNo: 4872159		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	52.1	1.0	50	0	104	81 - 120				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
LCS	Sample ID: VLCSW-1812018	Units: ug/L			Analysis Date: 18-Dec-2018 13:58					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872160	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	21.7	1.0	20	0	109	77 - 118				
1,1,1-Trichloroethane	20.06	1.0	20	0	100	70 - 130				
1,1,2,2-Tetrachloroethane	21.61	1.0	20	0	108	70 - 120				
1,1,2-Trichlor-1,2,2-trifluoroethane	21.66	1.0	20	0	108	70 - 130				
1,1,2-Trichloroethane	21.9	1.0	20	0	109	77 - 113				
1,1-Dichloroethane	20.96	1.0	20	0	105	71 - 122				
1,1-Dichloroethene	20.63	1.0	20	0	103	70 - 130				
1,1-Dichloropropene	22.62	1.0	20	0	113	78 - 118				
1,2,3-Trichlorobenzene	22.19	1.0	20	0	111	70 - 130				
1,2,3-Trichloropropane	21.99	1.0	20	0	110	70 - 127				
1,2,4-Trichlorobenzene	21.92	1.0	20	0	110	77 - 126				
1,2,4-Trimethylbenzene	23.08	1.0	20	0	115	73 - 121				
1,2-Dibromo-3-chloropropane	19.69	1.0	20	0	98.4	70 - 130				
1,2-Dibromoethane	22.19	1.0	20	0	111	76 - 123				
1,2-Dichlorobenzene	21.19	1.0	20	0	106	77 - 113				
1,2-Dichloroethane	20.79	1.0	20	0	104	70 - 124				
1,2-Dichloropropane	23.16	1.0	20	0	116	72 - 119				
1,3,5-Trimethylbenzene	23.2	1.0	20	0	116	75 - 118				
1,3-Dichlorobenzene	21.7	1.0	20	0	109	78 - 118				
1,3-Dichloropropane	22.28	1.0	20	0	111	75 - 116				
1,4-Dichlorobenzene	22.92	1.0	20	0	115	79 - 113				S
2,2-Dichloropropane	21.28	1.0	20	0	106	70 - 130				
2-Butanone	42.42	2.0	40	0	106	70 - 130				
2-Chlorotoluene	22.49	1.0	20	0	112	70 - 128				
2-Hexanone	46.58	2.0	40	0	116	70 - 130				
4-Chlorotoluene	22.67	1.0	20	0	113	74 - 126				
4-Isopropyltoluene	23.85	1.0	20	0	119	74 - 126				
4-Methyl-2-pentanone	46.52	2.0	40	0	116	70 - 130				
Acetone	42.82	2.0	40	0	107	70 - 130				
Benzene	22.64	1.0	20	0	113	74 - 120				
Bromobenzene	21.59	1.0	20	0	108	78 - 113				
Bromochloromethane	22.21	1.0	20	0	111	76 - 124				
Bromodichloromethane	20.86	1.0	20	0	104	74 - 122				
Bromoform	19.3	1.0	20	0	96.5	73 - 128				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
LCS	Sample ID: VLCSW-1812018	Units: ug/L			Analysis Date: 18-Dec-2018 13:58					
Client ID:	Run ID: VOA9_329535	SeqNo: 4872160	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Bromomethane	23.28	1.0	20	0	116	70 - 130				
Carbon disulfide	43.3	2.0	40	0	108	70 - 130				
Carbon tetrachloride	21.74	1.0	20	0	109	71 - 125				
Chlorobenzene	21.78	1.0	20	0	109	76 - 113				
Chloroethane	23.06	1.0	20	0	115	70 - 130				
Chloroform	20	1.0	20	0	100	71 - 121				
Chloromethane	24.03	1.0	20	0	120	70 - 129				
cis-1,2-Dichloroethene	20.82	1.0	20	0	104	75 - 122				
cis-1,3-Dichloropropene	22.79	1.0	20	0	114	73 - 127				
Dibromochloromethane	21.85	1.0	20	0	109	77 - 122				
Dibromomethane	21.43	1.0	20	0	107	78 - 121				
Dichlorodifluoromethane	21.72	1.0	20	0	109	70 - 130				
Ethylbenzene	22.49	1.0	20	0	112	77 - 117				
Hexachlorobutadiene	26.61	1.0	20	0	133	70 - 130				S
Isopropylbenzene	23.13	1.0	20	0	116	73 - 127				
m,p-Xylene	45.63	2.0	40	0	114	77 - 122				
Methylene chloride	22.57	2.0	20	0	113	70 - 127				
Naphthalene	22.83	1.0	20	0	114	70 - 130				
n-Butylbenzene	24.18	1.0	20	0	121	72 - 130				
n-Propylbenzene	23.41	1.0	20	0	117	73 - 124				
o-Xylene	22.9	1.0	20	0	114	75 - 119				
sec-Butylbenzene	23.6	1.0	20	0	118	73 - 128				
Styrene	23.39	1.0	20	0	117	72 - 126				
tert-Butylbenzene	23.13	1.0	20	0	116	73 - 124				
Tetrachloroethene	22.36	1.0	20	0	112	76 - 119				
Toluene	22.56	1.0	20	0	113	77 - 118				
trans-1,2-Dichloroethene	20.73	1.0	20	0	104	72 - 127				
trans-1,3-Dichloropropene	19.97	1.0	20	0	99.9	77 - 119				
Trichloroethene	21.82	1.0	20	0	109	77 - 121				
Trichlorofluoromethane	20.41	1.0	20	0	102	70 - 130				
Vinyl chloride	23.07	1.0	20	0	115	70 - 130				
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>44.22</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>88.4</i>	<i>70 - 130</i>				
<i>Surr: 4-Bromofluorobenzene</i>	<i>51.18</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>102</i>	<i>82 - 115</i>				
<i>Surr: Dibromofluoromethane</i>	<i>44.56</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>89.1</i>	<i>73 - 126</i>				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
LCS	Sample ID: VLCSW-1812018	Units: ug/L		Analysis Date: 18-Dec-2018 13:58						
Client ID:	Run ID: VOA9_329535	SeqNo: 4872160		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	52.7	1.0	50	0	105	81 - 120				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MS	Sample ID: HS18120278-03MS	Units: ug/L			Analysis Date: 18-Dec-2018 17:41					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872166	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	23.88	1.0	20	0	119	70 - 120				
1,1,1-Trichloroethane	22.68	1.0	20	0	113	70 - 130				
1,1,2,2-Tetrachloroethane	23.94	1.0	20	0	120	70 - 123				
1,1,2-Trichlor-1,2,2-trifluoroethane	22.2	1.0	20	0	111	70 - 130				
1,1,2-Trichloroethane	24.23	1.0	20	0	121	70 - 117				S
1,1-Dichloroethane	23.28	1.0	20	0	116	70 - 127				
1,1-Dichloroethene	23.18	1.0	20	0.7106	112	70 - 130				
1,1-Dichloropropene	25.8	1.0	20	0	129	70 - 129				S
1,2,3-Trichlorobenzene	24.51	1.0	20	0	123	70 - 130				
1,2,3-Trichloropropane	23.25	1.0	20	0	116	70 - 130				
1,2,4-Trichlorobenzene	23.82	1.0	20	0	119	70 - 125				
1,2,4-Trimethylbenzene	25.68	1.0	20	0	128	70 - 125				S
1,2-Dibromo-3-chloropropane	21.8	1.0	20	0	109	70 - 130				
1,2-Dibromoethane	24.17	1.0	20	0	121	70 - 124				
1,2-Dichlorobenzene	23.25	1.0	20	0	116	70 - 115				S
1,2-Dichloroethane	22.65	1.0	20	0	113	70 - 127				
1,2-Dichloropropane	25.31	1.0	20	0	127	70 - 122				S
1,3,5-Trimethylbenzene	25.78	1.0	20	0	129	70 - 126				S
1,3-Dichlorobenzene	23.52	1.0	20	0	118	70 - 119				
1,3-Dichloropropane	24.4	1.0	20	0	122	70 - 121				S
1,4-Dichlorobenzene	25.75	1.0	20	0	129	70 - 114				S
2,2-Dichloropropane	21.93	1.0	20	0	110	70 - 130				
2-Butanone	45.49	2.0	40	0	114	70 - 130				
2-Chlorotoluene	24.73	1.0	20	0	124	70 - 130				
2-Hexanone	51.07	2.0	40	0	128	70 - 130				
4-Chlorotoluene	25.15	1.0	20	0	126	70 - 130				
4-Isopropyltoluene	26.9	1.0	20	0	134	70 - 130				S
4-Methyl-2-pentanone	50.44	2.0	40	0	126	70 - 130				
Acetone	46	2.0	40	0	115	70 - 130				
Benzene	25.3	1.0	20	0	126	70 - 127				
Bromobenzene	23.78	1.0	20	0	119	70 - 115				S
Bromochloromethane	24.54	1.0	20	0	123	70 - 127				
Bromodichloromethane	22.91	1.0	20	0	115	70 - 124				
Bromoform	20.79	1.0	20	0	104	70 - 129				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MS	Sample ID: HS18120278-03MS	Units: ug/L			Analysis Date: 18-Dec-2018 17:41					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872166	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Bromomethane	12.09	1.0	20	0	60.5	70 - 130				S
Carbon disulfide	44.81	2.0	40	0	112	70 - 130				
Carbon tetrachloride	24.05	1.0	20	0	120	70 - 130				
Chlorobenzene	25.62	1.0	20	1.542	120	70 - 114				S
Chloroethane	24.72	1.0	20	0	124	70 - 130				
Chloroform	22.07	1.0	20	0	110	70 - 125				
Chloromethane	15.68	1.0	20	0	78.4	70 - 130				
cis-1,2-Dichloroethene	44.9	1.0	20	21.65	116	70 - 128				
cis-1,3-Dichloropropene	24.3	1.0	20	0	121	70 - 125				
Dibromochloromethane	23.74	1.0	20	0	119	70 - 124				
Dibromomethane	22.87	1.0	20	0	114	70 - 124				
Dichlorodifluoromethane	10.42	1.0	20	0	52.1	70 - 130				S
Ethylbenzene	25.15	1.0	20	0	126	70 - 124				S
Hexachlorobutadiene	28.15	1.0	20	0	141	70 - 130				S
Isopropylbenzene	26.04	1.0	20	0	130	70 - 130				S
m,p-Xylene	50.97	2.0	40	0	127	70 - 130				
Methylene chloride	24.38	2.0	20	0	122	70 - 128				
Naphthalene	25.37	1.0	20	0	127	70 - 130				
n-Butylbenzene	26.87	1.0	20	0	134	70 - 130				S
n-Propylbenzene	26.34	1.0	20	0	132	70 - 130				S
o-Xylene	25.44	1.0	20	0	127	70 - 124				S
sec-Butylbenzene	26.76	1.0	20	0	134	70 - 130				S
Styrene	25.73	1.0	20	0	129	70 - 130				
tert-Butylbenzene	26.06	1.0	20	0	130	70 - 130				S
Tetrachloroethene	25.53	1.0	20	0	128	70 - 130				
Toluene	25.16	1.0	20	0	126	70 - 123				S
trans-1,2-Dichloroethene	23.47	1.0	20	0	117	70 - 130				
trans-1,3-Dichloropropene	21.37	1.0	20	0	107	70 - 121				
Trichloroethene	72.45	1.0	20	47.56	124	70 - 129				
Trichlorofluoromethane	20.92	1.0	20	0	105	70 - 130				
Vinyl chloride	20.18	1.0	20	0	101	70 - 130				
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>44.17</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>88.3</i>	<i>70 - 126</i>				
<i>Surr: 4-Bromofluorobenzene</i>	<i>51.45</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>103</i>	<i>81 - 113</i>				
<i>Surr: Dibromofluoromethane</i>	<i>45.03</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>90.1</i>	<i>77 - 123</i>				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MS	Sample ID: HS18120278-03MS	Units: ug/L			Analysis Date: 18-Dec-2018 17:41					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872166		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	53.03	1.0	50	0	106	82 - 127				

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MSD	Sample ID: HS18120278-03MSD	Units: ug/L			Analysis Date: 18-Dec-2018 18:06					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872167	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	22.94	1.0	20	0	115	70 - 120	23.88	3.99	20	
1,1,1-Trichloroethane	21.46	1.0	20	0	107	70 - 130	22.68	5.56	20	
1,1,2,2-Tetrachloroethane	23.01	1.0	20	0	115	70 - 123	23.94	3.93	20	
1,1,2-Trichlor-1,2,2-trifluoroethane	20.91	1.0	20	0	105	70 - 130	22.2	6	20	
1,1,2-Trichloroethane	23.16	1.0	20	0	116	70 - 117	24.23	4.52	20	
1,1-Dichloroethane	22.19	1.0	20	0	111	70 - 127	23.28	4.8	20	
1,1-Dichloroethene	21.54	1.0	20	0.7106	104	70 - 130	23.18	7.34	20	
1,1-Dichloropropene	24.29	1.0	20	0	121	70 - 129	25.8	6.04	20	
1,2,3-Trichlorobenzene	22.99	1.0	20	0	115	70 - 130	24.51	6.4	20	
1,2,3-Trichloropropane	21.98	1.0	20	0	110	70 - 130	23.25	5.62	20	
1,2,4-Trichlorobenzene	22.88	1.0	20	0	114	70 - 125	23.82	4.03	20	
1,2,4-Trimethylbenzene	24.38	1.0	20	0	122	70 - 125	25.68	5.16	20	
1,2-Dibromo-3-chloropropane	20.49	1.0	20	0	102	70 - 130	21.8	6.15	20	
1,2-Dibromoethane	23.25	1.0	20	0	116	70 - 124	24.17	3.85	20	
1,2-Dichlorobenzene	22.08	1.0	20	0	110	70 - 115	23.25	5.16	20	
1,2-Dichloroethane	21.98	1.0	20	0	110	70 - 127	22.65	3	20	
1,2-Dichloropropane	24.76	1.0	20	0	124	70 - 122	25.31	2.19	20	S
1,3,5-Trimethylbenzene	24.4	1.0	20	0	122	70 - 126	25.78	5.52	20	
1,3-Dichlorobenzene	22.5	1.0	20	0	112	70 - 119	23.52	4.46	20	
1,3-Dichloropropane	23.35	1.0	20	0	117	70 - 121	24.4	4.39	20	
1,4-Dichlorobenzene	24.57	1.0	20	0	123	70 - 114	25.75	4.7	20	S
2,2-Dichloropropane	20.71	1.0	20	0	104	70 - 130	21.93	5.72	20	
2-Butanone	45.19	2.0	40	0	113	70 - 130	45.49	0.66	20	
2-Chlorotoluene	23.65	1.0	20	0	118	70 - 130	24.73	4.47	20	
2-Hexanone	49.4	2.0	40	0	124	70 - 130	51.07	3.32	20	
4-Chlorotoluene	23.94	1.0	20	0	120	70 - 130	25.15	4.92	20	
4-Isopropyltoluene	25.24	1.0	20	0	126	70 - 130	26.9	6.37	20	
4-Methyl-2-pentanone	48.64	2.0	40	0	122	70 - 130	50.44	3.64	20	
Acetone	43.54	2.0	40	0	109	70 - 130	46	5.49	20	
Benzene	24.46	1.0	20	0	122	70 - 127	25.3	3.38	20	
Bromobenzene	22.37	1.0	20	0	112	70 - 115	23.78	6.13	20	
Bromochloromethane	23.65	1.0	20	0	118	70 - 127	24.54	3.69	20	
Bromodichloromethane	22.31	1.0	20	0	112	70 - 124	22.91	2.64	20	
Bromoform	20.15	1.0	20	0	101	70 - 129	20.79	3.11	20	

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260							
MSD		Sample ID: HS18120278-03MSD		Units: ug/L		Analysis Date: 18-Dec-2018 18:06					
Client ID: 12WW24-181204		Run ID: VOA9_329535		SeqNo: 4872167		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual	
Bromomethane	13.33	1.0	20	0	66.6	70 - 130	12.09	9.71	20	S	
Carbon disulfide	42.21	2.0	40	0	106	70 - 130	44.81	5.98	20		
Carbon tetrachloride	23.31	1.0	20	0	117	70 - 130	24.05	3.09	20		
Chlorobenzene	24.33	1.0	20	1.542	114	70 - 114	25.62	5.14	20		
Chloroethane	20.67	1.0	20	0	103	70 - 130	24.72	17.8	20		
Chloroform	21.16	1.0	20	0	106	70 - 125	22.07	4.18	20		
Chloromethane	14.94	1.0	20	0	74.7	70 - 130	15.68	4.84	20		
cis-1,2-Dichloroethene	42.81	1.0	20	21.65	106	70 - 128	44.9	4.77	20		
cis-1,3-Dichloropropene	23.85	1.0	20	0	119	70 - 125	24.3	1.86	20		
Dibromochloromethane	23.1	1.0	20	0	115	70 - 124	23.74	2.73	20		
Dibromomethane	22.89	1.0	20	0	114	70 - 124	22.87	0.084	20		
Dichlorodifluoromethane	9.87	1.0	20	0	49.3	70 - 130	10.42	5.41	20	S	
Ethylbenzene	23.86	1.0	20	0	119	70 - 124	25.15	5.24	20		
Hexachlorobutadiene	26.61	1.0	20	0	133	70 - 130	28.15	5.63	20	S	
Isopropylbenzene	24.62	1.0	20	0	123	70 - 130	26.04	5.58	20		
m,p-Xylene	48.45	2.0	40	0	121	70 - 130	50.97	5.06	20		
Methylene chloride	22.91	2.0	20	0	115	70 - 128	24.38	6.23	20		
Naphthalene	24.2	1.0	20	0	121	70 - 130	25.37	4.71	20		
n-Butylbenzene	25.57	1.0	20	0	128	70 - 130	26.87	4.97	20		
n-Propylbenzene	24.76	1.0	20	0	124	70 - 130	26.34	6.18	20		
o-Xylene	24	1.0	20	0	120	70 - 124	25.44	5.82	20		
sec-Butylbenzene	25.1	1.0	20	0	125	70 - 130	26.76	6.41	20		
Styrene	24.44	1.0	20	0	122	70 - 130	25.73	5.15	20		
tert-Butylbenzene	24.73	1.0	20	0	124	70 - 130	26.06	5.27	20		
Tetrachloroethene	24.06	1.0	20	0	120	70 - 130	25.53	5.9	20		
Toluene	23.77	1.0	20	0	119	70 - 123	25.16	5.68	20		
trans-1,2-Dichloroethene	21.99	1.0	20	0	110	70 - 130	23.47	6.52	20		
trans-1,3-Dichloropropene	20.89	1.0	20	0	104	70 - 121	21.37	2.3	20		
Trichloroethene	68.98	1.0	20	47.56	107	70 - 129	72.45	4.9	20		
Trichlorofluoromethane	19.31	1.0	20	0	96.6	70 - 130	20.92	8.01	20		
Vinyl chloride	18.76	1.0	20	0	93.8	70 - 130	20.18	7.31	20		
<i>Surr: 1,2-Dichloroethane-d4</i>	43.56	1.0	50	0	87.1	70 - 126	44.17	1.38	20		
<i>Surr: 4-Bromofluorobenzene</i>	51.09	1.0	50	0	102	81 - 113	51.45	0.704	20		
<i>Surr: Dibromofluoromethane</i>	44.93	1.0	50	0	89.9	77 - 123	45.03	0.218	20		

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
 Project: LHAARP - 12
 WorkOrder: HS18120278

QC BATCH REPORT

Batch ID: R329535		Instrument: VOA9		Method: SW8260						
MSD	Sample ID: HS18120278-03MSD	Units: ug/L			Analysis Date: 18-Dec-2018 18:06					
Client ID: 12WW24-181204	Run ID: VOA9_329535	SeqNo: 4872167		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
<i>Surr: Toluene-d8</i>	52.59	1.0	50	0	105	82 - 127	53.03	0.822	20	

The following samples were analyzed in this batch:

HS18120278-01	HS18120278-02	HS18120278-03	HS18120278-04
HS18120278-05			

ALS Houston, US

Date: 19-Dec-18

Client: Aptim Environmental & Infrastructure, Inc.
Project: LHAARP - 12
WorkOrder: HS18120278

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
North Carolina	624-2018	31-Dec-2018
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	22-Dec-2018
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019

Sample Receipt Checklist

Client Name: CBI-Houston
 Work Order: HS18120278

Date/Time Received: **05-Dec-2018 09:30**
 Received by: **RPG**

Checklist completed by: Raegen Giga 6-Dec-2018 Reviewed by: Sonia West 10-Dec-2018
 eSignature Date eSignature Date

Matrices: **GW** Carrier name: **ALS.HS**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- Chain of custody present? Yes No
- Chain of custody signed when relinquished and received? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- TX1005 solids received in hermetically sealed vials? Yes No N/A
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.8c/2.1c uc/c IR 25
 Cooler(s)/Kit(s): 25741
 Date/Time sample(s) sent to storage: 12/05/2018 19:55

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

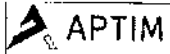
Login Notes:

Client Contacted: Date Contacted: Person Contacted:

Contacted By: Regarding:

Comments:

Corrective Action:



COC ID: **LHAAP12-DEC2018-ALSHT-1812-** TURNAROUND TIME: normal RUSH: Page 1 of


PROJECT/CLIENT INFO				LABORATORY				OTHER INFO			
Facility Name	Longhorn AAP			Lab Name	ALS Laboratories			Email Invoice To	Fedinvoices@aptim.com		
Project Number	501032			Lab Contact	RJ Modashia			Email Report To	Susan.Huang@aptim.com		
Address	LHAAP-12 1203-B East Grand Avenue PMB 202			Email	RJ.Modashia@alsglobal.com			Mail Reports To	Susan Huang		
City	Marshall	State	TX	Address	10450 Stanchiff Rd., Suite 210			Address	4005 Port Chicago Highway, Suit 200		
Postal Code	75670	Country	USA	City	Houston	State	TX	City	Concord	State	CA
Phone Number	713.243.7264			Postal Code	77099	Country	USA	Postal Code	94520	Country	USA
Project Manager	Praveen Srivastav			Phone Number	281.575.2279 or 281.530.5656			Shipping Company			

SAMPLE DETAILS										ANALYSIS REQUESTED
Sample ID	Location	Start Depth	End Depth	Depth Unit	Field Matrix	Date	Time (24hr)	# Of Cont.	ANALYSIS	Vials by 8260B (3-6ml Voa Vials with CD)
12WW21-181204	LHAAP12	22.20	22.43		WG	12/4/18	1005	3	X	
12WW21-181204-FD	LHAAP12	22.20	22.43		WG	12/4/18	1005	3	X	
12WW24-181204	LHAAP12	22.87	23.12		WG	12/4/18	110	3	X	
12WW24-181204-MS	LHAAP12	22.87	23.12		WG	12/4/18	1110	3	X	
12WW24-181204-MSD	LHAAP12	22.87	23.12		WG	12/4/18	1110	3	X	
12WW20-181204	LHAAP12	18.35	18.58		WG	12/4/18	1205	3	X	
Trip BLANK					W	12/4/18		2	X	

HS18120278
Aptim Environmental & Infrastructure, Inc.
LHAARP - 12



ADDITIONAL COMMENTS/SPECIAL INSTRUCTIONS	RELINQUISHED BY/AFFILIATION	DATE/TIME	ACCEPTED BY/AFFILIATION	DATE/TIME
	<i>Srinivas Beesaga</i> / BHAAR	12/4/18 1400	R Ciga	12/5/18 09:30 am
				cooler # 25741
				TEMP - 1.8c
				12/25 CF +0.3c

 ALS 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5687	CUSTODY SEAL		Seal Broken By: <i>SM</i>
	Date: <i>12/11/18</i>	Time: <i>14:00</i>	Date: <i>12/05/18</i>
	Name: <i>Scott Beesinger</i>		Comp any: <i>STATE</i>

25741 DEC 05 2018



Must Deliver Next Business Day
Time and Tempature Sensitive!

25741

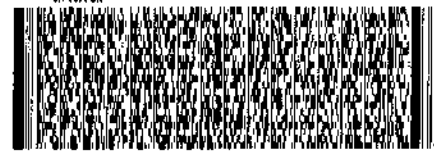
ORIGIN ID: SGRA (903) 930-6193
 ATT: SCOTT BEESINGER
 APTIM ENVIRONMENTAL & INFRASTR. INC
 1203-B EAST GRAND AVE PMB202
 MARSHALL, TX 75870
 UNITED STATES US

SHIP DATE: 29NOV18
 ACTWT: 1.00 LB MAN
 CAD: 300130/CAF3211
 DIMS: 19x15x13 IN

TO **CLIENT SERVICES**
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON TX 77099

(281) 530-6656
 REF: LHAAP-BO 62490-RJ

RMA: III III III

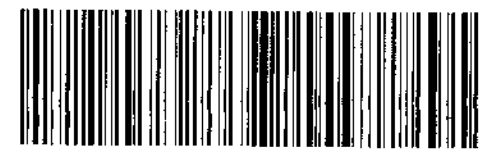


FedEx
 TRK#
 0221 4380 9535 0865

WED - 05 DEC 10:30A
 PRIORITY OVERNIGHT

AB SGRA

77099
 TX-US
 IAH



FID 162705 04DEC18 666A 553C1/71FF/BC8A

Chemical Analytical Data Review/Validation Form

Report Type: { } Preliminary {X} Final

Project Number: 501032

Project Name: LHAAP-12, Dec 2018 event

Analysis/Method: VOCs/SW8260

Laboratory: ALS

SDG # HS18120278

Evaluated By: Shuang

Date Evaluated: 1/7/2019

Sample Number(s): 12MW21-181204, 12MW21-181204FD, 12MW24-181204, 12MW20-181204, AND trip blank

REVIEW QUESTIONS	YES	NO	COMMENTS
1. Were holding times met?	X		Sampled on 12/4/2018, analyzed on 12/18/2018
2. Were sample preservation requirements met? (Sample condition, preservation, containers, temperature, etc.)	X		1.8 deg C, preserved with HCL
3. Were QAPP specified PQLs achieved?	X		LOQ>LOD>DL, ND at LOD
4. Were measurement results for all QAPP-specific target analytes reported?	X		
5. Was a method blank prepared and analyzed with each batch?	X		
6. Were target analytes reported in the method blank above the PQL?		X	VOCs ND in the blank
7. Was an equipment blank collected and analyzed at the required frequency stated in the site-specific FSP?		X	
8. Were target analytes reported in field blank analyses (trip or equipment) above the PQL?		X	Acetone was detected at 2.7 ppb>LOQ (2 ppb). Acetone was ND in all samples, no impact
9. Was a field duplicate analyzed? Were RPDs within QAPP specifications?	X		12MW21/FD, VOCS ND in the pair
10. Surrogate Recoveries – Were all samples spiked prior to purging or preparation?	X		
11. Were surrogate recoveries within QAPP specifications?	X		
12. Was an LCD/LCSD pair prepared and analyzed with each batch?	X		
13. Were LCS/LCSD RPDs within recoveries within QAPP specifications?		X	1,4-dicholobenzene 115% (79-113% lab, 79-118% DoD), hexachlorobutadiene at 133% (73-127% lab, 66-134% DoD), both are within the DoD limits. anlyates ND, no impact
14. Was a MS/MSD pair prepared with each batch?	X		

REVIEW QUESTIONS	YES	NO	COMMENTS
15. Is the MS/MSD parent sample a project sample?		X	12MW24, chlorobenzene 120/114% (DoD 82-118%, lab 70-114%), J/8A in all detected sample The lab report indicated high bias MS/MSD recoveries for other VOCs, however these analytes were ND and were not affected by high biased MS/MSD. Bromomethane and dichlorofluoromethane in MS/MSD were within the DoD LCS control limits but below lab in house limits. no impact since the DoD requirements were met.
16. Were initial calibration standards analyzed at the QAPP-specific frequency for each instrument?	X		
17. Were these results within project specifications?	X		
18. Were continuing calibration standards analyzed at the QAPP-specific frequency for each instrument?	X		
19. Were these results within project specifications?	X		
20. Were laboratory-generated Quality Control Exception Reports (i.e., QCERs) issued? If yes, summarize contents.			
21. Were lab comments included in the report? If yes, summarize contents			

12MW24:

1,1-dichloroethane: 0.71 ppb

Chlorobenzene: 1.5 ppb

Cis-1,2dichloroethene: 22 ppb

Trichloroethene: 48 ppb

All other VOCs ND in this sample.

VOCs ND all other samples

1/7/2019: reviewed EDD. Results in EDD matched the hardcopy

Appendix E

Mann-Kendall Trend Test Results

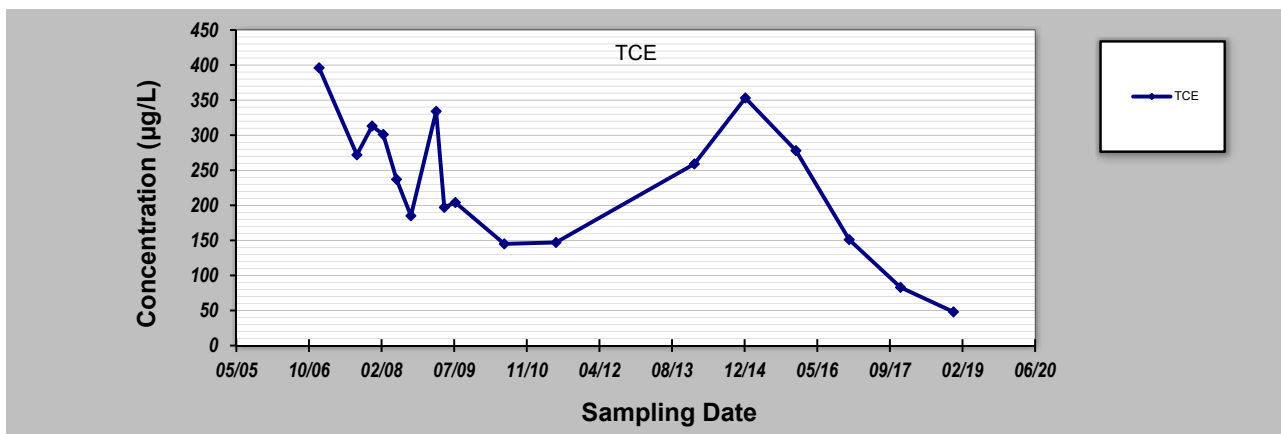
GSI MANN-KENDALL TOOLKIT for Constituent Trend Analysis

Evaluation Date: **11-Feb-19** Job ID: **501032**
 Facility Name: **LHAAP-12** Constituent: **12WW24**
 Conducted By: **Robert Mayer** Concentration Units: **µg/L**

Sampling Point ID: **TCE**

Sampling Event	Sampling Date	12WW24 CONCENTRATION (µg/L)					
1	19-Dec-06	396					
2	5-Sep-07	272					
3	19-Dec-07	313					
4	5-Mar-08	301					
5	4-Jun-08	237					
6	11-Sep-08	185					
7	4-Mar-09	334					
8	29-Apr-09	197					
9	14-Jul-09	204					
10	16-Jun-10	145					
11	7-Jun-11	147					
12	15-Jan-14	259					
13	1-Jan-15	353					
14	15-Dec-15	278					
15	16-Dec-16	151					
16	4-Dec-17	83					
17	4-Dec-18	48					
18							
19							
20							

Coefficient of Variation: **0.42**
 Mann-Kendall Statistic (S): **-60**
 Confidence Factor: **99.3%**
 Concentration Trend: **Decreasing**



Notes:

- At least four independent sampling events per well are required for calculating the trend. *Methodology is valid for 4 to 40 samples.*
- Confidence in Trend = Confidence (in percent) that constituent concentration is increasing (S>0) or decreasing (S<0): >95% = Increasing or Decreasing; ≥ 90% = Probably Increasing or Probably Decreasing; < 90% and S>0 = No Trend; < 90%, S≤0, and COV ≥ 1 = No Trend; < 90% and COV < 1 = Stable.
- Methodology based on "MAROS: A Decision Support System for Optimizing Monitoring Plans", J.J. Aziz, M. Ling, H.S. Rifai, C.J. Newell, and J.R. Gonzales, *Ground Water*, 41(3):355-367, 2003.

DISCLAIMER: The GSI Mann-Kendall Toolkit is available "as is". Considerable care has been exercised in preparing this software product; however, no party, including without limitation GSI Environmental Inc., makes any representation or warranty regarding the accuracy, correctness, or completeness of the information contained herein, and no such party shall be liable for any direct, indirect, consequential, incidental or other damages resulting from the use of this product or the information contained herein. Information in this publication is subject to change without notice. GSI Environmental Inc., disclaims any responsibility or obligation to update the information contained herein.

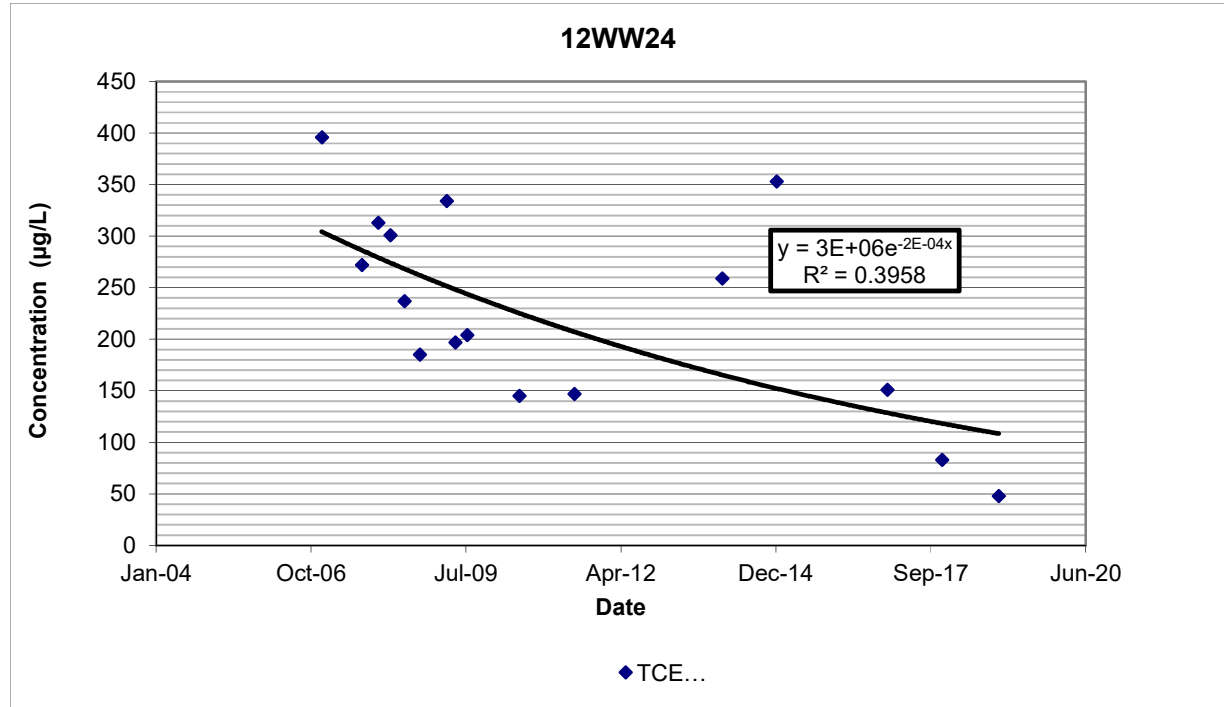
GSI Environmental Inc., www.gsi-net.com

Appendix F

Degradation Rates

**Time-Dependent Attenuation Rate Constant and Estimated Cleanup Time
12WW24**

Date	TCE (µg/L)
19-Dec-06	396
5-Sep-07	272
19-Dec-07	313
5-Mar-08	301
4-Jun-08	237
11-Sep-08	185
4-Mar-09	334
29-Apr-09	197
14-Jul-09	204
16-Jun-10	145
7-Jun-11	147
15-Jan-14	259
1-Jan-15	353
15-Dec-15	278
16-Dec-16	151
4-Dec-17	83
4-Dec-18	48



Chemical: TCE Well ID	Attenuation Rate Constant (day ⁻¹)	Attenuation Half-life (days)	Attenuation Half-life (years)	Current Conc. (µg/L)	Target Concentration for TCE (µg/L)	Estimated Cleanup Time (years)
12WW24	0.0002	2937.1	8.0	48	5	26.3

Notes:

The estimated cleanup time was calculated as the time it would take the most recent detected TCE concentration to reach the MCL using the site-specific attenuation rate, and assuming first order degradation kinetics.

µg/L - micrograms per liter

DCA - dichloroethane

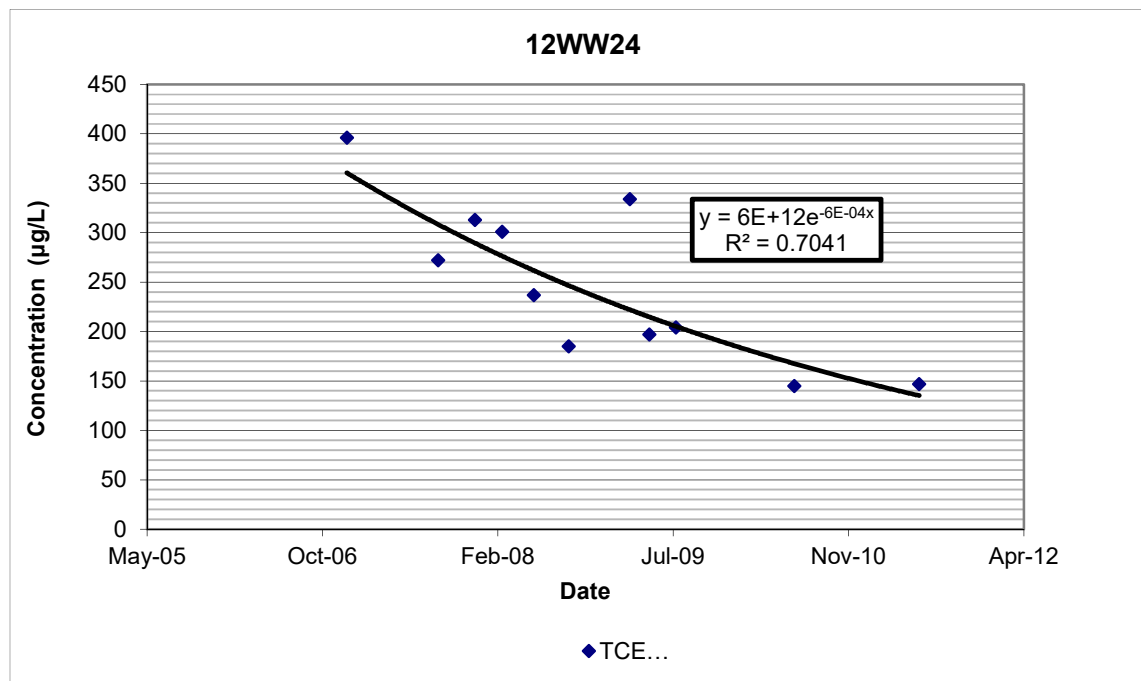
DCE - dichloroethene

MCL - maximum contaminant level

TCE - trichloroethene

Time-Dependent Attenuation Rate Constant and Estimated Cleanup Time 12WW24

Date	TCE (µg/L)
19-Dec-06	396
5-Sep-07	272
19-Dec-07	313
5-Mar-08	301
4-Jun-08	237
11-Sep-08	185
4-Mar-09	334
29-Apr-09	197
14-Jul-09	204
16-Jun-10	145
7-Jun-11	147
15-Jan-14	259
1-Jan-15	353
15-Dec-15	278
16-Dec-16	151
4-Dec-17	83
4-Dec-18	48



Chemical: TCE Well ID	Attenuation Rate Constant (day ⁻¹)	Attenuation Half-life (days)	Attenuation Half-life (years)	Current Conc. (µg/L)	Target Concentration for TCE (µg/L)	Estimated Cleanup Time (years)
12WW24	0.0006	1155.2	3.2	147	5	15.4

Notes:

The estimated cleanup time was calculated as the time it would take the most recent detected TCE concentration to reach the MCL using the site-specific attenuation rate, and assuming first order degradation kinetics.

µg/L - micrograms per liter

DCA - dichloroethane

DCE - dichloroethene

MCL - maximum contaminant level

TCE - trichloroethene

Subject: Final Minutes, Monthly Managers' Meeting (MMM),
Longhorn Army Ammunition Plant (LHAAP)

Location of Meeting: Teleconference

Date of Meeting: 20 June 2019– 10:00 AM Central Standard Time (CDT)

Attendees:

Army BRAC: Rose Zeiler (RMZ)
 EPA: Dorelle Harrison
 TCEQ: April Palmie (AP)
 USACE: Aaron Williams (AW)
 AEC: Andrew Maly and Amanda Sherman
 USGS: Kent Becher
 Bhate: Kim Nemmers (KN)
 APTIM: Bill Foss (BF) and Susan Watson (SW)

Action Items

Bhate/APTIM: BF stated that a round of groundwater elevation readings at LHAAP-16 was collected at the end of May, and potentiometric maps were prepared to compare the May 2019 readings to the October 2018 readings. Bill stated that the flow direction does not appear to differ much from the October maps, but that the ground water elevation is about 3 to 4 feet higher. BF stated that a list of wells for the baseline event is also being developed and will be sent with the potentiometric maps for discussion.

For the Oil and Gas Water Well Drilling Impact Update, United States Fish and Wildlife (USFWS) was not on the call. RMZ stated that she thought it can be as simple as a protocol for new developments.

Defense Environmental Restoration Program (DERP) Performance Based Remediation (PBR) Update

KN stated that the leak at LHAAP-18/24 had been repaired but that the line out to 18WW17 was capped. When the 2-inch well was repaired, then water was entering from the line that runs to extraction well 18WW17. The entire line to the wood will need to be excavated to find the leak but the soil is too wet to complete this work at this time. KN stated that the water line for the ICTs is online and working but the repair to the line out to 18WW17 will not be done until July sometime. RMZ asked that the repairing of the line to extraction well 18WW17 be included as an action item that is tracked.

KN explained that the groundwater treatment plant (GWTP) was not operating currently due to the pump for the air stripper feed seizing up. KN stated that bearings and seals needed to be replaced and that the same repairs occurred about 1.5 years ago. So, KN stated that the reason for the repairs is being evaluated to determine if it can be avoided or if regular proactive management of the pump is needed in the future.

AW stated that the award for the transformer replacement will be this week or early next week, and then the installation will be about a month out. So the transformer is about one month out and should be installed the end of July and or beginning of August 2019. AW explained that the transformer was being improved with increased impedance, bayonet fusing and current limiting fusing to protect the transformer from a storm current surge.

RMZ asked if the revised schedule will work. BF stated that the sequencing of the work is flexible, and the excavation at LHAAP-17 can start ahead of the injections, and well installation at LHAAP-16. BF said that the site conditions are being evaluated, and then the transformer installation plans will be considered.

BF stated that the field work includes excavation at LHAAP-17 and LHAAP-03, baseline sampling at LHAAP-16, and injections at LHAAP-04, LHAAP-16, and eventually LHAAP-50. BF stated that the same field personnel are planned for most of the work. BF stated that the priority is the well installation at LHAAP-16. BF stated that well installation is needed at LHAAP- -12, -16, -50 and -67. BF said that a bulldozer will be obtained for Scott Beesinger (onsite GWTP operator) to create access for the installation of the new wells. BF plans to be onsite with Rob Mayer (bioremediation expert) July 1 and 2, 2019 to look over the site conditions. BF said that the planned well installations will be completed the week of July 15th if site conditions allow. BF said that the excavation at LHAAP-03 is planned to piggy-back on the LHAAP-17 excavation given how small of a dig it is. RMZ asked if things look good, how long will the field work last. BF said that the schedule calls for two months with a week and a half of float time.

AP asked when the UIC injection information will be submitted for LHAAP-04 and LHAAP-50. BF said that the information has not been put together, but would be shortly. RMZ pointed out that it is a 30 day out notification. AP said that what was done for LHAAP-16 was a good example.

RMZ asked about the potential well installation at LHAAP-03. AP asked about LHAAP-03 and excavating around the well versus abandonment of the well. BF said that the well is still being used for performance monitoring for LHAAP-58. AP said that her concern is that material will be left around the well could still be detected in the well if everything isn't removed. RMZ stated that she has similar concerns and had the depth of the well evaluated. BF said that 8 feet of the annular seal (grout) will be left in the ground above the top of the filter pack below the excavation, which will include hand-digging around the well. AP stated that it seems like a lot of effort hand digging to save a well. BF said that if the well gets damaged during the excavation, then the well will be replaced.

KN asked everyone to refer to the Document and Issues Tracking Table dated June 20, 2019.

- **Task 1** (Project Management) -
 - KN stated that the May 2019 MMM Minutes were final.
 - KN stated that the Restoration Advisory Board (RAB) Meeting was being held the following month.
- **Task 3** (LHAAP-03) – BF stated that response to comments (RTCs) from the Regulators on the Remedial Design (RD)/Remedial Action Work Plan (RAWP) for LHAAP-03 are prepared and will be going to the Army for review. The RTCs will then be issued to the Regulators.
- **Task 4** (LHAAP-04) – BF stated that there are no current documents.
- **Task 5** (LHAAP-12) – BF stated that the Regulators had no comments on the Draft 2018 Annual Remedial Action Operation (RA-O) Report. BF stated that a final cover will be provided along with a transmittal letter that indicates no comments were received on the draft report. BF asked if new pages need to be issued because the footers say “Draft” or if a letter say that the document is final with no comments would be sufficient. BF said that the version going into the administrative record (AR) will have the word “final” on the cover and will include the letter. New compact discs (CDs) will be send out with the word Final but the hard copy will have only a new cover sheet and letter. RMZ said that the Federal Facility Agreement needs to be followed. AP also said that a CD is not necessary if it can

be downloaded. AP wants to move away from CDs because they just get thrown away. AP said she would provide something in writing regarding elimination of CDs.

- **Task 6** (LHAAP-16) – BF stated that May 2019 potentiometric maps will be provided along with the October 2018 maps plus the proposed baseline sampling event wells.
- **Task 7** (LHAAP-17) – BF stated that no documents are currently in process for LHAAP-17.
- **Task 9** (LHAAP-37) – BF stated that LHAAP-37 Year 2, Quarter 3 was completed in May. BF stated that the validated data will be provided at the July 2019 MMM.
- **Task 10** (LHAAP-46) – BF stated that the 2nd Semi-Annual event is planned for August 2019.
- **Task 11** (LHAAP-50) – BF stated that sampling was completed in May and the validated data would be available for the July 2019 meeting. BF stated that the Regulatory comments on the Explanation of Significant Difference (ESD) for a contingency remedy were received and that responses to the ESD comments are in Army review; due to the Regulators by July 8, 2019. BF also stated that the RD/RAWP is under Army review.
- **Task 12** (LHAAP-58) – KN stated that the LHAAP-58 sampling was completed and that validated data will be provided at the July 2019 MMM. KN stated that the Year 5 RA-O Report will be prepared once the data is received and validated.
- **Task 13** (LHAAP-67) – BF stated that the 2nd semi-annual Year 5 sampling was completed in May. BF stated that validated data from the May 2019 event will be available for the July 2019 MMM.
- **Task 14** (LHAAP-001-R and –003-R) - KN stated that no documents are in process.
- **Task 16** (GWTP) – KN stated that the 4th Quarter GWTP Report is being delivered on June 20, 2019 via Federal Express. AP asked if an email was being sent to which KN stated that an email would be provided after the meeting. KN stated that the 1st Quarter 2019 GWTP Report is under internal review.
- **Task 17** (LHAAP-18/24) – KN stated that the LHAAP-18/24 analytical data from December 2018 is presented within the 4th Quarter GWTP Report.
- **Task 18** (Surface Water) – KN stated that surface water would be collected in June or early July 2019.
- **AR** – SW said that the December 2018 AR should be sent out next week. SW said that the list of documents for the AR is being completed for the 1st quarter.

Update on other DERP Sites

- **LHAAP 18/24** – AW explained that the Final Proposed Plan (PP) for LHAAP-18/24 will be placed into the AR. AW stated that the Record of Decision (ROD) is being prepared for submittal to the Regulators on September 6, 2019.
- **LHAAP-29** – AW stated that the LHAAP-29 PP is ready for AR. Regulator comments on the ROD have been addressed and are under Army legal review. RTCs will be released by July 15, 2019.
- **LHAAP-47** – AW stated that the Post-Screening Investigation (PSI) Report is ready for the AR. AW explained that the Addendum to the PSI Report has been through a few, internal iterations but should be submitted next week. The draft ROD should be submitted by August 17, 2019.
- **Five Year Review (FYR)** – RMZ said that the FYR has an errata sheet, and the Army is determining if figures can be revised. AP said that she doesn't think the FYR should go into the AR until the changes are made. RMZ concurred with AP.

Schedule Next Managers' Meeting

The next MMM will be held on 25 July 2019 at 10:30 am CDT at the Karnack Community Center if power is not restored to the trailer at the LHAAP GWTP.

Call concluded at 10:47 am CDT.

ACRONYM LIST

AP	April Palmie
APTIM	APTIM Federal Services, LLC
AR	Administrative Record
AW	Aaron Williams
BF	Bill Foss
Bhate	Bhate Environmental Associates, Inc.
BRAC	Base Realignment and Closure
CD	Compact Disc
CDT	Central Daylight Time
DERP	Defense Environmental Restoration Program
EPA	United States Environmental Protection Agency
ESD	Explanation of Significant Differences
FYR	Five Year Review
GWTP	Ground Water Treatment Plant
ICT	Interception-Collection Trench
KN	Kim Nemmers
LHAAP	Longhorn Army Ammunition Plant
MMM	Monthly Managers' Meeting
PBR	Performance-Based Remediation
PP	Proposed Plan
PSI	Post-Screening Investigation
RAB	Restoration Advisory Board
RA-O	remedial action – operation
RAWP	Remedial Action Work Plan
RD	Remedial Design
RMZ	Rose M. Zeiler
ROD	Record of Decision
RTC	Response to Comment
SW	Susan Watson
TCEQ	Texas Commission on Environmental Quality
USACE	United States Army Corps of Engineers
USFWS	United States Fish and Wildlife Service

**LHAAP Data Validated
May 2019**

GWTP Effluent	<i>Weekly Perchlorate Sampling – May 2019</i> Perchlorate (6850)
GWTP Effluent	<i>Weekly, Bi-Weekly, and Monthly Sampling – May 2019</i> Ammonia (350.3) Ortho-Phosphate (365.3) Organic Carbon (415.1) VOC (8260C) Metals (6020A) Hexavalent Chromium (7196A) 1,4-Dioxane (8270D-SIM) Anions (9056)
GWTP Influent	<i>Monthly Sampling – May 2019</i> Metals (6020A) Perchlorate (6850) Hexavalent Chromium (7196A)
GWTP Quarterly	<i>Influent and Effluent – May 2019</i> Oil and Grease (1664A) Perchlorate (6850) Metals (6020A) 1,4-Dioxane (8270D-SIM) Chemical Oxygen Demand (410.4) VOC (8260C) Anions (9056)

GWTP Weekly/Effluent Perchlorate Sampling - May 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24- SP650_050119_BIX 5/1/19	LH18/24- SP650_050719_BIX 5/7/19	LH18/24- SP650_050719_BIX 5/7/19	LH18/24- SP650_051419_BIX 5/14/19	LH18/24- SP650_052119_BIX 5/21/19	LH18/24- SP650_052919_BIX 5/29/19	LH18/24- SP650_052919_BIX 5/29/19
Location Description		Collected from a spigot on the discharge of effluent TK-650.							
			Weekly	Weekly	Monthly EFF	Weekly	Weekly	Weekly	Quarterly EFF
Perchlorate (6850)									
Perchlorate	µg/L	589	1.1 J	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U

µg/L - micrograms per liter

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

BIX - before ion exchange

GWTP Weekly Sampling - May 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24- SP650_050119 5/1/19	LH18/24- SP650_050719 5/7/19	LH18/24- SP650_051419 5/14/19	LH18/24- SP650_052119 5/21/19	LH18/24- SP650_052919 5/29/19
Location Description			GWTP—Collected from a spigot on the discharge of effluent TK-650. Sampled Weekly.				
Ammonia as N (350.3)							
Ammonia as N	mg/L	NV	24	21	14	11	10
Ortho-Phosphate (365.3)							
Ortho-Phosphate	mg/L	NV	6.2	4.95	4.22	2.47	2.21
Organic Carbon (415.1)							
Total Organic Carbon (TOC)	mg/L	NV	1.51	1.46	1.64	1.65	2.12

mg/L - milligrams per liter

NV - No Value

GWTP Bi-Weekly Sampling - May 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_050119 5/1/19	LH 18/24-SP650_051419 5/14/19	LH18/24-SP650_052919 5/29/19
Location Description			GWTP – Collected from a spigot on the discharge of effluent TK-650. Sampled Biweekly.		
Volatile Organic Compounds (8260C)					
1,1,1-Trichloroethane	µg/L	7,230	< 0.5 U	< 0.5 U	< 0.5 U
1,1,2-Trichloroethane	µg/L	216.9	< 0.5 U	< 0.5 U	< 0.5 U
1,1-Dichloroethane	µg/L	14,032	< 0.5 U	< 0.5 U	< 0.5 U
1,1-Dichloroethene	µg/L	253	< 0.5 U	< 0.5 U	< 0.5 U
1,2-Dichloroethane	µg/L	181	0.51 J	< 0.5 U	< 0.5 U
1,2-Dichloropropane	µg/L	5	< 0.5 U	< 0.5 U	< 0.5 U
Acetone	µg/L	2,395	< 1.0 U	< 1.0 U	< 1.0 U
Benzene	µg/L	181	< 0.5 U	< 0.5 U	< 0.5 U
Carbon tetrachloride	µg/L	181	< 0.5 U	< 0.5 U	< 0.5 U
Chlorobenzene	µg/L	47,180	< 0.5 U	< 0.5 U	< 0.5 U
Chloroform	µg/L	3,615	< 0.5 U	< 0.5 U	< 0.5 U
Ethylbenzene	µg/L	57,025	< 0.5 U	< 0.5 U	< 0.5 U
m,p-Xylene	µg/L	83.6	< 1.0 U	< 1.0 U	< 1.0 U
Methylene chloride	µg/L	1,699	< 1.0 U	< 1.0 U	< 1.0 U
o-Xylene	µg/L	83.6	< 0.5 U	< 0.5 U	< 0.5 U
Styrene	µg/L	5,987	< 0.5 U	< 0.5 U	< 0.5 U
Tetrachloroethene	µg/L	180.7	< 0.5 U	< 0.5 U	< 0.5 U
Toluene	µg/L	4,189	< 0.5 U	< 0.5 U	< 0.5 U
Trichloroethene	µg/L	181	0.65 J	0.56 J	0.85 J
Vinyl chloride	µg/L	72	< 0.5 U	< 0.5 U	< 0.5 U
Anions (9056)					
Chloride	mg/L	NV	302	231	252
Sulfate	mg/L	NV	15	24.1	34

µg/L - micrograms per liter

mg/L - milligrams per liter

U - Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

NV - No Value

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

GWTP Monthly Effluent Sampling - May 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_050719 5/7/19
Location Description			GWTP – Collected from a spigot on the discharge of effluent TK-650. Sampled monthly
Volatile Organic Compounds (8260C)			
1,1,1-Trichloroethane	µg/L	7,230	< 0.5 U
1,1,2-Trichloroethane	µg/L	216.9	< 0.5 U
1,1-Dichloroethane	µg/L	14,032	< 0.5 U
1,1-Dichloroethene	µg/L	253	< 0.5 U
1,2-Dichloroethane	µg/L	181	0.51 J
1,2-Dichloropropane	µg/L	5	< 0.5 U
Acetone	µg/L	2,395	< 1.0 U
Benzene	µg/L	181	< 0.5 U
Carbon tetrachloride	µg/L	181	< 0.5 U
Chlorobenzene	µg/L	47,180	< 0.5 U
Chloroform	µg/L	3,615	< 0.5 U
Ethylbenzene	µg/L	57,025	< 0.5 U
m,p-Xylene	µg/L	83.6	< 1.0 U
Methylene chloride	µg/L	1,699	< 1.0 U
o-Xylene	µg/L	83.6	< 0.5 U
Styrene	µg/L	5,987	< 0.5 U
Tetrachloroethene	µg/L	180.7	< 0.5 U
Toluene	µg/L	4,189	< 0.5 U
Trichloroethene	µg/L	181	0.65 J
Vinyl chloride	µg/L	72	< 0.5 U
Metals (6020A)			
Barium	mg/L	2	0.136
Lead	mg/L	0.0046	< 0.00100 U
Selenium	mg/L	0.012	< 0.00250 U
Silver	mg/L	0.003	< 0.000500 U
Hexavalent Chromium (7196A)			
Hexavalent Chromium	mg/L	0.1244	< 0.0100 U
Semi-Volatile Organic Compounds (8270D SIM)			
1,4-Dioxane	µg/L	134.2	21

µg/L - micrograms per liter

mg/L - milligrams per liter

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

GWTP Monthly Influent Sampling - May 2019

Location ID: Sample Date:	Units	LH18/24-SP140_050719 5/7/19
Location Description		GWTP – Collected from a spigot on the influent to TK-140. Sampled Monthly.
Metals (6020A)		
Selenium	mg/L	< 0.00250 U
Silver	mg/L	< 0.000500 U
Hexavalent Chromium (7196A)		
Hexavalent Chromium	mg/L	< 0.0100 U
Perchlorate (6850)		
Perchlorate	µg/L	4,200

mg/L - milligrams per liter

µg/L - micrograms per liter

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

GWTP Quarterly Effluent Sampling - May 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_052919 5/29/19
Location Description			GWTP – Collected from a spigot on the discharge of effluent TK-650. Sampled Quarterly.
Oil and Grease (1664A)			
Oil & Grease	mg/L	15	1.46 J
Chemical Oxygen Demand (410.4)			
Chemical Oxygen Demand	mg/L	200	16
Volatile Organic Compounds (8260C)			
1,1,1-Trichloroethane	µg/L	7,230	< 0.5 U
1,1,2-Trichloroethane	µg/L	216.9	< 0.5 U
1,1-Dichloroethane	µg/L	14,032	< 0.5 U
1,1-Dichloroethene	µg/L	253	< 0.5 U
1,2-Dichloroethane	µg/L	181	< 0.5 U
1,2-Dichloropropane	µg/L	5	< 0.5 U
Acetone	µg/L	2,395	< 2.0 U
Benzene	µg/L	181	< 0.5 U
Carbon tetrachloride	µg/L	181	< 0.5 U
Chlorobenzene	µg/L	47,180	< 0.5 U
Chloroform	µg/L	3,615	< 0.5 U
Ethylbenzene	µg/L	57,025	< 0.5 U
m,p-Xylene	µg/L	83.6	< 1.0 U
Methylene chloride	µg/L	1,699	< 0.5 U
o-Xylene	µg/L	83.6	< 0.5 U
Styrene	µg/L	5,987	< 0.5 U
Tetrachloroethene	µg/L	180.7	< 0.5 U
Toluene	µg/L	4,189	< 0.5 U
Trichloroethene	µg/L	181	0.92 J
Vinyl chloride	µg/L	72	< 0.5 U
Metals (6020A)			
Aluminum	mg/L	1.644	0.0422
Antimony	mg/L	NV	< 0.000500 U
Arsenic	mg/L	0.722	0.000699 J
Barium	mg/L	2	0.114
Beryllium	mg/L	NV	< 0.000500 U
Cadmium	mg/L	0.0034	< 0.000500 U
Calcium	mg/L	NV	8.42
Chromium	mg/L	0.752	0.00105 J
Cobalt	mg/L	11.495	0.00105 J
Iron	mg/L	2.395	0.195 J
Lead	mg/L	0.0046	< 0.00100 U
Magnesium	mg/L	NV	10.7

Manganese	mg/L	15.494	0.0833
Nickel	mg/L	0.184	0.00257
Potassium	mg/L	NV	1.46
Selenium	mg/L	0.012	< 0.00250 U
Silver	mg/L	0.003	< 0.000500 U
Sodium	mg/L	NV	280
Thallium	mg/L	NV	< 0.000500 U
Vanadium	mg/L	3.592	0.000969 J
Zinc	mg/L	0.31	0.0228
Mercury	mg/L	NV	0.0000900 J
Anions (9056)			
Chloride	mg/L	NV	253
Sulfate	mg/L	NV	33
Semi-Volatile Organic Compounds (8270D SIM)			
1,4-Dioxane	µg/L	134.2	5.1

µg/L - micrograms per liter

mg/L - milligrams per liter

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

NV - No Value

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

GWTP Quarterly Influent Sampling - May 2019

Location ID: Sample Date:	Units	LH18/24-SP140_052919 5/29/19
Location Description		GWTP – Collected from a spigot on the influent to TK-140. Sampled Quarterly.
Oil and Grease (1664A)		
Oil & Grease	mg/L	6.56
Chemical Oxygen Demand (410.4)		
Chemical Oxygen Demand	mg/L	15
Perchlorate (6850)		
Perchlorate	µg/L	4,900
Volatile Organic Compounds (8260C)		
1,1,1,2-Tetrachloroethane	µg/L	< 50 U
1,1,1-Trichloroethane	µg/L	< 50 U
1,1,2,2-Tetrachloroethane	µg/L	< 50 U
1,1,2-Trichloroethane	µg/L	< 50 U
1,1-Dichloroethane	µg/L	< 50 U
1,1-Dichloroethene	µg/L	< 50 U
1,1-Dichloropropene	µg/L	< 50 U
1,2,3-Trichlorobenzene	µg/L	< 50 U
1,2,3-Trichloropropane	µg/L	< 50 U
1,2,4-Trichlorobenzene	µg/L	< 50 U
1,2,4-Trimethylbenzene	µg/L	< 50 U
1,2-Dibromo-3-chloropropane	µg/L	< 50 U
1,2-Dibromoethane	µg/L	< 50 U
1,2-Dichlorobenzene	µg/L	< 50 U
1,2-Dichloroethane	µg/L	53 J
1,2-Dichloropropane	µg/L	< 50 U
1,3,5-Trimethylbenzene	µg/L	< 50 U
1,3-Dichlorobenzene	µg/L	< 50 U
1,3-Dichloropropane	µg/L	< 50 U
1,4-Dichlorobenzene	µg/L	< 50 U
2,2-Dichloropropane	µg/L	< 50 U
2-Butanone	µg/L	< 100 U
2-Chlorotoluene	µg/L	< 50 U
2-Hexanone	µg/L	< 100 U
4-Chlorotoluene	µg/L	< 50 U
4-Isopropyltoluene	µg/L	< 50 U
4-Methyl-2-pentanone	µg/L	< 100 U
Acetone	µg/L	< 100 U
Benzene	µg/L	< 50 U
Bromobenzene	µg/L	< 50 U
Bromochloromethane	µg/L	< 50 U
Bromodichloromethane	µg/L	< 50 U

Bromoform	µg/L	< 50 U
Bromomethane	µg/L	< 50 U
Carbon disulfide	µg/L	< 100 U
Carbon tetrachloride	µg/L	< 50 U
Chlorobenzene	µg/L	< 50 U
Chloroethane	µg/L	< 50 U
Chloroform	µg/L	< 50 U
Chloromethane	µg/L	< 50 U
cis-1,2-Dichloroethene	µg/L	2,400
cis-1,3-Dichloropropene	µg/L	< 50 U
Dibromochloromethane	µg/L	< 50 U
Dibromomethane	µg/L	< 50 U
Dichlorodifluoromethane	µg/L	< 50 U
Ethylbenzene	µg/L	< 50 U
Hexachlorobutadiene	µg/L	< 100 U
Isopropylbenzene	µg/L	< 50 U
m,p-Xylene	µg/L	< 100 U
Methylene chloride	µg/L	3,900
Naphthalene	µg/L	< 50 U
n-Butylbenzene	µg/L	< 50 U
n-Propylbenzene	µg/L	< 50 U
o-Xylene	µg/L	< 50 U
sec-Butylbenzene	µg/L	< 50 U
Styrene	µg/L	< 50 U
tert-Butylbenzene	µg/L	< 50 U
Tetrachloroethene	µg/L	62 J
Toluene	µg/L	< 50 U
trans-1,2-Dichloroethene	µg/L	< 50 U
trans-1,3-Dichloropropene	µg/L	< 50 U
Trichloroethene	µg/L	5,600
Trichlorofluoromethane	µg/L	< 50 U
Vinyl chloride	µg/L	< 50 U
Metals (6020A)		
Aluminum	mg/L	0.165
Antimony	mg/L	< 0.000500 U
Arsenic	mg/L	0.00138 J
Barium	mg/L	0.448
Beryllium	mg/L	< 0.000500 U
Cadmium	mg/L	< 0.000500 U
Calcium	mg/L	22.2
Chromium	mg/L	0.00112 J
Cobalt	mg/L	0.00904
Iron	mg/L	1.04
Lead	mg/L	< 0.00100 U
Magnesium	mg/L	20.1
Manganese	mg/L	0.428

Nickel	mg/L	0.00947
Potassium	mg/L	1.41
Selenium	mg/L	< 0.00250 U
Silver	mg/L	< 0.000500 U
Sodium	mg/L	140
Thallium	mg/L	< 0.000500 U
Vanadium	mg/L	0.00202 J
Zinc	mg/L	0.0260
Mercury	mg/L	< 0.000100 U
Anions (9056)		
Chloride	mg/L	240
Sulfate	mg/L	37.4
Semi-Volatile Organic Compounds (8270D SIM)		
1,4-Dioxane	µg/L	8.6

µg/L - micrograms per liter

mg/L - milligrams per liter

J -estimated value between the detection limit and limit of quantitation and/or due to quality control issues

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

**QUARTERLY EVALUATION REPORT
1ST QUARTER (JANUARY - MARCH) 2019
GROUNDWATER TREATMENT PLANT
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS**

JULY 2019

Prepared For:



**U.S. Army Corps of Engineers
Tulsa District**

**Contract No. W9128F-13-D-0012
Task Order No. W912BV17F0150
Bhate Project No. NWO1312.0150.016.0001.03**

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GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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Appendices

Appendix A ICT Layout and GWTP Process Flow Diagram
Appendix B Groundwater Elevation Contour Maps
Appendix C GWTP Water Sampling Laboratory Analytical Results (Provided on CD Only)
Appendix D Laboratory Analytical Results for LHAAP-16 (Provided on CD Only)
Appendix E Quality Control Summary Report
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GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

ACRONYMS AND ABBREVIATIONS

AMCV	Air Monitoring Comparison Value
amsl	Above mean sea level
bgs	Below ground surface
Bhate	Bhate Environmental Associates, Inc.
CD	Compact disc
COD	Chemical oxygen demand
DCE	Dichloroethene
ESD	Explanation of Significant Difference
ESL	Effects Screening Level
FBR	Fluidized bed reactor
ft	Feet or foot
gpd	Gallons per day
gph	Gallons per hour
gpm	Gallons per minute
GWTP	Groundwater Treatment Plant
HCl	Hydrochloric acid
HDPE	High density polyethylene
ICT	Interception-collection trench
IRA	Interim Remedial Action
J	Estimated concentration
lbs/hr	Pounds per hour
LHAAP	Longhorn Army Ammunition Plant
MCL	Maximum Contaminant Level
µg/L	Micrograms per liter
Mg(OH) ₂	Magnesium hydroxide
MSC	Medium Specific Concentration
mV	Millivolts
NA	Not applicable
NaOH	Sodium hydroxide
No.	Number

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ORP	Oxidation-reduction potential
PCL	Protective Concentration Level
PID	Photoionization detector
ppmv	Parts per million by volume
psi	Pounds per square inch
ROD	Record of Decision
TAC	Texas Administrative Code
TCE	Trichloroethene
TCEQ	Texas Commission on Environmental Quality
tpy	Tons per year
TRRP	Texas Risk Reduction Program
UEP	Unlined Evaporation Pond
USEPA	United States Environmental Protection Agency
VOC	Volatile Organic Compound

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EXECUTIVE SUMMARY

The operation of the Groundwater Treatment Plant (GWTP) is part of the Interim Remedial Action (IRA) at Burning Ground Number (No.) 3, also referred to as Longhorn Army Ammunition Plant (LHAAP)-18/24. A historical pilot test for nearby landfill LHAAP-16 resulted in the installation of eight extraction wells which also contribute groundwater to the GWTP. Groundwater extraction, treatment, and monitoring activities consist of:

- Continuous extraction of groundwater from multiple interception-collection trenches (ICTs) and extraction wells at both LHAAP-18/24 and LHAAP-16;
- Treatment of extracted groundwater for heavy metals, chlorinated compounds, and perchlorate using precipitation, air stripping, and biological methods, respectively;
- Evaluation of the hydraulic effectiveness of the extraction system by groundwater monitoring;
- Monitoring of treated groundwater to ensure compliance with the discharge limits; and
- Discharge of treated water to Harrison Bayou, or to a holding pond (INF Pond), or the treated water may be released as irrigation water on LHAAP-18/24.

The location of the extraction wells and ICTs are shown on **Figure A-1** in **Appendix A**. The process flow diagram of the GWTP is shown on **Figure A-2** in **Appendix A**.

Figure ES-1 depicts the monthly total volume of groundwater that was extracted from the ICTs and extraction wells at LHAAP-18/24 and LHAAP-16 between September 2012 and March 2019.

The GWTP was not operational during June, July, and August 2012. This was related to meltdown of the scrubber system, associated with the catalytic oxidizer, due to system overheating. Overheating occurred when the blower became inoperable after the bearing on the scrubber blower unit was shattered and damaged the blower. This occurred around 1:00 PM on May 21, 2012.

After developing an interim air monitoring plan and obtaining concurrence from the Texas Commission on Environmental Quality (TCEQ) and the United States Environmental Protection Agency (USEPA) to operate the GWTP without use of air abatement equipment, a pilot run of the GWTP was conducted on September 6, 2012. In that first pilot run, 85,170 gallons of water that had been stored in the influent equalization tank (TK-140) were treated. The treated water was re-circulated through the fluidized bed reactor (FBR) to revive the FBR after 3 months of dormancy. Treated groundwater and air samples were collected and analyzed respectively for perchlorate, metals, and Volatile Organic Compounds (VOCs); and VOCs only. On September 19, 2012, a second pilot run was performed at the GWTP and 107,264 gallons of water were treated. Based on the successful re-start of the GWTP, continuous groundwater extraction began again on September 24, 2012. While groundwater extraction occurs on a continuous basis, operation of the GWTP occurs intermittently due to the low volume of water available for treatment with respect to the design capacity of the GWTP. During the 3rd quarter of 2012, groundwater extraction occurred only from LHAAP-18/24. Groundwater extraction from LHAAP-16 was not

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performed due to equipment failure. However, extraction from LHAAP-16 began in October 2012 and the extraction volumes increased steadily throughout the 4th quarter of 2012, as pumping equipment was gradually repaired/replaced. The GWTP operated under normal conditions until September 2015.

On September 14, 2015, at 11:15 AM, the blower on the air stripper (BL-340) malfunctioned during routine operation. The wiring on the blower was repaired and the blower operated for less than 2 hours on September 17, 2015, when the blower malfunctioned again. It was determined that the blower needed to be replaced, and groundwater extraction and operation of the GWTP ceased beginning September 18, 2015, as the influent equalization tank (TK-140) became full. Beginning on October 2, 2015, it was determined that the GWTP could operate without the blower at a reduced extraction rate. The operation of the GWTP allowed extraction of groundwater from ICTs 12E, 13A, 13B, and 13C (13C was changed to ICT 13E on October 12, 2015), which were considered critical ICTs to prevent migration of contaminants to Harrison Bayou. Groundwater extraction was switched frequently between ICTs 12E, 13A, 13B, and 13E to ICTs 14B, 14C, and 14D beginning on December 14, 2015.

On December 12, 2016, flange bolts at TK-380 failed and allowed hydrochloric acid (HCl) to drain into the sump. The containment area was washed down and the sump contents were transferred into the equalization tank (TK-140). Because of the acid release, extraction of groundwater from the ICTs was halted, and the GWTP was put into recycle mode (effluent sent back as influent) until the acid was neutralized and perchlorate, metals, and VOCs were below discharge criteria on March 17, 2017.

On August 12, 2017, severe storms caused a power outage at LHAAP-18/24. When electrical service was restored, the main transformer failed due to a manufacturing defect. A portable emergency generator was brought on-site on August 21, 2017, to allow the FBR to operate in full recycle mode. After a replacement transformer was installed on September 12, 2017, extraction began from ICT-13B, 13C, 13D, 13E, 13F, 7, and EW01 and the FBR was put into normal operation. Beginning on September 21, 2017, groundwater was extracted from all of the ICTs.

On December 27, 2018, severe storms caused a power outage in Karnack, Texas including LHAAP. When electrical service was restored, the main transformer failed due to a manufacturing defect. A portable emergency generator was brought on-site on December 28, 2018, to allow the FBR to operate in full recycle mode. After a replacement transformer was connected to the well field on February 11, 2019, extraction began from LHAAP-18/24, and the GWTP was put into normal operation. No extraction from LHAAP-16 occurred during the 1st Quarter 2019 due to the main transformer being down.

As shown on **Figure ES-1**, the total extracted groundwater volume from LHAAP-18/24 during the 1st quarter of 2019 was less than normal due to the main transformer being down. No groundwater was extracted in January 2019. In February and March 2019, repairs to the ICT and extraction well pumps and motor increased flow at LHAAP-18/24 but no groundwater was extracted during this quarter from LHAAP-16. The extracted groundwater volume was measured on a monthly basis as the sum of the difference between the flow meter totalizer reading at each

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ICT between the beginning and end of each month. Extraction quantities in LHAAP-18/24 were 515,618 gallons in February 2019 and 1,063,734 gallons in March 2019.

No groundwater was extraction from LHAAP-16 in 1st Quarter 2019. Approximately 1,579,352 gallons of groundwater were extracted from LHAAP-18/24 during the 1st quarter of 2019 compared to approximately 1,672,185 gallons extracted during the 1st quarter 2018 (year prior).

No treated water was returned to ICTs 6 and 9 during the 1st quarter of 2019 because this practice was discontinued after system restart in September 2012.

The typical discharged flowrate from the GWTP was calculated as 11 gallons per minute (gpm) during the 1st quarter of 2019. Water discharge from the INF Pond varied from a flow rate of 18 gpm to 184 gpm. Approximately 588,518 gallons of groundwater was discharged from the GWTP to the Harrison Bayou, and 1,779,495 gallons was discharged from the INF Pond to the Harrison Bayou (see **Figure ES-2** below).

Grab perchlorate samples from the GWTP influent were collected monthly on February 28 and March 14, 2019, and the following concentrations were reported: 3,700 micrograms per liter ($\mu\text{g/L}$) and 8,300 $\mu\text{g/L}$, respectively. In addition, a quarterly influent sample was collected and analyzed for perchlorate with a result of 6,500 $\mu\text{g/L}$. The average perchlorate concentration using these three values from the GWTP influent during the quarter was 6,167 $\mu\text{g/L}$. With the exception of low detections in effluent samples collected on March 27, 2019, no perchlorate was detected in the effluent (TK-650) samples during the 1st quarter of 2019.

As shown in **Table ES-1**, all treated water was discharged directly from the GWTP to Harrison Bayou in 1st Quarter 2019. In addition, 1,779,495 gallons of treated water was discharged from the INF Pond to Harrison Bayou.

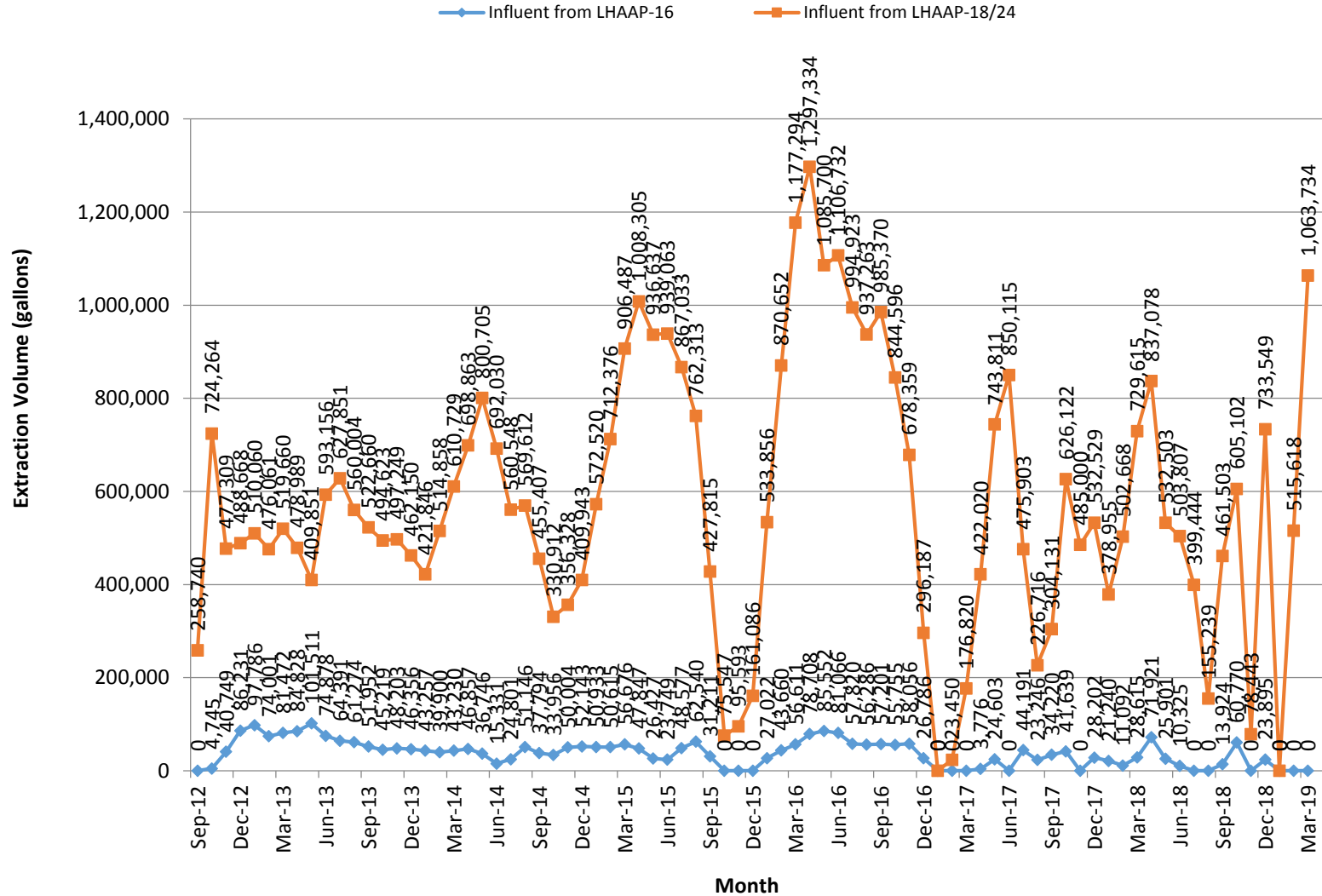
Groundwater was not extracted from LHAAP-18/24 in the 1st quarter of 2019 until February 11, 2019. After the GWTP resumed operation, the groundwater volume extracted for treatment at the LHAAP-18/24 increased from 515,618 gallons in February 2019 to 1,063,734 in March 2019. No groundwater was extracted from LHAAP-16 during the 1st quarter of 2019. The total water extracted for treatment by the GWTP for the 1st quarter of 2019 was approximately 1,579,352 gallons. The water quantities treated each month since June 2012 are shown on **Figure ES-2**. The total volume of water extracted from LHAAP-18/24 in the 1st quarter of 2019 (1,579,352 gallons) is higher than the volume of water discharged to the Harrison Bayou from the GWTP (588,518 gallons). The reason for the difference is the change in volume stored in the GWTP, the amount of water lost with the removed metals precipitation sludge, and the amount of evaporative water lost in the air stripper (which is included in the volume processed, but not in the volume discharged).

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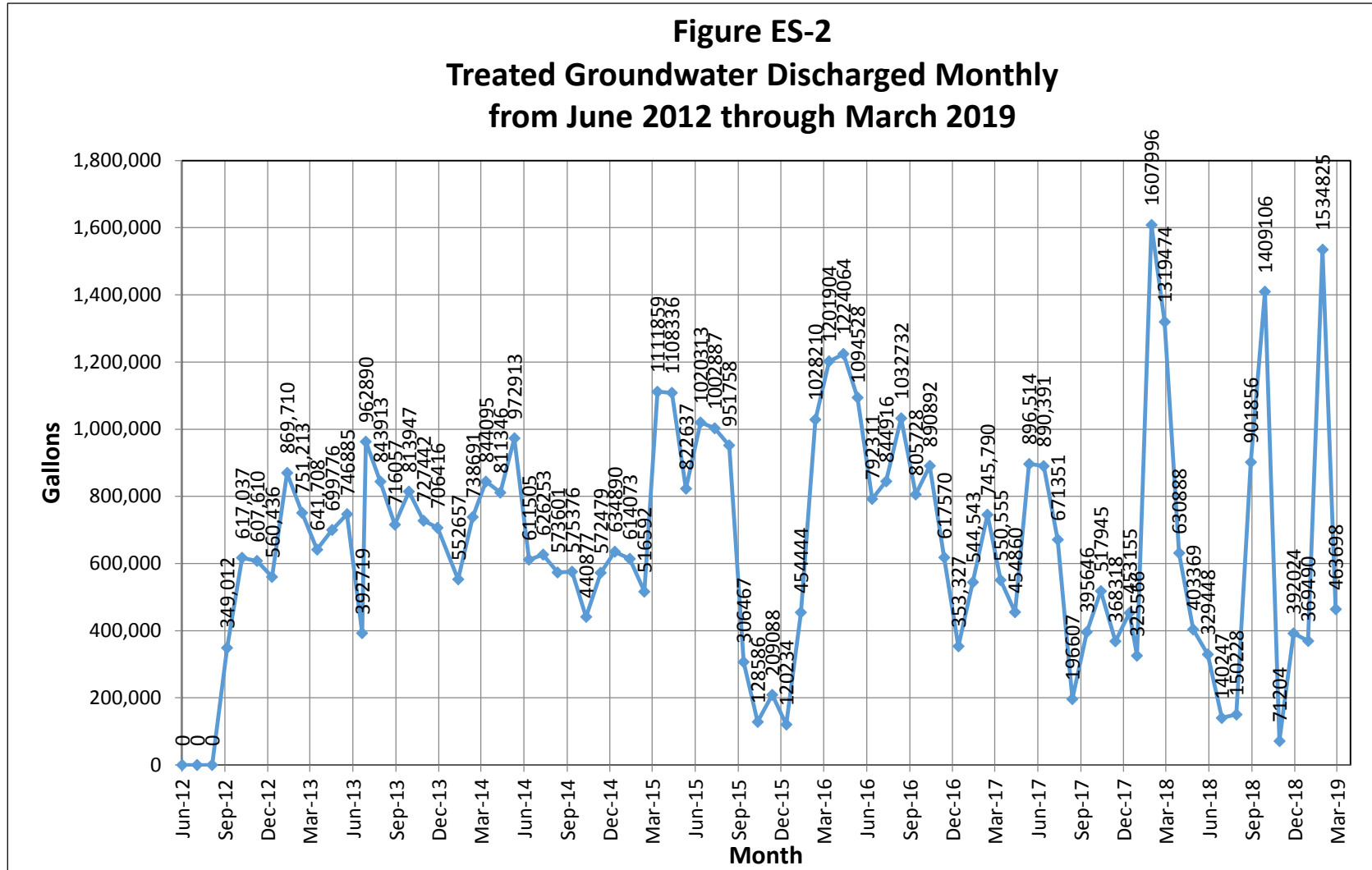
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Figure ES-1: Groundwater Recovery Between September 2012 & March 2019
LHAAP-18/24 & LHAAP-16



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Figure ES-2
Treated Groundwater Discharged Monthly
from June 2012 through March 2019



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Table ES-1: Discharge Information to Harrison Bayou During 1st Quarter 2019

DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
01/01/2019	Not Measured	Not Calculated	0	0	0	0	0	6.25
01/02/2019	Not Measured	Not Calculated	0	0	0	0	0	6.46
01/03/2019	Not Measured	Not Calculated	0	0	0	0	0	6.50
01/04/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/05/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/06/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/07/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/08/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/09/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/10/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/11/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/12/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/13/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/14/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/15/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/16/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/17/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/18/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/19/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/20/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/21/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/22/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/23/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89
01/24/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89
01/25/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89
01/26/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89

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DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
01/27/2019	Not Measured	Not Calculated	0	0	0	0	0	6.88
01/28/2019	Not Measured	Not Calculated	0	0	0	0	0	6.88
01/29/2019	16,975	305,560	0	176,319	0	0	176,319	6.45
01/30/2019	12,277	220,995	0	127,696	0	0	127,696	6.10
01/31/2019	8,224	148,042	0	65,475	0	0	65,475	5.92
02/01/2019	5,898	106,179	0	57,960	0	0	57,960	5.75
02/02/2019	Not Measured	Not Calculated	0	0	0	0	0	5.75
02/03/2019	Not Measured	Not Calculated	0	0	0	0	0	5.75
02/04/2019	Not Measured	Not Calculated	0	0	0	0	0	5.75
02/05/2019	Not Measured	Not Calculated	0	0	0	0	0	5.74
02/06/2019	Not Measured	Not Calculated	0	0	0	0	0	5.74
02/07/2019	Not Measured	Not Calculated	0	0	0	0	0	5.81
02/08/2019	Not Measured	Not Calculated	0	0	0	0	0	5.81
02/09/2019	Not Measured	Not Calculated	0	0	0	0	0	5.81
02/10/2019	Not Measured	Not Calculated	0	0	0	0	0	5.80
02/11/2019	11,579	3,383	893	0	0	893	893	5.83
02/12/2019	20,399	5,960	25,279	252,560	0	25,279	277,839	5.34
02/13/2019	13,411	3,918	15,361	221,340	0	15,361	236,701	4.78
02/14/2019	16,395	4,790	12,590	174,600	0	12,590	187,190	4.24
02/15/2019	15,454	4,515	9,109	174,240	0	9,109	183,349	3.69
02/16/2019	12,913	3,773	0	0	0	0	0	3.69
02/17/2019	10,373	3,031	21,479	0	0	21,479	21,479	3.69
02/18/2019	9,369	2,737	0	0	0	0	0	3.68
02/19/2019	8,366	2,444	17,960	0	0	17,960	17,960	3.68
02/20/2019	13,181	3,155	11,528	96,600	0	11,528	108,128	3.55
02/21/2019	20,096	4,810	11,444	28,025	0	11,444	39,469	3.50
02/22/2019	Flood Stage	Maximum	9,709	0	0	9,709	9,709	3.53
02/23/2019	Flood Stage	Maximum	0	0	0	0	0	3.60

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DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
02/24/2019	Flood Stage	Maximum	0	0	0	0	0	3.70
02/25/2019	Flood Stage	Maximum	38,835	0	0	38,835	38,835	3.70
02/26/2019	21,127	5,057	10,568	132,480	0	10,568	143,048	3.35
02/27/2019	19,533	4,675	11,891	97,680	0	11,891	109,571	3.15
02/28/2019	16,620	4,856	11,694	91,000	0	11,694	102,694	2.85
03/01/2019	16,125	4,711	11,607	83,520	0	11,607	95,127	2.47
03/02/2019	16,806	4,911	0	0	0	0	0	2.50
03/03/2019	Flood Stage	Maximum	0	0	0	0	0	2.50
03/04/2019	32,976	9,636	35,970	0	0	35,970	35,970	2.50
03/05/2019	23,207	6,781	10,521	0	0	10,521	10,521	2.50
03/06/2019	16,822	4,915	7,756	0	0	7,756	7,756	2.50
03/07/2019	10,652	3,564	4,826	0	0	4,826	4,826	2.50
03/08/2019	10,204	3,414	11,008	0	0	11,008	11,008	2.50
03/09/2019	18,255	5,464	0	0	0	0	0	2.50
03/10/2019	15,231	5,026	0	0	0	0	0	2.50
03/11/2019	14,601	4,885	34,669	0	0	34,669	34,669	2.49
03/12/2019	19,104	6,391	12,730	0	0	12,730	12,730	2.49
03/13/2019	16,809	5,624	12,255	0	0	12,255	12,255	2.58
03/14/2019	Flood Stage	Maximum	12,733	0	0	12,733	12,733	2.66
03/15/2019	Flood Stage	Maximum	10,626	0	0	10,626	10,626	2.66
03/16/2019	Flood Stage	Maximum	0	0	0	0	0	2.65
03/17/2019	20,598	5,723	0	0	0	0	0	2.65
03/18/2019	17,339	5,066	33,160	0	0	33,160	33,160	2.65
03/19/2019	7,625	2,228	14,714	0	0	14,714	14,714	2.64
03/20/2019	5,599	1,636	15,654	0	0	15,654	15,654	2.64
03/21/2019	6,950	2,031	8,703	0	0	8,703	8,703	2.64
03/22/2019	5,977	1,746	9,406	0	0	9,406	9,406	2.63
03/23/2019	5,326	1,544	0	0	0	0	0	2.63

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DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
03/24/2019	4,856	1,408	0	0	0	0	0	2.63
03/25/2019	4,431	1,295	18,556	0	0	18,556	18,556	2.66
03/26/2019	5,221	1,525	16,413	0	0	16,413	16,413	2.66
03/27/2019	5,592	1,634	16,818	0	0	16,818	16,818	2.66
03/28/2019	9,196	2,687	18,192	0	0	18,192	18,192	2.65
03/29/2019	7,977	2,331	15,769	0	0	15,769	15,769	2.65
03/30/2019	6,231	1,806	0	0	0	0	0	2.65
03/31/2019	5,426	1,573	48,092	0	0	48,092	48,092	2.65
Totals			588,518	1,779,495	0	588,518	2,368,013	

Notes: 1) Bayou flow is not measured on days when no release to the Bayou occurs from the GWTP or INF Pond.

2) When the bayou is in flood stage (over the banks), the flow is not measured and the maximum release is allowed.

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1 EVALUATION OF GROUNDWATER TREATMENT PLANT

The Groundwater Treatment Plant (GWTP) was constructed as part of the Interim Remedial Action (IRA) at Burning Ground Number (No.) 3, also referred to as Longhorn Army Ammunition Plant (LHAAP)-18/24, to treat groundwater extracted from interception-collection trenches (ICTs) and extraction wells. **Figure A-1** located in **Appendix A** presents the layout of the ICTs and extraction wells at LHAAP-18/24. The groundwater contamination at LHAAP-18/24 likely resulted from infiltration from an Unlined Evaporation Pond (UEP) that was used to store manufacturing wastewater, and from burning trenches and other industrial processes used to flash pyrotechnic, propellant, and explosive waste streams. The groundwater at LHAAP-18/24 is contaminated mainly with chlorinated ethenes and perchlorate, with lesser concentrations of 1,4-dioxane.

The GWTP also receives flow from eight extraction wells installed at LHAAP-16 as part of a historical treatability study. The extraction wells were installed in 1996 and 1997. The wells are located between the landfill at LHAAP-16 and Harrison Bayou. The groundwater at LHAAP-16 is also contaminated mainly with chlorinated ethenes and perchlorate.

1.1 Treatment Configuration

The process flow diagram for the GWTP is presented in **Appendix A, Figure A-2**. The GWTP was not operational between May 24, 2012, and September 6, 2012, due to malfunction of the scrubber unit associated with the catalytic oxidizer. Since September 6, 2012, the GWTP has operated without air abatement equipment. Although major repairs were conducted on the GWTP (e.g., replacement of level alarms, repair of the hydrochloric acid [HCl] tank, replacement of TK-650, replacement of malfunctioning valves and flow meters, replacement of metering pumps, repair or replacement of various system pumps, rust removal and repainting of various tanks, and replacement and repair of various extraction pumps, motors, and level switches), the GWTP treatment configuration has remained unchanged.

Malfunction of the blower on the air stripper (BL-340) on September 14, 2015, and on September 17, 2015, disrupted continuous extraction and routine operations of the GWTP, which lasted through January 7, 2016. Prior to this occurrence, the GWTP performed as designed and the GWTP was operated on an as needed basis (i.e., semi-continuous operational basis). During the 4th quarter of 2015, groundwater was extracted from a limited number of ICTs (ICTs 12E, 13A, 13B, 13C, and/or 13E, or ICTs 14B, 14C, and 14D). Operation of the GWTP occurred on a batch basis through the fluidized bed reactor (FBR). After replacement of the blower, attempts were made to restore continuous operations to the FBR but remained predominantly on a batch basis throughout January 2016.

In December 2016, a HCl spill caused plant operations to shut down until the issue could be properly addressed. The FBR performance was challenged by the increased chlorides in the neutralized wastewater, but performance gradually returned to normal in the 1st quarter of 2017. Groundwater extraction was gradually increased to full rates during the 2nd quarter of 2017.

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On August 12, 2017, severe storms caused a power outage at LHAAP-18/24. When electrical service was restored, the main transformer failed due to a manufacturing defect. A portable emergency generator was brought on-site on August 21, 2017, to allow the FBR to operate in full recycle mode. After a replacement transformer was installed on September 12, 2017, extraction began from ICT-13B, 13C, 13D, 13E, 13F, 7, and EW01 and the FBR was put into normal operation. Beginning on September 21, 2017, groundwater was extracted from all of the ICTs.

On December 27, 2018, severe storms caused a power out in Karnack, Texas. When electrical service was restored, the main was determined to have failed. A portable emergency generator was brought on-site on December 28, 2018, to allow the FBR to operate in full recycle mode. On February 5, 2019, the smaller generator mobilized in December 2018 was replaced with a larger generator capable of powering the LHAAP-18/24 well field as well as the entire GWTP. On February 8, 2019, the transformer at the GWTP was tested to ensure that it could handle backfeeding necessary to power the LHAAP-18/24 well field due to the necessary step-down in power from the generator. Following additional system modifications based upon the testing, the well field at LHAAP-18/24 had power restored on February 11, 2019, using the generator and transformer at the GWTP.

Flow rates for the treatment processes for metals and Volatile Organic Compounds (VOCs) ranged between 170 and 210 gallons per minute (gpm) with an average of approximately 185 gpm for the operating hours (i.e., this flow rate does not represent continuous flows). The GWTP operated for 64.75 hours during the quarter, with no operation of the GWTP in January and most of February 2019. The treatment configuration of the plant at these rates (with minor variations) is as follows:

GWTP Metals Precipitation Operating Parameters

Pretreatment Settings	Tank 200-A Mg(OH) ₂ Mixing	Tank 200-B NaOH Mixing	Tank 200-C Polymer Mixing	Tank 300 feed line to Air Stripper
pH Adjustment	9.0	10.5	NA	5.0 to meet ≤ 8.0 release from stripper
Feed Pump Settings	Speed 100% Stroke 100% 10 gph Mg(OH) ₂	Speed 100% Stroke 100% 9.0 gph NaOH	Speed 90% Stroke 100% 40 gph water	Speed 80% Stroke 80% 10 gph HCl
Notes: gph - gallons per hour, NaOH - sodium hydroxide, Mg(OH) ₂ - magnesium hydroxide, NA - not applicable				

GWTP Air Compressors Operating Parameters

Air Compressors	K-700A	K-700B	K-701
Air Pressure Settings	88 psi	88 psi	105 psi
Note: psi - pounds per square inch			

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GWTP Stripper Operating Parameters

pH Setting	7.4
Inlet Pressure Gauge	Not operational
Stripper Pressure Gauge	Not operational
Air Flow Rate	Not operational

GWTP Fluidized Bed Reactor Operating Parameters

Carbon Bed Height	12 feet & 8 to 11 inches
Recycle Flow Rate	200 gpm
pH	7.1 to 7.4
Recycle oxidation-reduction potential (ORP)	<-430 mV
Note: mV - millivolts	

1.2 Work Performed at the GWTP

Work performed at the GWTP during the 1st quarter of 2019 is described in the following subsections.

1.2.1 Major Maintenance

The major maintenance items that were completed at the GWTP during this quarterly reporting period are:

- January 15 and 16, 2019: Caddo Well Service was on site to install new bladder tank, pump, and piping in potable water well #1. This restored potable water pressure to the GWTP.
- February 5, 2019: Ark-La-Tex Electric was on site to connect wiring to 800 kilowatt generator.
- February 8, 2019: Transformer Testing Company was on site to check out pad mounted transformer at GWTP.
- February 11, 2019: Ark-La-Tex Electric was on site to connect wiring to allow back feed through the pad mounted transformer to the ICT wells.
- March 25, 2019: Bloc Design was on site to troubleshoot electrical problem with well EW01.
- March 25, 2019: The 800 kilowatt generator was repaired.
- March 28, 2019: Bloc Design was onsite to repair wiring problem with EW01 and repair MX-250 mixer.

1.2.2 Routine Maintenance

The following routine maintenance items were completed at the GWTP during this quarterly reporting period:

- Performed housekeeping in GWTP office

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- Performed housekeeping in Army trailer
- Performed housekeeping around GWTP and containment area
- Performed housekeeping in GWTP Shop
- Rebuilt grundfos pumps to be used in ICT wells
- Installed new weather station at GWTP
- Repaired 3-inch piping and 3-inch flange on discharge side of P-320
- Mowed grass around GWTP, Army trailer, decontamination pad, and storage connexes
- Replaced ¼-inch tubing on suction side of P-104
- Collected quarterly surface water samples

1.2.2.1 Safety

The GWTP Operators, Mr. Scott Beesinger and Mr. Kennie Moore, did not have any safety incidents. No safety training was completed.

1.2.2.2 Lubrication

No lubrication maintenance was conducted during the reporting period.

1.2.2.3 Air Compressors

During the 1st quarter of 2019, air compressor K-700A had routine maintenance completed that included replacement of the drive belt as well as the addition of oil.

1.2.2.4 Belt Press and Waste Disposal

No belt press or waste disposal was conducted during the reporting period.

1.2.2.5 Sand Filter

No maintenance or repairs were conducted on the sand filter during the reporting period.

1.2.2.6 Well Field at LHAAP-18/24

- Collected monthly flow meter readings
- Collected monthly water levels
- Freeze protected ICT wells
- Installed new flow meters on ICTs 2 and 13A
- Replaced a pump and motor in ICT 12D
- Replaced pump in ICT 13C
- Replaced pump in ICT 2
- Replaced pump and motor in ICT 14C
- Replaced pump in ICT 14D
- Replaced a leaking 1-inch union on ICT 2

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- Replaced pump, seal plate, and level probes in ICT 14E
- Replaced pump and motor in ICT 14A
- Repaired leaking 1-inch pipe on ICT 14A
- Replaced pump in EW01
- Replaced probe wire and level probes in 18WW17
- Lowered high level probes in ICTs 14A and 13C
- Cleaned level probes on ICT 13E and 18WW17

1.2.2.7 Miscellaneous Activities

- None

1.2.3 Routine Maintenance at LHAAP-16

- Checked site daily
- Collected monthly water levels
- Collected annual extraction well samples

1.2.4 Routine Maintenance (Potable Water Wells)

- Insulated the new piping at the potable water pump #1.
- Flushed potable water lines.

1.3 Filter Cake Operations and Management

No filter cake operations took place during this reporting period.

1.4 Fluidized Bed Reactor Operations

The FBR was operational during all of 1st Quarter 2019 though in recycle for the first half of the quarter. On February 10, 2019, the system sent a call out that the FBR feed was shut down. Upon arrival, it was discovered that P-641 or P-642 did not come on which allowed TK-650 to fill to capacity. The GWTP operator completed troubleshooting and then repaired the issue by resetting the electrical breaker. The FBR feed was then restarted.

The operating parameters for the GWTP FBR are presented in **Table 1**. With the exception of ORP on February 13, 2019, none of the operating parameters were outside of the optimal ranges in the 1st quarter of 2019. The ORP ranged between -403 mV and -516 mV, and the pH ranged between 7.1 and 7.4 standard units.

Table 1. Enhanced Fluidized Bed Reactor Operating Parameters – 1st Quarter 2019

Date	pH (7.1-7.4)	ORP (<-430mV)	Temperature (Degrees Fahrenheit)
1/1/2019	7.4	-492	55
1/2/2019	7.4	-493	55

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Date	pH (7.1-7.4)	ORP (<-430mV)	Temperature (Degrees Fahrenheit)
1/3/2019	7.4	-494	53
1/4/2019	7.4	-495	53
1/5/2019	7.4	-496	54
1/6/2019	7.4	-496	55
1/7/2019	7.3	-494	57
1/8/2019	7.2	-480	58
1/9/2019	7.2	-489	57
1/10/2019	7.3	-502	56
1/11/2019	7.3	-510	56
1/12/2019	7.2	-504	57
1/13/2019	7.2	-499	55
1/14/2019	7.2	-496	54
1/15/2019	7.2	-495	54
1/16/2019	7.2	-495	55
1/17/2019	7.2	-495	56
1/18/2019	7.2	-495	56
1/19/2019	7.3	-493	55
1/20/2019	7.3	-493	55
1/21/2019	7.3	-493	53
1/22/2019	7.2	-495	55
1/23/2019	7.3	-493	52
1/24/2019	7.3	-494	52
1/25/2019	7.2	-494	52
1/26/2019	7.2	-493	53
1/27/2019	7.2	-493	53
1/28/2019	7.2	-492	54
1/29/2019	7.2	-485	53
1/30/2019	7.2	-484	51
1/31/2019	7.2	-488	52
2/1/2019	7.2	-489	54
2/2/2019	7.2	-491	54
2/3/2019	7.2	-493	56
2/4/2019	7.2	-496	59
2/5/2019	7.2	-499	61
2/6/2019	7.1	-499	63
2/7/2019	7.1	-500	62
2/8/2019	7.2	-497	59
2/9/2019	7.3	-495	56
2/10/2019	7.3	-494	55

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Date	pH (7.1-7.4)	ORP (<-430mV)	Temperature (Degrees Fahrenheit)
2/11/2019	7.2	-498	58
2/12/2019	7.2	-499	58
2/13/2019	7.2	-403	57
2/14/2019	7.2	-482	58
2/15/2019	7.2	-485	58
2/16/2019	7.1	-486	57
2/17/2019	7.1	-487	56
2/18/2019	7.2	-486	56
2/19/2019	7.2	-486	53
2/20/2019	7.2	-445	54
2/21/2019	7.2	-470	52
2/22/2019	7.2	-475	52
2/23/2019	7.1	-475	55
2/24/2019	7.1	-474	53
2/25/2019	7.1	-471	55
2/26/2019	7.1	-468	55
2/27/2019	7.1	-466	57
2/28/2019	7.1	-470	55
3/1/2019	7.2	-476	54
3/2/2019	7.2	-482	55
3/3/2019	7.2	-483	53
3/4/2019	7.3	-484	52
3/5/2019	7.4	-473	51
3/6/2019	7.3	-474	51
3/7/2019	7.3	-445	52
3/8/2019	7.3	-454	56
3/9/2019	7.3	-483	58
3/10/2019	7.3	-482	58
3/11/2019	7.2	-494	59
3/12/2019	7.2	-500	60
3/13/2019	7.2	-489	61
3/14/2019	7.2	-493	63
3/15/2019	7.2	-493	62
3/16/2019	7.2	-494	62
3/17/2019	7.2	-495	61
3/18/2019	7.2	-500	62
3/19/2019	7.3	-497	63
3/20/2019	7.3	-500	63

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Date	pH (7.1-7.4)	ORP (<-430mV)	Temperature (Degrees Fahrenheit)
3/21/2019	7.3	-506	63
3/22/2019	7.3	-520	63
3/23/2019	7.3	-514	63
3/24/2019	7.3	-510	63
3/25/2019	7.4	-467	67
3/26/2019	7.4	-499	65
3/27/2019	7.4	-510	66
3/28/2019	7.4	-510	65
3/29/2019	7.4	-504	65
3/30/2019	7.3	-513	65
3/31/2019	7.3	-516	63

Note: Shaded value is outside of optimal operating range.

1.5 Process Chemical Usage at GWTP

Approximate chemical consumption and the quantity delivered during the 1st quarter of 2019 are shown in **Table 2**.

Table 2. Process Chemicals Delivered and Used

Chemical	Usage 1 st Quarter 2019	Quantity Delivered 1 st Quarter 2019
Hydrochloric acid	245 gallons	0
Sodium hydroxide (35%)	400 gallons	44,900 pounds (3,750 gallons)
Acetic acid (50%)	8 drums = 440 gallons	1,100 gallons
Phosphoric acid (75%)	30.1 liters	0
Magnesium hydroxide	105 gallons	0
Urea	223.3 pounds	500 pounds
Polymer (magnafloc 110-L)	4.1 liters	0

2 EVALUATION OF LHAAP-18/24 ICT EFFECTIVENESS

The ICT system at Burning Ground No. 3 is composed of 14 sections ranging in length from 100 feet (ft) to 1,300 ft. A total of approximately 5,000 linear ft of trench was installed within and around three sides of Burning Ground No. 3. The trench sections extend approximately 22 ft to 45 ft below ground surface (bgs). Most, but not all of the trenches are as deep as the confining clay layer of the shallow groundwater zone. High density polyethylene (HDPE) liners were installed in ICTs 12 and 13, located on the western and northern boundaries of LHAAP-18/24, respectively. The locations of the liners are shown on **Figure A-1** in **Appendix A**. **Table A-1** in **Appendix A** presents the depths of the ICTs.

2.1 Groundwater Elevation

Water levels from 95 monitoring wells and 11 piezometers (piezometer 12 was damaged and plugged and abandoned in May 2013) are measured monthly to generate potentiometric surface maps that assist in monitoring the effectiveness of the groundwater extraction system on plume containment. The groundwater contours are generated using the water levels from the shallow zone and Wilcox Formation wells. The water level data are presented in **Table 3**. No reinjection of treated groundwater or reapplication to LHAAP-18/24 grounds via the existing irrigation system occurred during the 1st quarter of 2019. Potentiometric surface maps are presented in **Appendix B** and groundwater elevations from the 1st quarter of 2019 are discussed in Section 2.2. No groundwater sampling was completed at LHAAP-18/24 in the 1st quarter of 2019.

2.2 Performance of Plume Capture

The intent of the ICTs is to control groundwater gradients, prevent off-site migration of contaminated groundwater, extract the most highly contaminated groundwater, and reduce the mass of contaminants in groundwater. Liners were installed in the ICTs on the northern (ICT 13) and western (ICT 12) site boundaries to limit migration of contaminated water from the site towards Harrison Bayou. At the same time, the liners reduce or prevent removal of contaminated groundwater that is outside the containment zone, between the site and Harrison Bayou. The ICTs are installed within the shallow subsurface at the site and capture primarily shallow groundwater (e.g., < 40 ft bgs).

In 2007 and 2008, in consultation with the Texas Commission on Environmental Quality (TCEQ) and the United States Environmental Protection Agency (USEPA), the Army ceased operations of ICTs 1, 3, 5, 10, and 12A for groundwater extraction (note that extraction from ICT 12A was resumed after pump replacement in December 2012). Two other ICTs (ICT 6 and ICT 9) were changed from extraction ICTs to re-injection ICTs. Groundwater extraction from well EW-1 located in the northeast central portion of the site began in October 2008 and well 18WW17 located to the northeast of the ICT containment area began in January 2008. **Table B-1** in **Appendix B** presents a summary of extraction equipment replacement since 2011, as dictated by poor extraction performance (malfunctioning pumps, poor pump positioning with respect to groundwater, non-operational level probes, scale build up, etc.). Further discussion of extraction performance of various ICTs and extraction wells is presented in Section 2.3.

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Potentiometric surface maps of the shallow zone groundwater in the vicinity of LHAAP-18/24, based on groundwater elevations measured on January 31, February 28, and March 26-27, 2019, are shown on **Figures B-1, B-2, and B-3 in Appendix B**, respectively. The potentiometric surface maps of the shallow zone were contoured using the Kriging geostatistical interpolation method included in the Golden Software Surfer® data analysis software.

The HDPE liners in the ICTs, where present, were interpreted as groundwater flow barriers. The potentiometric surface maps for January through March 2019 continue to reflect high groundwater elevations in the northern/northwestern portion of the site with groundwater flow occurring radially from groundwater highs at monitoring well AWD-2 (174.19 ft above mean sea level [amsl] in January 2019, 174.53 ft amsl in February 2019, and 174.66 ft amsl in March 2019) inside the ICT containment area.

The elevated potentiometric surface contours within the ICTs compared to the lower potentiometric surface contours on the outside of the ICTs is likely due to a no flow boundary condition caused by the ICT liners and groundwater extraction along the ICTs. From the groundwater high at monitoring well AWD-2, groundwater flows radially towards the surrounding ICTs which include ICT 13 to the north and northwest, and ICT 12 to the west and southwest.

Groundwater extraction rates from the ICTs were 515,618 gallons in February 2019 and 1,063,734 gallons in March 2019. Rainfall amounts recorded at the GWTP were 5.4 inches in January 2019, 3.58 inches in February 2019, and 2.75 inches in March 2019. This amount of rainfall resulted in almost 88,000 gallons of additional water treated and discharged but not metered with the influent totals.

During the reporting period, approximately 2.4-million gallons of treated groundwater was discharged to Harrison Bayou from either the GWTP or the INF Pond. No treated groundwater from the GWTP was returned to LHAAP-18/24 via the sprinkler system. Overall groundwater levels increased throughout the 1st quarter of 2019 with an average shallow zone groundwater elevation rise of 0.63 ft.

Groundwater levels in Wilcox Formation wells (generally > 40 to 50 ft bgs) were measured during the 1st quarter of 2019 groundwater gauging events. Wilcox Formation wells correspond generally to those wells previously identified as “Intermediate” and “Deep” wells. “Intermediate” wells are designated as Upper Wilcox Formation wells and “Deep” wells are designated as Lower Wilcox Formation wells. Generally, groundwater in the Upper and Lower Wilcox Formation wells are in hydraulic communication and so can be treated as a single hydrogeologic unit. Therefore, the groundwater elevations in Upper Wilcox wells were used to construct the potentiometric surface maps for the Wilcox Formation. **Figures B-4, B-5, and B-6 in Appendix B** show the locations of the Wilcox Formation monitoring wells and the potentiometric surface of the Wilcox Formation, based on static water levels measured during the January, February, and March 2019 gauging events, respectively. Groundwater in the Wilcox Formation generally flows in a northerly direction, towards Caddo Lake and there is a downward

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vertical gradient between the overlying shallow zone and the Wilcox Formation. However, a groundwater high in the Wilcox Formation occurs in the area of MW-14.

2.3 Quantity of Water Extracted from LHAAP-18/24

The average daily extraction rates from the ICTs were 18,415 gallons per day (gpd) in February 2019 and approximately 34,314 gpd in March 2019. In February 2019, the plant operated about half of the month whereas the system operated fully in March 2019.

The volume of groundwater removed from LHAAP-18/24 during the 1st Quarter 2019 measured approximately 1,579,352 gallons, based on total flow measured from the extraction wells and ICT wells. However, the influent totalizer (FIT-140) readings, as presented on **Figure 2-1**, only indicate 475,429 gallons were extracted. No groundwater was extracted from LHAAP-16 during the 1st quarter of 2019 due to the main transformer not being operational. **Figure 2-1** shows the historical trends of extracted volumes by quarter based upon the influent totalizer (FIT-140). It is noted that there is a significant difference between the totalizer readings at FIT-140 and the total obtained from the individual units observed this quarter, which will be further evaluated in the 2nd Quarter 2019.

In contrast to the approximate total extracted volume based on total flow measured at the GWTP, the total estimated volume discharged to Harrison Bayou following treatment by the GWTP (FIT-686) was 208,340 in February 2019 and 380,178 gallons in March 2019 for a total of 588,518 gallons discharged in 1st Quarter 2019 (**Table 4**). However, this volume does not account for water present in the decant tank at the end of March or the influent that was not treated after the GWTP last ran on March 28, 2019. The difference between the influent volume determined from the individual meters on the ICTs and extraction wells and effluent volume determined from FIT 686 is approximately 81%. However, considering the over 200,000 gallons of water within the treatment plant as of March 31, 2019, this percent difference is closer to 66% variation, which is contributable to variations in the flow meter recordings. However, this variation is larger than typically noted and should be further evaluate to determine if the issue is the flow meters or if there is a potential leak.

As indicated by **Table 5**, 23 of 27 ICTs and wells produced water during the 1st quarter of 2019. The table illustrates the power loss to LHAAP-18/24 as well as increase in flow in the ICTs and wells following repairs made throughout March 2019, which was completed on March 18, 2019.

2.4 Groundwater Treatment Plant Sampling and Analysis

As part of the GWTP operations, samples from various water streams are required to be collected and analyzed for the parameters cited in the Interim Record of Decision (ROD) and the TCEQ letter dated January 8, 2002 (see Administrative Record Volume 1 of 4 in 2002, Document A). Besides the ROD sampling requirement, additional sample analyses are typically performed on the influent and effluent samples to monitor the effectiveness of the perchlorate treatment (FBR and/or ion exchange vessels) process. Sections 2.4.1 through 2.4.4 present the results of analyses conducted during the 1st quarter of 2019. The complete laboratory results are provided on a compact disc (CD) (**Appendix C**).

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2.4.1 Perchlorate Sampling

Table 6 presents the weekly effluent perchlorate results for the 1st quarter of 2019. In January 2019, no effluent samples were collected due to the main transformer being non-operational. The generator placed at the GWTP was able to power the lights in the building and the FBR but not power the wells and ICTs at LHAAP-18/24. On February 5, 2019, a larger generator was brought onsite and the pumps for the ICTs and wells were backfed to the transformer located at the GWTP which then was powered by the larger generator. Power was officially resorted to LHAAP-18/24 on February 11, 2019. Therefore, the first effluent samples collected as part of the 1st Quarter 2019 for perchlorate occurred on February 21, 2019. The biweekly effluent results were non-detect for perchlorate in the 1st Quarter 2019. However, perchlorate was detected in the quarterly effluent sample on March 27, 2019, but the detection was just above the level of detection with a value of 2.3 micrograms per liter ($\mu\text{g/L}$). Based upon continued discharge to the Harrison Bayou, the ion exchange vessels were bypassed in the 1st quarter of 2019, per the discharge protocol. A total of three grab samples from the influent to the GWTP (TK-140) were collected, which included the quarterly sample. The perchlorate concentrations in these samples ranged from 3,700 $\mu\text{g/L}$ to 8,300 $\mu\text{g/L}$.

2.4.2 VOC Sampling

Tables 7 and **8** present the effluent VOC results for February and March 2019. Due to the power loss that occurred at the end of December 2018, no effluent was discharged in January 2019 such that no samples were collected. Sampling of the effluent for VOCs was conducted on a biweekly basis when GWTP was operational. The results, where applicable, were below the discharge limits. The tables also provide monthly influent concentrations for VOCs and perchlorate.

2.4.3 Monthly Metals Sampling

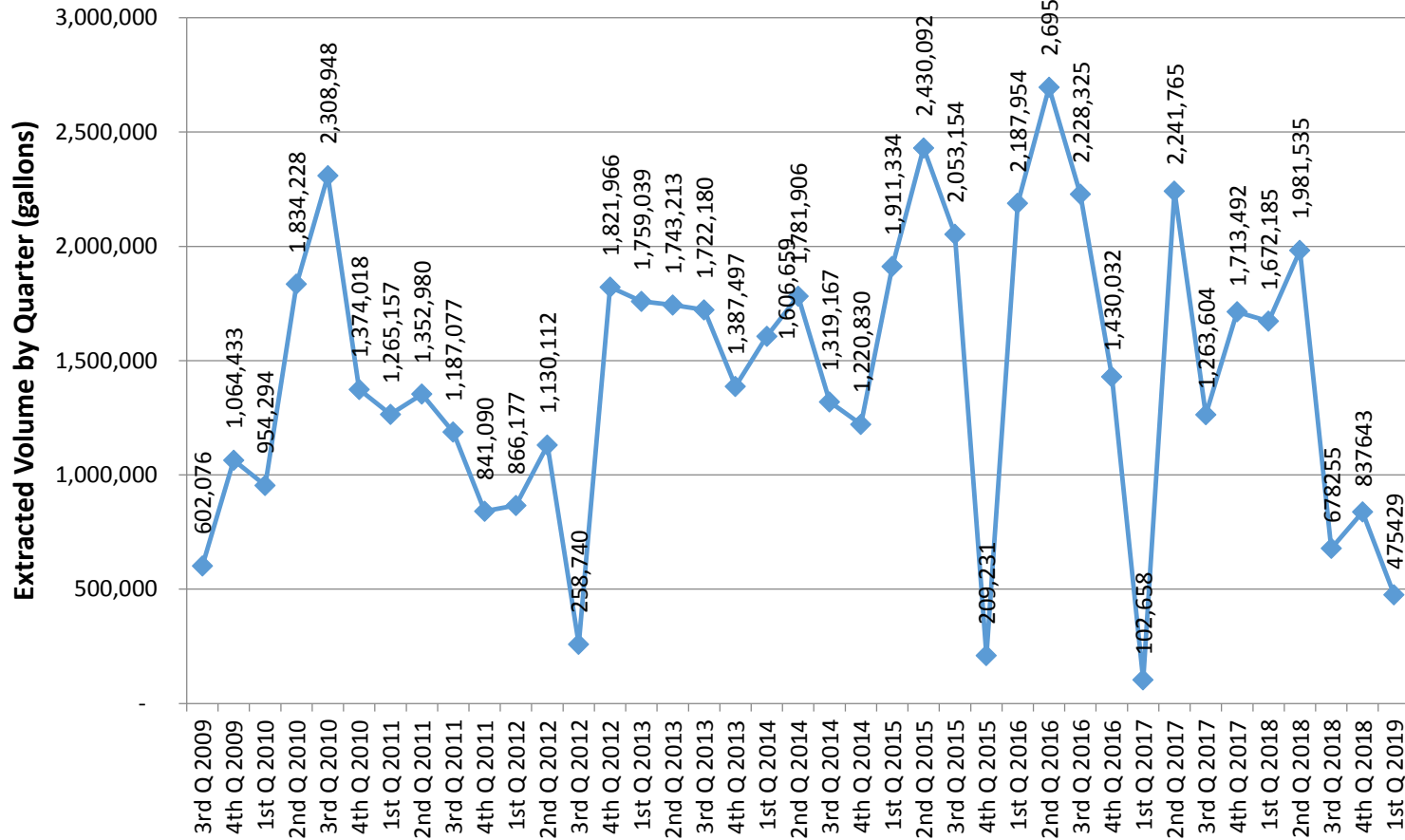
As per the Final Installation-Wide Work Plan (Bhate Environmental Associates, Inc. [Bhate], May 2018), the monthly metals sampling is reported in **Tables 7** and **8**. None of the metals exceeded the effluent discharge limits.

2.4.4 Quarterly Sampling

Sampling of the effluent for VOCs, anions, chemical oxygen demand (COD), oil and grease, perchlorate, and metals was conducted during this quarter and the results were below the discharge limits. **Table 9** presents the analytical results for the 1st quarter of 2019.

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**Figure 2-1
 Quarterly Extraction Rate**



Extraction rate beginning at 3rd Q 2013 is based in FIT-140 (influent totalizer)

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Table 3: Groundwater Elevations at LHAAP-18/24 Piezometers, Monitoring Wells, and Surface Water

Location Identification	Type	Reference Elevation (feet amsl)	Depth to Water (feet) 1/31/19	Groundwater Elevation (feet amsl) 1/31/19	Depth to Water (feet) 2/28/19	Groundwater Elevation (feet amsl) 2/28/19	Depth to Water (feet) 3/26/19-3/27/19	Groundwater Elevation (feet amsl) 3/26/16-3/27/19
BGPZ-1	Piezometer	184.99	6.12	178.87	5.91	179.08	5.57	179.42
BGPZ-2	Piezometer	184.39	13.57	170.82	13.45	170.94	13.02	171.37
BGPZ-3	Piezometer	180.35	7.96	172.39	7.79	172.56	6.91	173.44
BGPZ-4	Piezometer	177.77	7.89	169.88	7.70	170.07	6.84	170.93
BGPZ-5	Piezometer	180.76	12.05	168.71	11.89	168.87	10.95	169.81
BGPZ-6	Piezometer	197.82	25.77	172.05	25.85	171.97	27.05	170.77
BGPZ-7	Piezometer	195.96	28.15	167.81	28.02	167.94	26.86	169.10
BGPZ-8	Piezometer	197.08	29.93	167.15	29.80	167.28	28.93	168.15
BGPZ-9	Piezometer	196.45	28.69	167.76	28.55	167.90	27.44	169.01
BGPZ-10	Piezometer	197.00	27.95	169.05	27.80	169.20	27.53	169.47
BGPZ-11	Piezometer	196.99	27.83	169.16	27.91	169.08	27.64	169.35
BGPZ-12	Piezometer	188.17	NA	Plugged	NA	Plugged	NA	Plugged
AWD-1	Monitoring Well	182.27	9.16	173.11	9.25	173.02	9.13	173.14
AWD-2	Monitoring Well	186.78	12.59	174.19	12.25	174.53	12.12	174.66
AWD-3	Monitoring Well	200.13	27.90	172.23	27.78	172.35	27.59	172.54
AWD-4	Monitoring Well	193.89	24.29	169.60	24.35	169.54	24.11	169.78
MW-1	Monitoring Well	199.22	27.08	172.14	26.95	172.27	26.72	172.50
MW-2	Monitoring Well	196.73	25.49	171.24	26.36	170.37	26.12	170.61
MW-3	Monitoring Well	196.54	26.51	170.03	26.40	170.14	26.05	170.49
MW-4	Monitoring Well	197.27	26.73	170.54	26.65	170.62	26.33	170.94
MW-5	Monitoring Well	194.97	25.22	169.75	25.10	169.87	24.86	170.11
MW-6	Monitoring Well	192.18	23.45	168.73	23.31	168.87	23.09	169.09
MW-7	Monitoring Well	188.47	18.62	169.85	18.52	169.95	18.15	170.32
MW-8	Monitoring Well	187.13	17.43	169.70	17.29	169.84	16.88	170.25
MW-9	Monitoring Well	184.73	14.05	170.68	14.15	170.58	13.87	170.86
MW-10	Monitoring Well	178.12	8.10	170.02	7.97	170.15	7.31	170.81
MW-11	Monitoring Well	184.65	13.67	170.98	13.55	171.10	13.17	171.48

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Location Identification	Type	Reference Elevation (feet amsl)	Depth to Water (feet) 1/31/19	Groundwater Elevation (feet amsl) 1/31/19	Depth to Water (feet) 2/28/19	Groundwater Elevation (feet amsl) 2/28/19	Depth to Water (feet) 3/26/19-3/27/19	Groundwater Elevation (feet amsl) 3/26/16-3/27/19
MW-12	Monitoring Well	178.33	8.03	170.30	7.79	170.54	7.47	170.86
MW-13	Monitoring Well	176.72	7.67	169.05	7.57	169.15	6.92	169.80
MW-14	Monitoring Well	186.19	13.08	173.11	13.00	173.19	12.88	173.31
MW-16	Monitoring Well	178.59	7.91	170.68	7.71	170.88	7.45	171.14
MW-17	Monitoring Well	179.03	9.21	169.82	9.10	169.93	8.85	170.18
MW-18	Monitoring Well	178.58	8.71	169.87	8.60	169.98	8.38	170.20
MW-19	Monitoring Well	178.60	8.89	169.71	8.73	169.87	8.40	170.20
MW-20	Monitoring Well	186.64	9.59	177.05	9.47	177.17	9.11	177.53
MW-21	Monitoring Well	198.70	29.64	169.06	29.77	168.93	29.55	169.15
MW-22	Monitoring Well	197.51	27.65	169.86	27.53	169.98	27.28	170.23
MW-23	Monitoring Well	198.79	27.27	171.52	27.15	171.64	26.92	171.87
101	Monitoring Well	197.53	6.77	190.76	6.59	190.94	6.39	191.14
102	Monitoring Well	193.94	21.27	172.67	18.84	175.10	19.32	174.62
109	Monitoring Well	197.02	27.44	169.58	27.30	169.72	27.02	170.00
120	Monitoring Well	184.19	12.05	172.14	11.91	172.28	11.75	172.44
123	Monitoring Well	186.21	12.29	173.92	12.05	174.16	11.82	174.39
125	Monitoring Well	196.28	25.39	170.89	25.20	171.08	24.95	171.33
126	Monitoring Well	199.37	29.69	169.68	29.54	169.83	29.10	170.27
129	Monitoring Well	197.24	26.43	170.81	26.22	171.02	25.85	171.39
130	Monitoring Well	177.73	6.55	171.18	6.27	171.46	5.89	171.84
C-01	Monitoring Well	193.89	24.62	169.27	24.49	169.40	23.36	170.53
C-02	Monitoring Well	175.95	6.19	169.76	5.94	170.01	5.59	170.36
C-03	Monitoring Well	196.34	26.73	169.61	26.59	169.75	26.00	170.34
C-04	Monitoring Well	194.64	25.48	169.16	25.29	169.35	24.42	170.22
C-04A	Monitoring Well	194.61	25.17	169.44	25.01	169.60	24.19	170.42
C-05	Monitoring Well	180.74	12.47	168.27	11.97	168.77	11.53	169.21
C-06	Monitoring Well	192.22	25.62	166.60	25.50	166.72	23.33	168.89
C-07	Monitoring Well	196.80	28.57	168.23	28.67	168.13	26.71	170.09
C-08	Monitoring Well	193.10	24.11	168.99	24.00	169.10	23.90	169.20
C-09	Monitoring Well	202.35	33.03	169.32	32.90	169.45	32.65	169.70

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Location Identification	Type	Reference Elevation (feet amsl)	Depth to Water (feet) 1/31/19	Groundwater Elevation (feet amsl) 1/31/19	Depth to Water (feet) 2/28/19	Groundwater Elevation (feet amsl) 2/28/19	Depth to Water (feet) 3/26/19-3/27/19	Groundwater Elevation (feet amsl) 3/26/16-3/27/19
C-10	Monitoring Well	201.86	32.65	169.21	32.53	169.33	31.92	169.94
17WW08	Monitoring Well	179.72	9.25	170.47	8.95	170.77	8.27	171.45
18WW01	Monitoring Well	201.31	31.91	169.40	31.80	169.51	31.06	170.25
18WW02	Monitoring Well	179.30	9.02	170.28	8.85	170.45	8.29	171.01
18WW03	Monitoring Well	195.59	26.73	168.86	26.61	168.98	25.77	169.82
18WW04	Monitoring Well	183.74	15.62	168.12	15.22	168.52	14.89	168.85
18WW05	Monitoring Well	189.59	21.77	167.82	21.90	167.69	19.80	169.79
18WW06	Monitoring Well	179.70	9.49	170.21	9.30	170.40	8.71	170.99
18WW07	Monitoring Well	183.67	N/A	NM	N/A	NM	N/A	NM
18WW08	Monitoring Well	177.77	6.95	170.82	6.81	170.96	6.40	171.37
18WW09	Monitoring Well	177.51	7.13	170.38	7.02	170.49	6.79	170.72
18WW10	Monitoring Well	182.26	11.07	171.19	10.92	171.34	10.33	171.93
18WW11	Monitoring Well	182.29	12.52	169.77	12.39	169.90	11.81	170.48
18WW14	Monitoring Well	186.47	18.00	168.47	17.83	168.64	16.05	170.42
18WW15	Monitoring Well	186.24	17.39	168.85	17.24	169.00	15.46	170.78
18WW16	Monitoring Well	201.88	32.71	169.17	32.60	169.28	32.00	169.88
18WW18	Monitoring Well	196.82	26.53	170.29	26.84	169.98	26.80	170.02
18WW19	Monitoring Well	179.56	11.70	167.86	11.49	168.07	11.02	168.54
18WW20	Monitoring Well	180.42	12.45	167.97	12.29	168.13	11.86	168.56
18WW21	Monitoring Well	195.20	26.41	168.79	26.49	168.71	25.66	169.54
18WW22	Monitoring Well	195.37	25.94	169.43	25.78	169.59	25.24	170.13
18WW24	Monitoring Well	176.40	4.95	171.45	4.81	171.59	4.61	171.79
18WW25	Monitoring Well	175.15	4.55	170.60	4.48	170.67	4.29	170.86
18CPTMW01SW	Monitoring Well	198.20	27.35	170.85	27.44	170.76	27.21	170.99
18CPTMW01DW	Monitoring Well	197.92	27.29	170.63	27.39	170.53	27.20	170.72
18CPTMW03SW	Monitoring Well	198.53	28.43	170.10	28.31	170.22	28.10	170.43
18CPTMW04	Monitoring Well	196.60	24.15	172.45	24.03	172.57	23.79	172.81
18CPTMW04SW	Monitoring Well	196.42	26.84	169.58	26.70	169.72	26.51	169.91
18CPTMW06	Monitoring Well	198.12	27.61	170.51	27.49	170.63	27.24	170.88

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Location Identification	Type	Reference Elevation (feet amsl)	Depth to Water (feet) 1/31/19	Groundwater Elevation (feet amsl) 1/31/19	Depth to Water (feet) 2/28/19	Groundwater Elevation (feet amsl) 2/28/19	Depth to Water (feet) 3/26/19-3/27/19	Groundwater Elevation (feet amsl) 3/26/16-3/27/19
18CPTMW07	Monitoring Well	197.32	27.33	169.99	27.19	170.13	26.64	170.68
18CPTMW08SW	Monitoring Well	196.38	26.92	169.46	26.75	169.63	26.51	169.87
18CPTMW08DW	Monitoring Well	196.59	27.31	169.28	27.12	169.47	26.89	169.70
18CPTMW10SW	Monitoring Well	186.98	17.29	169.69	17.08	169.90	16.73	170.25
18CPTMW10DW	Monitoring Well	187.38	18.13	169.25	17.95	169.43	17.62	169.76
18CPTMW12SW	Monitoring Well	190.90	20.79	170.11	20.60	170.30	20.25	170.65
18CPTMW12DW	Monitoring Well	190.25	20.55	169.70	20.43	169.82	20.04	170.21
18CPTMW14	Monitoring Well	196.69	26.74	169.95	26.62	170.07	26.19	170.50
18CPTMW15	Monitoring Well	179.79	8.63	171.16	8.50	171.29	7.91	171.88
18CPTMW16	Monitoring Well	175.37	5.12	170.25	5.00	170.37	4.69	170.68
18CPTMW18	Monitoring Well	194.53	27.61	166.92	27.70	166.83	27.10	167.43
18CPTMW19	Monitoring Well	193.59	24.97	168.62	24.83	168.76	24.33	169.26
18CPTMW19SW	Monitoring Well	193.29	25.44	167.85	25.29	168.00	24.78	168.51
18CPTMW22SW	Monitoring Well	187.79	18.96	168.83	18.85	168.94	17.77	170.02
18CPTMW22R	Monitoring Well	187.23	5.77	181.46	5.70	181.53	5.60	181.63
18CPTMW22DW	Monitoring Well	188.00	19.21	168.79	19.07	168.93	17.65	170.35
18CPTMW23	Monitoring Well	177.47	6.92	170.55	6.77	170.70	6.49	170.98
18CPTMW23SW	Monitoring Well	177.43	6.74	170.69	6.59	170.84	6.30	171.13
18CPTMW24	Monitoring Well	194.89	26.79	168.10	26.60	168.29	25.95	168.94
18CPTMW26	Monitoring Well	182.60	17.13	165.47	17.00	165.60	14.41	168.19
18CPTMW26SW	Monitoring Well	182.00	12.07	169.93	11.91	170.09	11.73	170.27
1824HBSW7	Surface Water Sample	167.92	2.15	165.77	4.04	163.88	2.38	165.54

Notes: amsl - above mean sea level, NM-not measured

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Table 4: Treated Groundwater Discharged – January through March 2019

DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
01/01/2019	Not Measured	Not Calculated	0	0	0	0	0	6.25
01/02/2019	Not Measured	Not Calculated	0	0	0	0	0	6.46
01/03/2019	Not Measured	Not Calculated	0	0	0	0	0	6.50
01/04/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/05/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/06/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/07/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/08/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/09/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/10/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/11/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/12/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/13/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/14/2019	Not Measured	Not Calculated	0	0	0	0	0	6.69
01/15/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/16/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/17/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/18/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/19/2019	Not Measured	Not Calculated	0	0	0	0	0	6.68
01/20/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/21/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/22/2019	Not Measured	Not Calculated	0	0	0	0	0	6.67
01/23/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89
01/24/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89
01/25/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89
01/26/2019	Not Measured	Not Calculated	0	0	0	0	0	6.89
01/27/2019	Not Measured	Not Calculated	0	0	0	0	0	6.88

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DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
01/28/2019	Not Measured	Not Calculated	0	0	0	0	0	6.88
01/29/2019	16,975	305,560	0	176,319	0	0	176,319	6.45
01/30/2019	12,277	220,995	0	127,696	0	0	127,696	6.10
01/31/2019	8,224	148,042	0	65,475	0	0	65,475	5.92
02/01/2019	5,898	106,179	0	57,960	0	0	57,960	5.75
02/02/2019	Not Measured	Not Calculated	0	0	0	0	0	5.75
02/03/2019	Not Measured	Not Calculated	0	0	0	0	0	5.75
02/04/2019	Not Measured	Not Calculated	0	0	0	0	0	5.75
02/05/2019	Not Measured	Not Calculated	0	0	0	0	0	5.74
02/06/2019	Not Measured	Not Calculated	0	0	0	0	0	5.74
02/07/2019	Not Measured	Not Calculated	0	0	0	0	0	5.81
02/08/2019	Not Measured	Not Calculated	0	0	0	0	0	5.81
02/09/2019	Not Measured	Not Calculated	0	0	0	0	0	5.81
02/10/2019	Not Measured	Not Calculated	0	0	0	0	0	5.80
02/11/2019	11,579	3,383	893	0	0	893	893	5.83
02/12/2019	20,399	5,960	25,279	252,560	0	25,279	277,839	5.34
02/13/2019	13,411	3,918	15,361	221,340	0	15,361	236,701	4.78
02/14/2019	16,395	4,790	12,590	174,600	0	12,590	187,190	4.24
02/15/2019	15,454	4,515	9,109	174,240	0	9,109	183,349	3.69
02/16/2019	12,913	3,773	0	0	0	0	0	3.69
02/17/2019	10,373	3,031	21,479	0	0	21,479	21,479	3.69
02/18/2019	9,369	2,737	0	0	0	0	0	3.68
02/19/2019	8,366	2,444	17,960	0	0	17,960	17,960	3.68
02/20/2019	13,181	3,155	11,528	96,600	0	11,528	108,128	3.55
02/21/2019	20,096	4,810	11,444	28,025	0	11,444	39,469	3.50
02/22/2019	Flood Stage	Maximum	9,709	0	0	9,709	9,709	3.53
02/23/2019	Flood Stage	Maximum	0	0	0	0	0	3.60
02/24/2019	Flood Stage	Maximum	0	0	0	0	0	3.70
02/25/2019	Flood Stage	Maximum	38,835	0	0	38,835	38,835	3.70

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DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
02/26/2019	21,127	5,057	10,568	132,480	0	10,568	143,048	3.35
02/27/2019	19,533	4,675	11,891	97,680	0	11,891	109,571	3.15
02/28/2019	16,620	4,856	11,694	91,000	0	11,694	102,694	2.85
03/01/2019	16,125	4,711	11,607	83,520	0	11,607	95,127	2.47
03/02/2019	16,806	4,911	0	0	0	0	0	2.50
03/03/2019	Flood Stage	Maximum	0	0	0	0	0	2.50
03/04/2019	32,976	9,636	35,970	0	0	35,970	35,970	2.50
03/05/2019	23,207	6,781	10,521	0	0	10,521	10,521	2.50
03/06/2019	16,822	4,915	7,756	0	0	7,756	7,756	2.50
03/07/2019	10,652	3,564	4,826	0	0	4,826	4,826	2.50
03/08/2019	10,204	3,414	11,008	0	0	11,008	11,008	2.50
03/09/2019	18,255	5,464	0	0	0	0	0	2.50
03/10/2019	15,231	5,026	0	0	0	0	0	2.50
03/11/2019	14,601	4,885	34,669	0	0	34,669	34,669	2.49
03/12/2019	19,104	6,391	12,730	0	0	12,730	12,730	2.49
03/13/2019	16,809	5,624	12,255	0	0	12,255	12,255	2.58
03/14/2019	Flood Stage	Maximum	12,733	0	0	12,733	12,733	2.66
03/15/2019	Flood Stage	Maximum	10,626	0	0	10,626	10,626	2.66
03/16/2019	Flood Stage	Maximum	0	0	0	0	0	2.65
03/17/2019	20,598	5,723	0	0	0	0	0	2.65
03/18/2019	17,339	5,066	33,160	0	0	33,160	33,160	2.65
03/19/2019	7,625	2,228	14,714	0	0	14,714	14,714	2.64
03/20/2019	5,599	1,636	15,654	0	0	15,654	15,654	2.64
03/21/2019	6,950	2,031	8,703	0	0	8,703	8,703	2.64
03/22/2019	5,977	1,746	9,406	0	0	9,406	9,406	2.63
03/23/2019	5,326	1,544	0	0	0	0	0	2.63
03/24/2019	4,856	1,408	0	0	0	0	0	2.63
03/25/2019	4,431	1,295	18,556	0	0	18,556	18,556	2.66

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DATE	Harrison Bayou Flow (GPM)	Calculated Maximum Rate Allowable (GPM)	Released From GWTP To Harrison Bayou	Released From INF Pond to Harrison Bayou	Released From GWTP to INF Pond	Combined Total Released from GWTP	Combined Total Released to Harrison Bayou	INF Pond Staff Reading (6.20 = 3 ft. Freeboard)
03/26/2019	5,221	1,525	16,413	0	0	16,413	16,413	2.66
03/27/2019	5,592	1,634	16,818	0	0	16,818	16,818	2.66
03/28/2019	9,196	2,687	18,192	0	0	18,192	18,192	2.65
03/29/2019	7,977	2,331	15,769	0	0	15,769	15,769	2.65
03/30/2019	6,231	1,806	0	0	0	0	0	2.65
03/31/2019	5,426	1,573	48,092	0	0	48,092	48,092	2.65
Totals			588,518	1,779,495	0	588,518	2,368,013	

Notes: 1) Bayou flow is not measured on days when no release to the Bayou occurs from the GWTP or INF Pond.

2) When the bayou is in flood stage (over the banks), the flow is not measured and the maximum release is allowed.

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Table 5: Monthly Groundwater Extraction Quantities

ICT or Well Number	January 2019 (gallons)	February 2019 (gallons)	March 2019 (gallons)	Total
1	0	0	0	0
2	0	0	32,524	32,524
3	0	0	0	0
4	0	81,881	111,868	193,749
5	0	0	0	0
EW-1	0	0	368	368
7	0	16,983	20,394	37,377
8	0	100,461	170,110	270,571
18WW17	0	58	5,943	6,001
10	0	0	0	0
11	0	60,730	92,994	153,724
12A	0	5,937	9,785	15,722
12B	0	20,722	27,602	48,324
12C	0	30,102	40,083	70,185
12D	0	0	58,149	58,149
12E	0	29,473	28,942	58,415
13A	0	18,079	41,746	59,825
13B	0	74,529	135,849	210,378
13C	0	0	29,160	29,160
13D	0	30,988	40,081	71,069
13E	0	18,792	13,465	32,257
13F	0	8,752	14,438	23,190
14A	0	0	889	889
14B	0	17,996	22,458	40,454
14C	0	4	66,955	66,959
14D	0	0	55,422	55,422
14E	0	131	44,509	44,640
Total LHAAP-18/24	0	515,618	1,063,734	1,579,352
LHAAP-16	0	0	0	
Total LHAAP-16	0	0	0	0
TOTAL	0	515,618	1,063,734	1,579,352
Note: Power was lost to the GWTP and well field on December 27, 2018. Power was fully restored on February 11, 2019.				

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Table 6: Weekly Perchlorate Sample Results

Sample Identification	Lab Package	Date Sampled	Sample Location	Effluent Discharge Point	Harrison Bayou Maximum Allowable Daily Discharge Perchlorate Concentration (µg/L)	INF Pond Discharge Criteria for Perchlorate (µg/L)	Reporting Limit	Influent Perchlorate (6850)		Effluent Perchlorate (6850)		Does Concentration Meet Discharge Limit? (Yes/No)	No Daily Maximum Concentration		
								Result (µg/L)		Result (µg/L)	DVQ		Ammonia as N (350.3) (mg/L)	Ortho-Phosphate (365.3) (mg/L)	Organic Carbon (415.1) (mg/L)
LH18/24-SP650_022119/BIX	HS19021158	2/21/2019	TK-650	INF Pond	589	17	4	NA	< 2.0	U	Yes	15	4.2	8.4	
LH18/24-SP650_022819/BIX	HS19030014	2/28/2019	TK-650	INF Pond	589	17	4	NA	< 2.0	U	Yes	9.5	1.69	5.1	
LH18/24-SP650_022819_BIX (monthly)	HS19030012	2/28/2019	TK-650	INF Pond	589	17	4	NA	< 2.0	U	Yes	--	--	--	
LH18/24-SP140_022819 (monthly)	HS19030011	2/28/2019	TK-140	--	--	--	NA	3,700	NA		NA	--	--	--	
LH18/24-SP650_030619/BIX	HS19030298	3/6/2019	TK-650	INF Pond	589	17	4	NA	< 2.0	U	Yes	5.5	0.48	2.94	
LH18/24-SP650_031419/BIX	HS19030749	3/14/2019	TK-650	INF Pond	589	17	4	NA	< 2.0	U	Yes	18	4.65	2.96	
LH18/24-SP650_031419_BIX (monthly)	HS19030761	3/14/2019	TK-650	INF Pond	589	17	4	NA	< 2.0	U	Yes	--	--	--	
LH18/24-SP140_031419 (monthly)	HS19030763	3/14/2019	TK-140	--	--	--	NA	8,300	NA		NA	--	--	--	
LH18/24-SP650_032119/BIX	HS19031160	3/21/2019	TK-650	INF Pond	589	17	4	NA	< 2.0	U	NA	16	4.15	2.5	
LH18/24-SP650_032719/BIX	HS19031492	3/27/2019	TK-650	INF Pond	589	17	4	NA	2.6	J	Yes	11	2.42	2.29	
LH18/24-SP650_121818_BIX (quarterly)	HS19031511	3/27/2019	TK-650	INF Pond	589	17	4	NA	2.3	J	Yes	--	--	--	
LH18/24-SP140_121818 (quarterly)	HS19031508	3/27/2019	TK-140	--	--	--	NA	6,500	NA		NA	--	--	--	

Notes:

Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level (PCL)

SP140 samples are influent samples.

µg/L - micrograms per liter

DVQ - data validation qualifier

J - Estimated concentration between the detection limit and limit of quantitation and/or due to quality control discrepancies

NA - not applicable

U - non detect and reported to the limit of detection

BIX - before ion exchange

mg/L - milligrams per liter

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Table 7: Bi-Weekly GWTP Analytical Sampling Results for February 2019

Sample Location Sample Identification	EFFLUENT - Biweekly			EFFLUENT - Monthly		INFLUENT - Monthly*		Does Concentration Meet Effluent Discharge Limits? (Yes/No)	
	LH18/24-SP650_022119	LH18/24-SP650_022819/BIX†	LH18/24-SP140_022819	Result	DVQ	Result	DVQ		
Lab Package	HS19021165			HS19030012		HS19030011			
Sample Date	2/21/2019			2/28/2019		2/28/2019			
Sample Type	GRAB			GRAB		GRAB			
Effluent Limitation for Discharge (µg/L) per Table 2 of ROD	Daily Average Concentration	Daily Maximum Concentration	Detection Limit	Result	DVQ	Result	DVQ	Result	DVQ
	µg/L	µg/L	µg/L	µg/L		µg/L		µg/L	
VOLATILES	µg/L	µg/L	µg/L	µg/L		µg/L		µg/L	
1,1,1-Trichloroethane	3,417	7,230	1	< 0.5	U	< 0.5	U	NA	Yes
1,1,2-Trichloroethane	102.5	216.9	1	< 0.5	U	< 0.5	U	NA	Yes
1,1-Dichloroethane	6,633	14,032	1	< 0.5	U	< 0.5	U	NA	Yes
1,1-Dichloroethene	119	253	1	< 0.5	U	< 0.5	U	NA	Yes
1,2-Dichloroethane	85	181	1	< 0.5	U	< 0.5	U	NA	Yes
1,2-Dichloropropane	NA	NA	1	< 0.5	U	< 0.5	U	NA	Yes
Acetone	1,132	2,395	2	< 1.0	U	< 1.0	U	NA	Yes
Benzene	85	181	1	< 0.5	U	< 0.5	U	NA	Yes
Carbon Tetrachloride	85	181	1	< 0.5	U	< 0.5	U	NA	Yes
Chlorobezene	22,300	47,180	1	< 0.5	U	< 0.5	U	NA	Yes
Chloroform	1,708	3,615	1	< 0.5	U	< 0.5	U	NA	Yes
Ethylbenzene	26,954	57,025	1	< 0.5	U	< 0.5	U	NA	Yes
m,p-Xylenes	39.5	83.6	2	< 1.0	U	< 1.0	U	NA	Yes
Methylene Chloride	803	1,699	2	< 1.0	U	< 1.0	U	NA	Yes
o-Xylene	39.5	83.6	1	< 0.5	U	< 0.5	U	NA	Yes
Styrene	2,829	5,987	1	< 0.5	U	< 0.5	U	NA	Yes
Tetrachloroethene	85.4	180.7	1	< 0.5	U	< 0.5	U	NA	Yes
Toluene	1,980	4,189	1	< 0.5	U	< 0.5	U	NA	Yes
Trichloroethene	85	181	1	0.70	J	0.80	J	NA	Yes
Vinyl Chloride	34	72	1	< 0.5	U	< 0.5	U	NA	Yes
ANIONS	mg/L	mg/L	mg/L	mg/L		mg/L		mg/L	
Chloride	NA	NA	10	320		NA		NA	NA
Sulfate	NA	NA	10	29.8		NA		NA	NA
PERCHLORATE	µg/L	µg/L	µg/L	µg/L		µg/L		µg/L	
Perchlorate	278	589	4	NA		< 2.0	U	3,700	Yes
METALS	mg/L	mg/L	mg/L	mg/L		mg/L		mg/L	
Hexavalent Chromium	0.058	0.124	0.010	NA		< 0.0100	U	< 0.0100	U
Barium	1	2	0.004	NA		0.137		NA	Yes
Lead	0.0022	0.0046	0.002	NA		< 0.00100	U	NA	Yes
Selenium	0.0057	0.0120	0.002	NA		< 0.00250	U	< 0.00250	U
Silver	0.0014	0.0030	0.002	NA		< 0.000500	U	< 0.000500	U
SEMI-VOLATILES	µg/L	µg/L	µg/L	µg/L		µg/L		µg/L	
1,4-Dioxane**	NA	134.2	1	NA		4.3		NA	Yes

Notes:

µg/L - micrograms per liter

mg/L - milligrams per liter

DVQ - data validation qualifier

ROD - Record of Decision

GWTP - Groundwater Treatment Plant

*Influent sample not compared to discharge limits

NA - not applicable or not analyzed

†Perchlorate sample designated by BIX (before ion exchange)

Grab samples are compared to the daily maximum and composite samples to the daily average.

** Calculated Effluent Limit

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Table 8: Bi-Weekly Analytical GWTP Sampling Results for March 2019

Sample Location Sample Identification Lab Package Sample Date Sample Type	EFFLUENT - Biweekly			EFFLUENT - Monthly		INFLUENT - Monthly*		EFFLUENT - Biweekly		Does Concentration Meet Effluent Discharge Limits? (Yes/No)		
	LH18/24-SP650_030619			LH18/24-SP650_031419/BIX*		LH18/24-SP140_031419		LH18/24-SP650_032119				
	HS19030301			HS19030761		HS19030763		HS19031189				
	3/6/2019			3/14/2019		3/14/2019		3/21/2019				
	GRAB			GRAB		GRAB		GRAB				
Effluent Limitation for Discharge (µg/L) per Table 2 of ROD				Result	DVQ	Result	DVQ	Result	DVQ	Result	DVQ	
Daily Average Concentration	Daily Maximum Concentration	Detection Limit										
VOLATILES	µg/L	µg/L	µg/L	µg/L		µg/L						
1,1,1-Trichloroethane	3,417	7,230	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
1,1,2-Trichloroethane	102.5	216.9	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
1,1-Dichloroethane	6,633	14,032	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
1,1-Dichloroethene	119	253	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
1,2-Dichloroethane	85	181	1	< 0.5	U	0.53	J	NA		< 0.5	U	Yes
1,2-Dichloropropane	NA	NA	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
Acetone	1,132	2,395	2	< 2.0	U	< 2.0	U	NA		< 2.0	U	Yes
Benzene	85	181	1	< 0.5	U	< 0.5	U	NA		< 0.5	UJ	Yes
Carbon Tetrachloride	85	181	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
Chlorobenzene	22,300	47,180	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
Chloroform	1,708	3,615	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
Ethylbenzene	26,954	57,025	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
m,p-Xylenes	39.5	83.6	2	< 1.0	U	< 1.0	U	NA		< 1.0	U	Yes
Methylene Chloride	803	1,699	2	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
o-Xylene	39.5	83.6	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
Styrene	2,829	5,987	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
Tetrachloroethene	85.4	180.7	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
Toluene	1,980	4,189	1	< 0.5	U	< 0.5	U	NA		< 0.5	UJ	Yes
Trichloroethene	85	181	1	0.74	J	0.74	J	NA		0.71	J	Yes
Vinyl Chloride	34	72	1	< 0.5	U	< 0.5	U	NA		< 0.5	U	Yes
ANIONS	mg/L	mg/L	mg/L	mg/L		mg/L		mg/L		mg/L		
Chloride	NA	NA	10	312		NA		NA		314		NA
Sulfate	NA	NA	10	13.9		NA		NA		26.8		NA
PERCHLORATE	µg/L	µg/L	µg/L	µg/L		µg/L		µg/L		µg/L		
Perchlorate	278	589	4	NA		< 2.0	U	8,300		NA		Yes
METALS	mg/L	mg/L	mg/L	mg/L		mg/L		mg/L		mg/L		
Hexavalent Chromium	0.058	0.124	0.010	NA		< 0.0100	U	< 0.0100	U	NA		Yes
Barium	1	2	0.004	NA		0.139		NA		NA		Yes
Lead	0.0022	0.0046	0.002	NA		< 0.00100	U	NA		NA		Yes
Selenium	0.0057	0.0120	0.002	NA		< 0.00250	U	< 0.00250	U	NA		Yes
Silver	0.0014	0.0030	0.002	NA		< 0.000500	U	< 0.000500	U	NA		Yes
SEMI-VOLATILES	µg/L	µg/L	µg/L	µg/L		µg/L		µg/L		µg/L		
1,4-Dioxane**	NA	134.2	1	NA		3.1	J	NA		NA		Yes

Notes:

µg/L - micrograms per liter

DVQ - data validation qualifier

GWTP - Groundwater Treatment Plant

U - Non detect reported to the limit of detection

mg/L - milligrams per liter

*perchlorate sample designated by BIX (before ion exchange)

J - estimated concentration between the detection limit and limit of quantitation and/or due to quality control discrepancy

NA - not applicable or not analyzed

Grab samples are compared to the daily maximum and composite samples to the daily average

*Influent sample not compared to discharge limits

** Calculated Effluent Limit

BIX - before ion exchange vessel

UJ - estimated non-detect due to quality control issue

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Table 9: Quarterly GWTP Analytical Sampling Results

	Sample Location			EFFLUENT		INFLUENT*		Does Concentration Meet Discharge Limits? (Yes/No)
	Sample Identification			LH18/24-SP650_032719_BIX	LH18/24-SP140_032719			
	Lab Package			HS19031511	HS19031508			
	Sample Date			3/27/2019	3/27/2019			
	Sample Type			GRAB	GRAB			
	Effluent Limitation for Discharge (µg/L) per Protocol			Result	DVQ	Result	DVQ	
	Daily Average Concentration	Daily Maximum Concentration	Reporting Limit					
VOLATILES	µg/L	µg/L	µg/L	µg/L		µg/L		
1,1,1-Trichloroethane	3,417	7,230	1	< 0.5	U	< 5.0	U	
1,1,2-Trichloroethane	102.5	216.9	1	< 0.5	U	< 5.0	U	
1,1-Dichloroethane	6,633	14,032	1	< 0.5	U	6.4	J	
1,1-Dichloroethene	119	253	1	< 0.5	U	< 5.0	U	
1,2-Dichloroethane	85	181	1	0.57	J	58		
1,2-Dichloropropane	NA	NA	1	< 0.5	U	< 5.0	U	
Acetone	1,132	2,395	2	< 2.0	U	< 20	U	
Benzene	85	181	1	< 0.5	U	7.8	J	
Carbon Tetrachloride	85	181	1	< 0.5	U	25		
Chlorobenzene	22,300	47,180	1	< 0.5	U	< 5.0	U	
Chloroform	1,708	3,615	1	< 0.5	U	9.1	J	
Ethylbenzene	26,954	57,025	1	< 0.5	U	< 5.0	U	
m,p-Xylenes	39.5	83.6	2	< 1.0	U	< 5.0	U	
Methylene Chloride	803	1,699	2	< 0.5	U	110		
o-Xylene	39.5	83.6	1	< 0.5	U	< 5.0	U	
Styrene	2,829	5,987	1	< 0.5	U	< 5.0	U	
Tetrachloroethene	85.4	180.7	1	< 0.5	U	15		
Toluene	1,980	4,189	1	< 0.5	U	< 5.0	U	
Trichloroethene	85	181	1	0.85	J	5,600		
Vinyl Chloride	34	72	1	< 0.5	U	41		
ANIONS	mg/L	mg/L	mg/L	mg/L		mg/L		
Chloride	NA	NA	10	317		197		
Sulfate	NA	NA	10	28.5		32		
PERCHLORATE	µg/L	µg/L	µg/L	µg/L		µg/L		
Perchlorate	278	589	4	2.3	J	6,500		
METALS	mg/L	mg/L	mg/L	mg/L		mg/L		
Aluminum	0.777	1.644	0.0100	0.0311		0.0643		
Antimony	NA	NA	0.00200	0.00101	J	0.00216		
Arsenic	0.365	0.772	0.00200	0.000860	J	0.000971	J	

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Table 9: Quarterly GWTP Analytical Sampling Results

	Sample Location			EFFLUENT		INFLUENT*		Does Concentration Meet Discharge Limits? (Yes/No)
	Sample Identification			LH18/24-SP650_032719_BIX	LH18/24-SP140_032719			
	Lab Package			HS19031511	HS19031508			
	Sample Date			3/27/2019	3/27/2019			
	Sample Type			GRAB	GRAB			
	Effluent Limitation for Discharge (µg/L) per Protocol			Result	DVQ	Result	DVQ	
	Daily Average Concentration	Daily Maximum Concentration	Reporting Limit					
Barium	1	2	0.00400	0.132		0.223		Yes
Beryllium	NA	NA	0.00200	< 0.00250	U	< 0.00250	U	NA
Cadmium	0.0016	0.0034	0.00200	< 0.000500	U	< 0.000500	U	Yes
Calcium	NA	NA	0.500	14.5		16.8		NA
Chromium	0.355	0.752	0.00400	0.00142	J	0.00307	J	Yes
Cobalt	5.433	11.495	0.00500	0.00112	J	0.00548		Yes
Iron	1.132	2.395	0.200	0.125	J	0.317		Yes
Lead	0.0022	0.0046	0.00200	< 0.00100	U	< 0.00100	U	Yes
Magnesium	NA	NA	0.200	12.9		13.8		NA
Manganese	7.323	15.494	0.00500	0.0678		0.214		Yes
Nickel	0.087	0.184	0.00200	0.00344		0.0114		Yes
Potassium	NA	NA	0.200	1.27		1.30		NA
Selenium	0.0057	0.012	0.00200	< 0.00250	U	< 0.00250	U	Yes
Silver	0.0014	0.003	0.00200	< 0.000500	U	< 0.000500	U	Yes
Sodium	NA	NA	1.00	339		196		NA
Thallium	NA	NA	0.00200	< 0.00100	U	< 0.00100	U	NA
Vanadium	1.698	3.592	0.00500	0.00382	UB	0.00237	UB	Yes
Zinc	0.146	0.31	0.00400	0.0363		0.119		No
Mercury	NA	NA	0.000200	< 0.000100	U	< 0.000100	U	NA
1,4-DIOXANE	µg/L	µg/L	µg/L	µg/L		µg/L		
1,4-Dioxane	NA	134.2	1	11.0		16		Yes
CHEMICAL OXYGEN DEMAND (COD)	mg/L	mg/L	mg/L	mg/L		mg/L		
COD	NA	200	75	21		6.0	J	Yes
OIL AND GREASE (O&G)	mg/L	mg/L	mg/L	mg/L		mg/L		
O&G	NA	15	2	2.08		0.729	J	Yes

Notes:

µg/L - micrograms per liter

DVQ - data validation qualifier

Grab samples are compared to the daily maximum and composite samples to the daily average

* only Effluent sample is compared to discharge limits

J - Estimated concentration between the detection limit and limit of quantitation and/or due to quality control discrepancy

U - non detect and reported to the limit of detection

mg/L - milligrams per liter

NA - not applicable

GWTP - Groundwater Treatment Plant

BIX - before ion exchange (perchlorate sample)

UB - considered an artifact of blank contamination

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3 EVALUATION OF LHAAP-16 EXTRACTION SYSTEM

No groundwater was extracted from LHAAP-16 in the 1st quarter of 2019 due to the main transformer being down. However, annual compliance groundwater sampling was completed in this quarter as well as depth to water measurements, which are discussed in the following sections.

3.1 Groundwater Elevation

The groundwater elevations in the piezometers and monitoring wells at LHAAP-16 for January, February, and March 2019 are presented in **Table 10**. The potentiometric surface maps for the shallow and Upper Wilcox (intermediate) groundwater zones at LHAAP-16 for January, February, and March 2019 are presented on **Figures B-7** through **B-12** in **Appendix B**. Based on the potentiometric surface maps, the general groundwater flow direction in the shallow and intermediate zone is south-southeast. However, the March 2019 groundwater flow was more east-northeast towards the Harrison Bayou following 3 months without extraction.

3.2 Groundwater Sampling Activities

On February 26, 2019, a total of eight monitoring wells were sampled at Site LHAAP-16 for VOCs and perchlorate. The analytical results are presented in **Table 11** and the laboratory analytical report is presented in **Appendix D**. Parameters exceeding their respective USEPA Maximum Contaminant Level (MCL) or Texas Risk Reduction Program (TRRP) Groundwater Industrial Medium Specific Concentration (MSC) are 1,1,-dichlorethene (DCE); 1,2-dichloroethane; cis-1,2-DCE; trichloroethene (TCE); and vinyl chloride. Perchlorate was compared to the TRRP Protective Concentration Level (PCL) of 17 µg/L, which was exceeded at LHAAP-16. Two monitoring wells had perchlorate above the PCL with a detection of 51 J (estimated concentration) µg/L in 16EW01 and 230 µg/L (duplicate of 240 µg/L) in 16EW05. All eight of the monitoring wells sampled had exceedances of the MCL for TCE with the highest detection present in 16EW02 (34,000 µg/L). All other monitoring wells had TCE detected at less than 8,000 µg/L. Elevated detections of cis-1,2-DCE and vinyl chloride were also observed in groundwater sampled from 16EW02, indicating that monitored natural attenuation is occurring in addition to the extraction and treatment of LHAAP-16 groundwater. **Figure 3-1** depicts the groundwater sampling results on the LHAAP-16 map. No isoconcentration map was prepared due to the limited number of monitoring wells sampled.

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Gray highlighting indicates concentrations above the MCL/PCL/MSC
 MCL/PCL/MSC - Maximum Contaminant Level/Protective Concentration Level/Medium-Specific Concentrations
 µg/L - micrograms per liter
 J - estimated value between the limit of quantitation and the detection limit and/or estimated due to quality control discrepancies
 U- Undetected: The analyte was analyzed for, but not detected (value is the limit of detection)
 _a - Duplicate Sample
 *PCL – Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level



Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW05_ 022619	16EW05_ 022619_a
Perchlorate	µg/L	17*	230	240
1,1-Dichloroethene	µg/L	7	6.0	6.5
1,2-Dichloroethane	µg/L	5	6.0	6.4
cis-1,2-Dichloroethene	µg/L	70	1,000	1,200
Trichloroethene	µg/L	5	1,700	1,900
Vinyl chloride	µg/L	2	260 J	420 J

Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW01_ 022619
Perchlorate	µg/L	17*	51 J
1,1-Dichloroethene	µg/L	7	14
1,2-Dichloroethane	µg/L	5	12
cis-1,2-Dichloroethene	µg/L	70	3,700 J
Trichloroethene	µg/L	5	3,700
Vinyl chloride	µg/L	2	45

Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW02_ 022619
Perchlorate	µg/L	17*	<2.0 U
1,1-Dichloroethene	µg/L	7	86
1,2-Dichloroethane	µg/L	5	34 J
cis-1,2-Dichloroethene	µg/L	70	17,000
Trichloroethene	µg/L	5	34,000
Vinyl chloride	µg/L	2	51

Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW06_ 022619
Perchlorate	µg/L	17*	<2.0 U
1,1-Dichloroethene	µg/L	7	0.95 J
1,2-Dichloroethane	µg/L	5	0.56 J
cis-1,2-Dichloroethene	µg/L	70	180
Trichloroethene	µg/L	5	250
Vinyl chloride	µg/L	2	<0.50 U

Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW07_ 022619
Perchlorate	µg/L	17*	<2.0 U
1,1-Dichloroethene	µg/L	7	1.7
1,2-Dichloroethane	µg/L	5	<0.50 U
cis-1,2-Dichloroethene	µg/L	70	340
Trichloroethene	µg/L	5	81
Vinyl chloride	µg/L	2	25

Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW03_ 022619
Perchlorate	µg/L	17*	<2.0 U
1,1-Dichloroethene	µg/L	7	12
1,2-Dichloroethane	µg/L	5	6.2 J
cis-1,2-Dichloroethene	µg/L	70	2,000
Trichloroethene	µg/L	5	7,400
Vinyl chloride	µg/L	2	7.5 J

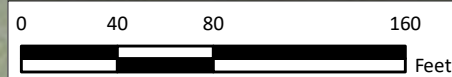
Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW08_ 022619
Perchlorate	µg/L	17*	<2.0 U
1,1-Dichloroethene	µg/L	7	0.76 J
1,2-Dichloroethane	µg/L	5	<0.50 U
cis-1,2-Dichloroethene	µg/L	70	170
Trichloroethene	µg/L	5	170
Vinyl chloride	µg/L	2	0.86 J

Location ID: Sample Date:	Units	MCL/PCL/MSC	16EW04_ 022619
Perchlorate	µg/L	17*	<2.0 U
1,1-Dichloroethene	µg/L	7	0.88 J
1,2-Dichloroethane	µg/L	5	0.55 J
cis-1,2-Dichloroethene	µg/L	70	260
Trichloroethene	µg/L	5	900
Vinyl chloride	µg/L	2	0.47 J

Unnamed Road

Harrison Bayou

LHAAP-16



Legend:

- Extraction Well Location
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Road Line
- Former Building Location and Number



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Table 10: Groundwater Elevations at LHAAP-16 Piezometers and Monitoring Wells-1st Quarter 2019

Location Identification	Type	Reference Elevation (feet amsl)	Depth to Water (feet) 1/30/19	Groundwater Elevation (feet amsl) 1/30/19	Depth to Water (feet) 2/27/19	Groundwater Elevation (feet amsl) 2/27/19	Depth to Water (feet) 3/26/19	Groundwater Elevation (feet amsl) 3/26/19
16PZ-1	Piezometer	199.44	27.55	171.89	27.43	172.01	26.67	172.77
16PZ-2	Piezometer	199.75	27.77	171.98	27.60	172.15	26.60	173.15
16PZ-3	Piezometer	198.61	26.64	171.97	26.49	172.12	25.47	173.14
16PZ-4	Piezometer	198.81	27.21	171.60	27.05	171.76	25.75	173.06
16PZ-5	Piezometer	198.31	26.35	171.96	26.21	172.10	25.05	173.26
16PZ-6	Piezometer	198.61	27.39	171.22	27.24	171.37	25.61	173.00
16PZ-7	Piezometer	200.10	28.70	171.40	28.50	171.60	26.63	173.47
16PZ-8	Piezometer	199.93	27.96	171.97	27.81	172.12	26.88	173.05
16PZ-9	Piezometer	196.49	24.73	171.76	24.58	171.91	23.50	172.99
16PZ-10	Piezometer	196.65	24.88	171.77	24.70	171.95	23.67	172.98
16PZ-11	Piezometer	198.88	26.79	172.09	26.65	172.23	25.68	173.20
16PZ-12	Piezometer	199.00	27.71	171.29	27.59	171.41	25.85	173.15
16PZ-13	Piezometer	196.58	24.75	171.83	24.60	171.98	23.45	173.13
16PZ-14	Piezometer	196.09	24.31	171.78	24.17	171.92	23.06	173.03
16PZ-15	Piezometer	191.93	19.64	172.29	19.49	172.44	18.69	173.24
16PZ-16	Piezometer	190.79	19.16	171.63	19.02	171.77	17.84	172.95
16PZ-17	Piezometer	186.67	16.47	170.20	16.30	170.37	13.59	173.08
16PZ-18	Piezometer	185.99	16.53	169.46	16.38	169.61	13.17	172.82
16PZ-19	Piezometer	183.98	11.29	172.69	10.92	173.06	11.07	172.91
16PZ-20	Piezometer	183.12	10.83	172.29	10.57	172.55	10.25	172.87
16WW12	Monitoring Well	188.81	16.65	172.16	16.52	172.29	16.25	172.56
16WW14	Monitoring Well	198.87	26.14	172.73	26.00	172.87	25.06	173.81
16WW22	Monitoring Well	200.13	28.34	171.79	28.25	171.88	26.92	173.21
16WW25	Monitoring Well	188.77	18.63	170.14	18.50	170.27	15.36	173.41
16WW26	Monitoring Well	188.83	17.09	171.74	16.97	171.86	14.90	173.93
16WW29	Monitoring Well	178.24	6.31	171.93	6.10	172.14	5.05	173.19
16WW30	Monitoring Well	178.47	6.60	171.87	6.41	172.06	5.30	173.17
16WW31	Monitoring Well	202.78	30.54	172.24	30.41	172.37	29.37	173.41
16WW33	Monitoring Well	203.09	30.50	172.59	30.37	172.72	29.50	173.59
16WW35	Monitoring Well	191.23	18.91	172.32	18.80	172.43	17.61	173.62
16WW36	Monitoring Well	190.94	18.15	172.79	18.04	172.90	17.10	173.84

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Table 11: LHAAP-16 Annual Sampling Results - February 2019

Location Identification: Sample Date:	Units	MCL/PCL/ MSC	16EW01_ 022619	16EW02_ 022619	16EW03_ 022619	16EW04_ 022619	16EW05_ 022619	16EW05_ 022619_a	16EW06_ 022619	16EW07_ 022619	16EW08_ 022619
Lab Package:			HS19021428								
Location Description			NE, middle region Sampled Annually	NE, middle region Sampled Annually	NE, middle region Sampled Annually	ENE, middle region Sampled Annually	NE, middle region Sampled Annually	NE, middle region Sampled Annually Duplicate.	NE, middle region Sampled Annually	NE, middle region Sampled Annually	ENE, middle region Sampled Annually
Perchlorate (6850)											
Perchlorate	µg/L	17*	51 J	< 2.0 U	< 2.0 U	< 2.0 U	230	240	< 2.0 U	< 2.0 U	< 2.0 U
Volatile Organic Compounds (8260C)											
1,1,1,2-Tetrachloroethane	µg/L	110	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,1-Trichloroethane	µg/L	200	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2,2-Tetrachloroethane	µg/L	14	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1,2-Trichloroethane	µg/L	5	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,1-Dichloroethane	µg/L	10,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	1.4	1.4	< 0.50 U	< 0.50 U	< 0.50 U
1,1-Dichloroethene	µg/L	7	14	86	12	0.88 J	6.0	6.5	0.95 J	1.7	0.76 J
1,1-Dichloropropene	µg/L	2.9	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,3-Trichlorobenzene	µg/L	310	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,3-Trichloropropane	µg/L	0.041	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-Trichlorobenzene	µg/L	70	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2,4-Trimethylbenzene	µg/L	5,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-Dibromo-3-chloropropane	µg/L	0.2	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-Dibromoethane	µg/L	0.05	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-Dichlorobenzene	µg/L	600	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,2-Dichloroethane	µg/L	5	12	34 J	6.2 J	0.55 J	6.0	6.4	0.56 J	< 0.50 U	< 0.50 U
1,2-Dichloropropane	µg/L	5	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3,5-Trimethylbenzene	µg/L	5,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-Dichlorobenzene	µg/L	3,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,3-Dichloropropane	µg/L	29	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
1,4-Dichlorobenzene	µg/L	75	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2,2-Dichloropropane	µg/L	42	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-Butanone	µg/L	61,000	< 5.0 U	< 25 U	< 10 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
2-Chlorotoluene	µg/L	2,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
2-Hexanone	µg/L	6,100	< 5.0 U	< 50 U	< 10 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U

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Table 11: LHAAP-16 Annual Sampling Results - February 2019

Location Identification: Sample Date:	Units	MCL/PCL/ MSC	16EW01_ 022619	16EW02_ 022619	16EW03_ 022619	16EW04_ 022619	16EW05_ 022619	16EW05_ 022619_a	16EW06_ 022619	16EW07_ 022619	16EW08_ 022619
Lab Package:			HS19021428								
Location Description			NE, middle region Sampled Annually	NE, middle region Sampled Annually	NE, middle region Sampled Annually	ENE, middle region Sampled Annually	NE, middle region Sampled Annually	NE, middle region Sampled Annually Duplicate.	NE, middle region Sampled Annually	NE, middle region Sampled Annually	ENE, middle region Sampled Annually
4-Chlorotoluene	µg/L	2,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
4-Isopropyltoluene	µg/L	10,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
4-Methyl-2-pentanone	µg/L	8,200	< 5.0 U	< 50 U	< 10 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
Acetone	µg/L	92,000	< 10 U	< 100 U	< 20 U	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U
Benzene	µg/L	5	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Bromobenzene	µg/L	2,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Bromochloromethane	µg/L	4,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Bromodichloromethane	µg/L	4.6	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Bromoform	µg/L	36	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Bromomethane	µg/L	140	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Carbon disulfide	µg/L	10,000	< 5.0 U	< 50 U	< 10 U	< 0.50 U	< 1.0 U	< 1.0 U	< 0.50 U	< 0.50 U	< 0.50 U
Carbon tetrachloride	µg/L	5	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Chlorobenzene	µg/L	100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Chloroethane	µg/L	41,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Chloroform	µg/L	1,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Chloromethane	µg/L	220	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
cis-1,2-Dichloroethene	µg/L	70	3,700 J	17,000	2,000	260	1,000	1,200	180	340	170
cis-1,3-Dichloropropene	µg/L	5.3	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Dibromochloromethane	µg/L	34	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Dibromomethane	µg/L	380	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Dichlorodifluoromethane	µg/L	20,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Ethylbenzene	µg/L	700	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Hexachlorobutadiene	µg/L	20	< 5.0 U	< 50 U	< 10 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U

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Table 11: LHAAP-16 Annual Sampling Results - February 2019

Location Identification: Sample Date:	Units	MCL/PCL/ MSC	16EW01_ 022619	16EW02_ 022619	16EW03_ 022619	16EW04_ 022619	16EW05_ 022619	16EW05_ 022619_a	16EW06_ 022619	16EW07_ 022619	16EW08_ 022619
Lab Package:			HS19021428								
Location Description			NE, middle region Sampled Annually	NE, middle region Sampled Annually	NE, middle region Sampled Annually	ENE, middle region Sampled Annually	NE, middle region Sampled Annually	NE, middle region Sampled Annually Duplicate.	NE, middle region Sampled Annually	NE, middle region Sampled Annually	ENE, middle region Sampled Annually
Isopropylbenzene	µg/L	10,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
m,p-Xylene	µg/L	10,000**	< 5.0 U	< 50 U	< 10 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U
Methylene chloride	µg/L	5	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Naphthalene	µg/L	2,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
n-Butylbenzene	µg/L	4,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
n-Propylbenzene	µg/L	4,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
o-Xylene	µg/L	10,000**	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
sec-Butylbenzene	µg/L	4,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Styrene	µg/L	100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
tert-Butylbenzene	µg/L	4,100	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Tetrachloroethene	µg/L	5	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Toluene	µg/L	1,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
trans-1,2-Dichloroethene	µg/L	100	8.7	24 J	< 5.0 U	0.72 J	3.7	3.9	< 0.50 U	0.84 J	0.64 J
trans-1,3-Dichloropropene	µg/L	29	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Trichloroethene	µg/L	5	3,700	34,000	7,400	900	1,700	1,900	250	81	170
Trichlorofluoromethane	µg/L	31,000	< 2.5 U	< 25 U	< 5.0 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U	< 0.50 U
Vinyl chloride	µg/L	2	45	51	7.5 J	0.47 J	260 J	420 J	< 0.50 U	25	0.86 J

Notes:

Blue Highlighting Indicates concentrations above the MCL/PCL/MSC

MCL/PCL/MSC - Maximum Contaminant Limit/Protective Concentration Level/Medium-Specific Concentration

µg/L - micrograms per liter

J - estimated value between the limit of quantitation and the detection limit and/or estimated due to quality control discrepancies

U - Undetected: The analyte was analyzed for, but not detected. Value presented is the limit of detection.

*PCL – Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level

** Value is for total xylenes

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4 QUALITY CONTROL

This report summarizes the data for samples collected during January, February, and March 2019. The samples were reviewed and validated in accordance with the guidelines in the *USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review* (USEPA, January 2017); *USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review* (USEPA, January 2017); and the quality control criteria specified in the *Basewide Uniform Federal Policy - Quality Assurance Project Plan Longhorn Army Ammunition Plant* which is in Appendix C of the *Final Installation-Wide Work Plan for Longhorn Army Ammunition Plant Karnack, Texas* (Bhate, May 2018).

The purpose of the sampling program is to evaluate the effectiveness of the groundwater pump and treat system, assess water quality within the capture zone, and assure compliance with the effluent discharge requirements of the Interim ROD. Quality control and quality assurance problems noted in the case narratives received from the laboratory are minor and do not affect the usability of the data for compliance at the GWTP. No sample results from the 1st quarter of 2019 were rejected due to quality control problems.

ALS Environmental analyzed the compliance samples collected from the GWTP. Independent data verification and validation was performed by the Bhate project chemist as described in the Quality Control Summary Report in **Appendix E**. The laboratory reports for the 1st quarter of 2019 are included in **Appendix C** on a CD and the laboratory results for sampling conducted at LHAAP-16 are included in **Appendix D** on CD. Air monitoring data is presented in **Appendix F** on CD.

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5 TREATED GROUNDWATER DISCHARGED

Reinjection of groundwater in ICT 6 and ICT 9 was discontinued as of July 15, 2012. The last injection occurred on May 24, 2012, immediately prior to the scrubber system malfunction which caused GWTP operation to cease temporarily.

Treated groundwater that met the perchlorate discharge criteria was discharged to Harrison Bayou or the INF Pond in accordance with the Protocol for Discharging GWTP Effluent (**Appendix G**). **Table 4** summarizes flow rates from the INF Pond to the Harrison Bayou, the maximum flow rate allowed by chloride and sulfate concentrations, and the approximated flow rate discharged for the 1st quarter of 2019. No treated groundwater was discharged to Harrison Bayou in January 2019 from the GWTP directly. Only water from the INF Pond was discharged to the Harrison Bayou in January 2019 because the main transformer was down such that the GWTP was not operational until February 2019. No treated groundwater was discharged to the INF Pond in the 1st quarter of 2019 due to the continuous flow of the Harrison Bayou.

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6 AIR MONITORING

6.1 Summary of Air Monitoring Approach

Operation of the GWTP without air abatement was approved by the TCEQ and USEPA conditioned on collection of air monitoring data to determine the effect of GWTP operation on ambient air quality with respect to potential human health exposure risk. An Interim Air Monitoring Plan was developed by AECOM in August 2012 and used to implement the air monitoring program. The air monitoring program included sampling emission concentrations from the air stripper, ambient air at the GWTP, and ambient air downwind of the GWTP. Collection of air data occurred on a weekly basis between September 2012 and September 2013, on a monthly basis between September 2013 and September 2014, and on a quarterly basis since that time. The sampling program includes use of Summa canisters and a photoionization detector (PID) to measure vapor phase concentrations. The air stripper emission sample is collected as a grab sample, while the ambient air samples are collected as composite samples. The GWTP sample is collected over 8 hours to represent a work day and the downwind sample is collected over 24 hours to represent potential exposure to an off-site receptor¹. The downwind sample is collected at the closest downwind property boundary, based on prevailing wind direction.

PID data (after system calibration) are collected each time the GWTP is operated and serve as a real-time indicator of ambient air conditions at and downwind of the GWTP. Correlations between definitive analytical air data and PID measurements were established and a means to calculate contaminant concentration from PID measurements was developed. A PID threshold of 0.4 parts per million by volume (ppmv) in ambient air was established, such that Summa canister measurements would occur when the PID threshold is exceeded.

The Summa canister samples are analyzed for VOCs using USEPA Method TO-15. The PID measurements are collected after instrument calibration. The air sampling results are summarized and reported to the USEPA and TCEQ in the GWTP quarterly reports; however, the air results are reviewed immediately upon receipt for the potential presence of any exceedances of ambient air concentrations. **Appendix H** (Tables 1 through 3) includes a comparison of ambient air concentrations with TCEQ Air Monitoring Comparison Values (AMCVs) or the short-term Effects Screening Levels (ESLs) for chemicals with no published AMCVs, calculations of emission rates from the emission point, and a compilation of PID results and calibration records. The air monitoring results to date indicate that all ambient air concentrations are lower than the AMCVs or ESLs. The stripper stack sample concentrations are used to calculate emission rates in

¹ Off-site receptor - Any recreational area, residence, commercial/industrial facility, or other normally occupied structures not used solely by the owner or operator of the facilities or the owner of the site upon which the facilities are located. Measurements of distances to determine compliance with this distance restriction must be taken toward structures that are in use as of the date that a notification is filed with the commission.

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pounds per hour (lbs/hr) and tons per year (tpy). The calculated emission rates in lbs/hr are then compared to the allowable emission rates per 30 Texas Administrative Code (TAC) §106.533(f)(1). The emission rates have been lower than the allowable emission rates to the conservatively-selected off-site receptor. The calculated emission rate in tpy is compared to the allowable limit of 5 tpy per chemical. All emission rates have been lower than the allowable emission rates.

The air monitoring results from the first few months of operation between September and November 2012 were compiled and submitted in a separate report (December 2012) (along with validated data) to TCEQ to demonstrate compliance with Texas Permit by Rule emission standards. Approval of the analytical results and concurrence that the site will continue to meet Title 30 TAC §106.533 without the use of air abatement using a catalytic oxidation system was obtained from the TCEQ via email on February 22, 2013.

On February 18, 2013, AECOM presented analysis of the approach applicable to obtaining a variance for operating the GWTP without air abatement equipment to the TCEQ and USEPA. The analysis indicated that the use of an Explanation of Significant Difference (ESD) was the appropriate approach for the site. Approval of use of an ESD was obtained from the USEPA via email on March 21, 2013. The ESD was developed, reviewed, and accepted by USEPA and TCEQ. The ESD was signed by the designated parties on April 3, 2014, and concurrence from the TCEQ was obtained in a letter dated April 16, 2014.

6.2 Air Monitoring Results for the 1st Quarter of 2019

During the 1st quarter of 2019, air sampling was completed on February 27 and 28, 2019. The laboratory data packages are presented in **Appendix F**. A summary of the air sampling results is presented in **Appendix H (Tables 1 through 3)**. All results met the criteria described in Section 6.1.

6.2.1 Summa Canister Monitoring Results

One sampling event was conducted on February 27 and 28, 2019, for presentation during the 1st Quarter 2019 reporting period using Summa canisters. The samples were collected and analyzed as described in Section 6.1 and per the approved air monitoring plan dated August 2012. The analytical results were then compiled in spreadsheets where calculations were completed and comparisons to applicable criteria were made as described in Section 6.1.

6.2.1.1 Ambient Air Results

Benzene, cis-1,2-dichloroethene, methylene chloride, trichloroethene, n-hexane, toluene, dichlorodifluoromethane, trichlorofluoromethane, and trichlorotrifluoroethane were detected in February 2019 in ambient air downwind of the GWTP.

Compounds originating at the GWTP would be expected to have lower concentrations in the downwind sampling location than at the GWTP sampling location. Likewise, compounds like dichlorodifluoromethane and trichlorofluoromethane with similar concentrations in both GWTP ambient air and downwind ambient air are suspected to be present in the ambient (background)

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air. The ambient air results during the quarter met the ambient air criteria, as presented in **Table 1** within **Appendix H**.

6.2.1.2 Air Stripper Effluent Results

The VOCs present in groundwater that are removed via the air stripper include 1,2-dichloroethane; cis-1,2-DCE; methylene chloride; TCE; and trichlorotrifluoroethane. The highest reported concentrations are for TCE; methylene chloride; cis-1,2-DCE; and trichlorotrifluoroethane. These compounds are frequently reported in groundwater at the site, with the exception of trichlorotrifluoroethane which is not typically a groundwater analyte at LHAAP. Trichlorotrifluoroethane, however, appears to be present in groundwater as indicated by limited analysis conducted in December 2013, where it was detected in several wells, and from historical accounts. Many of the chemicals that are reported in ambient air are not detected in the air stripper effluent. This is likely because the reporting limit for the air stripper effluent is higher than the reporting limit for the ambient air samples or the source for some of these chemicals are extraneous to groundwater.

The air stripper effluent concentrations were below the emission criteria, as presented in **Table 2** within **Appendix H**.

6.2.2 PID Results

Along with collection of Summa canister air samples, PID measurements from the same sources/areas are collected and recorded. These simultaneous measurements allowed establishing a correlation between PID readings and VOC concentrations in the Summa canister air samples. Conversion from PID to compound concentrations was established by TCEQ in 30 TAC §106.533(h). The TCEQ equation allows use of a PID to determine individual compound concentrations if the distribution of chemicals in the ambient air is known or assumed. This allows the use of a PID as a tool to measure VOC concentrations and convert the PID results to estimates of compound concentrations. All ambient air PID measurements during this quarter at the GWTP were reported at 0.0 parts per million. The results of the PID readings collected during GWTP operations are presented in **Table 3** within **Appendix H**.

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7 COMMENTS AND RESPONSES

The USEPA issued the following two comments on July 9, 2019.

Comment 1: Appendix D, Trend Analysis: For monitoring wells 18WW08, MW-2, MW-8, MW-109, and MW-120, please provide bigger charts with larger print for the contaminant concentrations. The print size for well AWD-3 is a good size to use. These are very useful charts.

Response to Comment 1: These charts will be reformatted and provided as change pages to the 4th Quarter 2018 Report.

Comment 2: General Comment: Have the pump replacements for ICTs 12D, 13C, 2, 14A, 14C, 14D, 14E, and WW-01 been completed?

Response to Comment 2: Section 1.2.2.6 of this report documents that these pumps were replaced in the 1st Quarter 2019.

The TCEQ issued the following comment on July 17, 2019.

Comment 3: Table ES-1 and 4 – Could “No Release” be moved to a different column? Isn’t bayou flow collected/calculated every day even when there is no release? If so, could this information be included in the Harrison Bayou Flow column. When there is no discharge, “No Release” could go in the calculated max rate column instead of NA.

Response to Comment 3: Bayou flow is not measured when there is no release to the bayou from the GWTP or the INF Pond. Tables ES-1 and 4 will be revised to note “Not Measured” in the “Harrison Bayou Flow” column and “Not Calculated” in the “Calculated Maximum Rate Allowable” column. The following notes will also be added to the tables:

- 1) *Bayou flow is not measured on days when no release to the Bayou occurs from the GWTP or INF Pond.*
- 2) *When the bayou is in flood stage (over the banks), the flow is not measured and the maximum release is allowed.*

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GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
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APPENDIX A
ICT LAYOUT AND GWTP PROCESS FLOW DIAGRAM

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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Table A-1: ICTs Completion Depths

ICT	TOC Elevation	Total Depth	Sump Elevation	Comment
1	186.07	22.5	163.57	Taken out of service in 2007.
2	185.02	29.5	155.52	
3	192.27	37.75	154.52	Taken out of service in 2007.
4	193.51	37.5	156.01	
5	192.67	35	157.67	Taken out of service in 2007.
6	197.30	40.75	156.55	Converted to infiltration in 2007. Ceased reinjection in July 2012.
7	198.03	32.33	165.7	
8	198.97	44.5	154.47	
9	197.64	45.5	152.14	Converted to infiltration in 2007. Ceased reinjection in July 2012.
10	198.07	45.42	152.65	Taken out of service in 2007.
11	198.01	43.33	154.68	
12A	189.06	31.5	157.56	Taken out of service in 2007. Reinstituted in December 2012.
12B	191.97	36.25	155.72	
12C	193.90	34.33	159.57	
12D	185.64	33.75	151.89	
12E	183.38	32.25	151.13	
13A	182.59	28.17	154.42	
13B	184.72	29.58	155.14	
13C	186.13	28.17	157.96	
13D	186.72	26.17	160.55	
13E	191.79	27.08	164.71	
13F	197.81	32.33	165.48	
13G	197.03	27.25	169.78	Taken out of service in 2008.
14A	196.8	43.00	153.8	
14B	197.61	43.42	154.19	
14C	197.86	41.33	156.53	
14D	198.47	44.25	154.22	
14E	198.47	43.08	155.39	

Note(s):

ICT - interception-collection trench

TOC - top of casing, measuring point for groundwater elevations

Elevations are reported as feet above mean sea level.

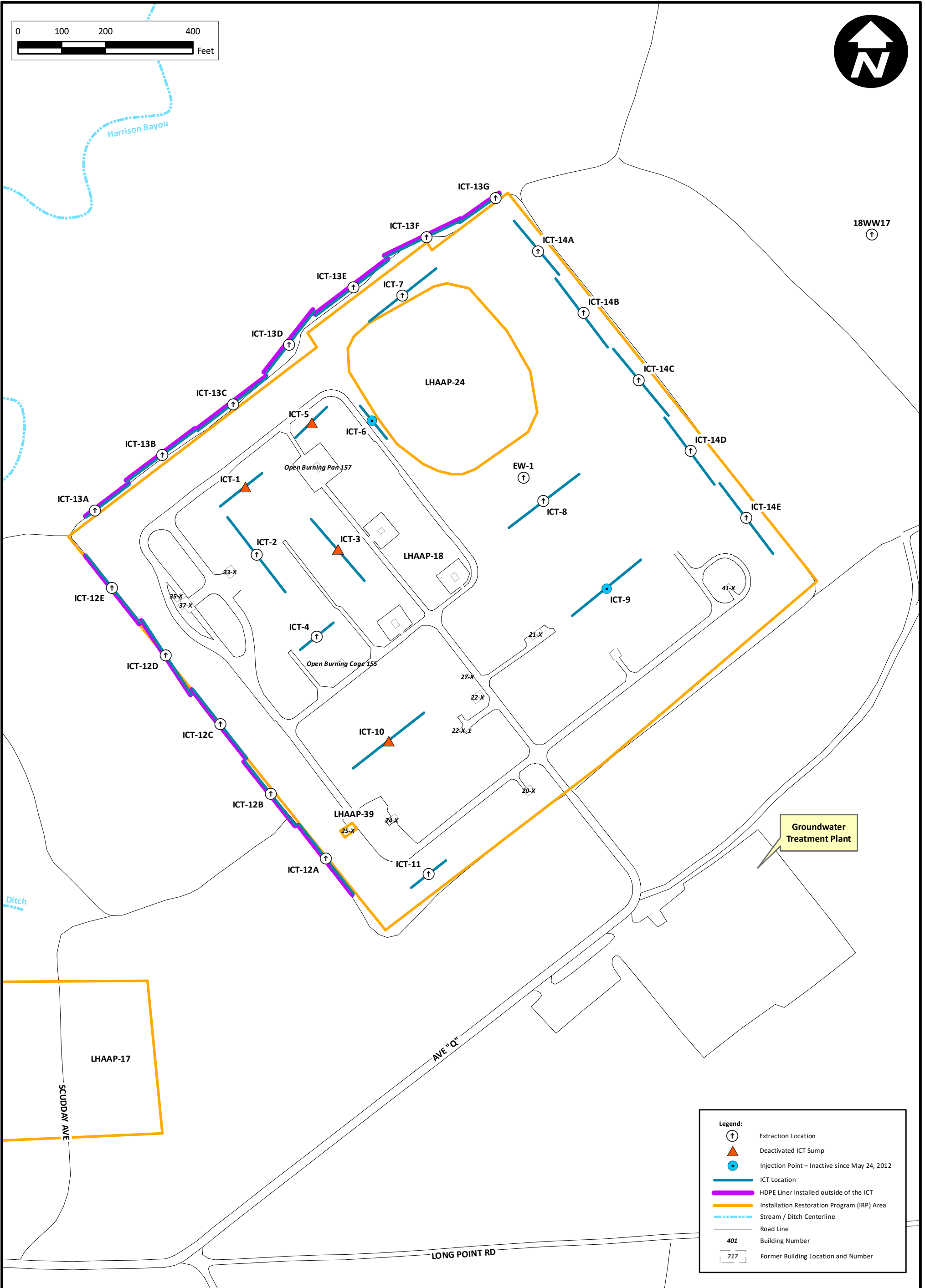
Total depths are reported as feet below TOC.

Sump elevation calculated by subtracting total depth from TOC elevation.

ICTs were installed in 1998.

ICT 12A was replaced on December 5, 2012, and extraction has resumed.

TOC Elevations and total depth measured in October 2003, 4th Quarter 2003, Groundwater Treatment Plant Report.



Legend:

- Extraction Location
- Deactivated ICT Sump
- Injection Point – Inactive since May 24, 2012
- ICT Location
- HDPE Liner Installed outside of the ICT
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Road Line
- Building Number
- Former Building Location and Number

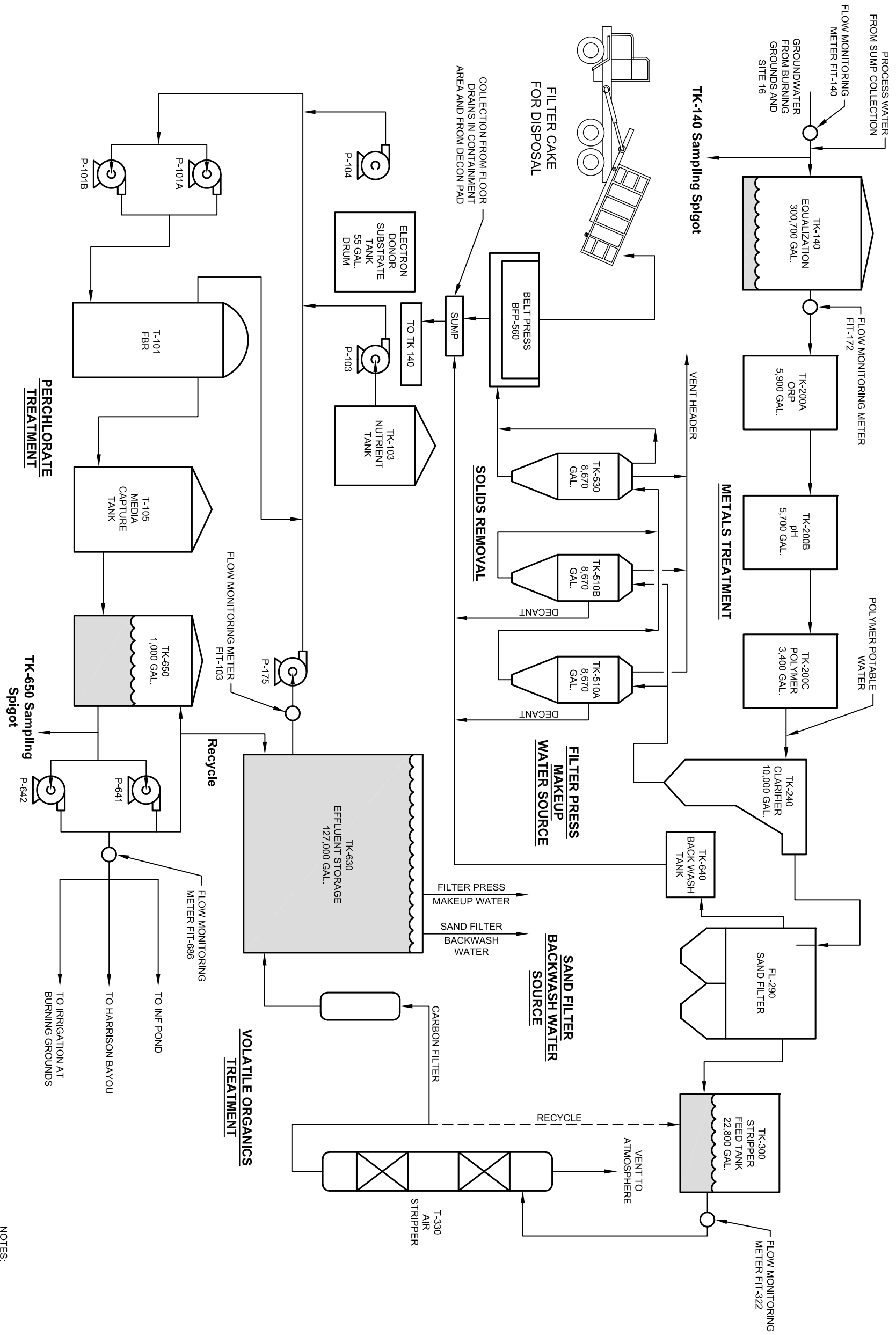


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ICT Layout Map

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Figure A-1



NOTES:
 GAL. GALLON
 TK or T TANK
 BFP BELT FILTER PRESS
 P PUMP
 FL FILTER
 FBR FLUIDIZED BED REACTOR

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NW01312.0150	Not to Scale	4/19/2019	MRM

Process Flow Diagram

Figure A-2



GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
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APPENDIX B
GROUNDWATER ELEVATION CONTOUR MAPS

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
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Table B-1: Extraction Equipment Maintenance Since 2011

LHAAP-18/24

Well I.D.	Replaced Parts	Date	Contractor
ICT 14E	Pump	3/15/2012	Shaw
ICT 14D	Pump, motor, level probes	3/16/2012	Shaw
ICT 14B	Pump, level probes, level probe wire	3/16/2012	Shaw
ICT 14A	Pump, motor, level probes, level probe wire	3/17/2012	Shaw
ICT 2	Pump, motor	3/17/2012	Shaw
ICT 13D	Pump	3/19/2012	Shaw
ICT 13B	Pump	3/20/2012	Shaw
ICT 14E	Pump, motor, broken piping	11/19/2012	AECOM
ICT 13C	Pump	11/20/2012	AECOM
ICT13E	Pump	11/20/2012	AECOM
ICT 12A	Pump, motor, wiring harness, level probes, level probe wire	12/5/2012	AECOM
ICT 7	Pump, motor, wiring harness, level probes	12/6/2012	AECOM
ICT2	Pump	6/10/2013	AECOM
ICT 13C	Pump	6/11/2013	AECOM
ICT 13D	Pump	6/12/2013	AECOM
ICT 14E	Pump rebuilt	6/15/2013	AECOM
ICT 14E	Replaced low level switch	6/19/2013	AECOM
ICT 13C	Pump, wiring harness, flow meter	4/15/2014	AECOM
ICT 14D	Repaired level probes	6/24/2014	AECOM
ICT 14E	Repaired level probes	6/24/2014	AECOM
ICT 14E	Pump and motor troubleshooting	6/26/2014	AECOM
ICT2, 13F, 14C, 14D, 14E	Repaired level probes	7/7/2014	AECOM
ICT 12E	Pump, motor	10/2/2014	AECOM
ICT 12E	Wiring harness, fixed leak	10/8/2014	AECOM
ICT 12E	Level probes	10/9/2014	AECOM
ICT13A	Pump, piping	10/15/2014	AECOM
ICT 12E	Repaired leaking fittings	10/16/2014	AECOM
ICT 11	1" tee and 1" elbow	1/13/2015	AECOM
ICT 12B	Flow meter	1/13/2015	AECOM
ICT 7	1" tee, Repaired 1" pipe	1/13/2015	AECOM
ICT 13A	Flow meter	1/15/2015	AECOM
ICT 13B	Pump	1/15/2015	AECOM
ICT 13C	Pump	1/16/2015	AECOM
ICT 7	Low level probe	1/16/2015	AECOM
ICT 13D	Pump, level probes	1/17/2015	AECOM
ICT 14C	Low level probe	1/17/2015	AECOM
ICT 14C	Low level probe	1/29/2015	AECOM
ICT 14D	Low level probe	1/29/2015	AECOM
ICT 13D	Level probes	1/29/2015	AECOM
ICT 2	Pump	1/30/2015	AECOM
ICT 8	Fuse	3/2/2015	AECOM
ICT 8	Fuse	3/9/2015	AECOM
ICT 12E	Flow meter	3/13/2015	AECOM
ICT 13D	Union	3/13/2015	AECOM
ICT 14C	Cleaned level probes	4/1/2015	AECOM
ICT 14D	Cleaned level probes	4/1/2015	AECOM
ICT 13A	Cleaned level probes	4/21/2015	AECOM
ICT 14C	Cleaned level probes	4/21/2015	AECOM
ICT 8	Low level probe	7/24/2015	AECOM
ICT 13C	Installed New Pump	7/28/2015	AECOM

Table B-1: Extraction Equipment Maintenance Since 2011

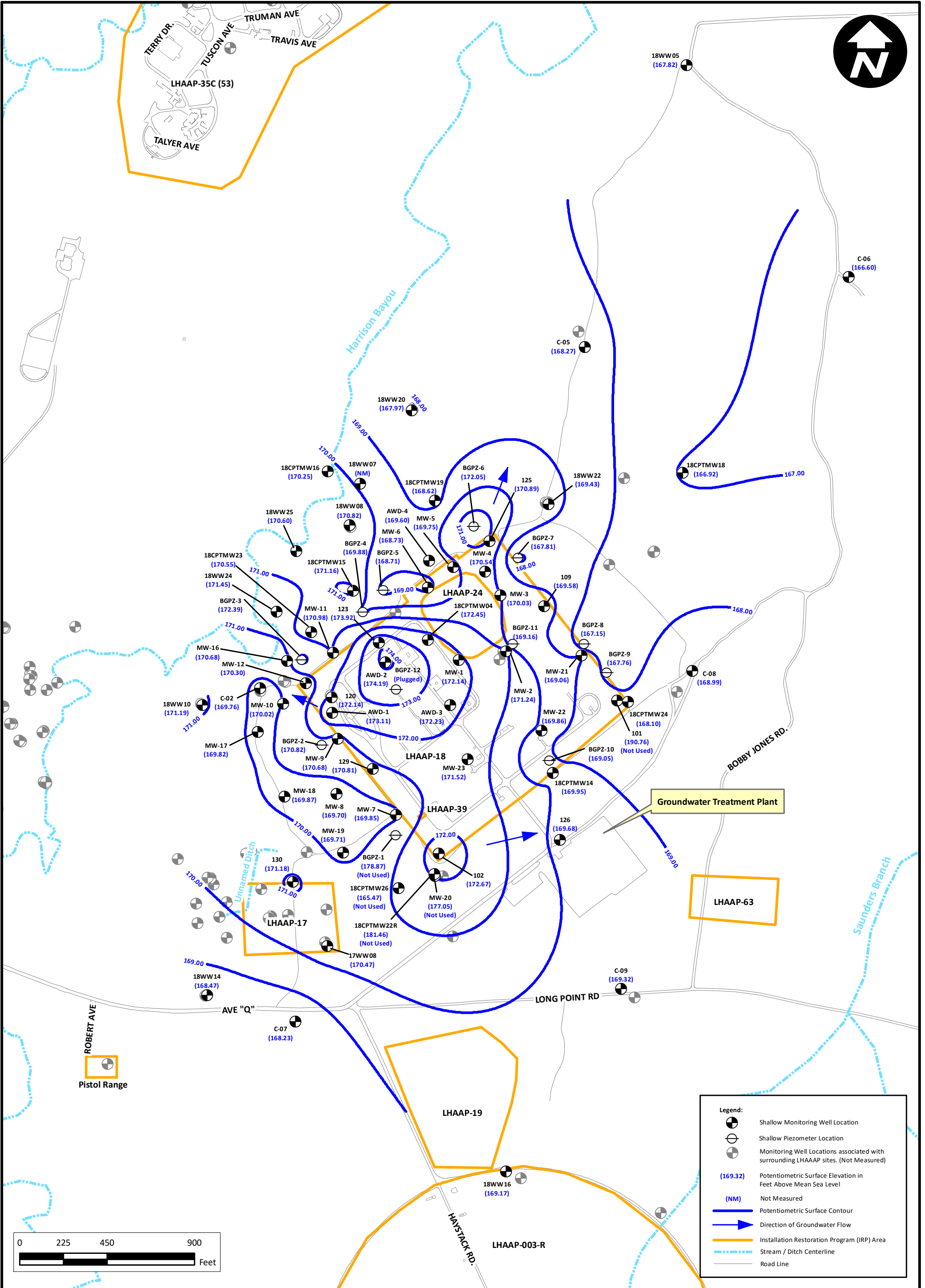
Well I.D.	Replaced Parts	Date	Contractor
ICT 14C	Installed New Pump and Electric Motor	7/29/2015	AECOM
ICT 14E	Installed New Pump and Electric Motor	7/31/2015	AECOM
ICT 12E	Repaired wiring	8/12/2015	AECOM
ICT 13E	Replaced high and low level probes	8/12/2015	AECOM
ICT 2	Installed Rebuilt Pump	9/9/2015	AECOM
ICT 12 E, ICT 14E	Replaced high level probe and wiring	9/15/2015	AECOM
ICT 13A	Installed new pump	12/23/2015	AECOM
ICT 13B	Installed new pump	12/24/2015	AECOM
ICT 13D	Replaced high level probe	2/11/2016	Aerotek
ICT 14C	Replaced low level probe on ICT 14C	2/15/2016	Aerotek
ICT 8	Installed new pump and electric motor	2/19/2016	Aerotek
ICT 14C	Repaired piping leak	3/10/2016	Aerotek
ICT 14E	Installed high and low level probes, level probe wire	3/22/2016	Aerotek
ICT 13D, ICT 14B	Installed high and low level probes, level probe wire	3/24/2016	Aerotek
ICT 14B	Installed new pump and electric motor	3/31/2016	Aerotek
ICT 14C	Installed new low level probe	4/20/2016	Aerotek
ICT 12B	Installed new mechanical flow meter	6/16/2016	Aerotek
ICT 13C	Installed rebuilt pump and new flow meter	8/10/2016	Aerotek
ICT 13A	Installed rebuilt pump, new flow meter, and new 1" unions	8/24/2016	Aerotek
ICT 14E	Installed new mechanical flow meter	8/26/2016	Aerotek
ICT 12C	Repaired flow meter	8/30/2016	Aerotek
ICT 2	Installed rebuilt pump and new flow meter	8/31/2016	Aerotek
ICT 14C	Cleaned and adjust level probes	9/7/2016	Aerotek
ICT 14C	Replaced level probes	9/12/2016	Aerotek
ICT 14C	Installed new level probe wire and level probes	9/21/2016	Aerotek
ICT 12C	Installed rebuilt pump, new electric motor, new wiring harness, new level probe wire, and new level probes	9/27/2016	Aerotek
ICT 14C	Cleaned and adjusted level probes	10/14/2016	Aerotek
ICT 13C	Cleaned and adjusted level probes	10/21/2016	Aerotek
ICT 13B	Installed rebuilt pump	10/25/2016	Aerotek
ICT 14D	Installed rebuilt pump	10/27/2016	Aerotek
ICT 13C	Replaced low level probe	11/8/2016	Aerotek
ICT 13B	Replaced relay base plate	11/8/2016	Aerotek
ICT 13E	Cleaned and adjusted low level probe	11/15/2016	Aerotek
ICT 13B	Replaced broken relay base plate and bad level probe wire	11/17/2016	Aerotek
ICT 13C	Cleaned & repaired leaking flow meter	11/18/2016	Aerotek
ICT 13B	Cleaned & adjusted low level probe	11/18/2016	Aerotek
ICT 13A, 13B, & 13E	Cleaned and adjusted low level probes	12/2/2016	Aerotek
ICT 13C & 14C	Pulled piping and pumps	2/8/2017	Aerotek
ICT 14C	Installed new electric motor	2/8/2017	Aerotek
ICT 13C & 14C	Installed rebuilt grundfos pumps	2/8/2017	Aerotek
ICT 7, 13A, & 14D	Repaired sample ports	2/9/2017	Aerotek
ICT 13B & 14E	Cleaned and adjusted low level probes	3/30/2017	Aerotek
ICT 13B & 13F	Installed new flow meters	3/30/2017	Aerotek
ICT 12B	Repaired flow meter	4/13/2017	Aerotek
ICT 12C	Replaced broken 1" tee	5/1/2017	Aerotek
ICT 11	Installed new manual flow meter	5/5/2017	Aerotek
ICT 2	Installed new flow meter	5/9/2017	Aerotek
ICT 14C & 14D	Cleaned and adjusted low level probes	5/31/2017	Aerotek
ICT 14C	Cleaned and adjusted low level probe	6/27/2017	Aerotek
ICT 8	Cleaned low level probe	7/11/2017	Aerotek
ICT 2 & 14D	Cleaned and replaced level probes	7/17/2017	Aerotek
ICT 14C	Cleaned low level probe	7/24/2017	Aerotek

Table B-1: Extraction Equipment Maintenance Since 2011

Well I.D.	Replaced Parts	Date	Contractor
ICT 13A	Installed new pump and flow meter	8/8/2017	Aerotek
ICT 13C & 13B	Installed new pump and flow meter	8/9/2017	Aerotek
ICT 13B	Installed new low level probe	8/10/2017	Aerotek
ICT 11	Installed new pump and flow meter	8/11/2017	Aerotek
ICT 4	Replaced low level probe	8/11/2017	Aerotek
ICT 2	Installed rebuilt pump	9/19/2017	Aerotek
ICT 13D	Adjusted level probes	9/22/2017	Aerotek
ICT 14C	Cleaned level probes	10/11/2017	Bhate
ICT 13E	Cleaned and adjusted level probes	10/24/2017	Bhate
ICT 12B	Adjusted level probes	11/15/2017	Bhate
ICT 14D	Cleaned level probes	11/15/2017	Bhate
ICT 8	Cleaned and adjusted level probes	11/28/2017	Bhate
ICT 2	Cleaned level probes	12/20/2017	Bhate
ICT 13C	Installed new flow meter	1/30/2018	Bhate
ICT 14C	Cleaned level probes	1/30/2018	Bhate
ICT 13C	Cleaned & adjusted level probes	2/1/2018	Bhate
ICT 13B	Repaired broken 1" union	2/1/2018	Bhate
ICT 14A	Repaired 2 broken 1" elbows & lower high level probe	2/1/2018	Bhate
ICT 14B	Repaired broken 1" tee & lower high level probe	2/1/2018	Bhate
ICT 14D	Installed new flow meter	2/8/2018	Bhate
ICT 8	Replaced broken 1" tee & cleaned level probes	2/8/2018	Bhate
ICT 14D	Cleaned level probes	2/28/2018	Bhate
ICT 14C	Replaced low level probe	2/28/2018	Bhate
ICT 13B	Cleaned level probes	2/28/2018	Bhate
ICT 13A	Installed new motor & replaced leaking 1" union	3/14/2018	Bhate
ICT 13C	Cleaned & adjusted level probes	3/21/2018	Bhate
ICT 12B	Replaced broken 1" elbow & installed new flow meter	3/21/2018	Bhate
ICT 2	Installed new pump	3/22/2018	Bhate
ICT 13B	Replaced level probes	3/30/2018	Bhate
ICT 14E	Lower high level probe	3/30/2018	Bhate
ICT 14C	Cleaned level probes	4/27/2018	Bhate
ICT 11	Installed new breaker	4/27/2018	Bhate
ICT 14E	Cleaned level probes	6/7/2018	Bhate
ICT 12C	Cleaned level probes	6/7/2018	Bhate
EW 01	Cleaned level probes	6/7/2018	Bhate
ICT 14E	Replaced level probes	6/8/2018	Bhate
ICT 11	Installed new electrical wire from breaker to well	6/14/2018	Bhate
ICT 12B	Replaced pump	6/25/2018	Bhate
ICT 14E	Cleaned level probes	6/26/2018	Bhate
ICT 8	Cleaned level probes	6/26/2018	Bhate
ICT 14C	Replaced pump	6/27/2018	Bhate
EW01	Replaced level probes	9/12/2018	Bhate
ICT 7	Cleaned level probes	9/12/2018	Bhate
ICT 12C	Replaced low level probe wire and probe	9/12/2018	Bhate
ICT 14D	Replaced high level probe wire & probe	9/13/2018	Bhate
ICT 12B	Replaced high and low level probe wires & probes	9/13/2018	Bhate
ICT 12B	Replaced electrical relay and relay base plate	9/13/2018	Bhate
ICT 14C	Cleaned level probes	9/13/2018	Bhate
ICT 13E	Replaced flow meter	9/14/2018	Bhate
ICT 14D	Replaced low level probe	10/31/2018	Bhate
ICT 8	Cleaned level probes	10/31/2018	Bhate
ICT 14C	Cleaned level probes	10/31/2018	Bhate
ICT 13A	Replaced broken 1" Tee	12/12/2018	Bhate
ICT 14C	Cleaned level probes	12/12/2018	Bhate

Table B-1: Extraction Equipment Maintenance Since 2011

Well I.D.	Replaced Parts	Date	Contractor
ICT 13A	Installed new flow meter	2/14/2019	Bhate
ICT 13E	Cleaned probes	2/25/2019	Bhate
18WW17	Cleaned probes	2/26/2019	Bhate
ICT 12D	Replaced pump & motor	3/6/2019	Bhate
ICT 13C	Replaced pump	3/8/2019	Bhate
ICT 2	Replaced pump & flow meter	3/11/2019	Bhate
ICT 14C	Replaced pump & motor	3/12/2019	Bhate
ICT 14D	Replaced pump	3/12/2019	Bhate
ICT 2	Replaced leaking 1" union	3/12/2019	Bhate
ICT 14E	Replaced pump, seal plate & level probes	3/14/2019	Bhate
ICT 14A	Replaced pump & motor and repaired leaking 1" pipe	3/15/2019	Bhate
18WW17	Replaced level prbes and level probe wire	3/18/2019	Bhate
ICT 13C	Lower high level probe	3/22/2019	Bhate
ICT 14A	Lower high level probe	3/22/2019	Bhate
EW-01	Replaced pump	3/29/2019	Bhate
LHAAP-16			
Well I.D.	Replaced Parts	Date	Contractor
EW08	New pump	2/28/2011	Shaw
EW01	Rebuild pump	8/25/2011	Shaw
EW06	Rebuild pump	8/25/2011	Shaw
EW02	Rebuild pump	2/12/2012	Shaw
EW03	Rebuild pump	2/12/2012	Shaw
EW08	Rebuild pump	11/8/2012	AECOM
EW01	Rebuild pump	11/8/2012	AECOM
EW04	Repaired pump	11/13/2012	AECOM
EW07	Rebuild pump	11/13/2012	AECOM
EW04	Installed New Pump	11/28/2012	AECOM
EW06	Installed New Pump	11/28/2012	AECOM
EW02	Installed New Pump	12/4/2012	AECOM
EW03	Installed New Pump	12/4/2012	AECOM
EW01	Installed New Pump	12/17/2012	AECOM
EW01	Replaced Low level probe	1/17/2015	AECOM
EW01	Cleaned and adjusted level probes	10/21/2016	Aerotek



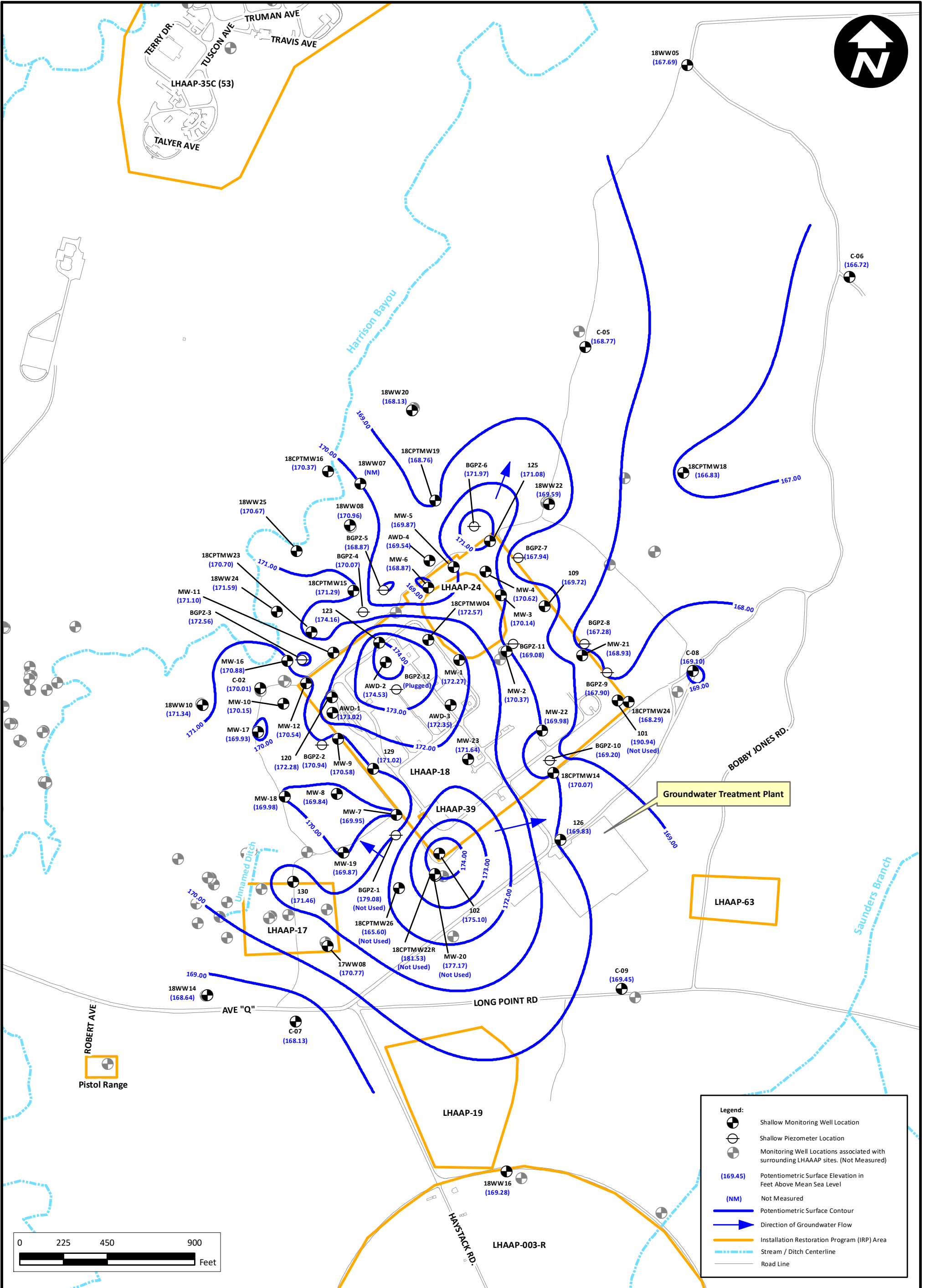
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 Groundwater Treatment Plant
 Longhorn Army Ammunition Plant, Karnack, Texas

Groundwater Potentiometric Surface Map
 Shallow Zone (January 31, 2019) LHAAP-18/24



PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	5/31/2019	MRM

Figure B-1



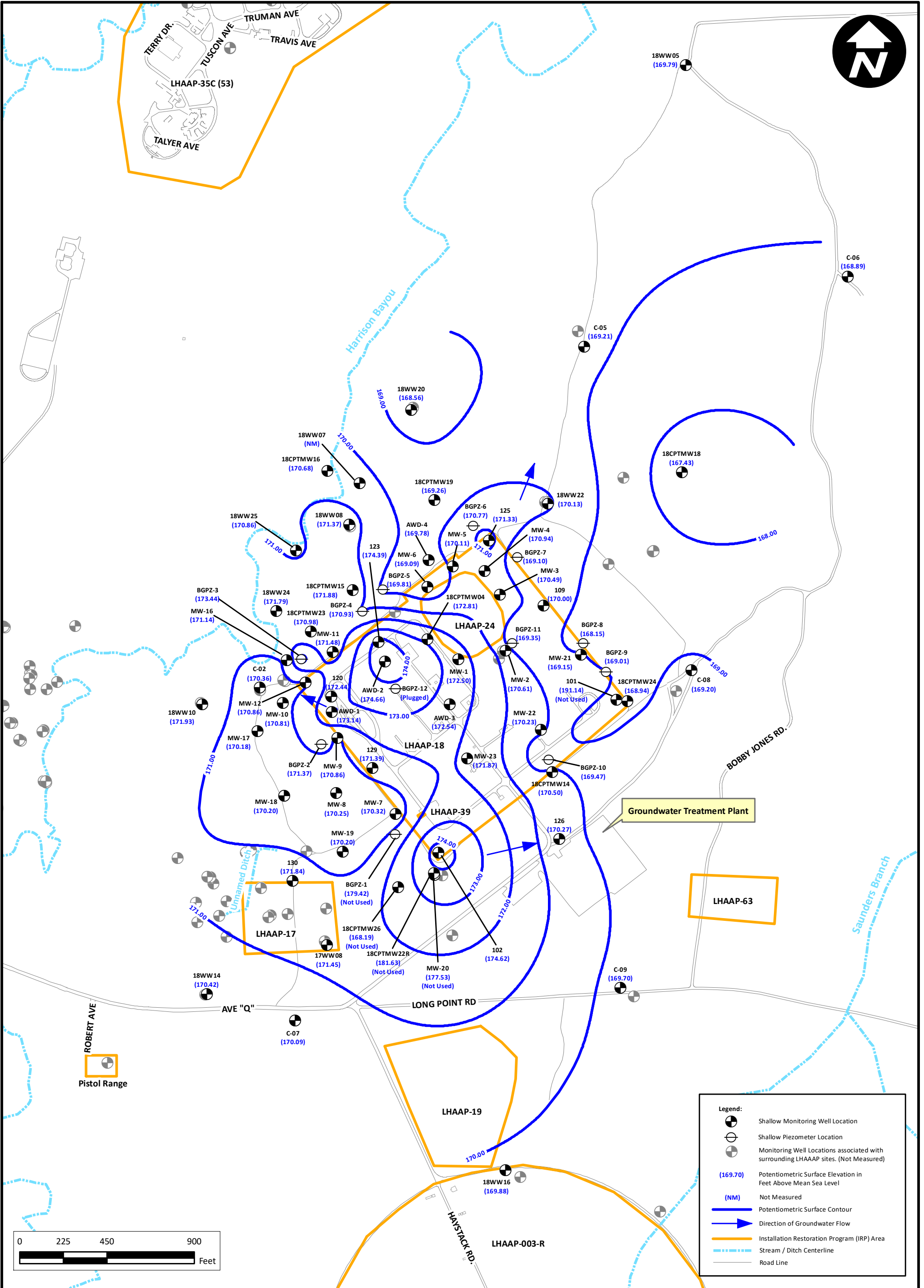
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Groundwater Potentiometric Surface Map
 Shallow Zone (February 28, 2019) LHAAP-18/24



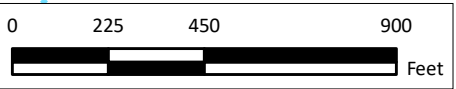
PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	5/31/2019	MRM

Figure B-2



Legend:

- Shallow Monitoring Well Location
- Shallow Piezometer Location
- Monitoring Well Locations associated with surrounding LHAAP sites. (Not Measured)
- (169.70)** Potentiometric Surface Elevation in Feet Above Mean Sea Level
- (NM)** Not Measured
- Potentiometric Surface Contour
- Direction of Groundwater Flow
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Road Line

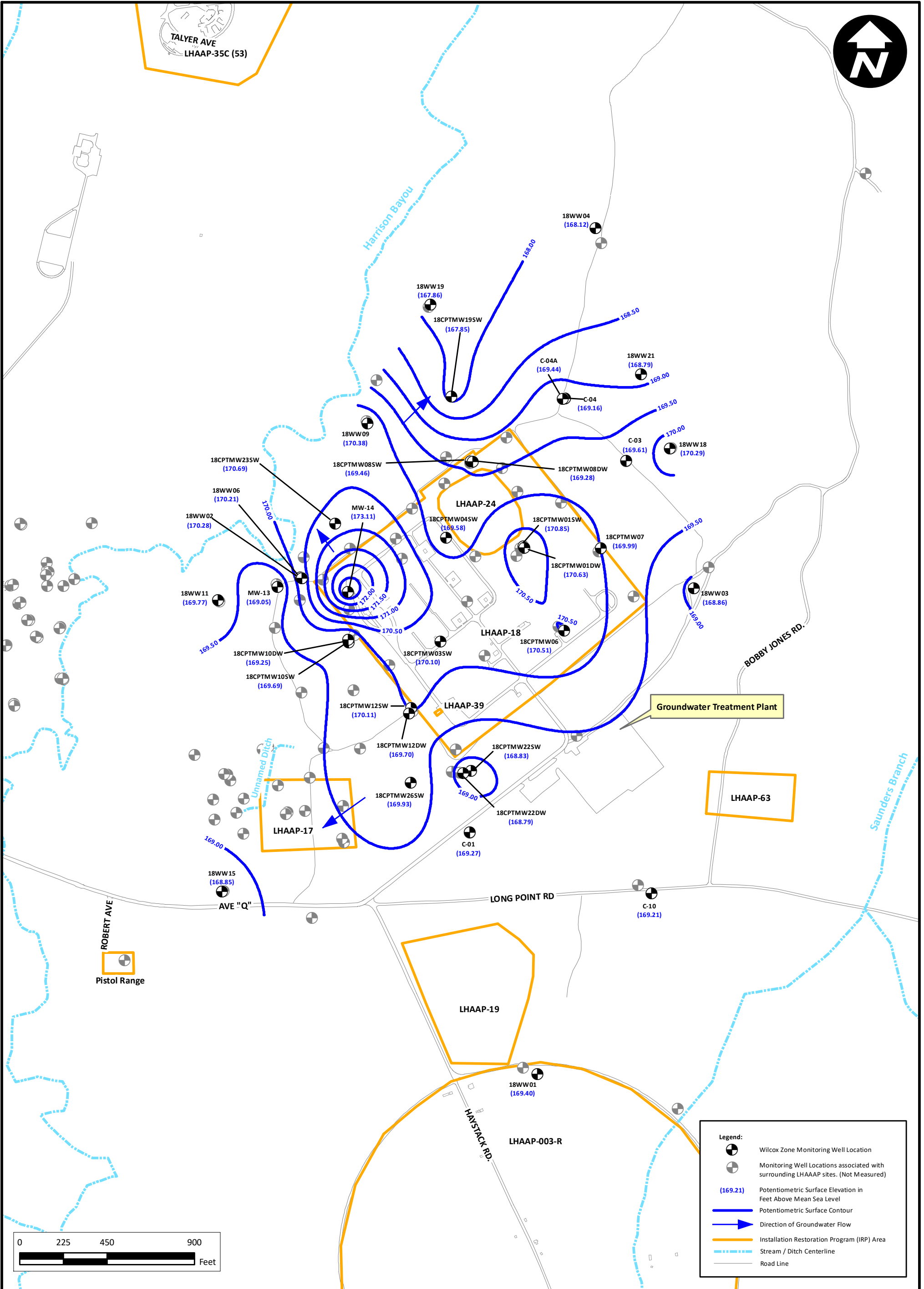


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**Groundwater Potentiometric Surface Map
 Shallow Zone (March 26-27, 2019) LHAAP-18/24**

Figure B-3



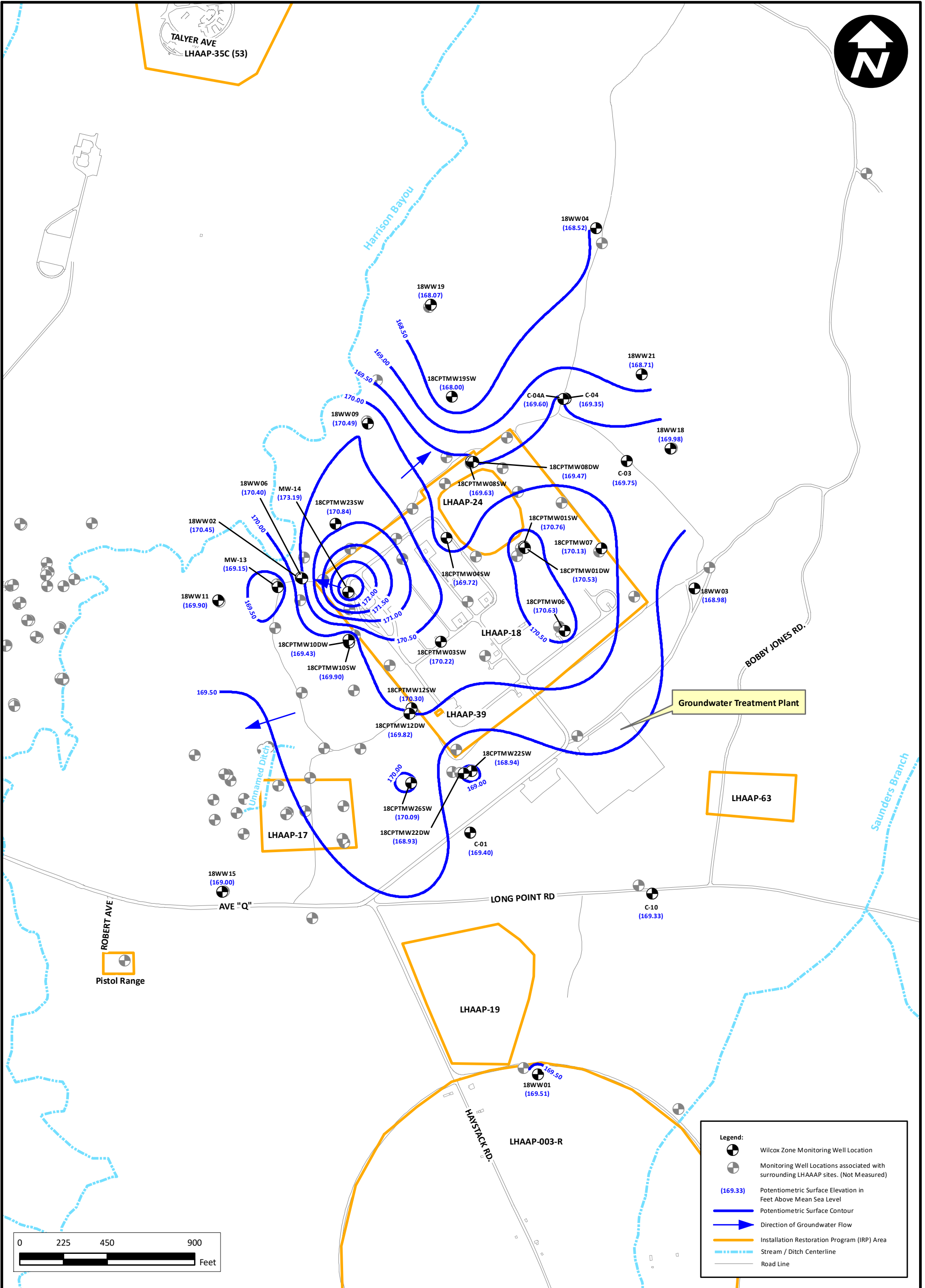
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Groundwater Potentiometric Surface Map
 Wilcox Zone (January 31, 2019) LHAAP-18/24



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NWO1312.0150	As Shown	4/24/2019	MRM

Figure B-4

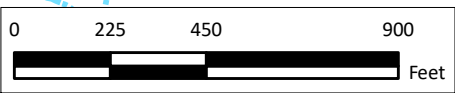
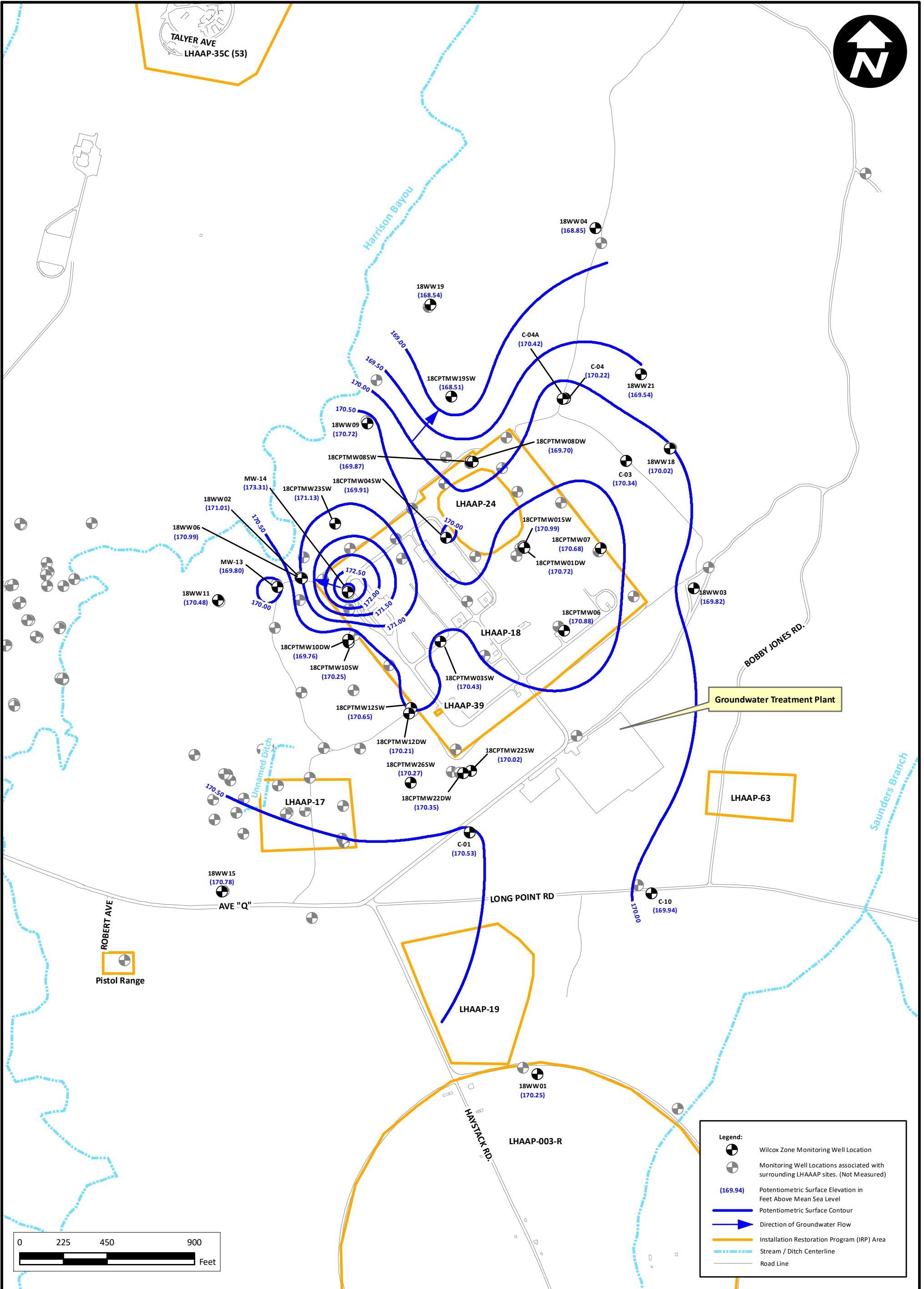


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Groundwater Potentiometric Surface Map
 Wilcox Zone (February 28, 2019) LHAAP-18/24

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	4/24/2019	MRM

Figure B-5

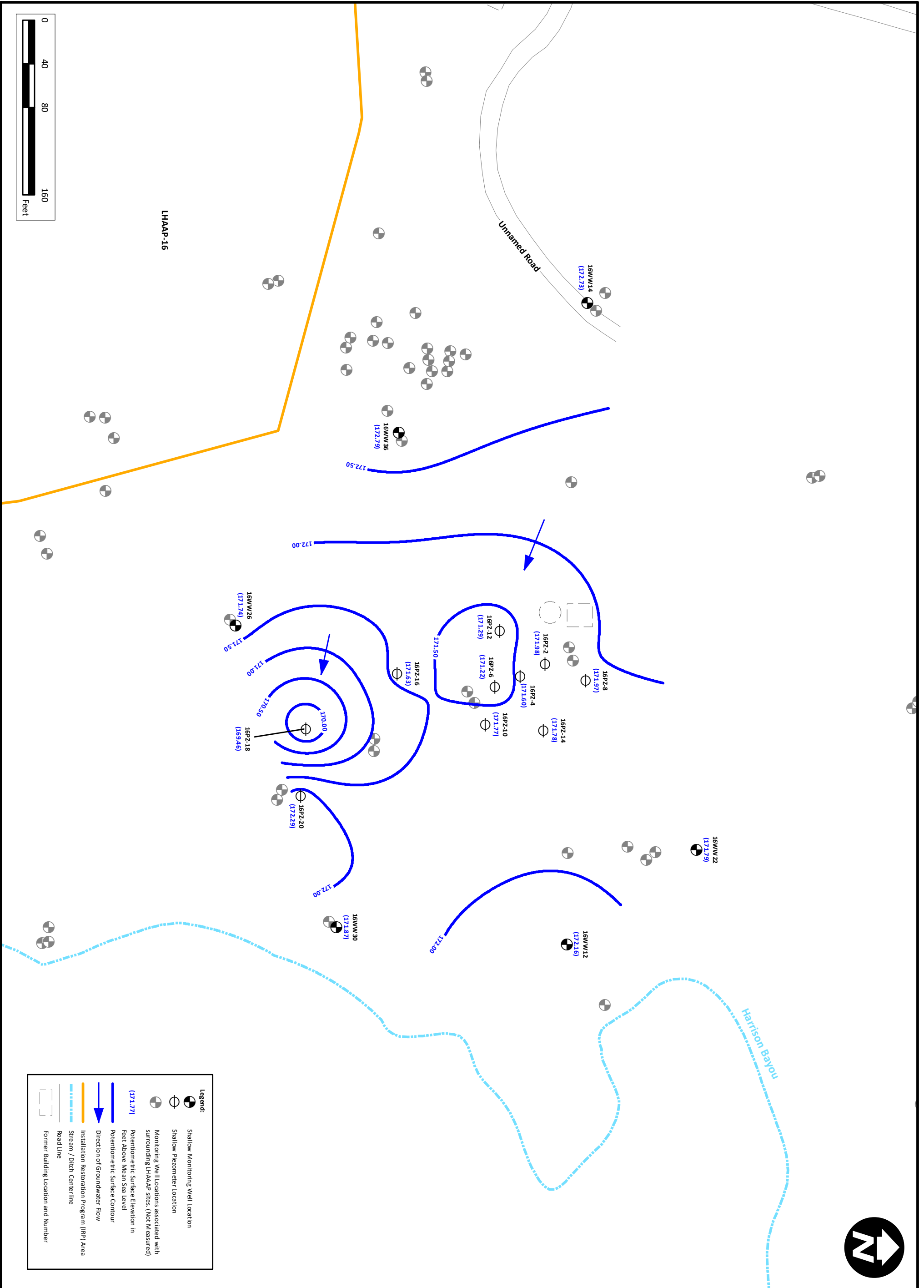


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Groundwater Potentiometric Surface Map
 Wilcox Zone (March 26-27, 2019) LHAAP-18/24

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	4/24/2019	MRM

Figure B-6



Legend:

- Shallow Monitoring Well Location
- Shallow Piezometer Location
- Monitoring Well Locations associated with surrounding LHAAP sites. (Not Measured)
- Potentiometric Surface Elevation in Feet Above Mean Sea Level
- Direction of Groundwater Flow
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Road Line
- Former Building Location and Number

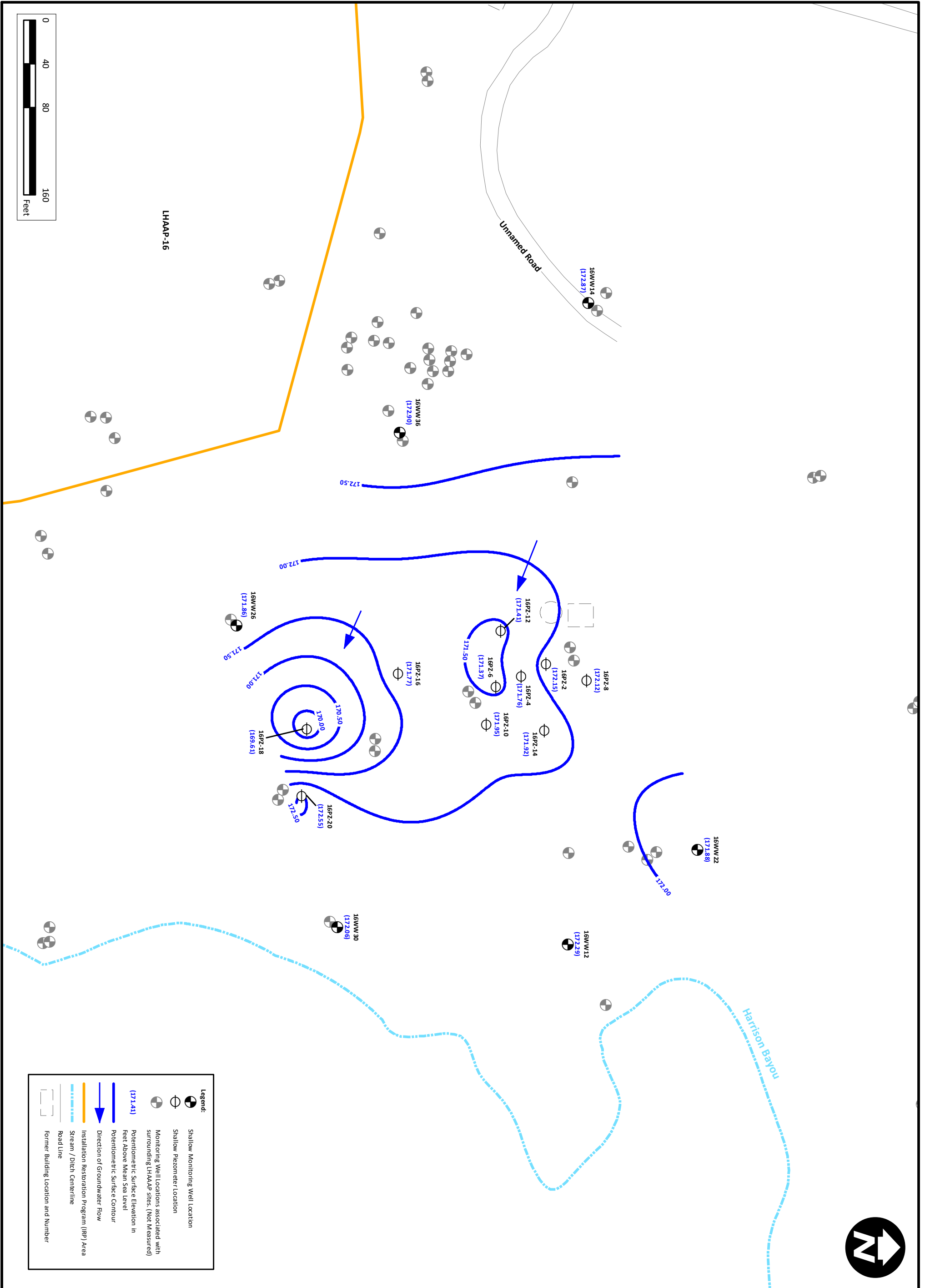


Quarterly Evaluation Report 1st Quarter (January – March) 2019
 Groundwater Treatment Plant
 Longhorn Army Ammunition Plant, Karnack, Texas

Groundwater Potentiometric Surface Map
 Shallow Zone (January 30, 2019) LHAAP-16

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NW01312.0150	As Shown	4/25/2019	MRM

Figure B-7



Legend:

- Shallow Monitoring Well Location
- Shallow Piezometer Location
- Monitoring Well Locations associated with surrounding LHAAP sites. (Not Measured)
- Potentiometric Surface Elevation in Feet Above Mean Sea Level
- Potentiometric Surface Contour
- Direction of Groundwater Flow
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Roadline
- Former Building Location and Number

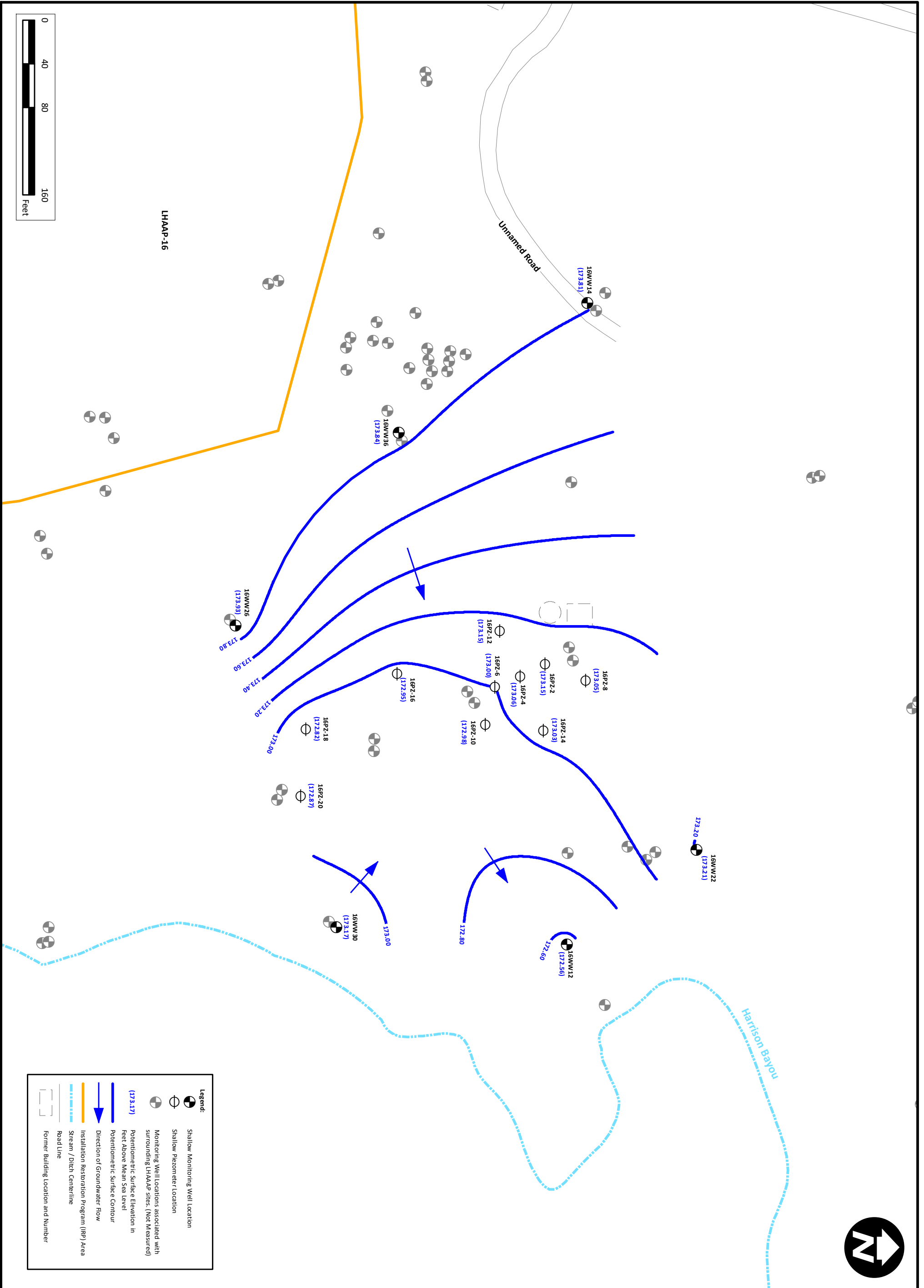


Quarterly Evaluation Report 1st Quarter (January – March) 2019
 Groundwater Treatment Plant
 Longhorn Army Ammunition Plant, Karnack, Texas

Groundwater Potentiometric Surface Map
 Shallow Zone (February 27, 2019) LHAAP-16

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	4/25/2019	MRM

Figure B-8



Legend:

- Shallow Monitoring Well Location
- Shallow Piezometer Location
- Monitoring Well Locations associated with surrounding LHAAP sites. (Not Measured)
- Potentiometric Surface Elevation in Feet Above Mean Sea Level
- Direction of Groundwater Flow
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Road Line
- Former Building Location and Number

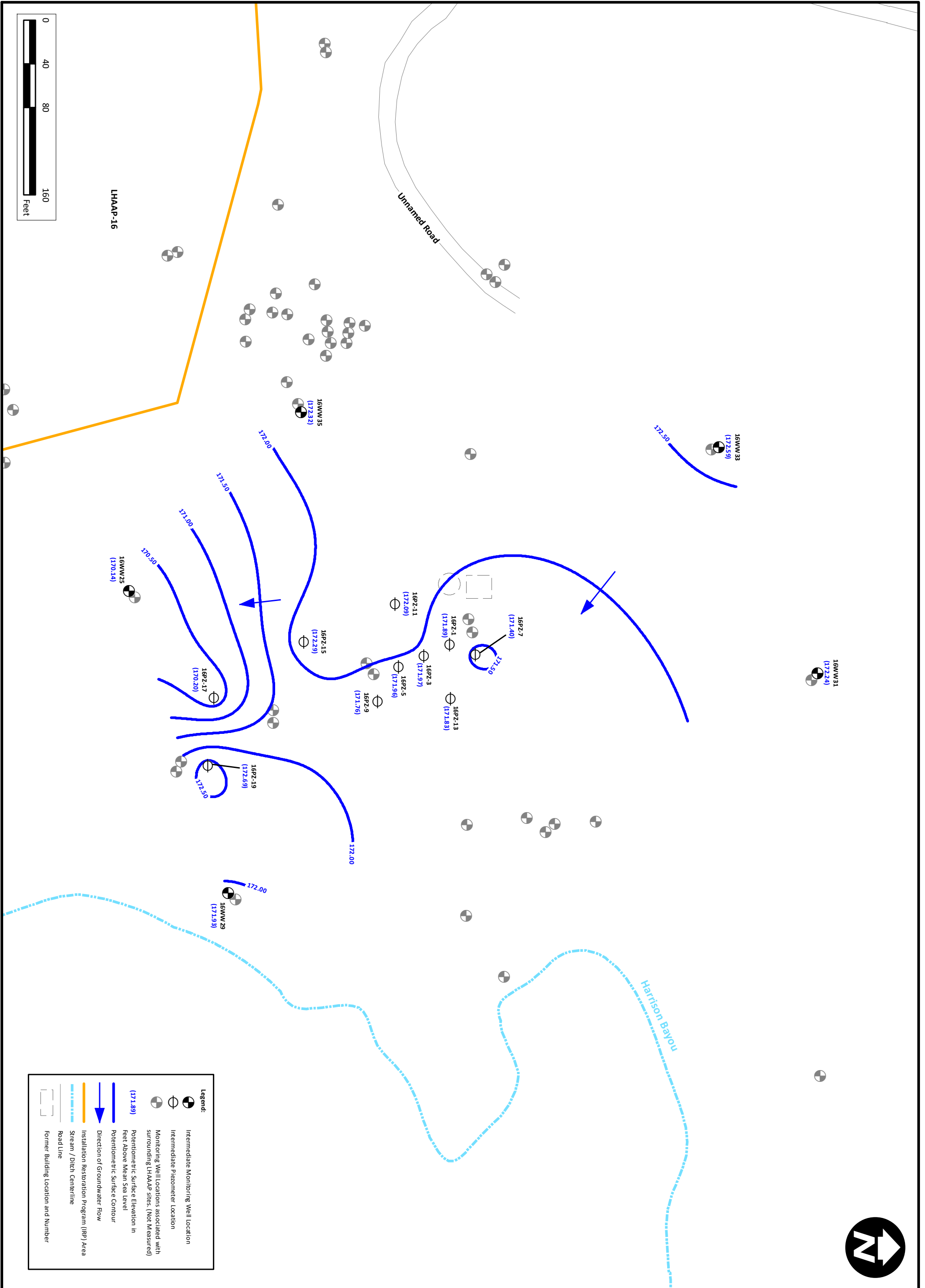


Quarterly Evaluation Report 1st Quarter (January – March) 2019
 Groundwater Treatment Plant
 Longhorn Army Ammunition Plant, Karnack, Texas

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	4/25/2019	MRM

**Groundwater Potentiometric Surface Map
 Shallow Zone (March 26, 2019) LHAAP-16**

Figure B-9



Legend:

- Intermediate Monitoring Well Location
- Intermediate Piezometer Location
- Monitoring Well Locations associated with surrounding LHAAP sites. (Not Measured)
- Potentiometric Surface Elevation in Feet Above Mean Sea Level
- Direction of Groundwater Flow
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Road Line
- Former Building Location and Number

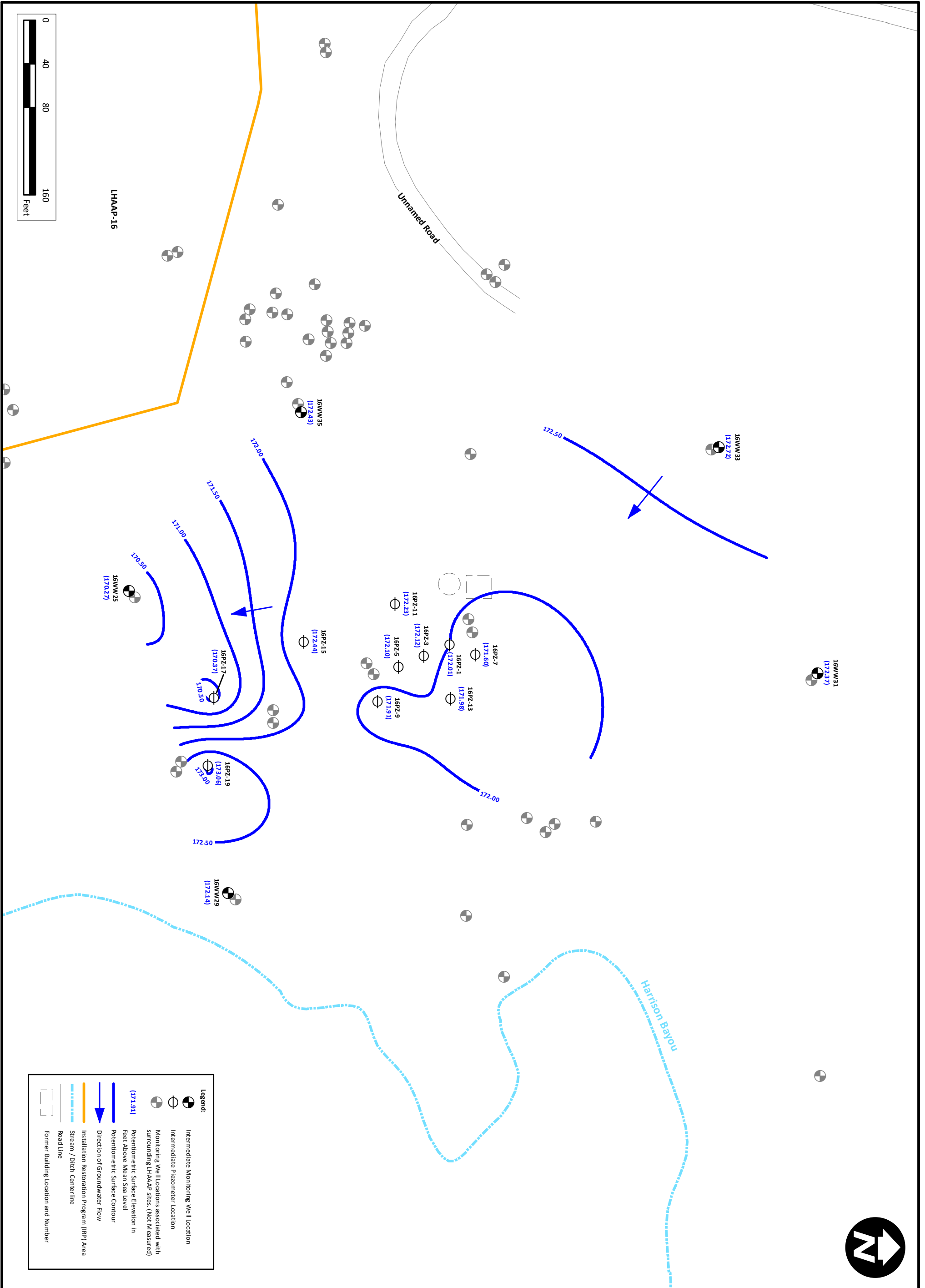


Quarterly Evaluation Report 1st Quarter (January – March) 2019
 Groundwater Treatment Plant
 Longhorn Army Ammunition Plant, Karnack, Texas

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	4/25/2019	MRM

**Groundwater Potentiometric Surface Map
 Intermediate Zone (January 30, 2019) LHAAP-16**

Figure B-10

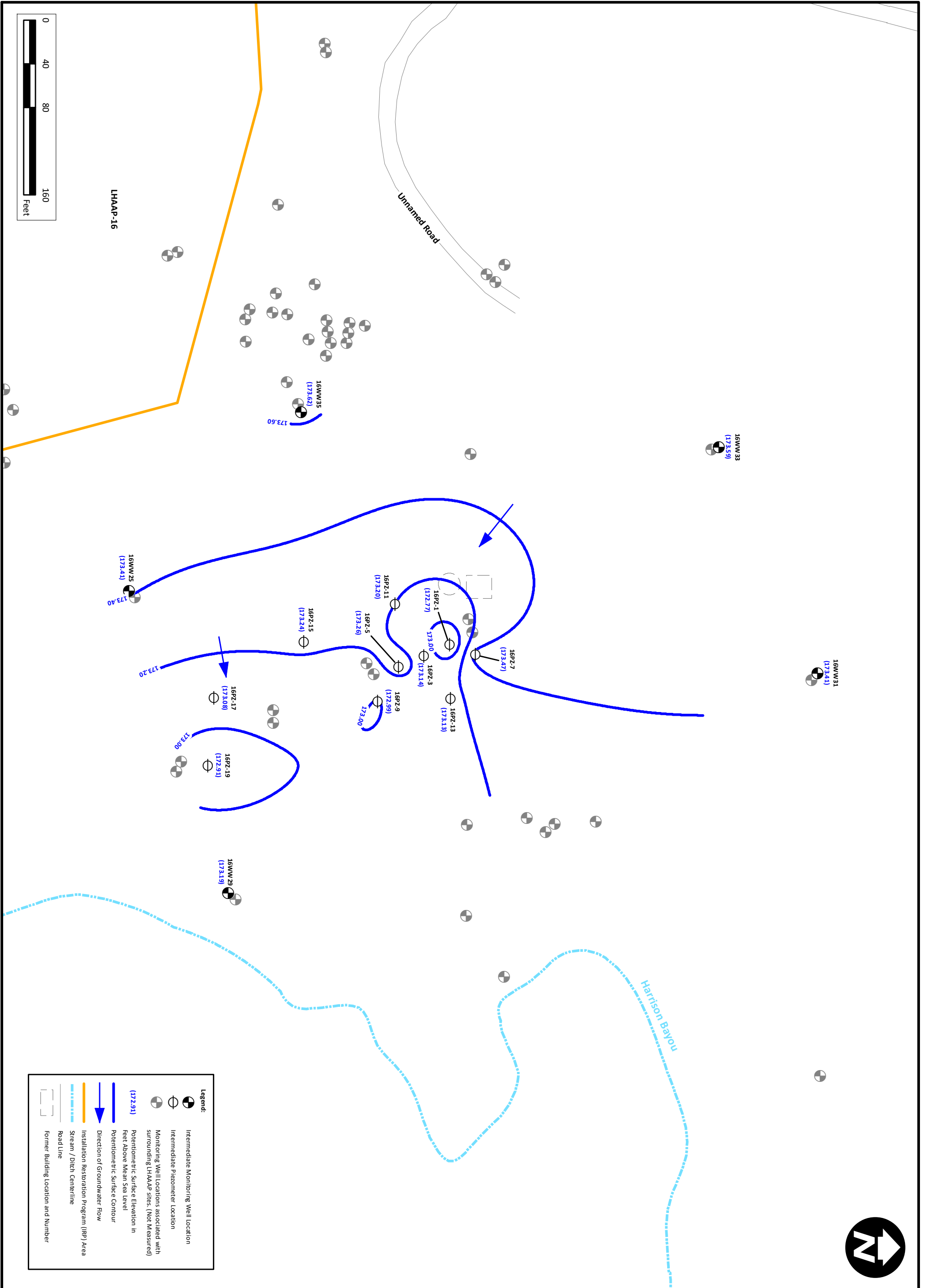


Quarterly Evaluation Report 1st Quarter (January – March) 2019
 Groundwater Treatment Plant
 Longhorn Army Ammunition Plant, Karnack, Texas

Groundwater Potentiometric Surface Map
 Intermediate Zone (February 27, 2019) LHAAP-16

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	4/25/2019	MRM

Figure B-11



Legend:

- Intermediate Monitoring Well Location
- Intermediate Piezometer Location
- Monitoring Well Locations associated with surrounding LHAAP sites. (Not Measured)
- Potentiometric Surface Elevation in Feet Above Mean Sea Level
- Direction of Groundwater Flow
- Installation Restoration Program (IRP) Area
- Stream / Ditch Centerline
- Road Line
- Former Building Location and Number



Quarterly Evaluation Report 1st Quarter (January – March) 2019
 Groundwater Treatment Plant
 Longhorn Army Ammunition Plant, Karnack, Texas

Groundwater Potentiometric Surface Map
 Intermediate Zone (March 26, 2019) LHAAP-16

PROJECT NO:	SCALE:	DATE:	DRAWN BY:
NWO1312.0150	As Shown	4/25/2019	MRM

Figure B-12

GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

APPENDIX C
GWTP WATER SAMPLING LABORATORY ANALYTICAL RESULTS
(PROVIDED ON CD ONLY)

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 14, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19021158**

Laboratory Results for: **Longhorn GW Treatment Plant**

Dear Marcia,

ALS Environmental received 2 sample(s) on Feb 22, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: DAYNA.FISHER
RJ Modashia
Project Manager

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
Work Order: HS19021158

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19021158-01	LH18/24-SP650_022119	Water		21-Feb-2019 09:06	22-Feb-2019 09:06	<input type="checkbox"/>
HS19021158-02	LH18/24-SP650_022119_BIX	Water		21-Feb-2019 14:00	22-Feb-2019 09:06	<input type="checkbox"/>

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.**CASE NARRATIVE****Project:** Longhorn GW Treatment Plant**Work Order:**

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
- The analysis for TOC was subcontracted to ALS Environmental in Kelso, WA. Final Report attached.

WetChemistry by Method E365.3**Batch ID: R333717**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

WetChemistry by Method E350.3**Batch ID: R333409**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: LH18/24-SP650_022119
 Collection Date: 21-Feb-2019 09:06

ANALYTICAL REPORT

WorkOrder:HS19021158
 Lab ID:HS19021158-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
AMMONIA AS N BY E350.3(ISE)								Analyst: KVL
Nitrogen, Ammonia (As N)	15		2.0	2.0	2.0	mg/L	10	25-Feb-2019 10:20
ORTHO PHOSPHATE (PO4) AS P BY E365.3								Analyst: MZD
Phosphorus, Total Orthophosphate (As P)	4.20		0.100	0.250	0.250	mg/L	10	22-Feb-2019 14:30
SUBCONTRACT ANALYSIS - TOC ANALYSIS								Analyst: SUBK
Subcontract Analysis	See Attached		0	0		NA	1	14-Mar-2019 14:02

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: LH18/24-SP650_022119_BIX
 Collection Date: 21-Feb-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19021158
 Lab ID:HS19021158-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021158

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R333409	Test Name : AMMONIA AS N BY E350.3(ISE)		Matrix: Water			
HS19021158-01	LH18/24-SP650_022119	21 Feb 2019 09:06			25 Feb 2019 10:20	10
Batch ID R333717	Test Name : ORTHO PHOSPHATE (PO4) AS P BY E365.3		Matrix: Water			
HS19021158-01	LH18/24-SP650_022119	21 Feb 2019 09:06			22 Feb 2019 14:30	10
Batch ID R334176	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19021158-02	LH18/24-SP650_022119_BIX	21 Feb 2019 14:00			07 Mar 2019 17:47	1
Batch ID R334558	Test Name : SUBCONTRACT ANALYSIS - TOC ANALYSIS		Matrix: Water			
HS19021158-01	LH18/24-SP650_022119	21 Feb 2019 09:06			14 Mar 2019 14:02	1

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021158

QC BATCH REPORT

Batch ID:	R333409	Instrument:	WetChem_HS	Method:	E350.3					
MBLK	Sample ID: MBLK-R333409	Units: mg/L	Analysis Date: 25-Feb-2019 10:20							
Client ID:	Run ID: WetChem_HS_333409	SeqNo: 4962077	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	0.20	0.20								U
LCS	Sample ID: LCS-R333409	Units: mg/L	Analysis Date: 25-Feb-2019 10:20							
Client ID:	Run ID: WetChem_HS_333409	SeqNo: 4962076	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	9.117	0.20	10	0	91.2	80 - 120				
MS	Sample ID: HS19021211-02MS	Units: mg/L	Analysis Date: 25-Feb-2019 10:20							
Client ID:	Run ID: WetChem_HS_333409	SeqNo: 4962079	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	9.246	0.20	10	0.3054	89.4	80 - 120				
MSD	Sample ID: HS19021211-02MSD	Units: mg/L	Analysis Date: 25-Feb-2019 10:20							
Client ID:	Run ID: WetChem_HS_333409	SeqNo: 4962078	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	9.272	0.20	10	0.3054	89.7	80 - 120	9.246	0.281	20	

The following samples were analyzed in this batch: HS19021158-01

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021158

QC BATCH REPORT

Batch ID: R333717		Instrument: UV-2450		Method: E365.3						
MBLK	Sample ID: MBLK-333717	Units: mg/L		Analysis Date: 22-Feb-2019 14:30						
Client ID:	Run ID: UV-2450_333717	SeqNo: 4969292		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.0250	0.0250							U	
LCS	Sample ID: LCS-333717	Units: mg/L		Analysis Date: 22-Feb-2019 14:30						
Client ID:	Run ID: UV-2450_333717	SeqNo: 4969293		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.222	0.0250	0.25	0	88.8	85 - 115				
MS	Sample ID: HS19021157-03MS	Units: mg/L		Analysis Date: 22-Feb-2019 14:30						
Client ID:	Run ID: UV-2450_333717	SeqNo: 4969296		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.23	0.0250	0.25	0.009	88.4	80 - 120				
MSD	Sample ID: HS19021157-03MSD	Units: mg/L		Analysis Date: 22-Feb-2019 14:30						
Client ID:	Run ID: UV-2450_333717	SeqNo: 4969297		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.239	0.0250	0.25	0.009	92.0	80 - 120	0.23	3.84	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 14-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021158

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19021158

Date/Time Received: **22-Feb-2019 09:06**
 Received by: **RPG**

Checklist completed by: Raegen Giga 22-Feb-2019
 eSignature Date

Reviewed by: RJ Modashia 22-Feb-2019
 eSignature Date

Matrices: **water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.6c/1.6c uc/c IR 11
 Cooler(s)/Kit(s): 24406
 Date/Time sample(s) sent to storage: 02/22/2019 11:59

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:



Contacted By: Regarding:



Comments:

Corrective Action:

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd, Suite 210 Houston, TX, 77099 (281) 530-5656 ATTN: R.J Modshia

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001			Analyses										HS19021158 Bhate Environmental Associates, Inc. Longhorn GW Treatment Plant 									
Job: GROUNDWATER TREATMENT PLANT WEEKLY SAMPLES						MS / MSD	No. OF CONTAINERS	AMMONIA-N	TOTAL ORGANIC CARBON	ORTHO-PHOSPHATE	PERCHLORATE											Remarks (Preservatives, etc.)	Lab I.D.#		
Prepared By: Scott Beesinger			P.O. Number																						
Field Sample I.D.		Sample Matrix		Date / Time																					
LH18/24-SP650_022119		Water		02/21/19 / 14:00		2		X		X						H2SO4									
LH18/24-SP650_022119		Water		02/21/19 / 14:00		1						X				NONE									
LH18/24-SP650_022119_BIX		Water		02/21/19 / 14:00		1						X				NONE									
Additional Remarks: Standard TAT on all parameters																									
Relinquished By: 		Date 02/21/19		Time 14:30		Received By: R. C. G...		Date 2/21/19		Time 29:06		Relinquished By:		Date		Time		Received By:		Date		Time			
For Lab Use Only																									
Received At Lab By:				Date		Time		Airbill No.				Opened By:				Date		Time		Temp of Container		Seal No.		Condition	
Remarks: cooler 24406 Temp 1-6c 12 11 CFO																									

	<p>ALS 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887</p>	<p align="center">CUSTODY SEAL</p> <p>Date: <u>2/21/99</u> Time: <u>1430</u> Name: <u>Scott B. King</u> Company: <u>BIBAC</u></p>	<p>Shipped By:  <u>2/21/99</u></p>
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TRK# 4380 9530 9467

FedEx.
 TRK# 4380 9530 9467

AB SGRA

**RETURNS MON - SAT
 PRIORITY OVERNIGHT**

**FRI - 22 FEB 10:30A
 PRIORITY OVERNIGHT**

**77099
 TX-US
 IAH**



ALS Environmental
ALS Group USA, Corp
1317 South 13th Avenue
Kelso, WA 98626
T : +1 360 577 7222
F : +1 360 636 1068
www.alsglobal.com

March 13, 2019

Analytical Report for Service Request No: K1901637

RJ Modashia
ALS Laboratory Group
10450 Stancliff Road
Suite 210
Houston, TX 77099-4338

RE: HS19021158 / HS19021158

Dear RJ,

Enclosed are the results of the sample(s) submitted to our laboratory February 23, 2019
For your reference, these analyses have been assigned our service request number **K1901637**.

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. The test results meet requirements of the current NELAP standards, where applicable, and except as noted in the laboratory case narrative provided. For a specific list of NELAP-accredited analytes, refer to the certifications section at www.alsglobal.com. All results are intended to be considered in their entirety, and ALS Group USA Corp. dba ALS Environmental (ALS) is not responsible for use of less than the complete report. Results apply only to the items submitted to the laboratory for analysis and individual items (samples) analyzed, as listed in the report.

Please contact me if you have any questions. My extension is 3350. You may also contact me via email at Kelley.Lovejoy@alsglobal.com.

Respectfully submitted,

ALS Group USA, Corp. dba ALS Environmental

Kelley Lovejoy
Project Manager



ALS Environmental
ALS Group USA, Corp
1317 South 13th Avenue
Kelso, WA 98626
T : +1 360 577 7222
F : +1 360 636 1068
www.alsglobal.com

Table of Contents

Acronyms

Qualifiers

State Certifications, Accreditations, And Licenses

Case Narrative

Chain of Custody

General Chemistry

Raw Data

 General Chemistry

Acronyms

ASTM	American Society for Testing and Materials
A2LA	American Association for Laboratory Accreditation
CARB	California Air Resources Board
CAS Number	Chemical Abstract Service registry Number
CFC	Chlorofluorocarbon
CFU	Colony-Forming Unit
DEC	Department of Environmental Conservation
DEQ	Department of Environmental Quality
DHS	Department of Health Services
DOE	Department of Ecology
DOH	Department of Health
EPA	U. S. Environmental Protection Agency
ELAP	Environmental Laboratory Accreditation Program
GC	Gas Chromatography
GC/MS	Gas Chromatography/Mass Spectrometry
LOD	Limit of Detection
LOQ	Limit of Quantitation
LUFT	Leaking Underground Fuel Tank
M	Modified
MCL	Maximum Contaminant Level is the highest permissible concentration of a substance allowed in drinking water as established by the USEPA.
MDL	Method Detection Limit
MPN	Most Probable Number
MRL	Method Reporting Limit
NA	Not Applicable
NC	Not Calculated
NCASI	National Council of the Paper Industry for Air and Stream Improvement
ND	Not Detected
NIOSH	National Institute for Occupational Safety and Health
PQL	Practical Quantitation Limit
RCRA	Resource Conservation and Recovery Act
SIM	Selected Ion Monitoring
TPH	Total Petroleum Hydrocarbons
tr	Trace level is the concentration of an analyte that is less than the PQL but greater than or equal to the MDL.

Inorganic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- E The result is an estimate amount because the value exceeded the instrument calibration range.
- J The result is an estimated value.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.
- H The holding time for this test is immediately following sample collection. The samples were analyzed as soon as possible after receipt by the laboratory.

Metals Data Qualifiers

- # The control limit criteria is not applicable. See case narrative.
- J The result is an estimated value.
- E The percent difference for the serial dilution was greater than 10%, indicating a possible matrix interference in the sample.
- M The duplicate injection precision was not met.
- N The Matrix Spike sample recovery is not within control limits. See case narrative.
- S The reported value was determined by the Method of Standard Additions (MSA).
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- W The post-digestion spike for furnace AA analysis is out of control limits, while sample absorbance is less than 50% of spike absorbance.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- + The correlation coefficient for the MSA is less than 0.995.
- Q See case narrative. One or more quality control criteria was outside the limits.

Organic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- A A tentatively identified compound, a suspected aldol-condensation product.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- C The analyte was qualitatively confirmed using GC/MS techniques, pattern recognition, or by comparing to historical data.
- D The reported result is from a dilution.
- E The result is an estimated value.
- J The result is an estimated value.
- N The result is presumptive. The analyte was tentatively identified, but a confirmation analysis was not performed.
- P The GC or HPLC confirmation criteria was exceeded. The relative percent difference is greater than 40% between the two analytical results.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a chromatographic interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.

Additional Petroleum Hydrocarbon Specific Qualifiers

- F The chromatographic fingerprint of the sample matches the elution pattern of the calibration standard.
- L The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of lighter molecular weight constituents than the calibration standard.
- H The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of heavier molecular weight constituents than the calibration standard.
- O The chromatographic fingerprint of the sample resembles an oil, but does not match the calibration standard.
- Y The chromatographic fingerprint of the sample resembles a petroleum product eluting in approximately the correct carbon range, but the elution pattern does not match the calibration standard.
- Z The chromatographic fingerprint does not resemble a petroleum product.

**ALS Group USA Corp. dba ALS Environmental (ALS) - Kelso
State Certifications, Accreditations, and Licenses**

Agency	Web Site	Number
Alaska DEH	http://dec.alaska.gov/eh/lab/cs/csapproval.htm	UST-040
Arizona DHS	http://www.azdhs.gov/lab/license/env.htm	AZ0339
Arkansas - DEQ	http://www.adeq.state.ar.us/techsvs/labcert.htm	88-0637
California DHS (ELAP)	http://www.cdph.ca.gov/certlic/labs/Pages/ELAP.aspx	2795
DOD ELAP	http://www.denix.osd.mil/edqw/Accreditation/AccreditedLabs.cfm	L16-58-R4
Florida DOH	http://www.doh.state.fl.us/lab/EnvLabCert/WaterCert.htm	E87412
Hawaii DOH	http://health.hawaii.gov/	-
ISO 17025	http://www.pjlabs.com/	L16-57
Louisiana DEQ	http://www.deq.louisiana.gov/page/la-lab-accreditation	03016
Maine DHS	http://www.maine.gov/dhhs/	WA01276
Minnesota DOH	http://www.health.state.mn.us/accreditation	053-999-457
Nevada DEP	http://ndep.nv.gov/bsdw/labservice.htm	WA01276
New Jersey DEP	http://www.nj.gov/dep/enforcement/oqa.html	WA005
New York - DOH	https://www.wadsworth.org/regulatory/elap	12060
North Carolina DEQ	https://deq.nc.gov/about/divisions/water-resources/water-resources-data/water-sciences-home-page/laboratory-certification-branch/non-field-lab-certification	605
Oklahoma DEQ	http://www.deq.state.ok.us/CSDnew/labcert.htm	9801
Oregon – DEQ (NELAP)	http://public.health.oregon.gov/LaboratoryServices/EnvironmentalLaboratoryAccreditation/Pages/index.aspx	WA100010
South Carolina DHEC	http://www.scdhec.gov/environment/EnvironmentalLabCertification/	61002
Texas CEQ	http://www.tceq.texas.gov/field/qa/env_lab_accreditation.html	T104704427
Washington DOE	http://www.ecy.wa.gov/programs/eap/labs/lab-accreditation.html	C544
Wyoming (EPA Region 8)	https://www.epa.gov/region8-waterops/epa-region-8-certified-drinking-water	-
Kelso Laboratory Website	www.alsglobal.com	NA

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. A complete listing of specific NELAP-certified analytes, can be found in the certification section at www.ALSGlobal.com or at the accreditation bodies web site.

Please refer to the certification and/or accreditation body's web site if samples are submitted for compliance purposes. The states highlighted above, require the analysis be listed on the state certification if used for compliance purposes and if the method/analyte is offered by that state.



Case Narrative

ALS Environmental—Kelso Laboratory
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Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com



Client: ALS Environmental - US
Project: HS19021158
Sample Matrix: Water

Service Request: K1901637
Date Received: 02/23/2019

CASE NARRATIVE

All analyses were performed consistent with the quality assurance program of ALS Environmental. This report contains analytical results for samples designated for Tier IV validation deliverables including summary forms and all of the associated raw data for each of the analyses. When appropriate to the method, method blank results have been reported with each analytical test.

Sample Receipt:

One water sample was received for analysis at ALS Environmental on 02/23/2019. The sample was received in good condition and consistent with the accompanying chain of custody form. The sample was stored in a refrigerator at 4°C upon receipt at the laboratory.

General Chemistry:

No significant anomalies were noted with this analysis.

Approved by Kelley Lovejoy

Date 03/13/2019



Chain of Custody

ALS Environmental—Kelso Laboratory
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10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

K190637

Subcontract Chain of Custody

COC ID: 10804

SUBCONTRACT TO:

ALS Environmental Kelso
1317 S. 13th Avenue
Kelso, WA 98626

Phone: +1 360 501 3312

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com


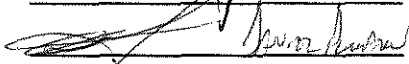
INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19021158
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19021158-01	LH18/24-SP650_022119	Water	21 Feb 2019 09:06
TOC Analysis for DOD Level IV			08 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: 
Received By: 
Cooler ID(s): _____

Date/Time: 2/22/19 1800
Date/Time: 2-28-19 1130
Temperature(s): _____



Cooler Receipt and Preservation Form

Client ALS-HOUSTON Service Request K19 01637
 Received: 2-23-19 Opened: 2-23-19 By: ASP Unloaded: 2-23-19 By: BP

1. Samples were received via? USPS Fed Ex UPS DHL PDX Courier Hand Delivered
2. Samples were received in: (circle) Cooler Box Envelope Other NA
3. Were custody seals on coolers? NA Y N If yes, how many and where? 2 TOP FRONT
 If present, were custody seals intact? Y N If present, were they signed and dated? Y N

Raw Cooler Temp	Corrected Cooler Temp	Raw Temp Blank	Corrected Temp Blank	Corr. Factor	Thermometer ID	Cooler/COC ID	Tracking Number	NA	Filed
4.4	4.2	1.4	1.2	-0.2	322	NA	4809 7831 0617	NA	

4. Packing material: Inserts Baggies Bubble Wrap Gel Packs Wet Ice Dry Ice Sleeves
5. Were custody papers properly filled out (ink, signed, etc.)? NA Y N
6. Were samples received in good condition (temperature, unbroken)? *Indicate in the table below.* NA Y N
 If applicable, tissue samples were received: Frozen Partially Thawed Thawed
7. Were all sample labels complete (i.e analysis, preservation, etc.)? NA Y N
8. Did all sample labels and tags agree with custody papers? *Indicate major discrepancies in the table on page 2.* NA Y N
9. Were appropriate bottles/containers and volumes received for the tests indicated? NA Y N
10. Were the pH-preserved bottles (*see SMO GEN SOP*) received at the appropriate pH? *Indicate in the table below* NA Y N
11. Were VOA vials received without headspace? *Indicate in the table below.* NA Y N
12. Was C12/Res negative? NA Y N

Sample ID on Bottle	Sample ID on COC	Identified by:

Sample ID	Bottle Count	Out of	Head-	Broke	pH	Reagent	Volume	Reagent Lot	Initials	Time
	Bottle Type	Temp	space				added	Number		

Notes, Discrepancies, & Resolutions: _____



General Chemistry

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www.alsglobal.com

Analytical Report

Client: ALS Environmental - US
Project: HS19021158/HS19021158
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1901637
Date Collected: 02/21/19
Date Received: 02/23/19
Units: mg/L
Basis: NA

Carbon, Total Organic

Sample Name	Lab Code	Result	LOQ	LOD	MDL	Dil.	Date Analyzed	Q
HS19021158 -01	K1901637-001	8.40	0.50	0.20	0.07	1	03/06/19 22:02	
Method Blank	K1901637-MB2	ND U	0.50	0.20	0.07	1	03/06/19 15:18	
Method Blank	K1901637-MB3	ND U	0.50	0.20	0.07	1	03/07/19 00:39	

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19021158/HS19021158
Sample Matrix: Water

Service Request: K1901637
Date Collected: 02/21/19
Date Received: 02/23/19

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA

Replicate Sample Summary
Carbon, Total Organic

Sample Name:	Lab Code:	LOQ	LOD	MDL	Sample Result	Duplicate Result	Average	RPD	RPD Limit	Date Analyzed
Batch QC	K1901602-003DUP	2.0	0.8	0.3	65.7	66.0	65.8	<1	10	03/06/19
HS19021158 -01	K1901637-001DUP2	0.50	0.20	0.07	8.40	8.29	8.35	1	10	03/06/19

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19021158/HS19021158
Sample Matrix: Water

Service Request: K1901637
Date Collected: N/A
Date Received: N/A
Date Analyzed: 03/6/19
Date Extracted: NA

Matrix Spike Summary
Carbon, Total Organic

Sample Name: Batch QC
Lab Code: K1901602-003
Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA

Matrix Spike
K1901602-003MS

<u>Analyte Name</u>	<u>Sample Result</u>	<u>Result</u>	<u>Spike Amount</u>	<u>% Rec</u>	<u>% Rec Limits</u>
Carbon, Total Organic	65.7	173	100	107	83-117

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19021158/HS19021158
Sample Matrix: Water

Service Request: K1901637
Date Analyzed: 03/06/19
Date Extracted: NA

Lab Control Sample Summary
Carbon, Total Organic

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA
Analysis Lot: 627457

Sample Name	Lab Code	Result	Spike Amount	% Rec	% Rec Limits
Lab Control Sample	K1901637-LCS2	27.5	25.0	110	83-117

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19021158/HS19021158
Sample Matrix: Water

Service Request: K1901637
Date Analyzed: 03/07/19
Date Extracted: NA

Lab Control Sample Summary
Carbon, Total Organic

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA
Analysis Lot: 627458

Sample Name	Lab Code	Result	Spike Amount	% Rec	% Rec Limits
Lab Control Sample	K1901637-LCS3	27.6	25.0	110	83-117

ALS Group USA, Corp.
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QA/QC Report

Client: ALS Environmental - US
Project: HS19021158/HS19021158

Service Request: K1901637

Continuing Calibration Verification (CCV) Summary

Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis		Date	True	Measured	Percent	Acceptance
	Lot	Lab Code	Analyzed	Value	Value	Recovery	Limits
CCV1	627457	KQ1902968-06	03/06/19 14:44	25.0	26.2	105	90-110
CCV2	627457	KQ1902968-07	03/06/19 18:45	25.0	25.8	103	90-110
CCV3	627457	KQ1902968-08	03/07/19 00:09	25.0	26.1	104	90-110
CCV4	627457	KQ1902968-09	03/07/19 04:52	25.0	26.3	105	90-110
CCV5	627457	KQ1902968-10	03/07/19 10:17	25.0	25.7	103	90-110
CCV6	627458	KQ1902969-05	03/07/19 04:52	25.0	26.3	105	90-110
CCV7	627458	KQ1902969-06	03/07/19 10:17	25.0	25.7	103	90-110
CCV8	627458	KQ1902969-07	03/07/19 15:00	25.0	25.9	103	90-110
CCV9	627458	KQ1902969-08	03/07/19 19:43	25.0	25.8	103	90-110

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19021158/HS19021158

Service Request:K1901637

Continuing Calibration Blank (CCB) Summary
Carbon, Total Organic

Analysis Method: SM 5310 C**Units:**mg/L

	Analysis Lot	Lab Code	Date Analyzed	LOQ	LOD	MDL	Result	Q
CCB1	627457	KQ1902968-01	03/06/19 15:01	0.50	0.20	0.07	ND	U
CCB2	627457	KQ1902968-02	03/06/19 18:59	0.50	0.20	0.07	ND	U
CCB3	627457	KQ1902968-03	03/07/19 00:24	0.50	0.20	0.07	ND	U
CCB4	627457	KQ1902968-04	03/07/19 05:07	0.50	0.20	0.07	ND	U
CCB5	627457	KQ1902968-05	03/07/19 10:31	0.50	0.20	0.07	ND	U
CCB6	627458	KQ1902969-01	03/07/19 05:07	0.50	0.20	0.07	ND	U
CCB7	627458	KQ1902969-02	03/07/19 10:31	0.50	0.20	0.07	ND	U
CCB8	627458	KQ1902969-03	03/07/19 15:15	0.50	0.20	0.07	0.17	J
CCB9	627458	KQ1902969-04	03/07/19 19:58	0.50	0.20	0.07	ND	U



Raw Data

ALS Environmental—Kelso Laboratory
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General Chemistry

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Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 627457 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1900530-026	Carbon, Total Organic	N/A		Water	0.43 mg/L	10 mL	0.50 mg/L U	1	0.07	0.50			3/6/19 16:23:00	N	I
K1901484-001	Carbon, Total Organic	N/A		Water	8.93 mg/L	10 ml	893 mg/L	100	7	50			3/7/19 01:08	N	II
K1901484-004	Carbon, Total Organic	N/A		Water	7.41 mg/L	10 ml	741 mg/L	100	7	50			3/7/19 01:36	N	II
K1901559-001	Carbon, Total Organic	N/A		Ground Water	1.04 mg/L	10 ml	1.04 mg/L	1	0.07	0.50			3/7/19 05:22	N	II
K1901564-002	Carbon, Total Organic	N/A		Water	4.27 mg/L	10 ml	4.27 mg/L	1	0.07	0.50			3/6/19 17:48	N	IV
K1901564-003	Carbon, Total Organic	N/A		Water	3.26 mg/L	10 ml	3.26 mg/L	1	0.07	0.50			3/6/19 18:16	N	IV
K1901564-019	Carbon, Total Organic	N/A		Water	20.14 mg/L	10 ml	80.6 mg/L	4	0.3	2.0			3/6/19 19:14	N	IV
K1901564-020	Carbon, Total Organic	N/A		Water	16.24 mg/L	10 ml	65.0 mg/L	4	0.3	2.0			3/6/19 19:42	N	IV
K1901602-003	Carbon, Total Organic	N/A		Water	16.41 mg/L	10 ml	65.7 mg/L	4	0.3	2.0			3/6/19 16:38	N	II
K1901612-001	Carbon, Total Organic	N/A		Water	111.94 mg/L	10 ml	1120 mg/L	10	0.7	5.0			3/6/19 20:10	N	II
K1901612-002	Carbon, Total Organic	N/A		Water	111.76 mg/L	10 ml	447 mg/L	4	0.3	2.0			3/6/19 20:38	N	II
K1901637-001	Carbon, Total Organic	N/A		Water	8.40 mg/L	10 ml	8.40 mg/L	1	0.07	0.50			3/6/19 22:02	N	IV
K1901638-001	Carbon, Total Organic	N/A		Ground Water	2.65 mg/L	10 ml	2.65 mg/L	1	0.07	0.50			3/6/19 22:30	N	IV
K1901638-002	Carbon, Total Organic	N/A		Ground Water	11.19 mg/L	10 ml	11.2 mg/L	1	0.07	0.50			3/6/19 22:58	N	IV
K1901638-003	Carbon, Total Organic	N/A		Ground Water	4.88 mg/L	10 ml	4.88 mg/L	1	0.07	0.50			3/6/19 23:26	N	IV
K1901651-001	Carbon, Total Organic	N/A		Water	9.25 mg/L	10 ml	925 mg/L	100	7	50			3/7/19 02:04	N	II
K1901651-002	Carbon, Total Organic	N/A		Water	8.89 mg/L	10 ml	889 mg/L	100	7	50			3/7/19 02:32	N	II
K1901651-003	Carbon, Total Organic	N/A		Water	6.58 mg/L	10 ml	658 mg/L	100	7	50			3/7/19 03:00	N	II
K1901651-004	Carbon, Total Organic	N/A		Water	7.90 mg/L	10 ml	790 mg/L	100	7	50			3/7/19 03:28	N	II
K1901651-005	Carbon, Total Organic	N/A		Water	7.24 mg/L	10 ml	724 mg/L	100	7	50			3/7/19 03:56	N	II
KQ1902968-01	Carbon, Total Organic	CCB		Water	0.01 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/6/19 15:01	N	I
KQ1902968-02	Carbon, Total Organic	CCB		Water	2.199999999999998E	10 ml	0.50 mg/L U	1	0.07	0.50			3/6/19 18:59	N	I
KQ1902968-03	Carbon, Total Organic	CCB		Water	3.400000000000001E	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 00:24	N	I
KQ1902968-04	Carbon, Total Organic	CCB		Water	-0.01 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 05:07	N	I
KQ1902968-05	Carbon, Total Organic	CCB		Water	-0.23 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 10:31	N	I
KQ1902968-06	Carbon, Total Organic	CCV		Water	26.22 mg/L	10 ml	26.2 mg/L	1			105		3/6/19 14:44	N	I
KQ1902968-07	Carbon, Total Organic	CCV		Water	25.78 mg/L	10 ml	25.8 mg/L	1			103		3/6/19 18:45	N	I
KQ1902968-08	Carbon, Total Organic	CCV		Water	26.12 mg/L	10 ml	26.1 mg/L	1			104		3/7/19 00:09	N	I
KQ1902968-09	Carbon, Total Organic	CCV		Water	26.35 mg/L	10 ml	26.3 mg/L	1			105		3/7/19 04:52	N	I
KQ1902968-10	Carbon, Total Organic	CCV		Water	25.73 mg/L	10 ml	25.7 mg/L	1			103		3/7/19 10:17	N	I
KQ1902968-11	Carbon, Total Organic	MB		Water	-0.02 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/6/19 15:18	N	I
KQ1902968-12	Carbon, Total Organic	LCS		Water	27.50 mg/L	10 ml	27.5 mg/L	1	0.07	0.50	110		3/6/19 15:39	N	I
KQ1902968-13	Carbon, Total Organic	LODV		Water	0.14 mg/L	10 mL	0.14 mg/L J	1	0.07	0.50			3/6/19 16:09:00	N	I
KQ1902968-14	Carbon, Total Organic	LOQV		Water	0.43 mg/L	10 mL	0.43 mg/L J	1	0.07	0.50			3/6/19 16:23:00	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

03/08/19

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 627457 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1902968-17	Carbon, Total Organic	DUP	K1901602-003	Water	16.49 mg/L	10 ml	66.0 mg/L	4	0.3	2.0		<1	3/6/19 16:38	N	II
KQ1902968-18	Carbon, Total Organic	DUP	K1901564-002	Water	4.23 mg/L	10 ml	4.23 mg/L	1	0.07	0.50		<1	3/6/19 17:48	N	IV
KQ1902968-19	Carbon, Total Organic	DUP	K1901564-003	Water	3.10 mg/L	10 ml	3.10 mg/L	1	0.07	0.50		5	3/6/19 18:16	N	IV
KQ1902968-20	Carbon, Total Organic	DUP	K1901564-019	Water	20.70 mg/L	10 ml	82.8 mg/L	4	0.3	2.0		3	3/6/19 19:14	N	IV
KQ1902968-21	Carbon, Total Organic	DUP	K1901564-020	Water	16.48 mg/L	10 ml	65.9 mg/L	4	0.3	2.0		1	3/6/19 19:42	N	IV
KQ1902968-22	Carbon, Total Organic	DUP	K1901612-001	Water	112.46 mg/L	10 ml	1120 mg/L	10	0.7	5.0		<1	3/6/19 20:10	N	II
KQ1902968-23	Carbon, Total Organic	DUP	K1901612-002	Water	119.75 mg/L	10 ml	479 mg/L	4	0.3	2.0		7	3/6/19 20:38	N	II
KQ1902968-24	Carbon, Total Organic	DUP	K1901637-001	Water	8.29 mg/L	10 ml	8.29 mg/L	1	0.07	0.50		1	3/6/19 22:02	N	IV
KQ1902968-25	Carbon, Total Organic	DUP	K1901638-001	Ground Water	2.55 mg/L	10 ml	2.55 mg/L	1	0.07	0.50		4	3/6/19 22:30	N	IV
KQ1902968-26	Carbon, Total Organic	DUP	K1901638-002	Ground Water	10.99 mg/L	10 ml	11.0 mg/L	1	0.07	0.50		2	3/6/19 22:58	N	IV
KQ1902968-27	Carbon, Total Organic	DUP	K1901638-003	Ground Water	4.80 mg/L	10 ml	4.80 mg/L	1	0.07	0.50		1	3/6/19 23:26	N	IV
KQ1902968-28	Carbon, Total Organic	DUP	K1901484-001	Water	8.99 mg/L	10 ml	899 mg/L	100	7	50		<1	3/7/19 01:08	N	II
KQ1902968-29	Carbon, Total Organic	DUP	K1901484-004	Water	7.54 mg/L	10 ml	754 mg/L	100	7	50		2	3/7/19 01:36	N	II
KQ1902968-30	Carbon, Total Organic	DUP	K1901651-001	Water	9.26 mg/L	10 ml	926 mg/L	100	7	50		<1	3/7/19 02:04	N	II
KQ1902968-31	Carbon, Total Organic	DUP	K1901651-002	Water	8.91 mg/L	10 ml	891 mg/L	100	7	50		<1	3/7/19 02:32	N	II
KQ1902968-32	Carbon, Total Organic	DUP	K1901651-003	Water	6.44 mg/L	10 ml	644 mg/L	100	7	50		2	3/7/19 03:00	N	II
KQ1902968-33	Carbon, Total Organic	DUP	K1901651-004	Water	7.88 mg/L	10 ml	788 mg/L	100	7	50		<1	3/7/19 03:28	N	II
KQ1902968-34	Carbon, Total Organic	DUP	K1901651-005	Water	7.21 mg/L	10 ml	721 mg/L	100	7	50		<1	3/7/19 03:56	N	II
KQ1902968-35	Carbon, Total Organic	DUP	K1901559-001	Ground Water	1.00 mg/L	10 ml	1.00 mg/L	1	0.07	0.50		5	3/7/19 05:22	N	II
KQ1902968-36	Carbon, Total Organic	MS	K1901602-003	Water	43.16 mg/L	10 ml	173 mg/L	4	0.3	2.0	107		3/6/19 17:06	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 627458 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1901559-002	Carbon, Total Organic	N/A		Ground Water	0.53 mg/L	10 ml	0.53 mg/L	1	0.07	0.50			3/7/19 05:50	N	II
K1901559-003	Carbon, Total Organic	N/A		Ground Water	1.09 mg/L	10 ml	1.09 mg/L	1	0.07	0.50			3/7/19 06:18	N	II
K1901594-001	Carbon, Total Organic	N/A		Water	1.52 mg/L	10 ml	1.52 mg/L	1	0.07	0.50			3/7/19 15:58	N	II
K1901594-002	Carbon, Total Organic	N/A		Water	29.13 mg/L	10 ml	29.1 mg/L	1	0.07	0.50			3/7/19 16:26	N	II
K1901594-003	Carbon, Total Organic	N/A		Water	31.09 mg/L	10 ml	31.1 mg/L	1	0.07	0.50			3/7/19 16:54	N	II
K1901594-004	Carbon, Total Organic	N/A		Water	1.34 mg/L	10 ml	1.34 mg/L	1	0.07	0.50			3/7/19 17:22	N	II
K1901594-005	Carbon, Total Organic	N/A		Water	1.87 mg/L	10 ml	1.87 mg/L	1	0.07	0.50			3/7/19 17:50	N	II
K1901674-001	Carbon, Total Organic	N/A		Water	3.39 mg/L	10 ml	169 mg/L	50	4	25			3/7/19 06:46	Y	II
K1901674-002	Carbon, Total Organic	N/A		Water	0.96 mg/L	10 ml	9.6 mg/L	10	0.7	5.0			3/7/19 07:57	N	II
K1901674-003	Carbon, Total Organic	N/A		Water	4.90 mg/L	10 ml	49.0 mg/L	10	0.7	5.0			3/7/19 08:25	N	II
K1901674-004	Carbon, Total Organic	N/A		Water	3.94 mg/L	10 ml	39.4 mg/L	10	0.7	5.0			3/7/19 08:53	N	II
K1901676-001	Carbon, Total Organic	N/A		Water	2.10 mg/L	10 ml	2.10 mg/L	1	0.07	0.50			3/7/19 11:15	N	II
K1901676-002	Carbon, Total Organic	N/A		Water	17.93 mg/L	10 ml	179 mg/L	10	0.7	5.0			3/7/19 11:43	N	II
K1901676-003	Carbon, Total Organic	N/A		Water	1.20 mg/L	10 ml	60 mg/L	50	4	25			3/7/19 12:11	N	II
K1901676-004	Carbon, Total Organic	N/A		Water	2.53 mg/L	10 ml	2.53 mg/L	1	0.07	0.50			3/7/19 12:39	N	II
K1901706-001	Carbon, Total Organic	N/A		Ground Water	3.15 mg/L	10 ml	3.15 mg/L	1	0.07	0.50			3/7/19 13:36	N	IV
K1901706-002	Carbon, Total Organic	N/A		Ground Water	14.68 mg/L	10 ml	14.7 mg/L	1	0.07	0.50			3/7/19 14:04	N	IV
K1901706-003	Carbon, Total Organic	N/A		Ground Water	9.39 mg/L	10 ml	9.39 mg/L	1	0.07	0.50			3/7/19 14:32	N	IV
K1901706-004	Carbon, Total Organic	N/A		Ground Water	6.28 mg/L	10 ml	6.28 mg/L	1	0.07	0.50			3/7/19 15:30	N	IV
K1901897-001	Carbon, Total Organic	N/A		Water	5.41 mg/L	10 ml	5.41 mg/L	1	0.07	0.50			3/7/19 18:18	N	II
KQ1902969-01	Carbon, Total Organic	CCB		Ground Water	-0.01 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 05:07	N	II
KQ1902969-02	Carbon, Total Organic	CCB		Ground Water	-0.23 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 10:31	N	II
KQ1902969-03	Carbon, Total Organic	CCB		Ground Water	0.17 mg/L	10 ml	0.17 mg/L J	1	0.07	0.50			3/7/19 15:15	N	II
KQ1902969-04	Carbon, Total Organic	CCB		Ground Water	-0.22 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 19:58	N	II
KQ1902969-05	Carbon, Total Organic	CCV		Ground Water	26.35 mg/L	10 ml	26.3 mg/L	1			105		3/7/19 04:52	N	II
KQ1902969-06	Carbon, Total Organic	CCV		Ground Water	25.73 mg/L	10 ml	25.7 mg/L	1			103		3/7/19 10:17	N	II
KQ1902969-07	Carbon, Total Organic	CCV		Ground Water	25.85 mg/L	10 ml	25.9 mg/L	1			104		3/7/19 15:00	N	II
KQ1902969-08	Carbon, Total Organic	CCV		Ground Water	25.81 mg/L	10 ml	25.8 mg/L	1			103		3/7/19 19:43	N	II
KQ1902969-09	Carbon, Total Organic	MS	K1901674-001	Water	31.30 mg/L	10 ml	1570 mg/L	50	4	25	112		3/7/19 07:14	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

03/08/19
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Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 627458 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1902969-10	Carbon, Total Organic	DUP	K1901559-002	Ground Water	0.42 mg/L	10 ml	0.42 mg/L J	1	0.07	0.50		24*	3/7/19 05:50	N	II
KQ1902969-11	Carbon, Total Organic	DUP	K1901559-003	Ground Water	0.96 mg/L	10 ml	0.96 mg/L	1	0.07	0.50		13*	3/7/19 06:18	N	II
KQ1902969-12	Carbon, Total Organic	DUP	K1901674-001	Water	3.33 mg/L	10 ml	166 mg/L	50	4	25		2	3/7/19 06:46	N	II
KQ1902969-13	Carbon, Total Organic	DUP	K1901674-002	Water	0.91 mg/L	10 ml	9.1 mg/L	10	0.7	5.0		5	3/7/19 07:57	N	II
KQ1902969-14	Carbon, Total Organic	DUP	K1901674-003	Water	4.35 mg/L	10 ml	43.5 mg/L	10	0.7	5.0		12*	3/7/19 08:25	N	II
KQ1902969-15	Carbon, Total Organic	DUP	K1901674-004	Water	3.80 mg/L	10 ml	38.0 mg/L	10	0.7	5.0		3	3/7/19 08:53	N	II
KQ1902969-16	Carbon, Total Organic	DUP	K1901676-001	Water	1.75 mg/L	10 mL	1.75 mg/L	1	0.07	0.50		18*	3/7/19 11:15:00	N	II
KQ1902969-17	Carbon, Total Organic	DUP	K1901676-002	Water	18.02 mg/L	10 ml	180 mg/L	10	0.7	5.0		<1	3/7/19 11:43	N	II
KQ1902969-18	Carbon, Total Organic	DUP	K1901676-003	Water	0.66 mg/L	10 ml	33 mg/L	50	4	25		58*	3/7/19 12:11	N	II
KQ1902969-19	Carbon, Total Organic	DUP	K1901676-004	Water	2.40 mg/L	10 ml	2.40 mg/L	1	0.07	0.50		5	3/7/19 12:39	N	II
KQ1902969-20	Carbon, Total Organic	DUP	K1901706-001	Ground Water	3.56 mg/L	10 ml	3.56 mg/L	1	0.07	0.50		12*	3/7/19 13:36	N	IV
KQ1902969-21	Carbon, Total Organic	DUP	K1901706-002	Ground Water	12.07 mg/L	10 ml	12.1 mg/L	1	0.07	0.50		19*	3/7/19 14:04	N	IV
KQ1902969-22	Carbon, Total Organic	DUP	K1901706-003	Ground Water	7.15 mg/L	10 ml	7.15 mg/L	1	0.07	0.50		27*	3/7/19 14:32	N	IV
KQ1902969-23	Carbon, Total Organic	DUP	K1901706-004	Ground Water	6.10 mg/L	10 ml	6.10 mg/L	1	0.07	0.50		3	3/7/19 15:30	N	IV
KQ1902969-24	Carbon, Total Organic	DUP	K1901594-001	Water	1.42 mg/L	10 ml	1.42 mg/L	1	0.07	0.50		7	3/7/19 15:58	N	II
KQ1902969-25	Carbon, Total Organic	DUP	K1901594-002	Water	29.44 mg/L	10 ml	29.4 mg/L	1	0.07	0.50		1	3/7/19 16:26	N	II
KQ1902969-26	Carbon, Total Organic	DUP	K1901594-003	Water	31.30 mg/L	10 ml	31.3 mg/L	1	0.07	0.50		<1	3/7/19 16:54	N	II
KQ1902969-27	Carbon, Total Organic	DUP	K1901594-004	Water	1.20 mg/L	10 ml	1.20 mg/L	1	0.07	0.50		11*	3/7/19 17:22	N	II
KQ1902969-28	Carbon, Total Organic	DUP	K1901594-005	Water	1.80 mg/L	10 ml	1.80 mg/L	1	0.07	0.50		4	3/7/19 17:50	N	II
KQ1902969-29	Carbon, Total Organic	DUP	K1901897-001	Water	5.36 mg/L	10 ml	5.36 mg/L	1	0.07	0.50		<1	3/7/19 18:18	N	II
KQ1902969-30	Carbon, Total Organic	LCS		Ground Water	27.56 mg/L	10 ml	27.6 mg/L	1	0.07	0.50	110		3/7/19 00:53	N	II
KQ1902969-31	Carbon, Total Organic	MB		Ground Water	0.02 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 00:39	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 627459 Method/Testcode: SM 5310 C/TOC D


Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1901553-001	Carbon, Dissolved Organic (DOC)	N/A		Water	7.55 mg/L	10 ml	15.1 mg/L	2	0.2	1.0			3/7/19 23:01	N	I
K1901553-002	Carbon, Dissolved Organic (DOC)	N/A		Water	3.06 mg/L	10 ml	3.06 mg/L	1	0.07	0.50			3/7/19 23:29	N	I
K1901703-001	Carbon, Dissolved Organic (DOC)	N/A		Water	1.99 mg/L	10 ml	1.99 mg/L	1	0.07	0.50			3/7/19 18:46	N	II
K1901703-002	Carbon, Dissolved Organic (DOC)	N/A		Water	1.83 mg/L	10 ml	1.83 mg/L	1	0.07	0.50			3/7/19 20:13	N	II
K1901703-003	Carbon, Dissolved Organic (DOC)	N/A		Water	3.41 mg/L	10 ml	3.41 mg/L	1	0.07	0.50			3/7/19 20:41	N	II
K1901703-004	Carbon, Dissolved Organic (DOC)	N/A		Water	6.14 mg/L	10 ml	6.14 mg/L	1	0.07	0.50			3/7/19 21:09	N	II
K1901703-005	Carbon, Dissolved Organic (DOC)	N/A		Water	7.31 mg/L	10 ml	7.31 mg/L	1	0.07	0.50			3/7/19 21:37	N	II
K1901703-006	Carbon, Dissolved Organic (DOC)	N/A		Water	7.26 mg/L	10 ml	7.26 mg/L	1	0.07	0.50			3/7/19 22:05	N	II
K1901703-007	Carbon, Dissolved Organic (DOC)	N/A		Water	0.69 mg/L	10 ml	0.69 mg/L	1	0.07	0.50			3/7/19 22:33	N	II
K1901728-001	Carbon, Dissolved Organic (DOC)	N/A		Water	1.43 mg/L	10 ml	1.43 mg/L	1	0.07	0.50			3/8/19 02:19	N	I
K1901728-002	Carbon, Dissolved Organic (DOC)	N/A		Water	0.25 mg/L	10 ml	25 mg/L J	100	7	50			3/8/19 02:47	N	I
K1901865-001	Carbon, Dissolved Organic (DOC)	N/A		Water	5.58 mg/L	10 ml	5.58 mg/L	1	0.07	0.50			3/8/19 00:27	N	II
K1901865-002	Carbon, Dissolved Organic (DOC)	N/A		Water	2.46 mg/L	10 ml	2.46 mg/L	1	0.07	0.50			3/8/19 00:55	N	II
KQ1902970-01	Carbon, Dissolved Organic (DOC)	CCB		Water	0.17 mg/L	10 ml	0.17 mg/L J	1	0.07	0.50			3/7/19 15:15	N	II
KQ1902970-02	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.22 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/7/19 19:58	N	II
KQ1902970-03	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.29 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/8/19 00:12	N	II
KQ1902970-04	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.32 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/8/19 03:30	N	II
KQ1902970-05	Carbon, Dissolved Organic (DOC)	CCV		Water	25.85 mg/L	10 ml	25.9 mg/L	1			104		3/7/19 15:00	N	II
KQ1902970-06	Carbon, Dissolved Organic (DOC)	CCV		Water	25.81 mg/L	10 ml	25.8 mg/L	1			103		3/7/19 19:43	N	II
KQ1902970-07	Carbon, Dissolved Organic (DOC)	CCV		Water	25.44 mg/L	10 ml	25.4 mg/L	1			102		3/7/19 23:57	N	II
KQ1902970-08	Carbon, Dissolved Organic (DOC)	CCV		Water	24.94 mg/L	10 ml	24.9 mg/L	1			100		3/8/19 03:15	N	II
KQ1902970-09	Carbon, Dissolved Organic (DOC)	MB		Water	-0.26 mg/L	10 mL	0.50 mg/L U	1	0.07	0.50			3/7/19 10:46:00	N	II
KQ1902970-10	Carbon, Dissolved Organic (DOC)	LCS		Water	26.60 mg/L	10 mL	26.6 mg/L	1	0.07	0.50	106		3/7/19 11:01:00	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Printed 3/8/19 10:46

Results Summary

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03/08/19


Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 627459 Method/Testcode: SM 5310 C/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1902970-11	Carbon, Dissolved Organic (DOC)	MS	K1901703-001	Water	30.19 mg/L	10 ml	30.2 mg/L	1	0.07	0.50	113		3/7/19 19:14	N	II
KQ1902970-12	Carbon, Dissolved Organic (DOC)	DUP	K1901703-001	Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50		7	3/7/19 18:46	N	II
KQ1902970-13	Carbon, Dissolved Organic (DOC)	DUP	K1901703-002	Water	1.85 mg/L	10 ml	1.85 mg/L	1	0.07	0.50		<1	3/7/19 20:13	N	II
KQ1902970-14	Carbon, Dissolved Organic (DOC)	DUP	K1901703-003	Water	3.35 mg/L	10 ml	3.35 mg/L	1	0.07	0.50		2	3/7/19 20:41	N	II
KQ1902970-15	Carbon, Dissolved Organic (DOC)	DUP	K1901703-004	Water	6.18 mg/L	10 ml	6.18 mg/L	1	0.07	0.50		<1	3/7/19 21:09	N	II
KQ1902970-16	Carbon, Dissolved Organic (DOC)	DUP	K1901703-005	Water	7.26 mg/L	10 ml	7.26 mg/L	1	0.07	0.50		<1	3/7/19 21:37	N	II
KQ1902970-17	Carbon, Dissolved Organic (DOC)	DUP	K1901703-006	Water	7.29 mg/L	10 ml	7.29 mg/L	1	0.07	0.50		<1	3/7/19 22:05	N	II
KQ1902970-18	Carbon, Dissolved Organic (DOC)	DUP	K1901703-007	Water	0.67 mg/L	10 ml	0.67 mg/L	1	0.07	0.50		4	3/7/19 22:33	N	II
KQ1902970-19	Carbon, Dissolved Organic (DOC)	DUP	K1901553-001	Water	7.64 mg/L	10 ml	15.3 mg/L	2	0.2	1.0		1	3/7/19 23:01	N	I
KQ1902970-20	Carbon, Dissolved Organic (DOC)	DUP	K1901553-002	Water	2.92 mg/L	10 ml	2.92 mg/L	1	0.07	0.50		5	3/7/19 23:29	N	I
KQ1902970-21	Carbon, Dissolved Organic (DOC)	DUP	K1901865-001	Water	5.66 mg/L	10 ml	5.66 mg/L	1	0.07	0.50		1	3/8/19 00:27	N	II
KQ1902970-22	Carbon, Dissolved Organic (DOC)	DUP	K1901865-002	Water	2.48 mg/L	10 ml	2.48 mg/L	1	0.07	0.50		<1	3/8/19 00:55	N	II
KQ1902970-23	Carbon, Dissolved Organic (DOC)	DUP	K1901728-001	Water	1.45 mg/L	10 ml	1.45 mg/L	1	0.07	0.50		1	3/8/19 02:19	N	I
KQ1902970-24	Carbon, Dissolved Organic (DOC)	DUP	K1901728-002	Water	0.22 mg/L	10 ml	22 mg/L	J 100	7	50		14*	3/8/19 02:47	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

TOC: 627457
627458
DOC: 627459

Schedule: Daily Run Method 010711

Version: 67

Instrument: Fusion1

Last Saved by: Fusion1 (Fusion1)

Last Saved on: 2019/03/06 16:23 - Wednesday

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Blank)	Blank	Reagent/Acid Blank		1	True	Ready
D	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
1	Sample	MB1	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
2	Sample	ICS	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
3	Sample	LOD	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
4	Sample	LOQ	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
5	Sample	K1901602-003.01 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
6	Sample	K1901602-003.01 ms 4x	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
7	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
8	Sample	K1901564-002.11	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
9	Sample	K1901564-003.11	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
10	Sample	K1901564-019.11 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
11	Sample	K1901564-020.11 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
12	Sample	K1901612-001.01 10x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
13	Sample	K1901612-002.01 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
14	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
15	Sample	K1901637-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
16	Sample	K1901638-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
17	Sample	K1901638-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
18	Sample	K1901638-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
19	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
20	Sample	MB2	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
21	Sample	K1901484-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
22	Sample	K1901484-004.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
23	Sample	K1901651-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
24	Sample	K1901651-002.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
25	Sample	K1901651-003.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
26	Sample	K1901651-004.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
27	Sample	K1901651-005.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
28	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
29	Sample	K1901559-001.09	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
30	Sample	K1901559-002.09	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
31	Sample	K1901559-003.09	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
32	Sample	K1901674-001.01 50x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
33	Sample	K1901674-001.01 ms 50x	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
34	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
35	Sample	K1901674-002.01 10x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
36	Sample	K1901674-003.01 10x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
37	Sample	K1901674-004.01 10x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
38	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready

Printed on: March 8, 2019 09:02:53

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B. [Signature]

Schedule: Daily Run Method 010711

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
39	Sample	MB3	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
40	Sample	K1901676-001.02	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
41	Sample	K1901676-002.02 10x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
42	Sample	K1901676-003.02 50x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
43	Sample	K1901676-004.02	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
44	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
45	Sample	K1901706-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
46	Sample	K1901706-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
47	Sample	K1901706-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
48	Sample	K1901706-004.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
49	Sample	K1901594-001.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
50	Sample	K1901594-002.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
51	Sample	K1901594-003.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
52	Sample	K1901594-004.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
53	Sample	K1901594-005.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
54	Sample	K1901897-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
55	Sample	K1901703-001.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
56	Sample	K1901703-001.01 ms doc	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
57	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
58	Sample	MB4	CAS_salt_010711 (CAS_salt_010711)	1	False	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	False	Ready
59	Sample	K1901703-002.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
60	Sample	K1901703-003.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
61	Sample	K1901703-004.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
62	Sample	K1901703-005.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
63	Sample	K1901703-006.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
64	Sample	K1901703-007.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
65	Sample	K1901553-001.06 doc 2x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
66	Sample	K1901553-002.05 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
67	Sample	K1901865-001.02 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
68	Sample	K1901865-002.02 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
69	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
70	Sample	FB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
71	Sample	K1901728-001.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
72	Sample	K1901728-002.01 doc 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
					False	

Fusion Report - Daily Run Method 010711 Wednesday, March 06, 2019 12:36 PM

(View - Repts, Unused Repts, Meta-Data, Signature, History)
Printed on 2019/03/08 09:03 - Friday

Report Summary Information

Company Location:	Gen Chem Lab	Engine	1.1.5.1
Schedule Name:	Daily Run Method 010711	Version:	
Instrument Name:	Fusion1	Firmware	1.2.0696
Report Version:	1 of 1	Version:	
Report Creation by Operators (schedule version):	Fusion1 (Fusion1) (v59) Fusion1 (Fusion1) (v60) Fusion1 (Fusion1) (v61) Fusion1 (Fusion1) (v62) Fusion1 (Fusion1) (v63) Fusion1 (Fusion1) (v64) Fusion1 (Fusion1) (v65) Fusion1 (Fusion1) (v66) Fusion1 (Fusion1) (v67)	Connection:	RS232 COM1

Comment:

Report Results

03/08/19
[Signature]

Sample Type: Clean		From Schedule Version 59				
Pos	Analysis Type	Sample ID			Start Time	
◆ (clean)		Clean			2019/03/06 12:36	
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	13.35	17.28	3.93	49.57	05:24
2	TC Clean	6.77	9.63	2.86	49.99	04:01
3	TC Clean	1.88	4.84	2.96	50.01	03:55
4	TC Clean	1.47	4.58	3.10	50.03	03:55

Sample Type: Clean		From Schedule Version 60				
Pos	Analysis Type	Sample ID			Start Time	
◆ (clean)		Clean			2019/03/06 12:59	
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	11.70	14.60	2.91	49.67	05:22

2	TC Clean	4.26	7.36	3.10	49.99	04:04
3	TC Clean	2.14	5.17	3.02	50.05	03:55
4	TC Clean	1.41	4.44	3.03	50.10	03:56

Sample Type: Clean From Schedule Version 61

Pos	Analysis Type	Sample ID			Start Time	
◊ (clean)		Clean			2019/03/06 13:33	

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	11.51	14.34	2.83	49.74	05:23
2	TC Clean	5.23	7.99	2.75	50.05	04:01
3	TC Clean	2.47	5.42	2.96	50.11	03:46
4	TC Clean	1.76	4.78	3.02	50.07	03:46

Sample Type: Blank (Creating v1229) From Schedule Version 62

Pos	Analysis Type	Sample ID			Start Time	
◊ (blank)		Reagent/Acid Blank			2019/03/06 13:55	

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	0.87	3.78	2.91	49.71	05:12
2	TC Clean	4.88	8.00	3.13	50.12	04:02
3	TC Clean	2.68	5.63	2.95	50.12	03:45
4	TC Clean	2.03	4.98	2.95	50.12	03:47
5	Reagent Blank	7.97	10.93	2.95	50.11	05:06
6	Acid Blank	1.37	4.40	3.03	49.55	05:29

Sample Type: Sample From Schedule Version 62

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ D	TOC	RB	0.9560 ppm	0.0000 ppm	0.0000%	2019/03/06 14:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.9560	9.5600	15.54	18.56	3.02	50.25	10:31

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 63

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.5547 ppm (PASS)	0.0000 ppm	0%	2019/03/06 14:44

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.5547	265.5467	189.71	192.74	3.02	50.29	10:29

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 64

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.3488 ppm (PASS)	0.0000 ppm	0%	2019/03/06 15:01

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.3488	3.4878	11.83	14.80	2.97	50.26	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 65

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 1	TOC	MB1	0.3156 ppm	0.0000 ppm	0.0000%	2019/03/06 15:18

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3156	3.1560	11.19	14.09	2.90	50.28	10:32

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS From Schedule Version 66

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Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	27.8336 ppm (PASS)	0.0000 ppm	0%	2019/03/06 15:39

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	27.8336	278.3355	198.40	201.37	2.98	50.24	10:28

Completion State	Success Action	Method	Calibration	STD Conc - Pos C
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	25 ppmC

Sample Type: Sample From Schedule Version 66

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 2	TOC	ICS	0.7157 ppm	0.0000 ppm	0.0000%	2019/03/06 15:54

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7157	7.1572	13.91	16.87	2.96	50.28	10:34

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 3	TOC	LOD	0.4741 ppm	0.0000 ppm	0.0000%	2019/03/06 16:09

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4741	4.7412	12.27	15.13	2.87	50.31	10:27

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Sample From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 4	TOC	LOQ	0.7682 ppm	0.0000 ppm	0.0000%	2019/03/06 16:23

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7682	7.6817	14.26	17.26	3.00	50.33	10:30

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time

5	TOC	K1901602-003.01 4x	16.7892 ppm	0.0542 ppm	0.3200%	2019/03/06 16:38
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Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	16.7509	167.5093	122.75	125.69	2.94	50.36	10:28
2	TOC	16.8275	168.2753	123.27	126.23	2.96	50.37	10:29

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
6	TOC	K1901602-003.01 ms 4x	43.4995 ppm	0.0000 ppm	0.0000%	2019/03/06 17:06

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	43.4995	434.9954	304.32	307.30	2.98	50.39	10:29

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
7	TOC	RB	0.3908 ppm	0.1172 ppm	29.9900%	2019/03/06 17:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4737	4.7368	12.26	15.25	2.99	50.39	10:29
2	TOC	0.3079	3.0794	11.14	14.21	3.07	50.41	10:30

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
8	TOC	K1901564-002.11	4.5904 ppm	0.0277 ppm	0.6000%	2019/03/06 17:48

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	4.6100	46.0998	40.34	43.18	2.84	50.42	10:26
2	TOC	4.5708	45.7080	40.07	43.06	2.98	50.40	10:26

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
9	TOC	K1901564-003.11	3.5189 ppm	0.1106 ppm	3.1400%	2019/03/06 18:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.5972	35.9716	33.47	36.50	3.04	50.43	10:30
2	TOC	3.4407	34.4070	32.40	35.36	2.96	50.37	10:27

Dilution **Blank Contribution** **Method** **Calibration**

1:10 (TC) 9.0477 (IC) CAS_salt_010711 CAS_salt_010711
(v1229) (v4) (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
◆	B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.1143 ppm (PASS)	0.0000 ppm	0%	2019/03/06 18:45

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.1143	261.1433	186.72	189.64	2.91	50.48	10:33

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
◆	D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.3404 ppm (PASS)	0.0000 ppm	0%	2019/03/06 18:59

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.3404	3.4038	11.77	14.71	2.94	50.50	10:29

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample

From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
◆	10	TOC	K1901564-019.11 4x	20.7624 ppm	0.3955 ppm	1.9100%	2019/03/06 19:14

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	20.4827	204.8269	148.08	151.03	2.95	50.51	10:27
2	TOC	21.0421	210.4207	151.88	154.91	3.03	50.53	10:30

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
◆	11	TOC	K1901564-020.11 4x	16.6990 ppm	0.1645 ppm	0.9900%	2019/03/06 19:42

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	16.5827	165.8269	121.61	124.60	2.99	50.55	10:29
2	TOC	16.8153	168.1530	123.19	126.24	3.05	50.56	10:27

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
12	TOC	K1901612-001.01 10x	112.5359 ppm	0.3656 ppm	0.3200%	2019/03/06 20:10

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	112.2774	1122.7739	771.18	774.16	2.98	50.57	10:27
2	TOC	112.7945	1127.9449	774.69	777.84	3.15	50.59	10:29

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
13	TOC	K1901612-002.01 4x	116.0911 ppm	5.6458 ppm	4.8600%	2019/03/06 20:38

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	112.0990	1120.9899	769.97	773.21	3.24	50.60	10:28
2	TOC	120.0833	1200.8330	824.17	828.26	4.10	50.60	10:28

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
14	TOC	RB	3.5314 ppm	4.1684 ppm	118.0400%	2019/03/06 21:06

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	9.7391	97.3908	75.16	79.27	4.12	50.63	10:26
2	TOC	2.1399	21.3987	23.57	26.84	3.27	50.63	10:25
3	TOC	1.2919	12.9189	17.82	20.87	3.05	50.65	10:29
4	TOC	0.9548	9.5482	15.53	18.48	2.95	50.61	10:24

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
15	TOC	K1901637-001.01	8.6847 ppm	0.0773 ppm	0.8900%	2019/03/06 22:02

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	8.7394	87.3937	68.37	71.34	2.97	50.59	10:28
2	TOC	8.6301	86.3005	67.63	70.64	3.01	50.50	10:28

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
16	TOC	K1901638-001.01	2.9411 ppm	0.0696 ppm	2.3700%	2019/03/06 22:30

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.9903	29.9035	29.35	32.52	3.18	50.49	10:28
2	TOC	2.8919	28.9194	28.68	31.60	2.92	50.48	10:26

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
17	TOC	K1901638-002.01	11.4260 ppm	0.1424 ppm	1.2500%	2019/03/06 22:58

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	11.5267	115.2666	87.29	90.10	2.81	50.44	10:28
2	TOC	11.3253	113.2527	85.92	88.83	2.91	50.42	10:27

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
18	TOC	K1901638-003.01	5.1774 ppm	0.0507 ppm	0.9800%	2019/03/06 23:26

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.2133	52.1326	44.44	47.54	3.11	50.37	10:28
2	TOC	5.1415	51.4151	43.95	46.97	3.02	50.36	10:28

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
19	TOC	RB	0.4492 ppm	0.0000 ppm	0.0000%	2019/03/06 23:55

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4492	4.4922	12.10	15.09	3.00	50.35	10:32

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 67

Concentration	Min / Max

Pos	BAT	(ppm)	Dil	Sample ID	(% dev)	Result	Std. Dev.	RSD	Start Time
♦ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.4560 ppm (PASS)	0.0000 ppm	0%	2019/03/07 00:09

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.4560	264.5596	189.04	191.98	2.94	50.32	10:32

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.3416 ppm (PASS)	0.0000 ppm	0%	2019/03/07 00:24

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.3416	3.4156	11.78	14.62	2.84	50.32	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 20	TOC	MB2	0.3558 ppm	0.0000 ppm	0.0000%	2019/03/07 00:39

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3558	3.5582	11.46	14.21	2.74	50.29	10:29

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	27.8960 ppm (PASS)	0.0000 ppm	0%	2019/03/07 00:53

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time

C	TOC	25.0 ppm	1	27.8960	278.9602	198.82	201.65	2.83	50.27	10:32
Completion State		Success Action		Method		Calibration		STD Conc - Pos C		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		25 ppmC		

Sample Type: Sample From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
21	TOC	K1901484-001.01 100x	9.2955 ppm	0.0429 ppm	0.4600%	2019/03/07 01:08

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	9.2652	92.6515	71.94	74.67	2.73	50.26	10:30
2	TOC	9.3258	93.2585	72.35	75.23	2.88	50.28	10:27

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
22	TOC	K1901484-004.01 100x	7.8135 ppm	0.0953 ppm	1.2200%	2019/03/07 01:36

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.7461	77.4613	61.63	64.55	2.92	50.28	10:26
2	TOC	7.8809	78.8093	62.54	65.32	2.77	50.27	10:27

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
23	TOC	K1901651-001.01 100x	9.5939 ppm	0.0064 ppm	0.0700%	2019/03/07 02:04

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	9.5894	95.8940	74.14	77.01	2.87	50.26	10:26
2	TOC	9.5984	95.9839	74.20	77.23	3.03	50.26	10:28

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
24	TOC	K1901651-002.01 100x	9.2408 ppm	0.0160 ppm	0.1700%	2019/03/07 02:32

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	9.2295	92.2950	71.70	74.56	2.86	50.26	10:31
2	TOC	9.2522	92.5219	71.85	74.63	2.78	50.24	10:25

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) **Method** CAS_salt_010711 **Calibration** CAS_salt_010711

(v1229)

(v4)

(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
25	TOC	K1901651-003.01 100x	6.8494 ppm	0.0944 ppm	1.3800%	2019/03/07 03:00

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.9161	69.1613	55.99	58.88	2.89	50.24	10:28
2	TOC	6.7827	67.8266	55.09	57.97	2.88	50.24	10:26

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
26	TOC	K1901651-004.01 100x	8.2291 ppm	0.0176 ppm	0.2100%	2019/03/07 03:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	8.2416	82.4157	64.99	67.94	2.95	50.22	10:26
2	TOC	8.2167	82.1667	64.82	67.64	2.81	50.23	10:27

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
27	TOC	K1901651-005.01 100x	7.5602 ppm	0.0206 ppm	0.2700%	2019/03/07 03:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.5748	75.7480	60.46	63.31	2.85	50.22	10:26
2	TOC	7.5456	75.4563	60.27	63.11	2.84	50.23	10:25

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
28	TOC	RB	0.5632 ppm	0.0731 ppm	12.9800%	2019/03/07 04:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6150	6.1496	13.22	16.24	3.02	50.19	10:26
2	TOC	0.5115	5.1154	12.52	15.25	2.73	50.20	10:29

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
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♦	B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.6839 ppm (PASS)	0.0000 ppm	0%	2019/03/07 04:52
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Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.6839	266.8387	190.59	193.38	2.78	50.20	10:28

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 67

♦	D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.3270 ppm (PASS)	0.0000 ppm	0%	2019/03/07 05:07
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Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.3270	3.2698	11.68	14.51	2.83	50.20	10:32

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 29	TOC	K1901559-001.09	1.3591 ppm	0.0340 ppm	2.5000%	2019/03/07 05:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.3831	13.8308	18.44	21.25	2.82	50.19	10:29
2	TOC	1.3351	13.3506	18.11	20.97	2.86	50.22	10:30

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 30	TOC	K1901559-002.09	0.8134 ppm	0.0800 ppm	9.8400%	2019/03/07 05:50

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.8700	8.6997	14.95	17.92	2.97	50.19	10:31
2	TOC	0.7568	7.5683	14.18	17.14	2.95	50.19	10:26

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0477 (IC) (v1229)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
31	TOC	K1901559-003.09	1.3630 ppm	0.0964 ppm	7.0700%	2019/03/07 06:18

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.4311	14.3111	18.76	21.55	2.79	50.19	10:29
2	TOC	1.2948	12.9484	17.84	20.78	2.95	50.17	10:24

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
32	TOC	K1901674-001.01 50x	3.6953 ppm	0.0453 ppm	1.2300%	2019/03/07 06:46

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.7274	37.2739	34.35	37.26	2.91	50.20	10:31
2	TOC	3.6633	36.6330	33.91	36.74	2.83	50.22	10:25

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
33	TOC	K1901674-001.01 ms 50x	31.6409 ppm	0.0000 ppm	0.0000%	2019/03/07 07:14

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	31.6409	316.4086	223.82	226.59	2.77	50.19	10:31

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
34	TOC	RB	0.3333 ppm	0.0729 ppm	21.8800%	2019/03/07 07:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3848	3.8484	11.66	14.60	2.94	50.18	10:30
2	TOC	0.2817	2.8172	10.96	13.85	2.89	50.19	10:29

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
35	TOC	K1901674-002.01 10x	1.2695 ppm	0.0338 ppm	2.6600%	2019/03/07 07:57

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.2934	12.9337	17.83	20.72	2.90	50.20	10:26

2	TOC	1.2456	12.4563	17.50	20.40	2.89	50.20	10:32
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Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
36	TOC	K1901674-003.01 10x	4.9624 ppm	0.3896 ppm	7.8500%	2019/03/07 08:25

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.2379	52.3786	44.60	47.46	2.86	50.18	10:29
2	TOC	4.6869	46.8688	40.86	43.91	3.05	50.20	10:27

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
37	TOC	K1901674-004.01 10x	4.2076 ppm	0.0953 ppm	2.2700%	2019/03/07 08:53

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	4.2750	42.7498	38.07	40.94	2.88	50.18	10:27
2	TOC	4.1402	41.4018	37.15	40.13	2.98	50.19	10:31

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
38	TOC	RB	0.1295 ppm	0.0214 ppm	16.5400%	2019/03/07 09:21

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1460	1.4604	10.04	12.97	2.93	50.21	10:29
2	TOC	0.1490	1.4898	10.06	12.92	2.86	50.17	10:23
3	TOC	0.1179	1.1790	9.85	12.83	2.98	50.24	10:28
4	TOC	0.1052	1.0523	9.76	12.62	2.86	50.27	10:25

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

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Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.0678 ppm (PASS)	0.0000 ppm	0%	2019/03/07 10:17

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
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B	TOC	25 ppm	1	26.0678	260.6777	186.41	189.20	2.79	50.31	10:30
Completion State		Success Action		Method		Calibration		STD Conc - Pos B		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		50 ppmC		

Sample Type: Check Standard --> CCB From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.1057 ppm (PASS)	0.0000 ppm	0%	2019/03/07 10:31

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.1057	1.0570	10.18	12.99	2.81	50.33	10:33

Completion State		Success Action		Method		Calibration		STD Conc - Pos D		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		0 ppmC		

Sample Type: Sample From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 39	TOC	MB3	0.0790 ppm	0.0000 ppm	0.0000%	2019/03/07 10:46

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0790	0.7901	9.58	12.38	2.79	50.36	10:32

Dilution		Blank Contribution		Method		Calibration	
1:10		(TC) 9.0477 (IC) (v1229)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)	

Sample Type: Check Standard --> LCS From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.9353 ppm (PASS)	0.0000 ppm	0%	2019/03/07 11:01

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.9353	269.3534	192.30	195.05	2.75	50.37	10:29

Completion State		Success Action		Method		Calibration		STD Conc - Pos C		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		25 ppmC		

Sample Type: Sample

From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
40	TOC	K1901676-001.02	2.2636 ppm	0.2458 ppm	10.8600%	2019/03/07 11:15

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.4375	24.3745	25.59	28.67	3.07	50.39	10:29
2	TOC	2.0898	20.8978	23.23	26.41	3.18	50.42	10:28

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
41	TOC	K1901676-002.02 10x	18.3144 ppm	0.0619 ppm	0.3400%	2019/03/07 11:43

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	18.2707	182.7068	133.07	136.14	3.07	50.42	10:27
2	TOC	18.3582	183.5819	133.66	136.63	2.96	50.43	10:26

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
42	TOC	K1901676-003.02 50x	1.2686 ppm	0.3821 ppm	30.1200%	2019/03/07 12:11

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.5388	15.3880	19.49	22.49	3.00	50.46	10:30
2	TOC	0.9984	9.9843	15.82	18.79	2.96	50.44	10:25

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
43	TOC	K1901676-004.02	2.8031 ppm	0.0879 ppm	3.1400%	2019/03/07 12:39

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.8653	28.6527	28.50	31.49	2.99	50.43	10:26
2	TOC	2.7409	27.4093	27.65	30.68	3.02	50.43	10:28

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
44	TOC	RB	0.3509 ppm	0.1574 ppm	44.8600%	2019/03/07 13:07

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time

1	TOC	0.4622	4.6219	12.18	15.13	2.94	50.45	10:26
2	TOC	0.2396	2.3959	10.67	13.63	2.96	50.46	10:28

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
45	TOC	K1901706-001.01	3.6934 ppm	0.2852 ppm	7.7200%	2019/03/07 13:36

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.4917	34.9168	32.75	35.76	3.01	50.46	10:28
2	TOC	3.8950	38.9504	35.49	38.44	2.95	50.47	10:29

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
46	TOC	K1901706-002.01	13.7127 ppm	1.8415 ppm	13.4300%	2019/03/07 14:04

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	15.0149	150.1491	110.97	113.95	2.98	50.45	10:26
2	TOC	12.4106	124.1058	93.29	96.46	3.17	50.43	10:26

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
47	TOC	K1901706-003.01	8.6082 ppm	1.5858 ppm	18.4200%	2019/03/07 14:32

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	9.7295	97.2950	75.09	78.13	3.03	50.41	10:30
2	TOC	7.4868	74.8685	59.87	62.82	2.95	50.38	10:27

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.1926 ppm (PASS)	0.0000 ppm	0%	2019/03/07 15:00

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.1926	261.9256	187.26	190.28	3.03	50.40	10:33

Completion State Success - Criteria met.	Success Action Do Nothing	Method CAS_salt_010711 (v4)	Calibration CAS_salt_010711 (v30)	STD Conc - Pos B 50 ppmC
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Sample Type: Check Standard --> CCB From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev. (ppm)	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.5100 ppm (PASS)	0.0000 ppm	0%	2019/03/07 15:15

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.5100	5.0995	12.92	15.76	2.84	50.44	10:29

Completion State Success - Criteria met.	Success Action Do Nothing	Method CAS_salt_010711 (v4)	Calibration CAS_salt_010711 (v30)	STD Conc - Pos D 0 ppmC
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Sample Type: Sample From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 48	TOC	K1901706-004.01	6.5273 ppm	0.1218 ppm	1.8700%	2019/03/07 15:30

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.6134	66.1339	53.94	56.91	2.97	50.39	10:30
2	TOC	6.4412	64.4117	52.77	55.73	2.96	50.45	10:26

Dilution 1:10	Blank Contribution (TC) 9.0477 (IC) (v1229)	Method CAS_salt_010711 (v4)	Calibration CAS_salt_010711 (v30)
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Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 49	TOC	K1901594-001.04	1.8063 ppm	0.0694 ppm	3.8400%	2019/03/07 15:58

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.8554	18.5539	21.64	24.59	2.94	50.39	10:26
2	TOC	1.7573	17.5728	20.98	23.88	2.90	50.42	10:26

Dilution 1:10	Blank Contribution (TC) 9.0477 (IC) (v1229)	Method CAS_salt_010711 (v4)	Calibration CAS_salt_010711 (v30)
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Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 50	TOC	K1901594-002.04	29.6254 ppm	0.2211 ppm	0.7500%	2019/03/07 16:26

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	29.4691	294.6907	209.08	211.97	2.89	50.42	10:28

2	TOC	29.7817	297.8168	211.20	214.26	3.05	50.44	10:28
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Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
51	TOC	K1901594-003.04	31.5352 ppm	0.1514 ppm	0.4800%	2019/03/07 16:54

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	31.4281	314.2813	222.38	225.28	2.90	50.45	10:25
2	TOC	31.6422	316.4219	223.83	226.87	3.04	50.47	10:26

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
52	TOC	K1901594-004.04	1.6091 ppm	0.1022 ppm	6.3500%	2019/03/07 17:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.6814	16.8141	20.46	23.60	3.14	50.42	10:27
2	TOC	1.5369	15.3689	19.48	22.47	2.99	50.45	10:26

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
53	TOC	K1901594-005.04	2.1766 ppm	0.0484 ppm	2.2300%	2019/03/07 17:50

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.2109	22.1088	24.06	27.02	2.96	50.43	10:27
2	TOC	2.1424	21.4237	23.59	26.83	3.24	50.44	10:28

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
54	TOC	K1901897-001.01	5.7203 ppm	0.0368 ppm	0.6400%	2019/03/07 18:18

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.7463	57.4626	48.05	50.98	2.93	50.46	10:28
2	TOC	5.6943	56.9426	47.70	50.53	2.83	50.41	10:28

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
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55	TOC	K1901703-001.01 doc	2.2652 ppm	0.0938 ppm	4.1400%	2019/03/07 18:46
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Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.3315	23.3153	24.87	27.84	2.96	50.49	10:27
2	TOC	2.1989	21.9894	23.97	27.02	3.05	50.49	10:23

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
56	TOC	K1901703-001.01 ms doc	30.5280 ppm	0.0000 ppm	0.0000%	2019/03/07 19:14

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	30.5280	305.2801	216.27	219.40	3.13	50.50	10:30

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
57	TOC	RB	0.2627 ppm	0.0000 ppm	0.0000%	2019/03/07 19:29

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2627	2.6271	10.83	13.87	3.04	50.50	10:29

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.1528 ppm (PASS)	0.0000 ppm	0%	2019/03/07 19:43

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.1528	261.5278	186.99	189.86	2.88	50.56	10:30

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time

◆	D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.1190 ppm (PASS)	0.0000 ppm	0%	2019/03/07 19:58
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.1190	1.1896	10.27	13.16	2.89	50.53	10:32
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>		<u>Calibration</u>		<u>STD Conc - Pos D</u>		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		0 ppmC		

Sample Type: Sample From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◆ 59	TOC	K1901703-002.01 doc	2.1764 ppm	0.0127 ppm	0.5800%	2019/03/07 20:13		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.1674	21.6742	23.76	26.74	2.98	50.48	10:26
2	TOC	2.1854	21.8539	23.88	26.87	2.99	50.45	10:24
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0477 (IC) (v1229)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◆ 60	TOC	K1901703-003.01 doc	3.7162 ppm	0.0463 ppm	1.2400%	2019/03/07 20:41		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.7489	37.4890	34.49	37.39	2.90	50.42	10:29
2	TOC	3.6835	36.8349	34.05	36.98	2.93	50.40	10:26
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0477 (IC) (v1229)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◆ 61	TOC	K1901703-004.01 doc	6.4987 ppm	0.0318 ppm	0.4900%	2019/03/07 21:09		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.4762	64.7623	53.01	55.91	2.90	50.39	10:25
2	TOC	6.5212	65.2117	53.31	56.18	2.86	50.39	10:26
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0477 (IC) (v1229)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◆ 62	TOC	K1901703-005.01 doc	7.6223 ppm	0.0299 ppm	0.3900%	2019/03/07 21:37

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.6435	76.4345	60.93	63.69	2.76	50.36	10:28
2	TOC	7.6012	76.0117	60.64	63.26	2.62	50.37	10:28

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
63	TOC	K1901703-006.01 doc	7.6095 ppm	0.0211 ppm	0.2800%	2019/03/07 22:05

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.5945	75.9454	60.60	63.50	2.91	50.33	10:28
2	TOC	7.6244	76.2445	60.80	63.72	2.92	50.29	10:27

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
64	TOC	K1901703-007.01 doc	1.0166 ppm	0.0185 ppm	1.8200%	2019/03/07 22:33

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.0297	10.2966	16.04	18.95	2.91	50.12	10:27
2	TOC	1.0034	10.0344	15.86	18.52	2.66	50.21	10:29

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
65	TOC	K1901553-001.06 doc 2x	7.9336 ppm	0.0659 ppm	0.8300%	2019/03/07 23:01

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.8870	78.8697	62.58	65.48	2.89	50.40	10:30
2	TOC	7.9802	79.8022	63.22	66.07	2.85	50.43	10:26

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
66	TOC	K1901553-002.05 doc	3.3275 ppm	0.0993 ppm	2.9800%	2019/03/07 23:29

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.3977	33.9769	32.11	35.01	2.90	50.41	10:29
2	TOC	3.2573	32.5729	31.16	33.95	2.79	50.39	10:31

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

1:10 (TC) 9.0477 (IC) CAS_salt_010711 CAS_salt_010711
(v1229) (v4) (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.7792 ppm (PASS)	0.0000 ppm	0%	2019/03/07 23:57

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.7792	257.7917	184.45	187.32	2.87	50.36	10:30

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0491 ppm (PASS)	0.0000 ppm	0%	2019/03/08 00:12

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0491	0.4913	9.80	12.77	2.97	50.32	10:31

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample

From Schedule Version 67

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 67	TOC	K1901865-001.02 doc	5.9540 ppm	0.0577 ppm	0.9700%	2019/03/08 00:27

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.9132	59.1318	49.19	52.18	2.99	50.29	10:30
2	TOC	5.9948	59.9479	49.74	52.50	2.76	50.30	10:27

Dilution 1:10
Blank Contribution (TC) 9.0477 (IC) (v1229)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 68	TOC	K1901865-002.02 doc	2.8095 ppm	0.0151 ppm	0.5400%	2019/03/08 00:55

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.7988	27.9883	28.05	30.99	2.94	50.29	10:24
2	TOC	2.8202	28.2019	28.19	31.23	3.04	50.28	10:29

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
69	TOC	RB	0.1225 ppm	0.0605 ppm	49.3900%	2019/03/08 01:23

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1653	1.6534	10.17	12.88	2.71	50.28	10:27
2	TOC	0.0797	0.7974	9.59	12.49	2.90	50.28	10:26

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
70	TOC	FB	0.0226 ppm	0.0198 ppm	87.6400%	2019/03/08 01:51

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0366	0.3658	9.30	12.23	2.94	50.27	10:28
2	TOC	0.0086	0.0859	9.11	11.91	2.80	50.26	10:25

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
71	TOC	K1901728-001.01 doc	1.7817 ppm	0.0129 ppm	0.7200%	2019/03/08 02:19

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.7726	17.7260	21.08	23.90	2.82	50.26	10:25
2	TOC	1.7909	17.9087	21.20	23.95	2.75	50.25	10:30

Dilution 1:10 **Blank Contribution** (TC) 9.0477 (IC) (v1229) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
72	TOC	K1901728-002.01 doc 100x	0.5750 ppm	0.0235 ppm	4.0900%	2019/03/08 02:47

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5917	5.9168	13.06	15.82	2.75	50.24	10:28
2	TOC	0.5584	5.5839	12.84	15.78	2.94	50.23	10:27

Dilution **Blank Contribution** **Method** **Calibration**

1:10 (TC) 9.0477 (IC) CAS_salt_010711 CAS_salt_010711
(v1229) (v4) (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.2814 ppm (PASS)	0.0000 ppm	0%	2019/03/08 03:15

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.2814	252.8138	181.07	183.95	2.88	50.22	10:28

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 67

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0213 ppm (PASS)	0.0000 ppm	0%	2019/03/08 03:30

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0213	0.2129	9.61	12.42	2.81	50.22	10:31

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Meta Data Used in this Report

Blanks

Version	Reagent (Abs)	Acid (Abs)	DI IC (Abs)	DI TC (Abs)	DI TOC (Abs)	Save Time	Operator
v1228	1.4157	0.7880	0.0000	0.0000	0.0000	2019/03/05 15:06	Fusion1 (Fusion1)
v1229	2.6573	1.3700	0.0000	0.0000	0.0000	2019/03/06 14:28	Fusion1 (Fusion1)

Calibrations

Name: CAS_salt_010711 (TOC)

Version: v30 Calibration curve formula: TOC: $y = 6.788x + 9.463$
 Ver Creation: 2019/03/05 17:42 r^2 value: TOC: $r^2 = 0.99963$
 Comment:
 Operator: Fusion1 (Fusion1)
 Basic Analysis Type: TOC

Basic Analysis Type: TOC

Sample ID	Y Raw Value	X Expected	Message	End Time
DI Water	7.8970	0.0000		2019/03/05 16:15
0.500 ppm	11.5280	0.5000		2019/03/05 16:29
1.0 ppm	14.9760	1.0000		2019/03/05 16:44
5.0 ppm	43.6500	5.0000		2019/03/05 16:58
10 ppm	79.6020	10.0000		2019/03/05 17:12
25 ppm	183.3580	25.0000		2019/03/05 17:26
50 ppm	346.3230	50.0000		2019/03/05 17:40

Methods

Name: CAS_salt_010711 (TOC)

Version: v4 Operator: Fusion1 (Fusion1)
 Ver Creation: 2019/02/21 17:57
 Comment:

Parameter	Value	Advanced Parameter	Value
SampleVolume	10.0 mL	NeedleRinseVolume	5.0 ml
Dilution	1:10	VialPrimeVolume	2.0 ml
AcidVolume	0.5 ml	ICSamplePrimeVolume	2.0 ml
ReagentVolume	2.0 ml	ICSpurgeRinseVolume	12.0 ml
UVReactorPrerinse	Off	BaselineStabilizeTime	0.70 min
UVReactorPrerinseVolume	5.0	DetectorPressureFlow	150 ml/min
NumberOfUVReactorPrerinses	1	SyringeSpeedWaste	10
ICSpurgeTime	1.00 mins	SyringeSpeedAcid	7
DetectorSweepFlow	500 ml/min	SyringeSpeedReagent	7
PreSpurgeTime	2.00 mins	SyringeSpeedDIWater	7
SystemFlow	500 ml/min	NDIRPressurization	60 psig
		SyringeSpeedSampleDispense	5
		SyringeSpeedSampleAspirate	4
		SyringeSpeedUVDispense	5
		SyringeSpeedUVAspirate	5
		SyringeSpeedICDispense	5
		SyringeSpeedICAspirate	5
		NDIRPressureStabilize	1.75 min
		SampleMixing	Off
		SampleMixingCycles	1
		SampleMixingVolume	10.0

LowLevelFilterNDIR	Off
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Acceptance / Approval

Electronic Signatures				
Report Version	User Name	Acceptance	Reason	Date

Report History

Report History				
Report Version	User Name	System Reason	User Reason	Date
1	Fusion1 (Fusion1)	Schedule completed	Schedule completed	2019/03/08 03:48

StarLIMS Run: 627457, 627458, 627459
 Analysis: TOC
 Method: 9060, 415.1, SM 5310 C, 9060A

CCV: 11-GEN-05-76B 50 ppm LCS: 11-GEN-05-74L 25.0 ppm

ICAL Date: 3/6/19

ICAL ID: 11-GEN-05-76H

ICS ID: 11-GEN-05-74A

ICS TV: 25.0 ppm ICS % R = 2

Spike ID: 11-GEN-05-700 0.05 ml of 5000 ppm stock ---> 10.0 ml = 25.0 ppm x dilution factor

Sodium Persulfate: 11-GEN-05-76F

21 % H3PO4: 11-GEN-05-76G

Equipment ID: K-TOC-03

PIPETTE ID: 124276B, 129001F, N11314F, Marge

FILTER ID: NA

Analyzed By: <i>BCD</i>	Date Analyzed: <i>3/6/19</i>
Reviewed By: <i>[Signature]</i>	Date Reviewed: <i>07/08/19</i>



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1905651; 1906112; 1906330;
1906332; 1906334

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2223 (233911)

General Set Information: There were thirteen field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ¹⁸O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50μL of an ¹⁸O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Method QC data: The method blank (LMB 642099) was less than 1/2 the CRDL. The recovery for the LCS (642100) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.µg/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in µg/L. Results were calculated in µg/L by the equation $(A) \times (B)$,

where: A = Analyte concentration from the standard curve (µg/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level of 4.0µg/L. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported. Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEBI04) along with datafiles 05MARD07-10.

Thomas Bosch March 06, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 07, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1905651**

Project ID: 10803

Purchase Order: 10803

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
HS19021158-02/LH18/24-SP650_022119_BIX	1905651001	02/21/19	02/23/19	



ANALYTICAL REPORT

Workorder: 34-1905651

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: HS19021158-02/LH18/24-SP650_022119_BIX	Sampling Site: NA	Collected: 02/21/2019				
Lab ID: 1905651001	Media: 250 mL Nalgene	Received: 02/23/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 09:40	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 233911)

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/06/2019 09:17	/S/ Stephen Brose 03/07/2019 11:04

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1905651

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00934387

Analysis Information

Workorder: 1905651

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2223 (HBN: 233911)
Analyzed By: Thomas Bosch

Blank

LMB: 642099 Analyzed: 03/05/2019 09:26 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 642100 Analyzed: 03/05/2019 09:00 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.11	4.00	103	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1906112001 Analyzed: 03/05/2019 09:54 Dilution: 1 Units: ug/L		MS: 1906112002 Analyzed: 03/05/2019 10:07 Dilution: 1 Units: ug/L				MSD: 1906112003 Analyzed: 03/05/2019 10:20 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	51.0	53.9	4	▲ 65.7	78.8 123.8	53.7	▲ 61.2	0.337	0.0 20.0

Continuing Calibration Verification

CCV: 642096 Analyzed: 03/05/2019 08:44 Units: ug/L Criteria: ± 15%			CCV: 642101 Analyzed: 03/05/2019 12:18 Units: ug/L Criteria: ± 15%			CCV: 642102 Analyzed: 03/05/2019 14:07 Units: ug/L Criteria: ± 15%			
Analyte	Result	Target	% Rec.	Result	Target	% Rec.	Result	Target	% Rec.
Perchlorate	25.0	25.0	100	25.0	25.0	100	24.5	25.0	97.8

Interference Check Sample

ICSA: 642098 Analyzed: 03/05/2019 09:13 Units: ug/L Criteria: ± 30%			
Analyte	Result	Target	% Rec.
Perchlorate	3.93	4.00	98.2

Comments

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.



Quality Control Sample Batch Report

00934388

Analysis Information

Workorder: 1905651

Limits: Client SOW/Contract Specified

Preparation: NA

Analysis: EPA 6850, DoD QSM

Basis: DoD QSM

Batch: NA

Batch: ELMS/2223 (HBN: 233911)

Prepared By: NA

Analyzed By: Thomas Bosch

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/06/2019 13:45	/S/ Stephen Brose 03/07/2019 11:04

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



1905651

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18698/#2

Subcontract Chain of Custody

COC ID: 10803

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

1905051

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19021158
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19021158-02	LH18/24-SP650_022119_BIX	Water	21 Feb 2019 14:00
SUB_Perch-6850			08 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: [Signature]
Received By: [Signature]
Cooler ID(s): 9181

Date/Time: 2/22/19 18:00
Date/Time: 2/23/19 850
Temperature(s): 2°



ALS Environmental
CHAIN-OF-CUSTODY

Project / Job / Task: 10803		Split:		Workorder ID: 1905651		Level: ENV_LVL4		Requested Analysis	
Client: ALS Environmental (Houston)				Account: 8101		Type: 250Poly			
Comments:									
Item	Collect Date/Time	Sample ID	Lab ID	QC	Matrix	ID(s)	Containers	Count	
1	02/21/2019 14:00	HS19021158-02/LH18/24-SP650_0	1905651001		Water	A	COOL	1	A
2									
3									
4									
5									
6									
7									
8									
9									
10									

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY					SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY				
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location	Sample Prep / Analysis for: _____	Lab Notebook No.: _____	Prepared / Analyzed by: _____	Date / Time: _____	Received By: (Signature)	Reason for Transfer / Storage Location
Schmith, Marianne	02/23/2019 08:50	ALS Sample Receiving	Sample Login						
<i>[Signature]</i>	03/01/19 13:10	R-33-1 10 B	Storage						
R-33-1		T. Bush	6850						

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: All Houston Project/Task/Site: 1905051
 Date/Time of Receipt: 2/23/19 850 Number of Coolers Received: 1

Condition of Coolers: Acceptable/Unacceptable Temperature Control: Present/Not Included
 Cooler Custody Seals: Present/Absent/NA Location Temp Taken: Control/Between Samples
 Container Custody Seals: Present/Absent/NA Intact/Broken/NA
 Intact/Broken/NA
 Ice Present: Yes/No/NA Are all temperatures within project specific guidelines? Yes/No/NA
 Frozen/Melted/NA VOA Headspace Present? Yes/No/NA *NA/No 2/23/19*

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9181</u>	<u>2</u> °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: *Meredith Lewis* Signature Printed Name Meredith Lewis Date 2/23/19

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Must Deliver Next Business Day
Time and Tempature Sensitive!

ORIGIN ID:SGRA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

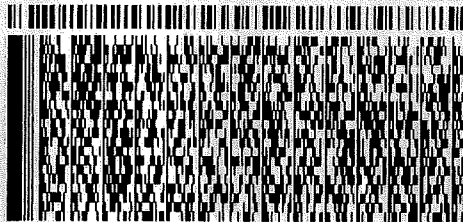
SHIP DATE: 22FEB19
ACTWGT: 8.65 LB
CAD: 300130/CAFE3211

BILL THIRD PARTY

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(801) 288-7700
REF: HS19021158 - RJ



FedEx
Express



2007/05/07/20155

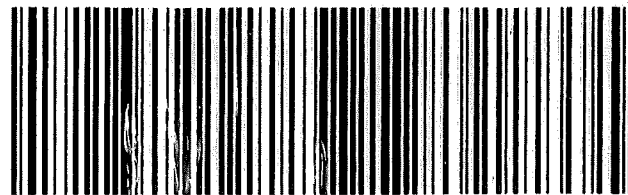
Part # 159469-434 RT2 EXP 11/19

TRK# 4809 7831 0661
0201

SATURDAY 12:00P
PRIORITY OVERNIGHT

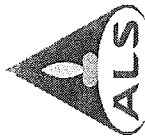
XO BTFA

84123
UT-US SLC



Saturday Delivery

12:00
0661
B
8
FedEx Express
FedEx



Batch Worklist

HBN: 233911

Instrument:



Status: WP

Created: 3/5/2019 08:21

Analyst: T. Bosch

Batch: ELMS/ 2223
 Rule: EPA 6850, DoD QSM Water

- Workorder: 1905651 [ENV_LVL4]
- Workorder: 1906112 [ENV_LVL4]
- Workorder: 1906330 [ENV_LVL4]
- Workorder: 1906332 [ENV_LVL4]
- Workorder: 1906334 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mix	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	642096	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
2	642097	RLVS for HBN 233911 [ELMS/2223]				RLVS	3		E685041C3Q	5311		3/7/2019	
3	642098	ICS for HBN 233911 [ELMS/2223]				ICS	3		E6850.D3Q	5311		3/7/2019	
4	642099	LMB for HBN 233911 [ELMS/2223]				LMB	3		E6850Q413Q	5311		3/7/2019	
5	642100	LCS for HBN 233911 [ELMS/2223]				LCS	3		E6850Q413Q	5311		3/7/2019	
6	1905651001	HS19021158-02/LH18/24-SP650_02				SAMPLE	3	1905651001-A	E6850Q41.3	5480	3/21/2019	3/7/2019	
7	1906112001	EW01_022619				SAMPLE	3	1906112001-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
8	1906112002	EW01_022619MS				MS	3	1906112002-A	E6850Q413Q	5480		3/7/2019	
9	1906112003	EW01_022619MSD				MSD	3	1906112003-A	E6850Q413Q	5480		3/7/2019	
10	1906112004	EW05_022619				SAMPLE	3	1906112004-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
11	1906112005	EW05_022619_FD				FLDDUP	3	1906112005-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
12	1906112006	EW02_022619				SAMPLE	3	1906112006-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
13	1906112007	EW06_022619				SAMPLE	3	1906112007-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
14	1906112008	EW03_022619				SAMPLE	3	1906112008-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
15	1906112009	EW07_022619				SAMPLE	3	1906112009-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
16	642101	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
17	1906112010	EW04_022619				SAMPLE	3	1906112010-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
18	1906112011	EW08_022619				SAMPLE	3	1906112011-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
19	1906330001	LH18/24-SP140_022819				SAMPLE	3	1906330001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
20	1906332001	LH18/24-SP650_022819_BIX				SAMPLE	3	1906332001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
21	1906334001	LH18-24-SP650_022819_BIX				SAMPLE	3	1906334001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
22	642102	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1905651 (001); 1906112 (001-11); 1906330 (001); 1906332 (001); 1906334 (001)
 ELMS Batch/HBN ID: 2223 (233911)
 Prep Date: 03/04/2019 Analysis Date: 03/05/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\05MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by TNB. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 02/15/2019, sequence 15FEB19D.s Offline Quantitation Method: CLO4-DP1.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 3 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 642100; Target = 4.0µg/L. ASTM type II water was used for LMB 642099.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\23911-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEB104) along with datafiles 05MARD07-10.

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
Batch(es)/SDG: <u>E LMS: 2223 HBN: 233911</u>		
Sample Set IDs if Applicable: <u>1905651 / 190912 / 1906330</u> <u>1906332 / 1906334</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850.WKG Std 100.ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK			Description - 6850 Stock AccStd 1,000ug/mL
Standard: 43659		Created By: Thomas Bosch	
MFG: AccuStandard		Create Date: 09/17/2018 09:09AM	
MFG Lot: 218065075		Amount: 100 mL	
Part ID: IC-PER-10X-1		Expires: 07/25/2020	
		Usable: No	
		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT			Description - 6850 Intermdt AccStd 10.ug/mL		
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK			Description: 6850 QC WKG STD 100ug/L		
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdf Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type: II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK			Description: -6850 QC Stock STD 1,000ug/mL
Standard: 36748		Created By: Thomas Bosch	
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	
MFG Lot: CP-0860		Amount: 100 mL	
Part ID: ICC-013		Expires: 03/31/2020	
		Usable: Yes	
		Lab Lot: CLO4 QC STOCK	
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730		Created By: Thomas Bosch		Amount: 25 mL	
MFG: ALS/SLC		Create Date: 09/20/2018 09:09AM		Expires: 09/20/2019	
MFG Lot: TNB: 05/09/2018		Verified By: Thomas Bosch		Usable: Yes	
Pipette ID: Not Provided		Verify Date:		Lab Lot: CLO4ISTDWRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	.CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

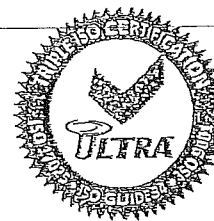
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:
This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:
Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:
The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:
This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:
This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:
Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

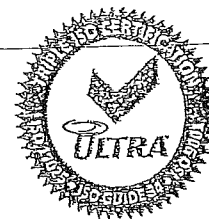
Hazards:
Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:
The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis

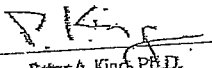


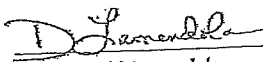
ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:
The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QAVRA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleared Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

For use in routine laboratory analysis.

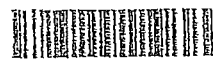


Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:

ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: NaCl*O4

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 µg/mL
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 ± 2.8 µg/mL (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount	
*	642096	CCV@25	Vial 71	1	Control	1	2.57589e6	8.017	25.04585
*	642100	QC@4.0	Vial 72	1	Control	2	4.23307e5	8.139	4.10749
*	642098	ICS@4.0	Vial 73	1	Control	3	3.17619e5	7.883	3.92708
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	7.71127e6	7.438	51.25151
*	1906112002	MS	Vial 77	1	Sample	7	8.02696e6	7.464	53.87972
*	1906112003	MSD	Vial 78	1	Sample	8	7.94242e6	7.442	53.69940
*	1906112004		Vial 79	1	Sample	9	2.33017e7	7.227	172.36522
*	1906112005		Vial 80	1	Sample	10	2.30991e7	7.229	180.38568
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.33611e6	8.114	25.01987
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	2.53126e6	7.736	226.48155
*	1906112005	10X	Vial 91	1	Sample	22	2.68237e6	7.739	241.03710
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	3.77363e6	8.090	3708.65666
*	642102	CCV@25	Vial 71	1	Control	25	2.15787e6	8.144	24.45533

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount	
*	642096	CCV@25	Vial 71	1	Control	1	6.79028e5	8.036	25.15984
*	642100	QC@4.0	Vial 72	1	Control	2	1.27412e5	8.156	4.45187
*	642098	ICS@4.0	Vial 73	1	Control	3	9.68721e4	7.906	4.29888
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	2.02786e6	7.452	51.10329
*	1906112002	MS	Vial 77	1	Sample	7	2.14637e6	7.478	54.46797
*	1906112003	MSD	Vial 78	1	Sample	8	2.10991e6	7.456	53.98100
*	1906112004		Vial 79	1	Sample	9	6.79668e6	7.243	180.45962
*	1906112005		Vial 80	1	Sample	10	6.82834e6	7.246	190.34382
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	6.28961e5	8.128	25.63410
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	6.46870e5	7.754	220.91905
*	1906112005	10X	Vial 91	1	Sample	22	6.95643e5	7.756	238.44674
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	1.01021e6	8.102	3771.67475
*	642102	CCV@25	Vial 71	1	Control	25	5.83711e5	8.160	25.16652

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount	
*	642096	CCV@25	Vial 71	1	Control	1	3.12207e5	8.043	5.00000
*	642100	QC@4.0	Vial 72	1	Control	2	3.41038e5	8.157	5.00000
*	642098	ICS@4.0	Vial 73	1	Control	3	2.68237e5	7.902	5.00000
*	642099	LMB	Vial 74	1	Control	4	3.53313e5	8.102	5.00000
*	1905651001		Vial 75	1	Sample	5	3.26356e5	7.780	5.00000
*	1906112001		Vial 76	1	Sample	6	4.26473e5	7.464	5.00000
*	1906112002	MS	Vial 77	1	Sample	7	4.19549e5	7.491	5.00000
*	1906112003	MSD	Vial 78	1	Sample	8	4.16709e5	7.467	5.00000
*	1906112004		Vial 79	1	Sample	9	2.95705e5	7.253	5.00000
*	1906112005		Vial 80	1	Sample	10	2.75946e5	7.263	5.00000
*	1906112006		Vial 81	1	Sample	11	3.73575e5	7.480	5.00000
*	1906112007		Vial 82	1	Sample	12	5.00533e5	7.256	5.00000
*	1906112008		Vial 83	1	Sample	13	4.65121e5	7.237	5.00000
*	1906112009		Vial 84	1	Sample	14	2.43675e5	7.672	5.00000
*	1906112010		Vial 85	1	Sample	15	2.51865e5	7.710	5.00000
*	1906112011		Vial 86	1	Sample	16	2.80792e5	7.876	5.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.83460e5	8.140	5.00000
*	1906332001		Vial 88	1	Sample	19	2.64674e5	7.827	5.00000
*	1906334001		Vial 89	1	Sample	20	2.65662e5	7.808	5.00000
*	1906112004	10X	Vial 90	1	Sample	21	3.41608e5	7.756	50.00000
*	1906112005	10X	Vial 91	1	Sample	22	3.38724e5	7.764	50.00000
*	1906112007	RE	Vial 82	1	Sample	23	4.31135e5	7.332	5.00000
*	1906330001	100	Vial 92	1	Sample	24	2.98985e5	8.111	500.00000
*	642102	CCV@25	Vial 71	1	Control	25	2.68305e5	8.165	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

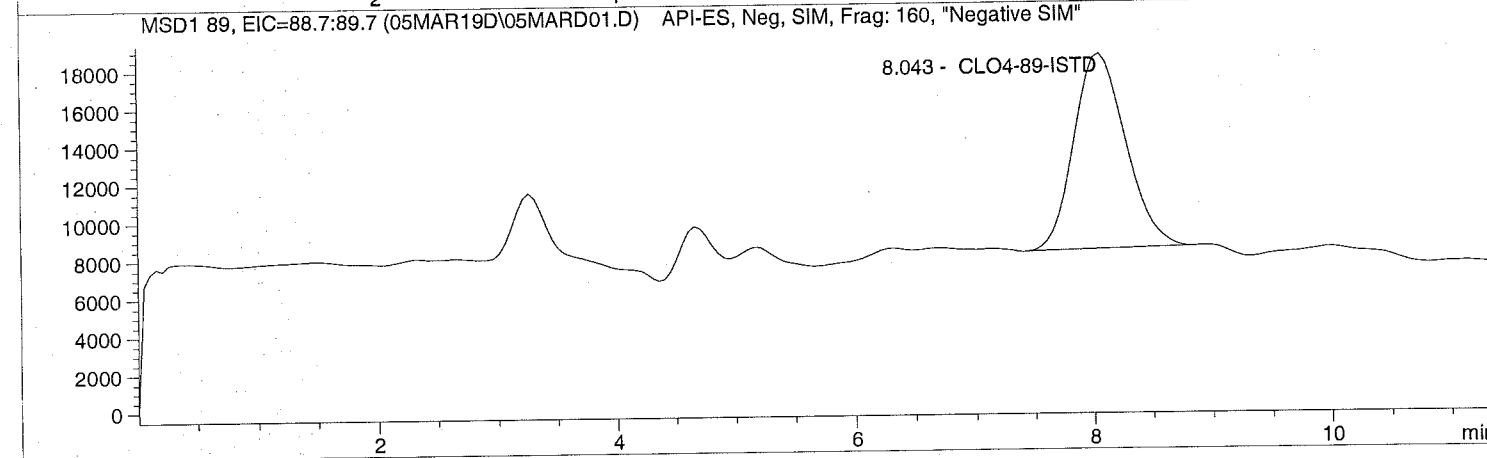
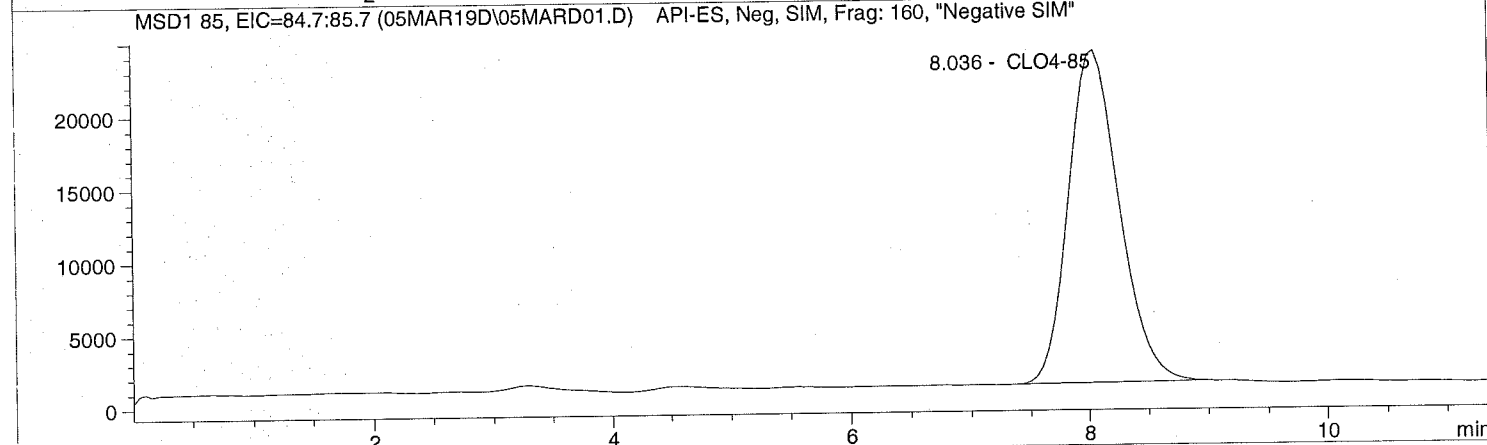
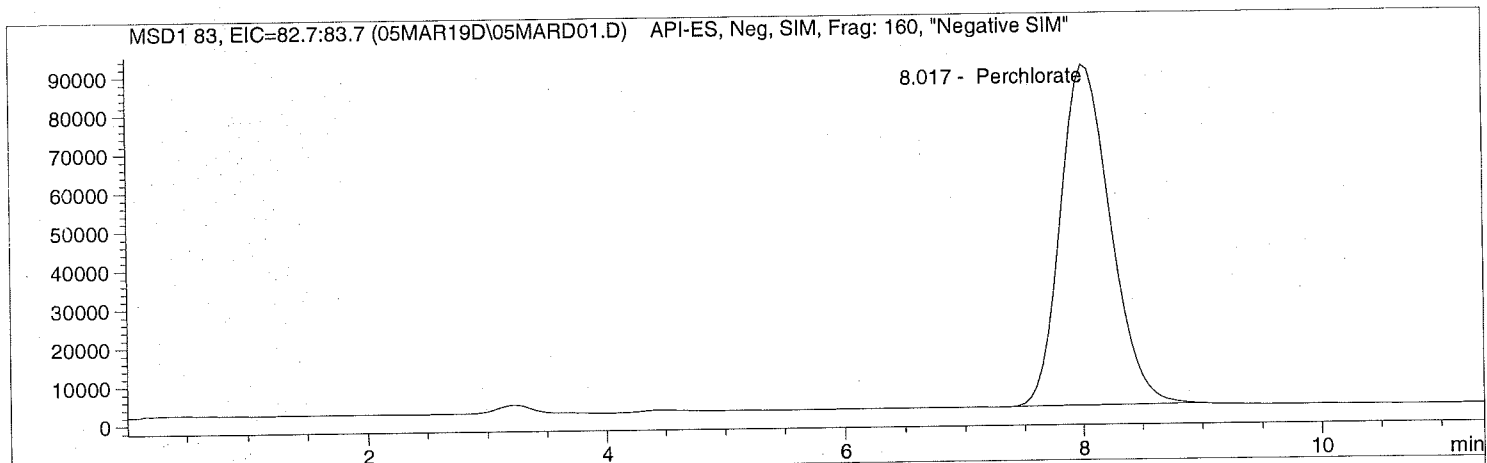
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	642096	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	642100	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	642098	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	642099	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1905651001		CLO4-AQN	1	Sample	
6	Vial 76	1906112001		CLO4-AQN	1	Sample	
7	Vial 77	1906112002	MS	CLO4-AQN	1	Sample	
8	Vial 78	1906112003	MSD	CLO4-AQN	1	Sample	
9	Vial 79	1906112004		CLO4-AQN	1	Sample	
10	Vial 80	1906112005		CLO4-AQN	1	Sample	
11	Vial 81	1906112006		CLO4-AQN	1	Sample	
12	Vial 82	1906112007		CLO4-AQN	1	Sample	
13	Vial 83	1906112008		CLO4-AQN	1	Sample	
14	Vial 84	1906112009		CLO4-AQN	1	Sample	
15	Vial 85	1906112010		CLO4-AQN	1	Sample	
16	Vial 86	1906112011		CLO4-AQN	1	Sample	
17	Vial 71	642101	CCV@25	CLO4-AQN	1	Ctrl Samp	
18	Vial 87	1906330001	1K	CLO4-AQN	1	Sample	
19	Vial 88	1906332001		CLO4-AQN	1	Sample	
20	Vial 89	1906334001		CLO4-AQN	1	Sample	
21	Vial 90	1906112004	10X	CLO4-AQN	1	Sample	
22	Vial 91	1906112005	10X	CLO4-AQN	1	Sample	
23	Vial 82	1906112007	RE	CLO4-AQN	1	Sample	
24	Vial 92	1906330001	100	CLO4-AQN	1	Sample	
25	Vial 71	642102	CCV@25	CLO4-AQN	1	Ctrl Samp	

Injection Date: 3/05/2019 08:44:45
Sample Name: 642096 CCV@25
Acq Operator: TNB

Seq Line: 1
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 08:44:45 Seq Line: 1
Sample Name: 642096 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.017	PBA	2575886.3	25.0459	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.036	PBA	679028.4	25.1598	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.043	PBA	312206.9	5.0000	CLO4-89-ISTD

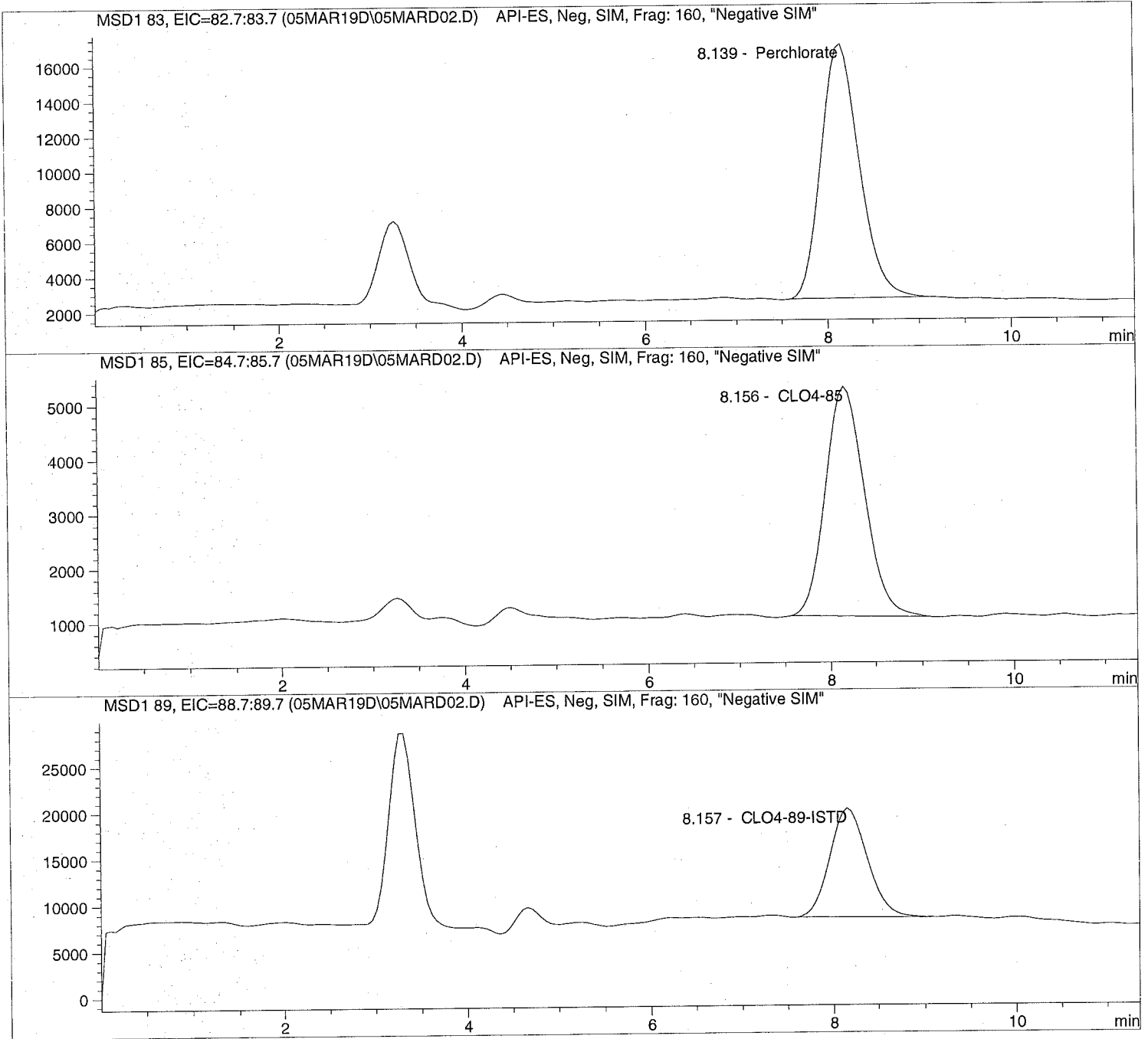
*** End of Report ***

Injection Date: 3/05/2019 09:00:30
Sample Name: 642100 QC@4.0
Acq Operator: TNB

Seq Line: 2
Location: Vial 72
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 09:00:30      Seq Line: 2
Sample Name: 642100 QC@4.0              Location: Vial 72
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.139	PBA	423307.1	4.1075	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.156	PBA	127412.0	4.4519	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

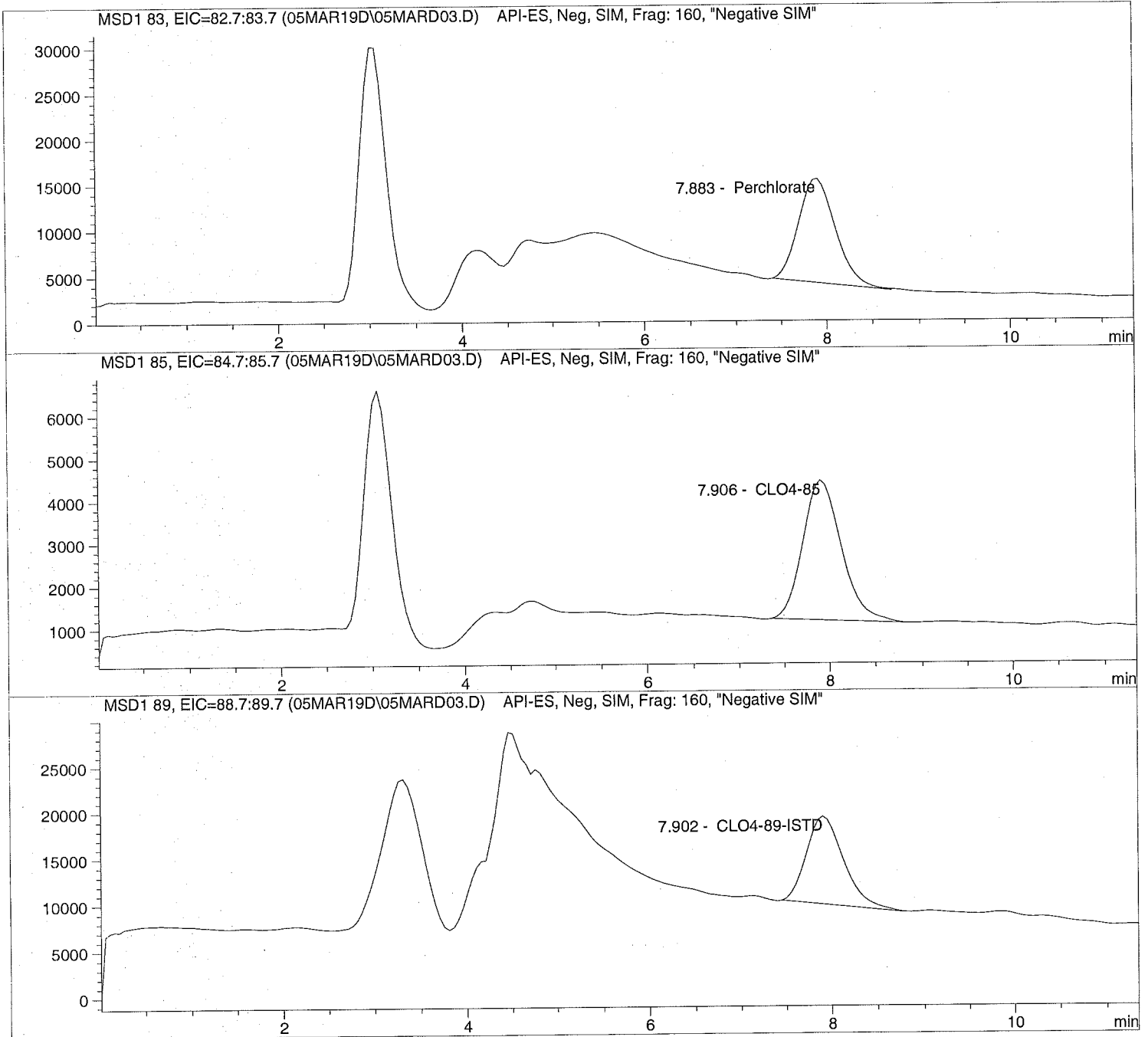
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.157	PBA	341038.3	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

Injection Date: 3/05/2019 09:13:34 Seq Line: 3
Sample Name: 642098 ICS@4.0 Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 09:13:34 Seq Line: 3
Sample Name: 642098 ICS@4.0 Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.883	PBA	317618.9	3.9271	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.906	PBA	96872.1	4.2989	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.902	PBA	268236.7	5.0000	CLO4-89-ISTD

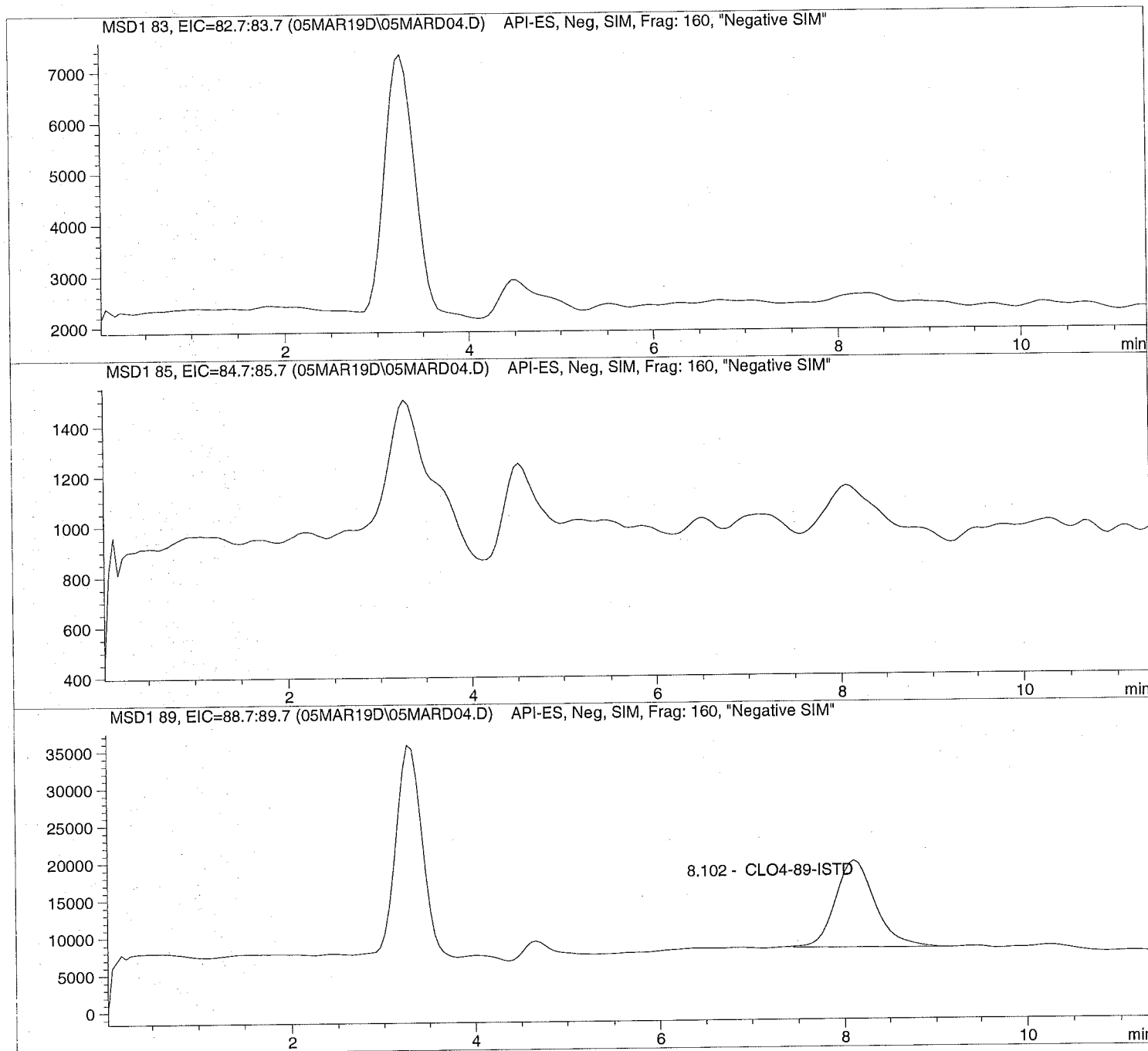
*** End of Report ***

Injection Date: 3/05/2019 09:26:40
Sample Name: 642099 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 09:26:40 Seq Line: 4
Sample Name: 642099 LMB Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	BBA	353313.1	5.0000	CLO4-89-ISTD

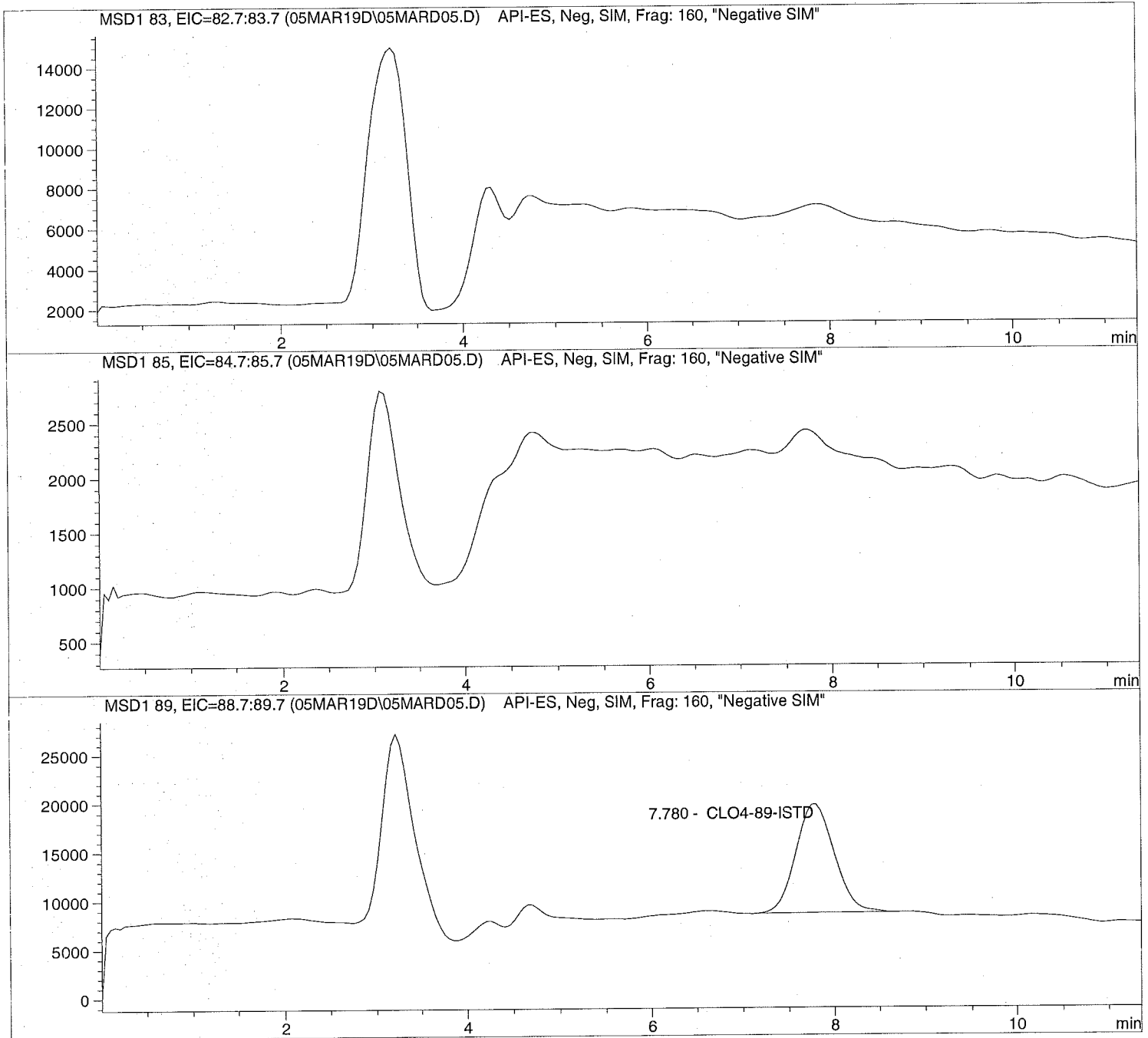
*** End of Report ***

Injection Date: 3/05/2019 09:40:58
Sample Name: 1905651001
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 09:40:58      Seq Line: 5
Sample Name: 1905651001                 Location: Vial 75
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.780	PBA	326356.2	5.0000	CLO4-89-ISTD

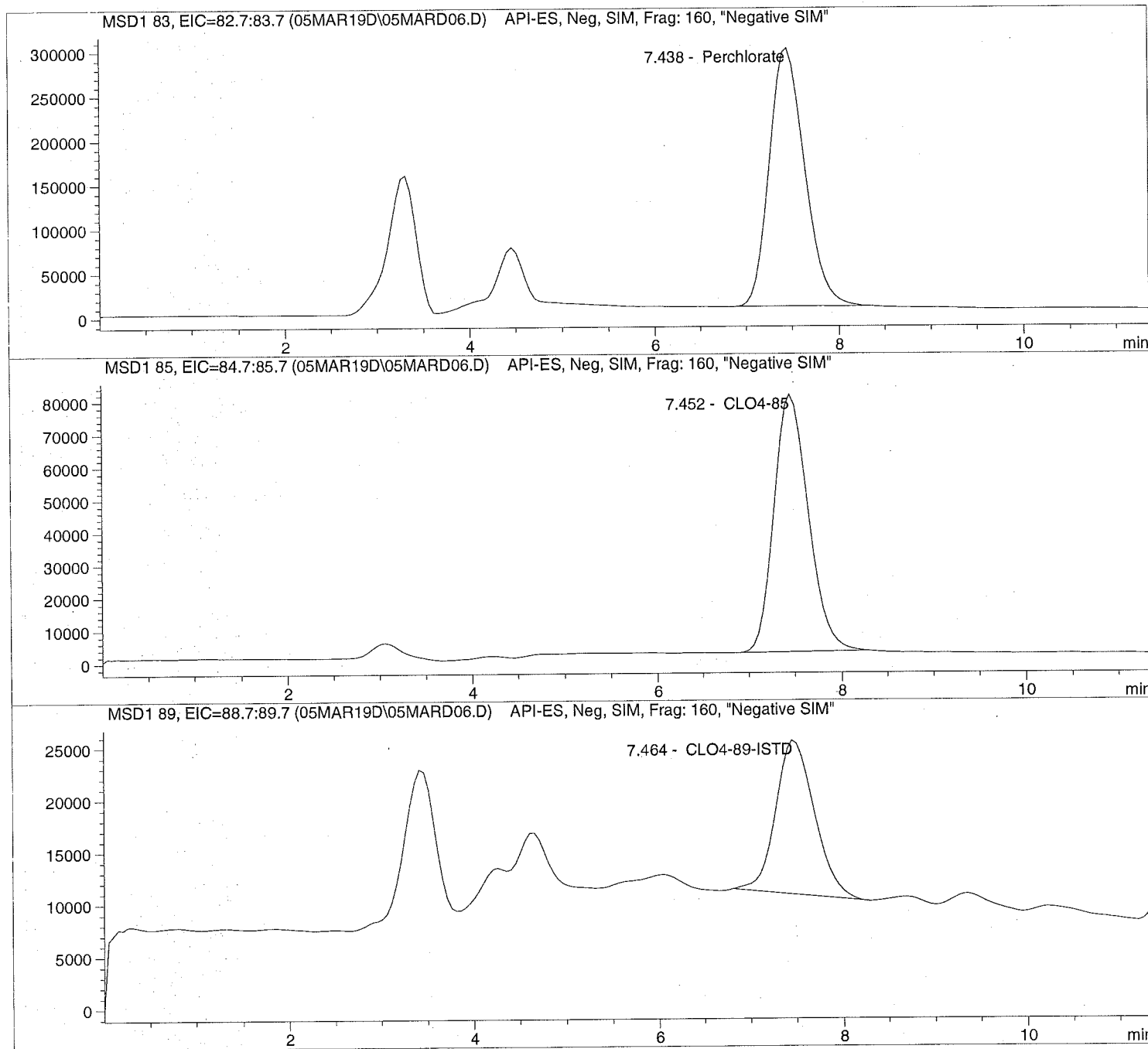
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*** End of Report ***

Injection Date: 3/05/2019 09:54:04
Sample Name: 1906112001
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 09:54:04      Seg Line: 6  
Sample Name: 1906112001                 Location: Vial 76  
Acq Operator: TNB                       Inj. No.: 1  
                                           Inj. Vol.: 20 µl
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.438	PBA	7711270.5	51.2515	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.452	PBA	2027855.1	51.1033	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PB	426473.5	5.0000	CLO4-89-ISTD

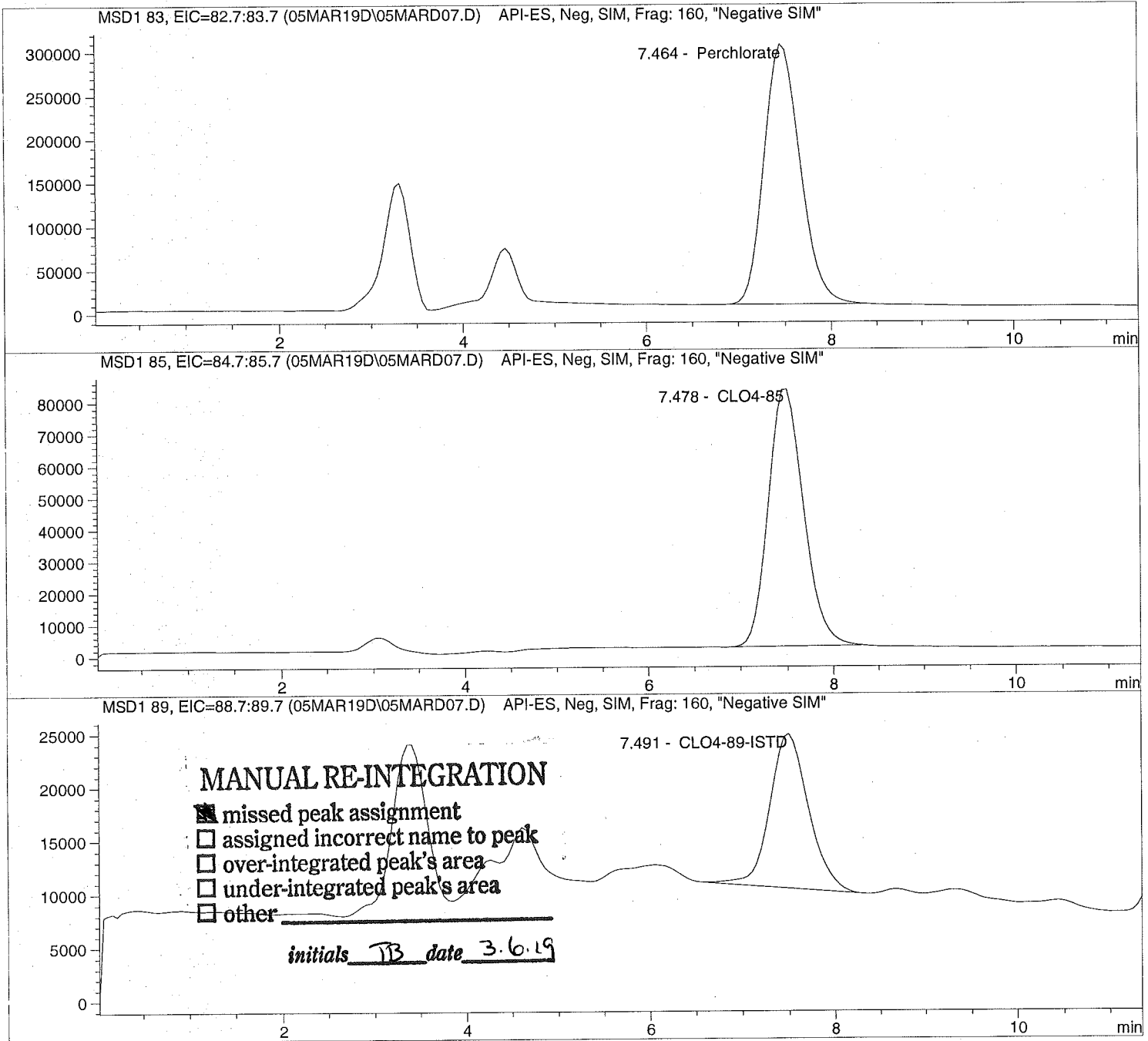
=====
*** End of Report ***

Injection Date: 3/05/2019 10:07:11
Sample Name: 1906112002 MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis




```
=====
Injection Date: 3/05/2019 10:07:11      Seq Line: 7
Sample Name: 1906112002 MS              Location: Vial 77
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	53.8797	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	54.4680	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.491	MM	419549.0	5.0000	CLO4-89-ISTD

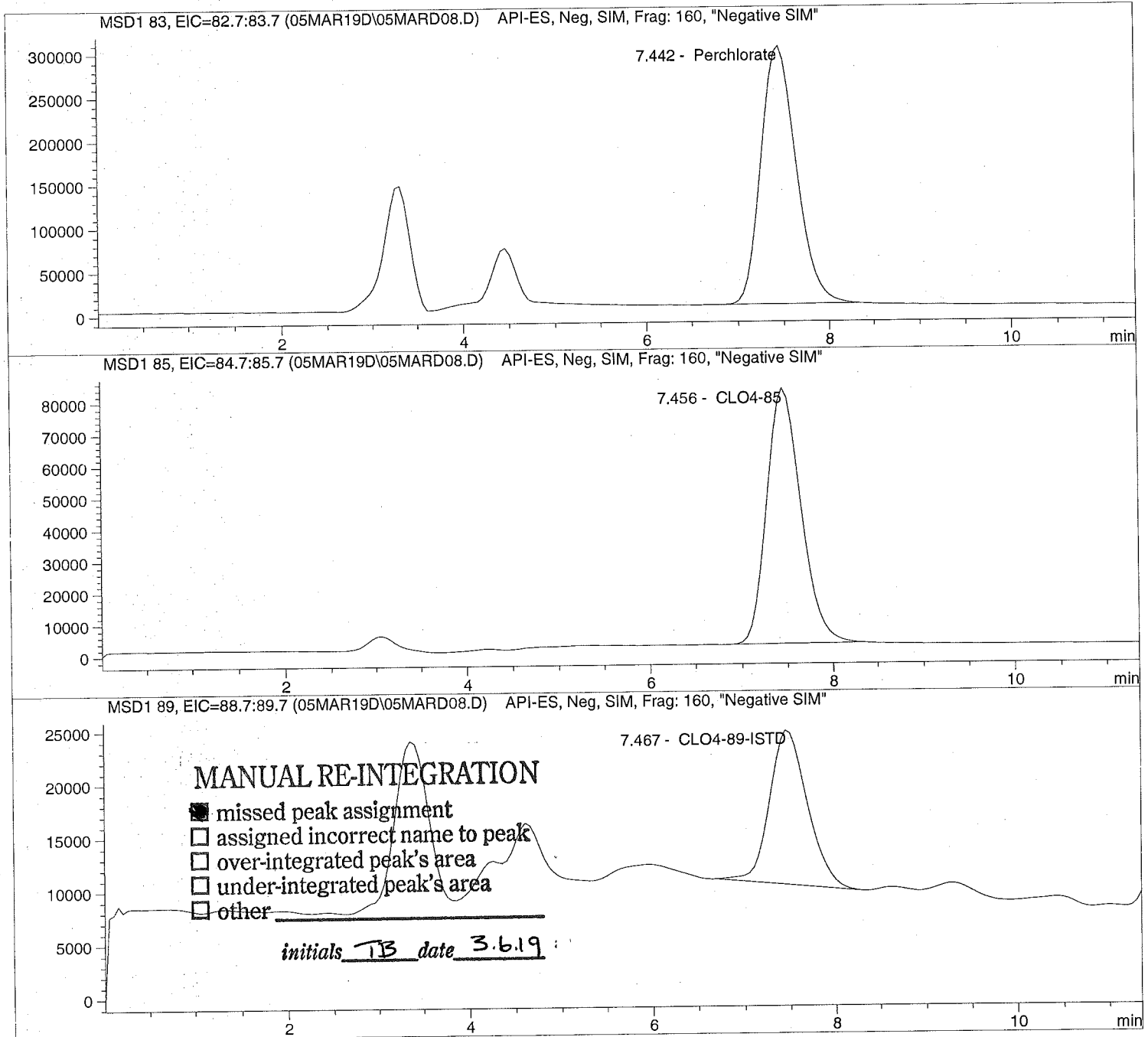
=====
*** End of Report ***

Injection Date: 3/05/2019 10:20:17
Sample Name: 1906112003 MSD
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	53.6994	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	53.9810	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.467	MM	416709.1	5.0000	CLO4-89-ISTD

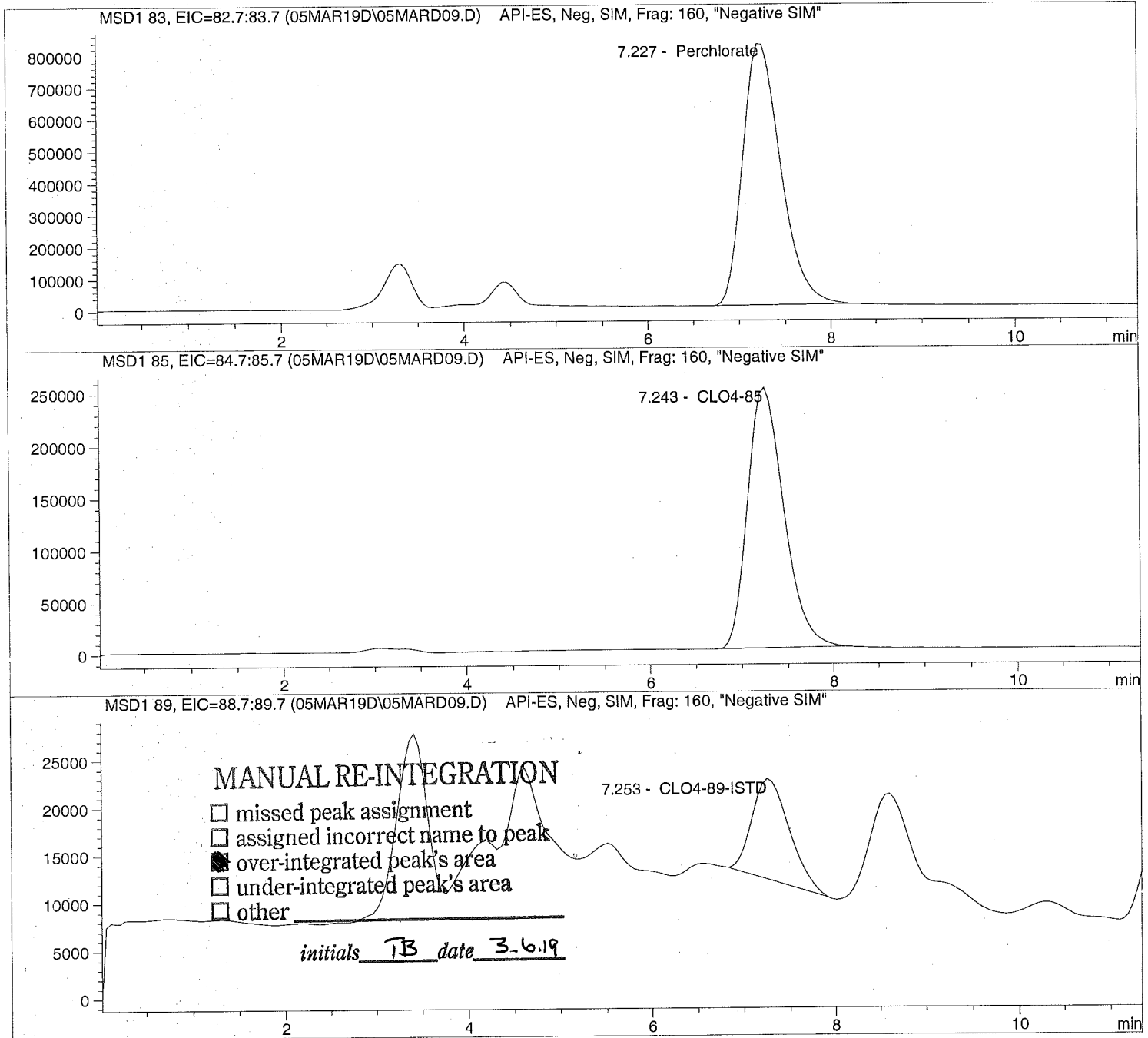
*** End of Report ***

Injection Date: 3/05/2019 10:33:21
Sample Name: 1906112004
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 10:33:21      Seq Line:          9
Sample Name:    1906112004              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	172.3652	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	180.4596	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.253	MM	295705.2	5.0000	CLO4-89-ISTD

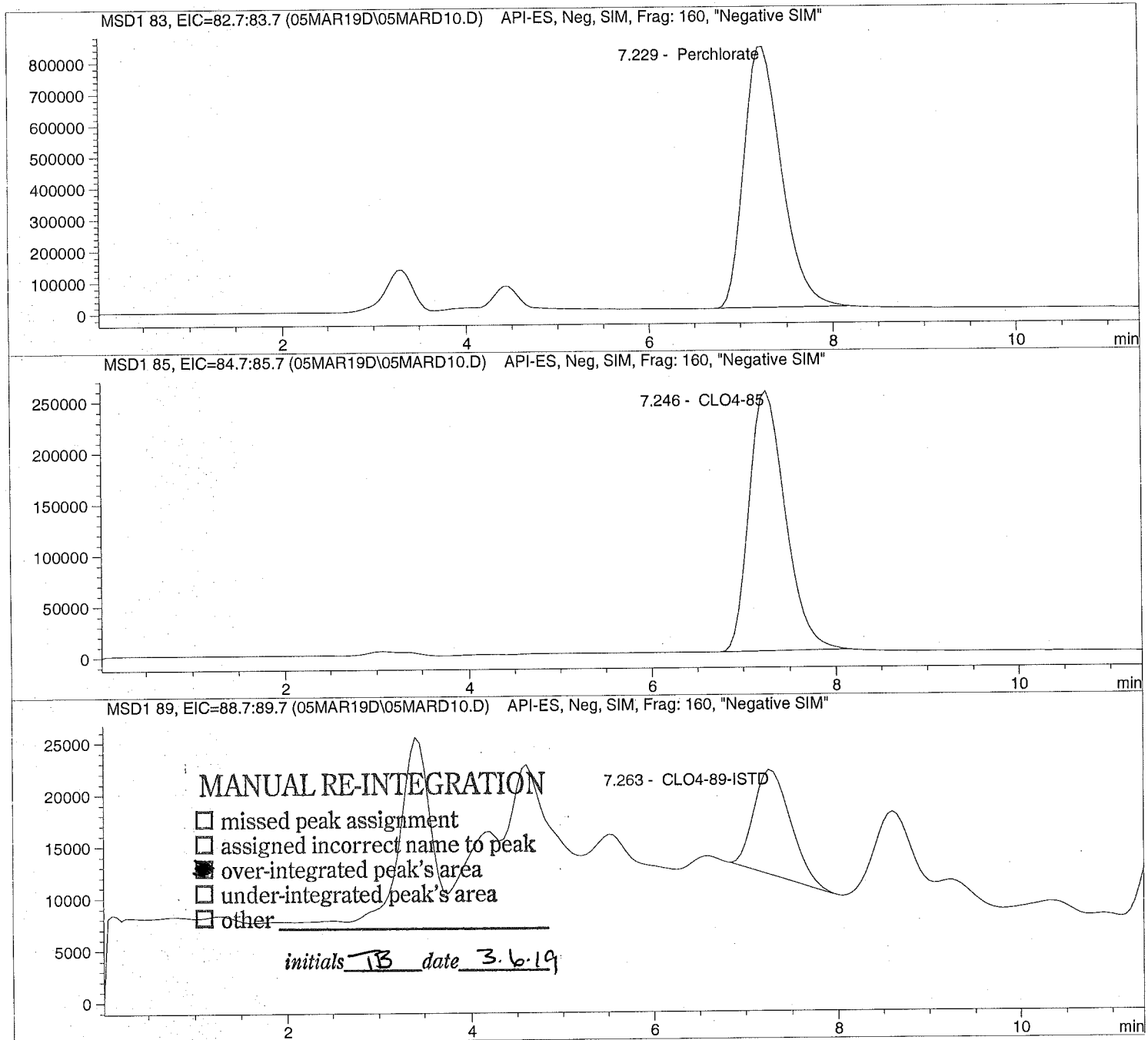
=====
*** End of Report ***

Injection Date: 3/05/2019 10:46:26
Sample Name: 1906112005
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```

=====
Injection Date: 3/05/2019 10:46:26      Seq Line: 10
Sample Name: 1906112005                  Location: Vial 80
Acq Operator: TNB                         Inj. No.: 1
                                           Inj. Vol.: 20 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	180.3857	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	190.3438	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	MM	275946.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

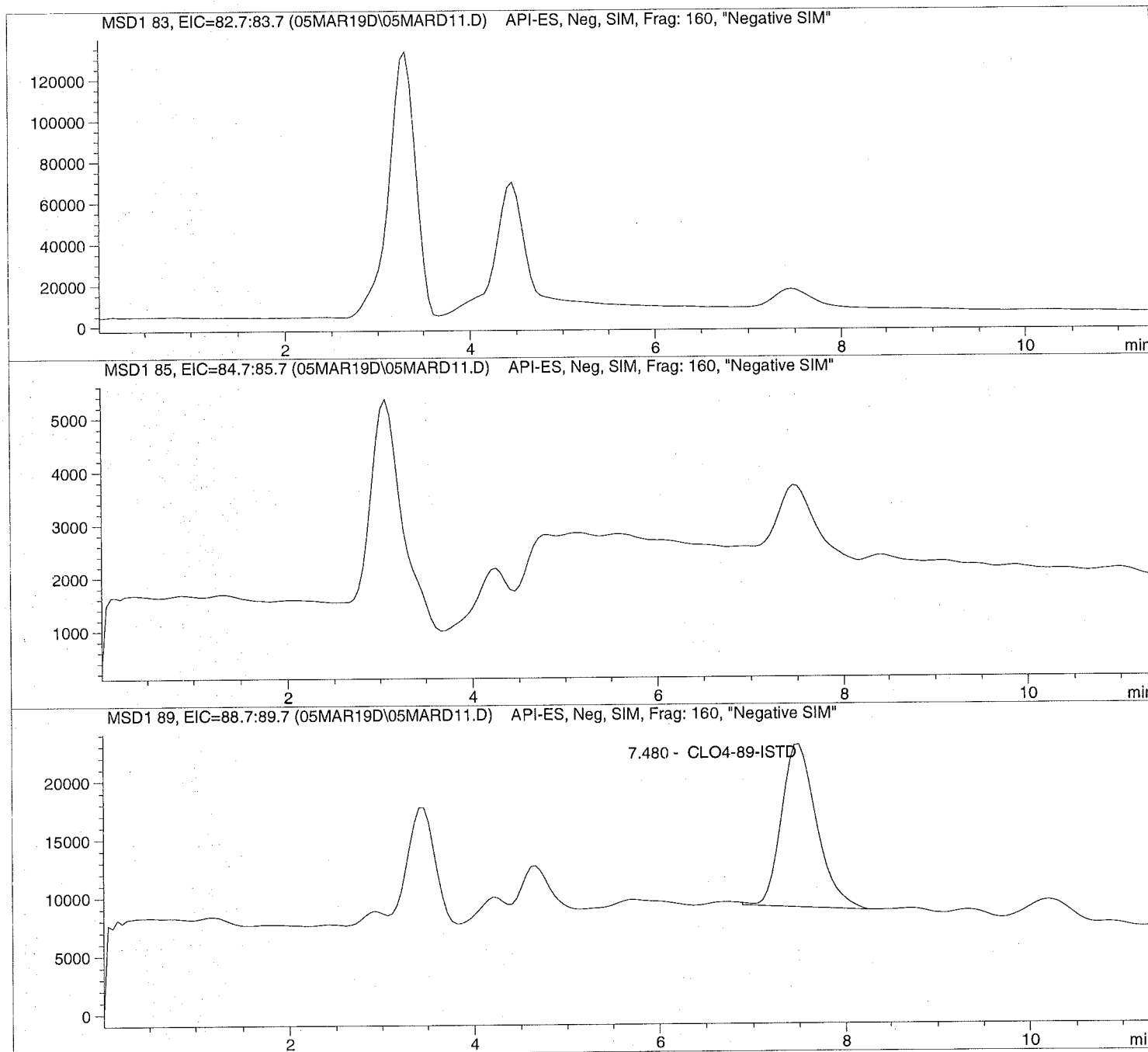
```

Injection Date: 3/05/2019 10:59:36
Sample Name: 1906112006
Acq Operator: TNB

Seq Line: 11
Location: Vial 81
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis




```
=====
Injection Date:  3/05/2019  10:59:36      Seq Line:      11
Sample Name:    1906112006                Location:      Vial 81
Acq Operator:   TNB                       Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  12:13:46
=====
```

Perchlorate analysis

```
=====
Sample Information
=====
```

```
Sorted By:      Signal
Calib. Data Modified:  Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====
```

```
=====
LCMS Results
=====
```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.480	BBA	373575.4	5.0000	CLO4-89-ISTD

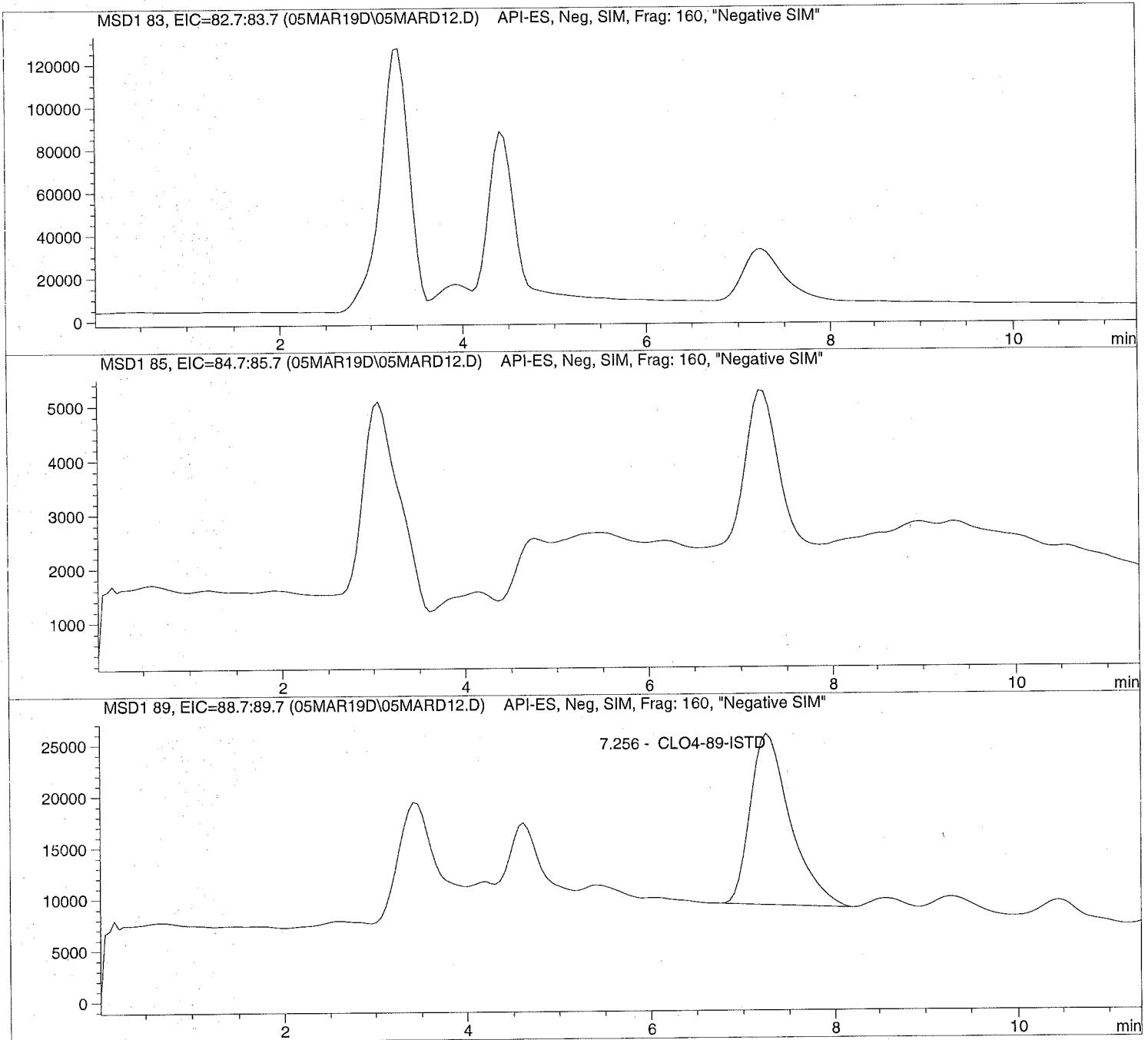
```
=====
*** End of Report ***
=====
```

Injection Date: 3/05/2019 11:12:28
Sample Name: 1906112007
Acq Operator: TNB

Seq Line: 12
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 11:12:28      Seq Line: 12  
Sample Name: 1906112007                Location: Vial 82  
Acq Operator: TNB                      Inj. No.: 1  
                                         Inj. Vol.: 20 µl
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.256	PB	500532.9	5.0000	CLO4-89-ISTD

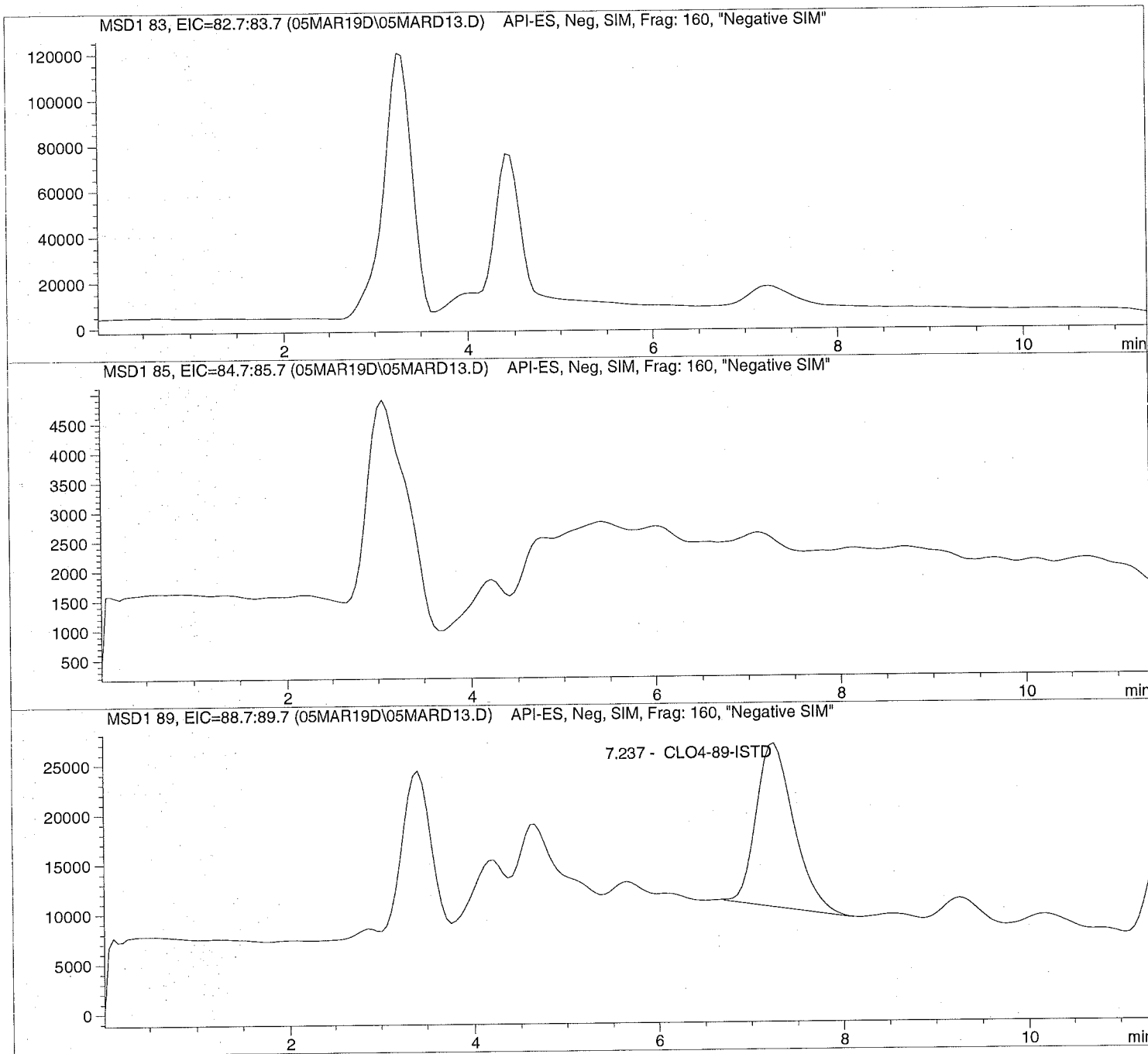
=====
*** End of Report ***

Injection Date: 3/05/2019 11:26:23
Sample Name: 1906112008
Acq Operator: TNB

Seq Line: 13
Location: Vial 83
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```

=====
Injection Date: 3/05/2019 11:26:23      Seq Line: 13
Sample Name: 1906112008                 Location: Vial 83
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.237	BBA	465121.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

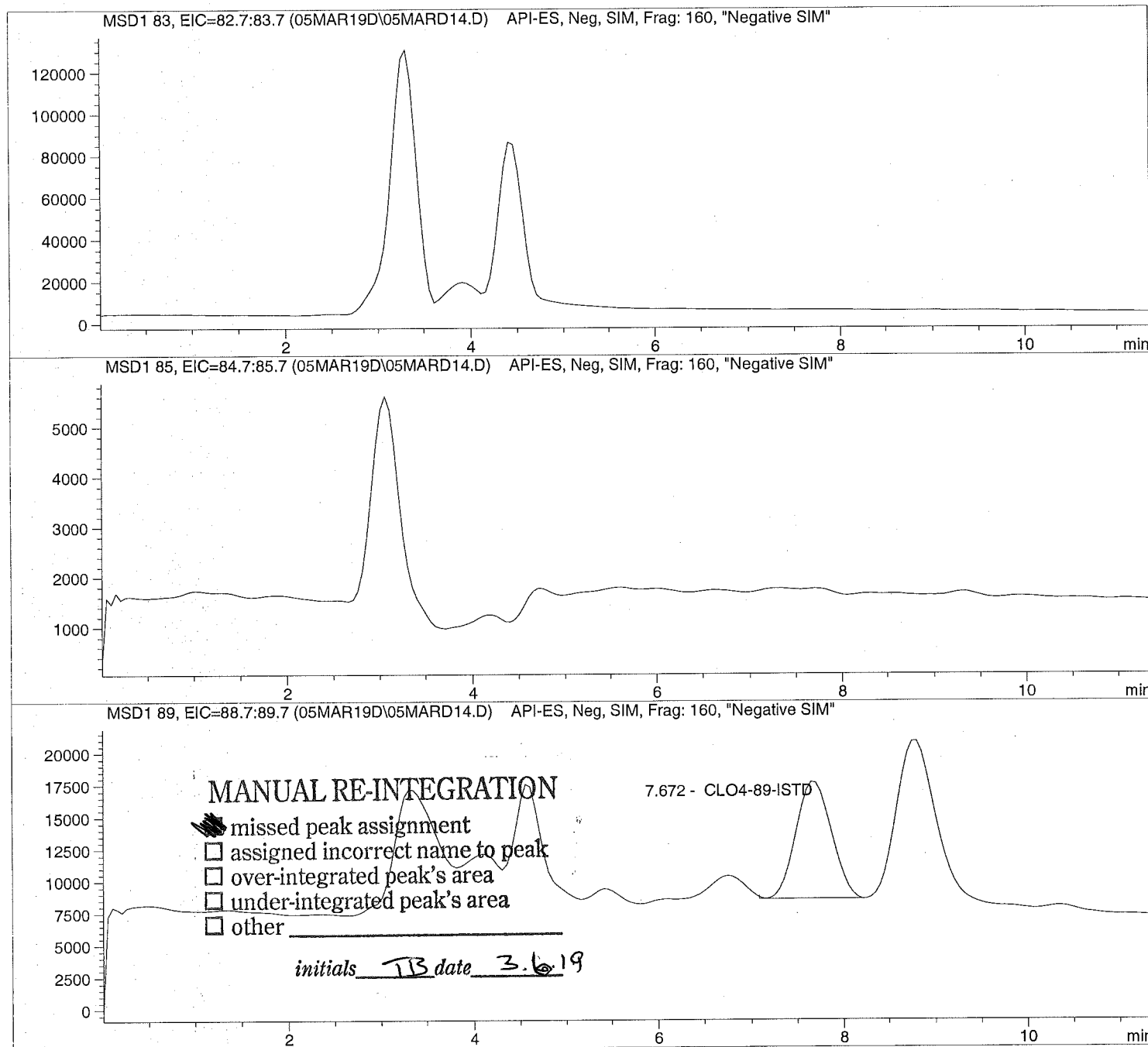
```

Injection Date: 3/05/2019 11:39:24
Sample Name: 1906112009
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 11:39:24 Seq Line: 14
Sample Name: 1906112009 Location: Vial 84
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	5.0000	CLO4-89-ISTD

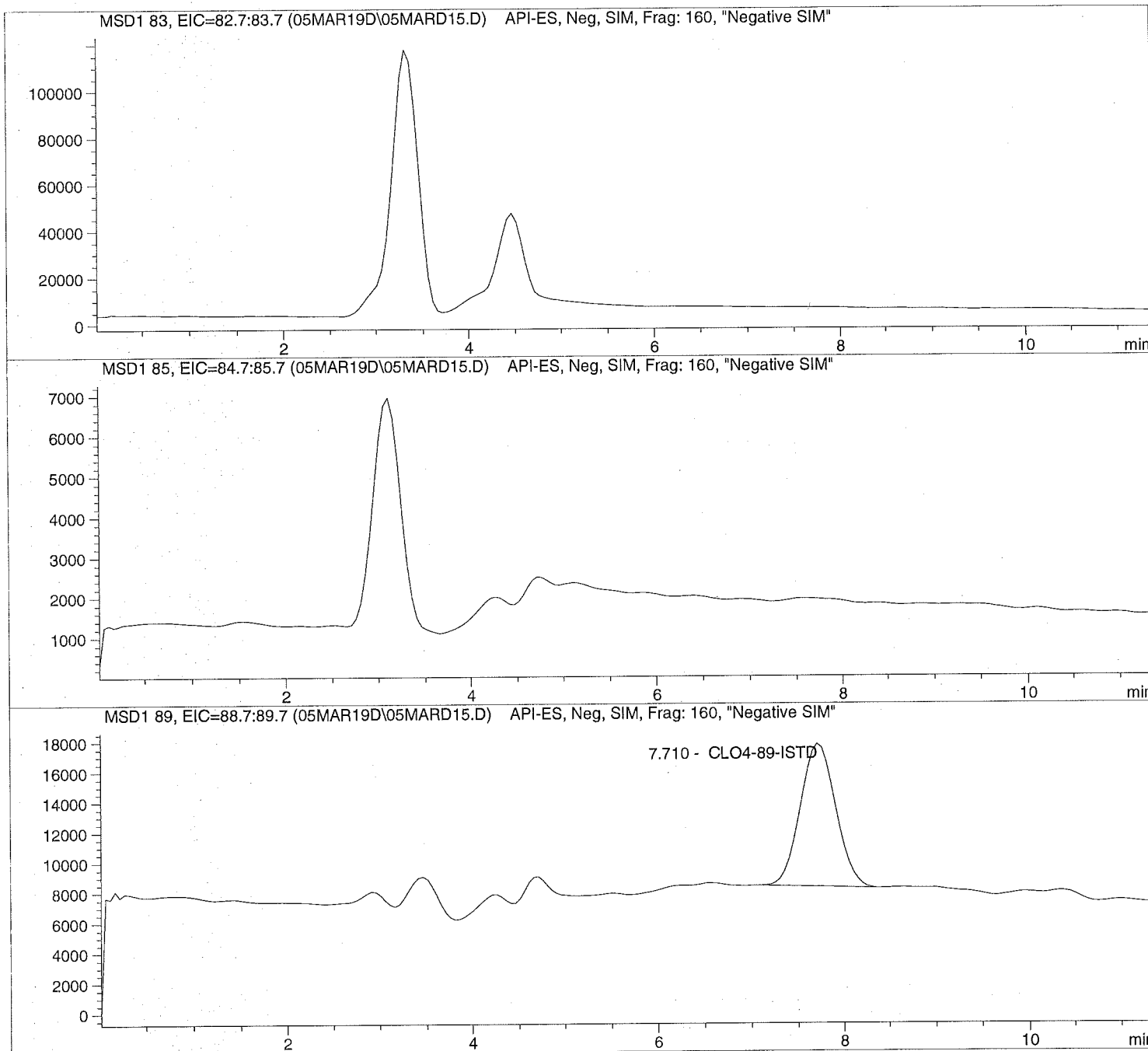
*** End of Report ***

Injection Date: 3/05/2019 11:52:27
Sample Name: 1906112010
Acq Operator: TNB

Seq Line: 15
Location: Vial 85
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis




```
=====
Injection Date: 3/05/2019 11:52:27      Seq Line: 15
Sample Name: 1906112010                 Location: Vial 85
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.710	BBA	251865.4	5.0000	CLO4-89-ISTD

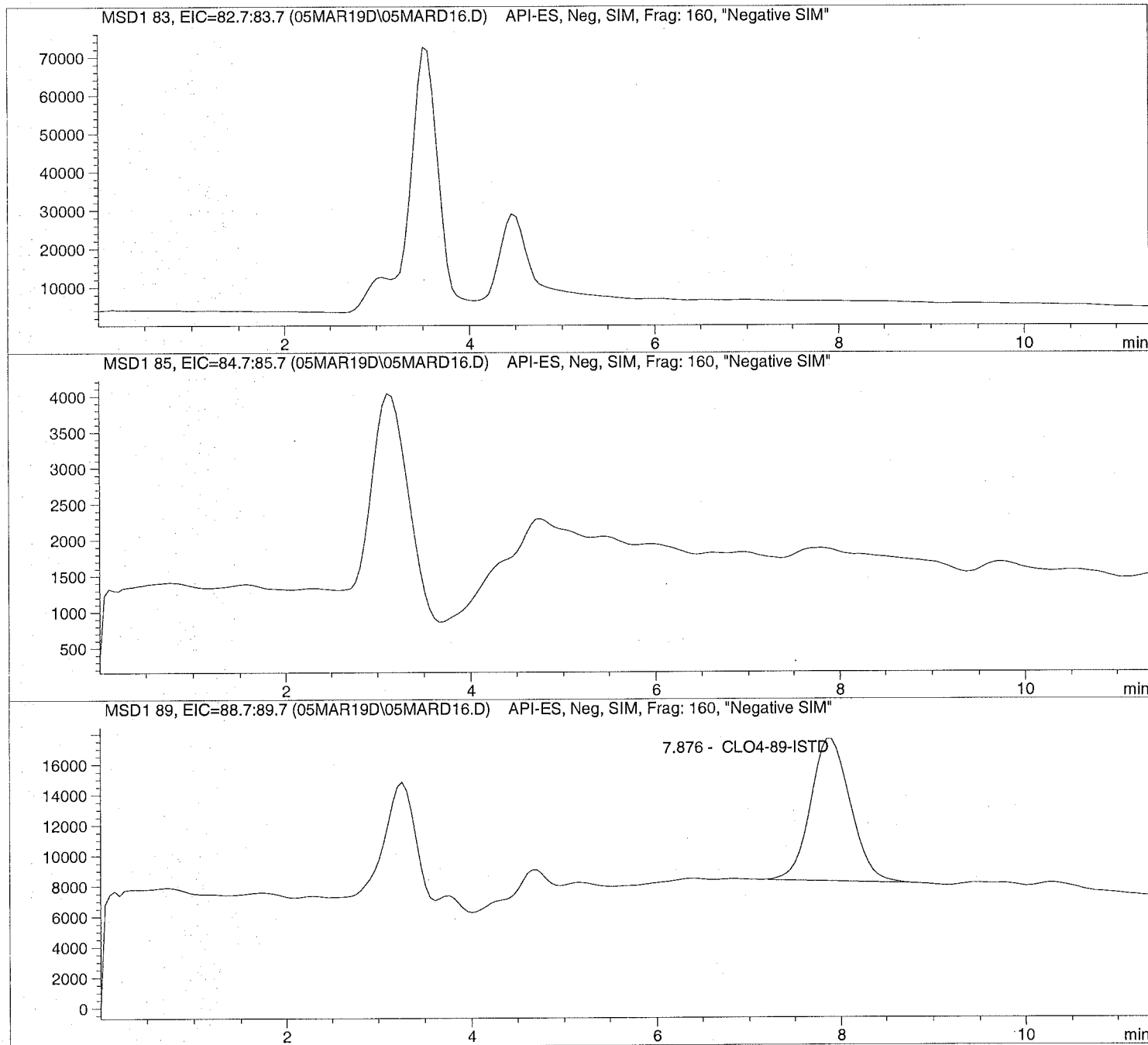
=====
*** End of Report ***

Injection Date: 3/05/2019 12:05:39
Sample Name: 1906112011
Acq Operator: TNB

Seq Line: 16
Location: Vial 86
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 12:05:39      Seq Line: 16
Sample Name: 1906112011                 Location: Vial 86
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

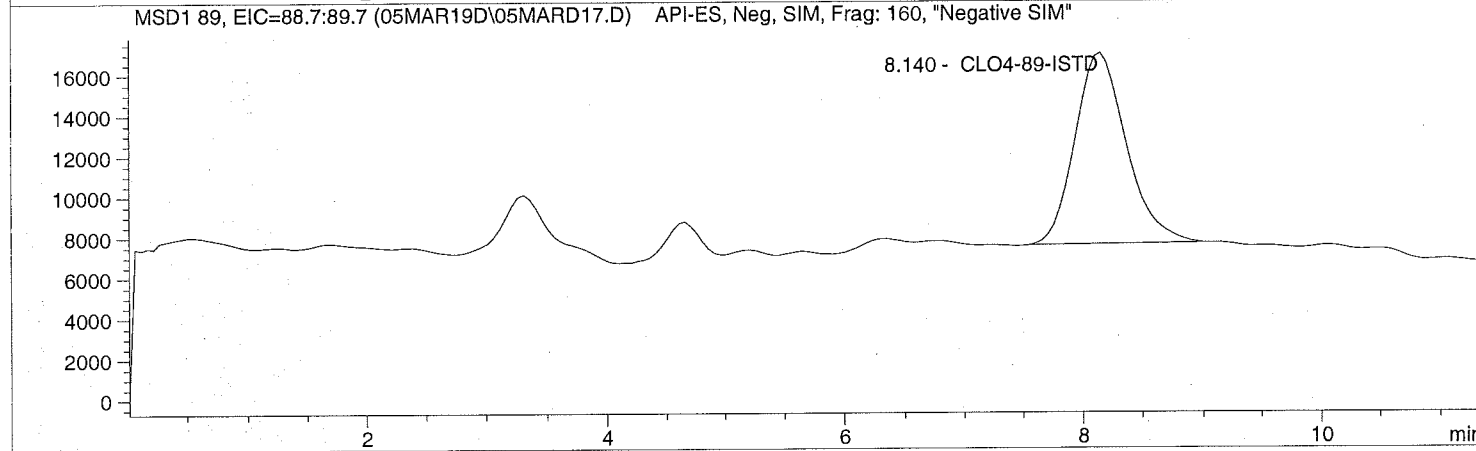
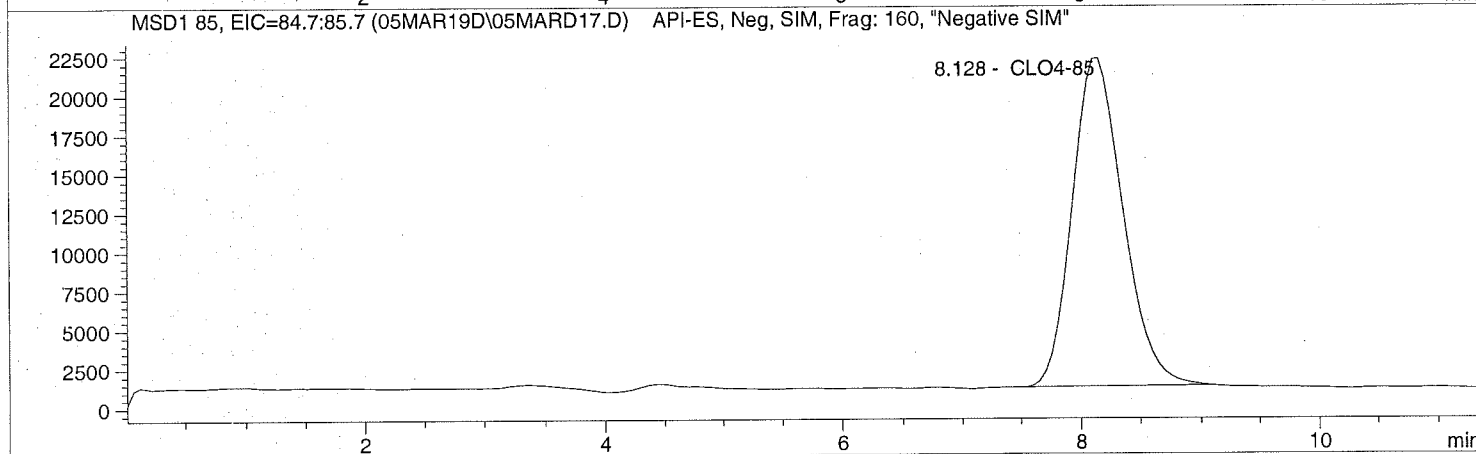
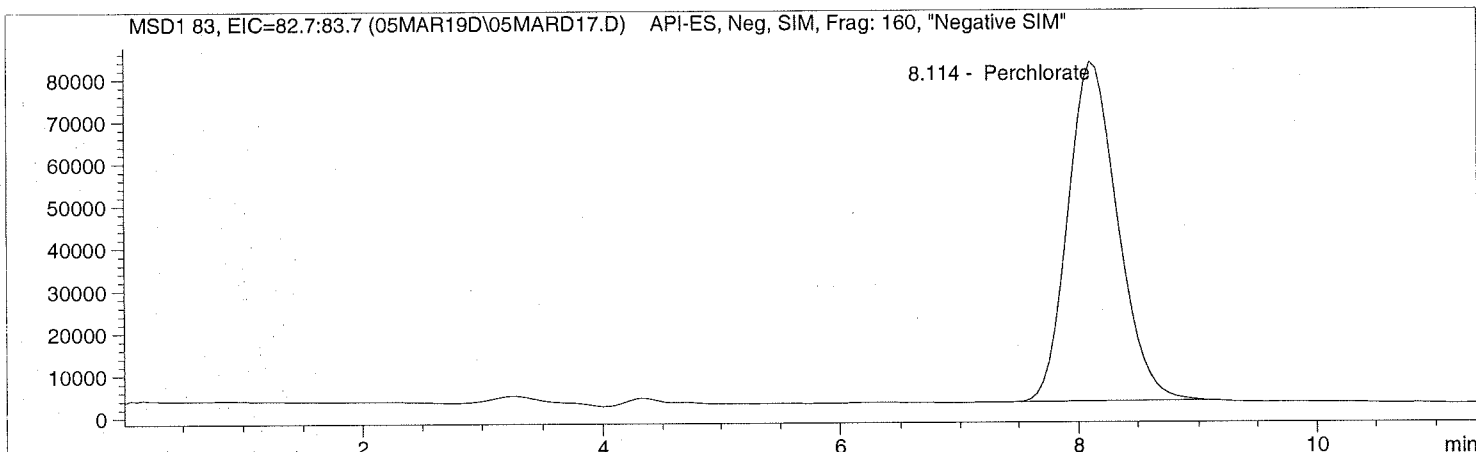
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.876	BBA	280792.2	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

Injection Date: 3/05/2019 12:18:41 Seq Line: 17
Sample Name: 642101 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 12:18:41 Seq Line: 17
Sample Name: 642101 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	2336112.7	25.0199	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.128	PBA	628961.2	25.6341	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.140	PBA	283460.4	5.0000	CLO4-89-ISTD

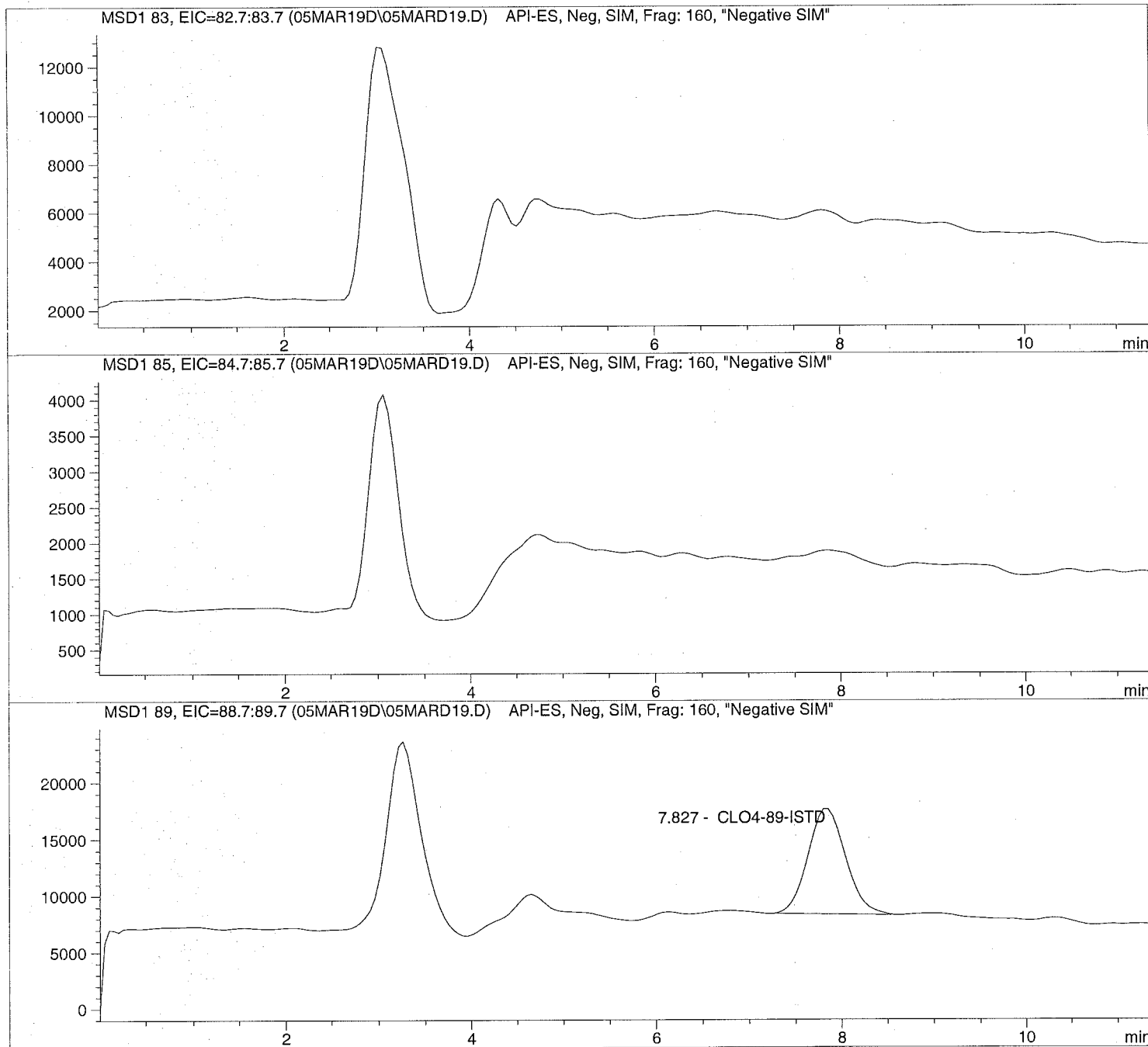
*** End of Report ***

Injection Date: 3/05/2019 12:49:07
Sample Name: 1906332001
Acq Operator: TNB

Seq Line: 19
Location: Vial 88
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 12:49:07      Seq Line:      19
Sample Name:    1906332001              Location:     Vial 88
Acq Operator:   TNB                     Inj. No.:    1
                                           Inj. Vol.:   20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:      1.000000
Sample Amount:  0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.827	PBA	264674.3	5.0000	CLO4-89-ISTD

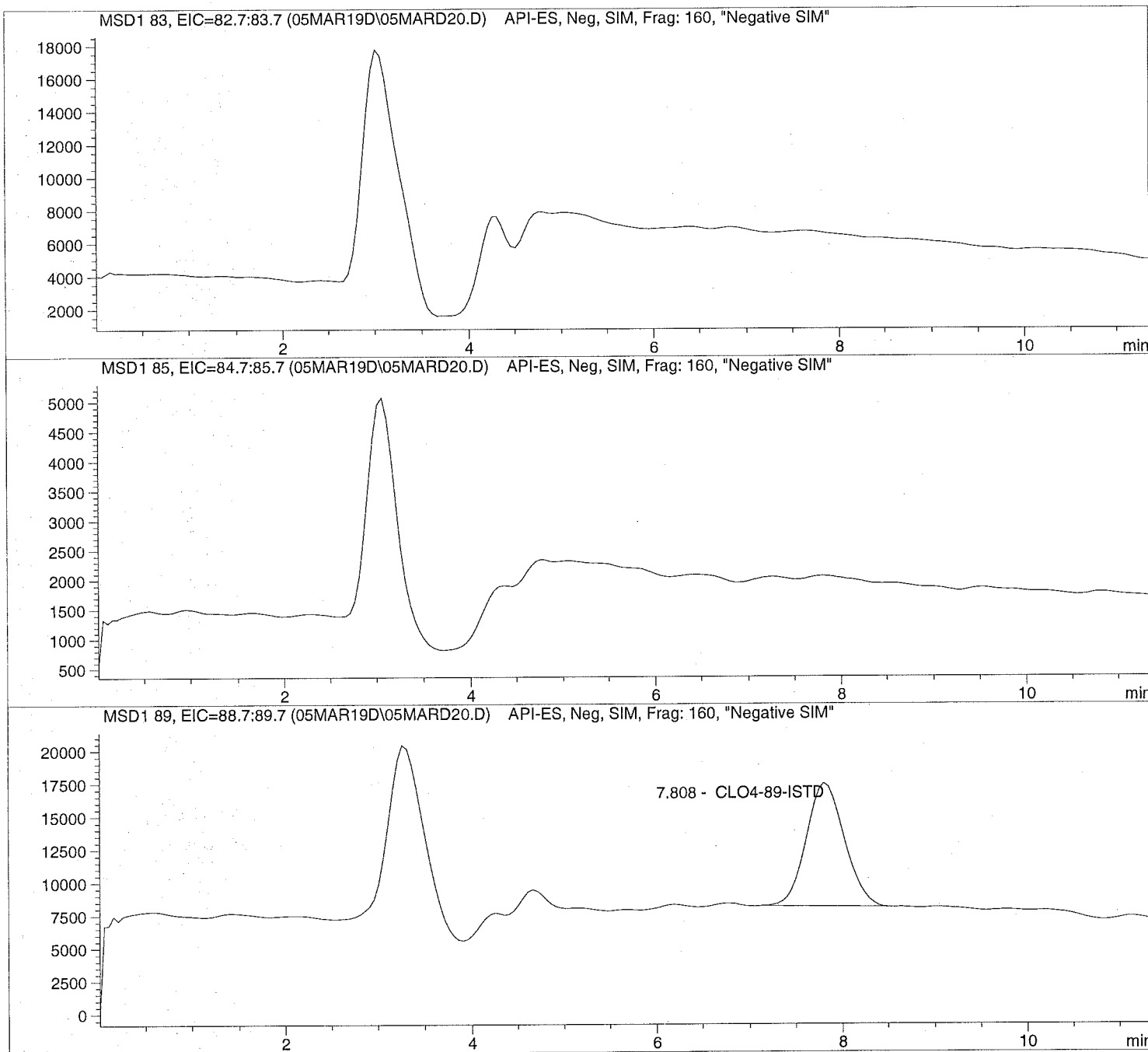
=====
*** End of Report ***

Injection Date: 3/05/2019 13:02:15
Sample Name: 1906334001
Acq Operator: TNB

Seq Line: 20
Location: Vial 89
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis




```
=====
Injection Date: 3/05/2019 13:02:15      Seq Line: 20
Sample Name: 1906334001                 Location: Vial 89
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.808	PBA	265662.0	5.0000	CLO4-89-ISTD

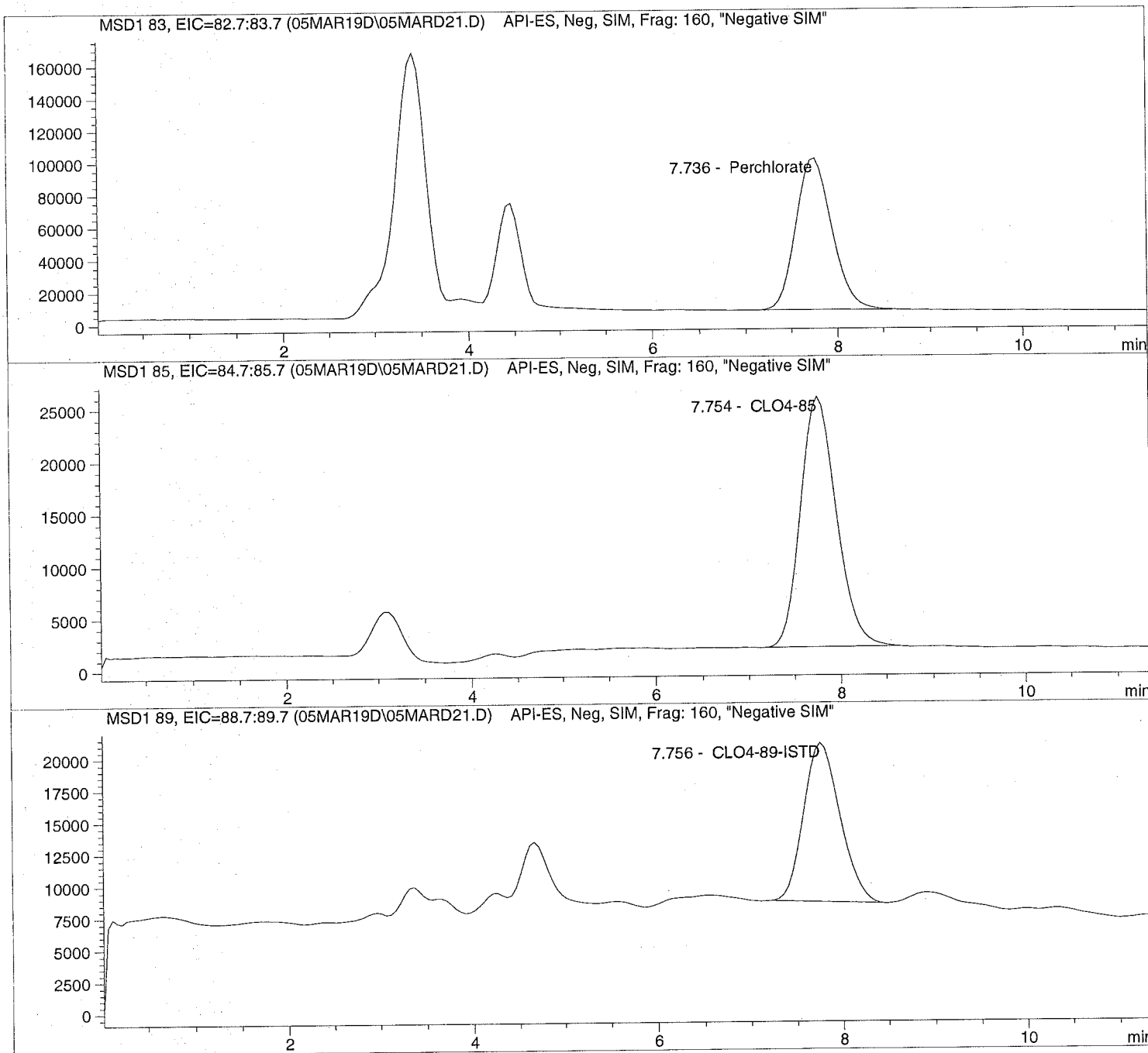
=====
*** End of Report ***

Injection Date: 3/05/2019 13:15:18
Sample Name: 1906112004 10X
Acq Operator: TNB

Seq Line: 21
Location: Vial 90
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 13:15:18 Seq Line: 21
Sample Name: 1906112004 10X Location: Vial 90
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 10.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.736	PBA	2531259.3	226.4815	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.754	PBA	646870.4	220.9191	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	PB	341607.9	50.0000	CLO4-89-ISTD

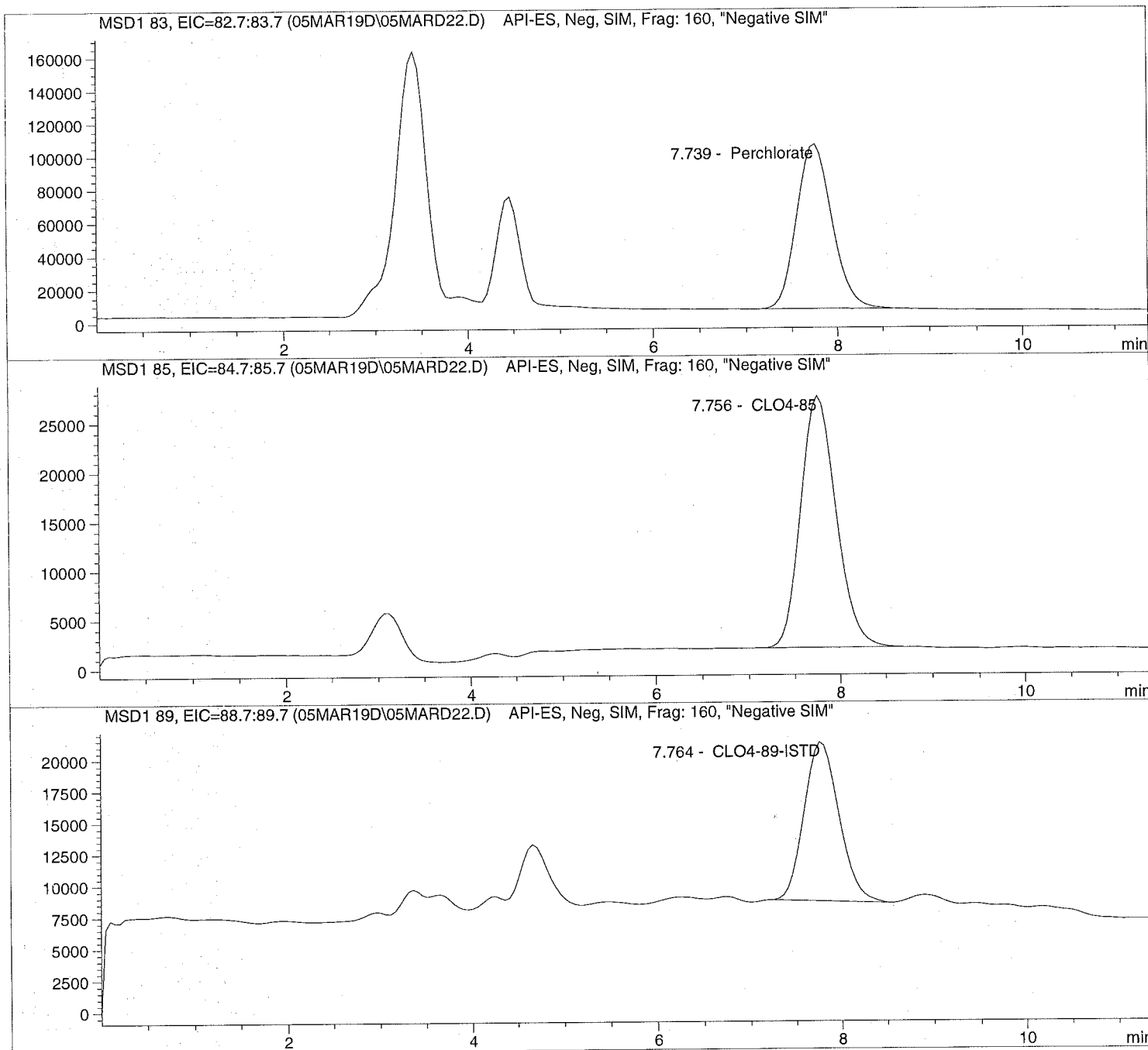
*** End of Report ***

Injection Date: 3/05/2019 13:28:32
Sample Name: 1906112005 10X
Acq Operator: TNB

Seq Line: 22
Location: Vial 91
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



=====
Injection Date: 3/05/2019 13:28:32 Seq Line: 22
Sample Name: 1906112005 10X Location: Vial 91
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

=====
Sample Information
=====

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 10.000000
Sample Amount: 0.000

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.739	PBA	2682370.5	241.0371	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	BBA	695643.1	238.4467	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.764	PB	338724.1	50.0000	CLO4-89-ISTD

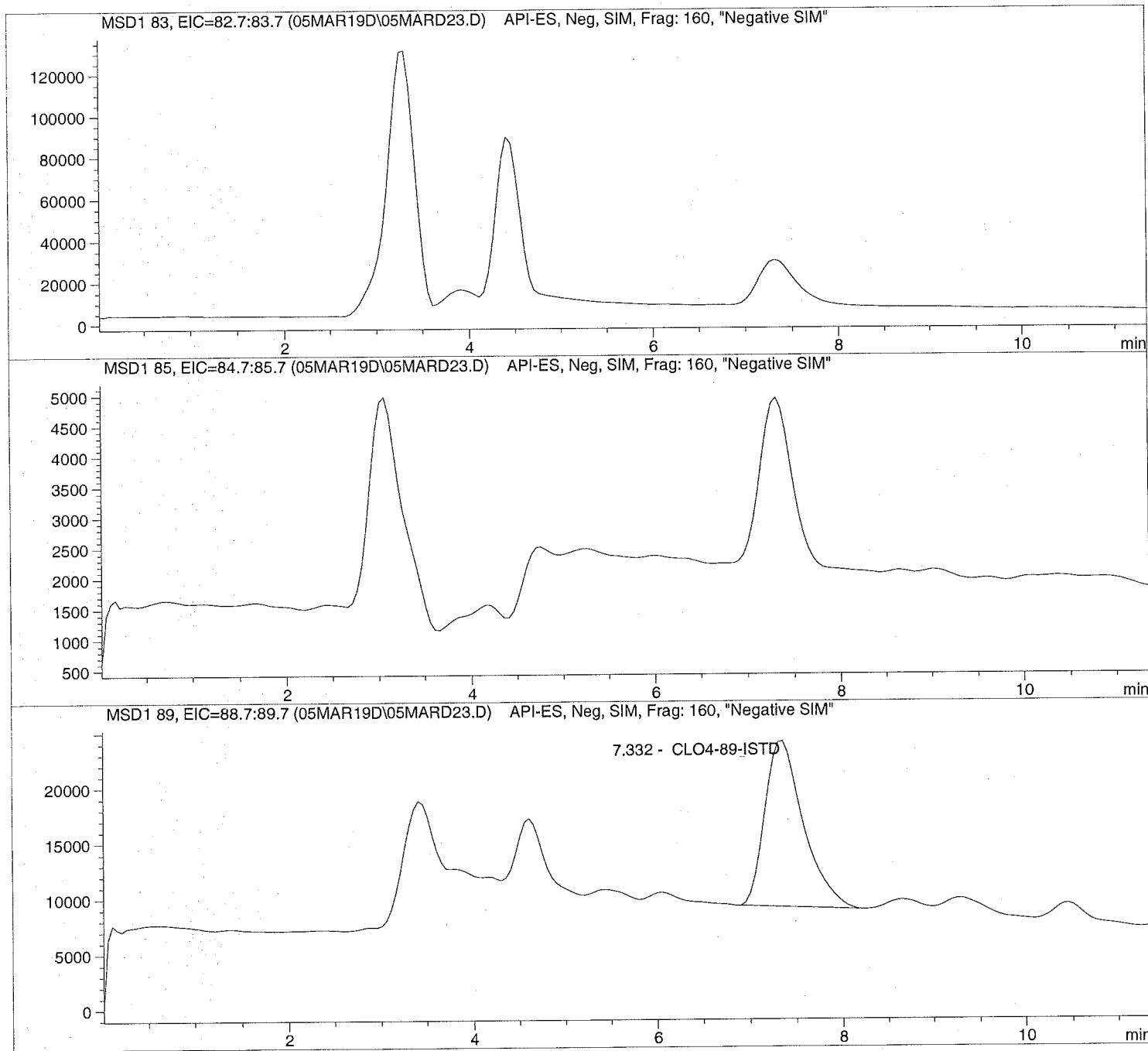
=====
*** End of Report ***
=====

Injection Date: 3/05/2019 13:41:34
Sample Name: 1906112007 RE
Acq Operator: TNB

Seq Line: 23
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date:  3/05/2019  13:41:34      Seq Line:      23
Sample Name:    1906112007  RE            Location:      Vial 82
Acq Operator:   TNB                               Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.332	PB	431134.6	5.0000	CLO4-89-ISTD

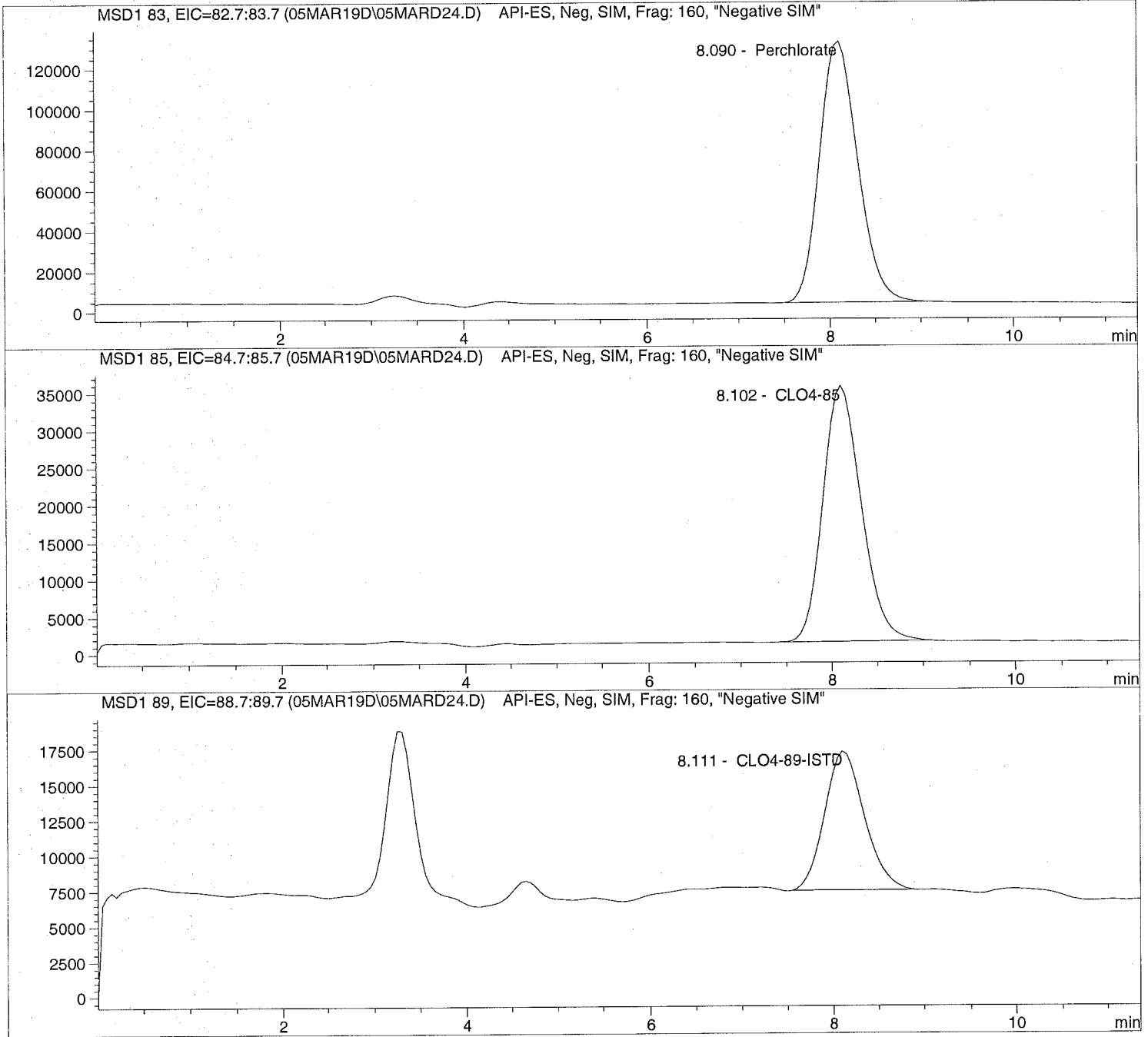
=====
*** End of Report ***

Injection Date: 3/05/2019 13:54:34
Sample Name: 1906330001 100
Acq Operator: TNB

Seq Line: 24
Location: Vial 92
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis




```
=====
Injection Date: 3/05/2019 13:54:34      Seq Line:      24
Sample Name:    1906330001 100          Location:      Vial 92
Acq Operator:   TNB                    Inj. No.:      1
                                           Inj. Vol.:     20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       100.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.090	PBA	3773628.7	3708.6567	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	PBA	1010205.2	3771.6748	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.111	PBA	298984.6	500.0000	CLO4-89-ISTD

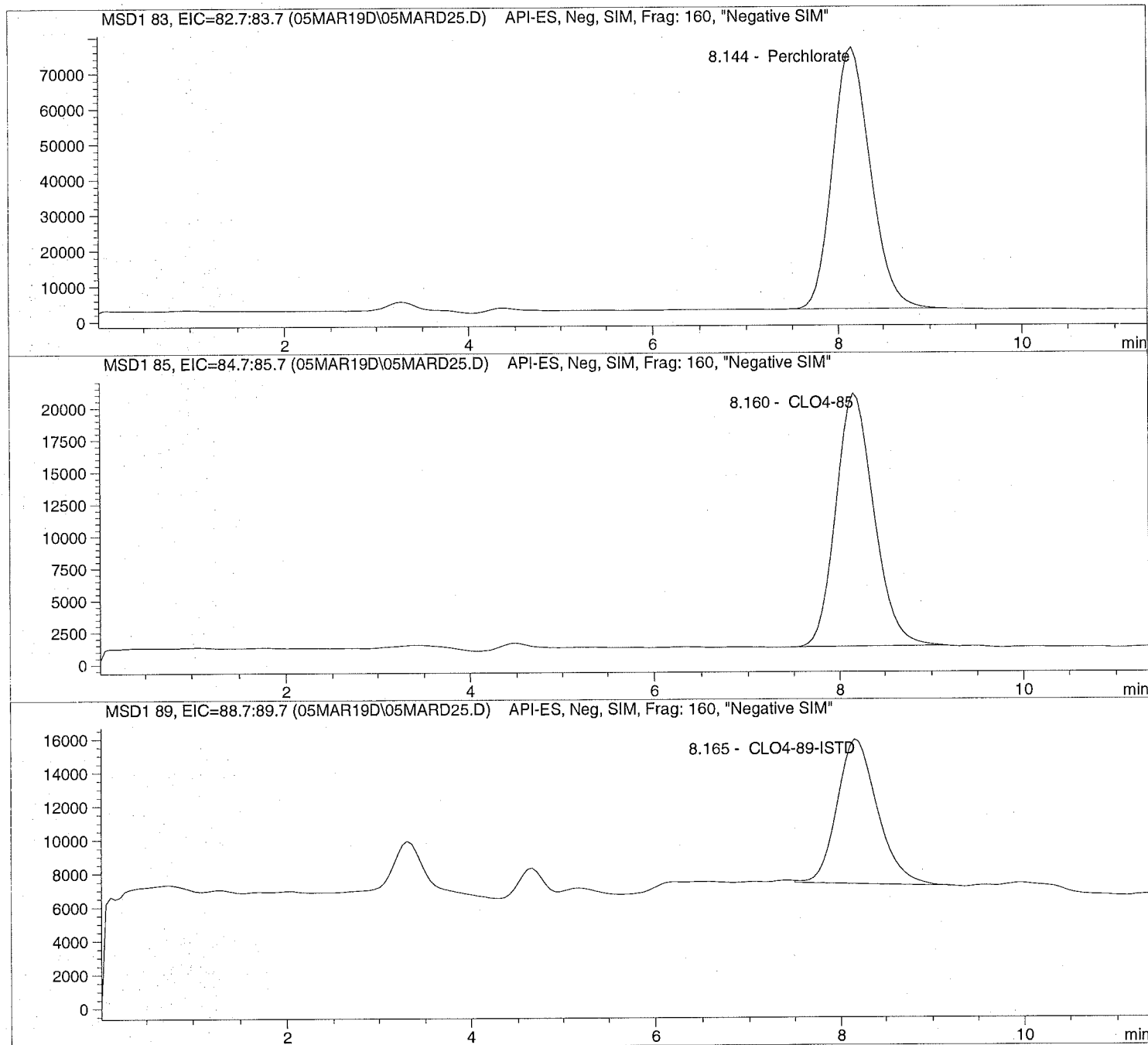
=====
*** End of Report ***

Injection Date: 3/05/2019 14:07:36
Sample Name: 642102 CCV@25
Acq Operator: TNB

Seq Line: 25
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 14:07:36      Seq Line:          25  
Sample Name:    642102  CCV@25          Location:          Vial 71  
Acq Operator:  TNB                      Inj. No.:         1  
                                           Inj. Vol.:        20 µl
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 25.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.144	PBA	2157871.5	24.4553	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.160	PBA	583711.5	25.1665	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.165	BBA	268305.4	5.0000	CLO4-89-ISTD

=====
*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

**Initial
Calibration**

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	8.94006e4	7.889	9.89924e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.97443e5	8.114	2.26028
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	4.79370e5	7.828	4.65688
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	9.30136e5	7.904	9.14998
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.81067e6	7.793	25.52636
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	5.66830e6	7.976	51.07439
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	8.69624e6	7.886	74.30603
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	1.01141e6	7.988	9.46019

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.26121e4	7.914	9.98836e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	5.53134e4	8.127	2.11360
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.39247e5	7.842	4.91261
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.54396e5	7.923	9.39034
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	7.35969e5	7.811	25.48268
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.47152e6	7.993	50.35774
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.32809e6	7.900	74.72233
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.81230e5	8.007	9.87858

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.41443e5	7.900	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.99651e5	8.132	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.38646e5	7.853	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	3.25154e5	7.925	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	3.33799e5	7.819	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	3.14712e5	7.999	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	3.13909e5	7.908	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	3.41503e5	8.005	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

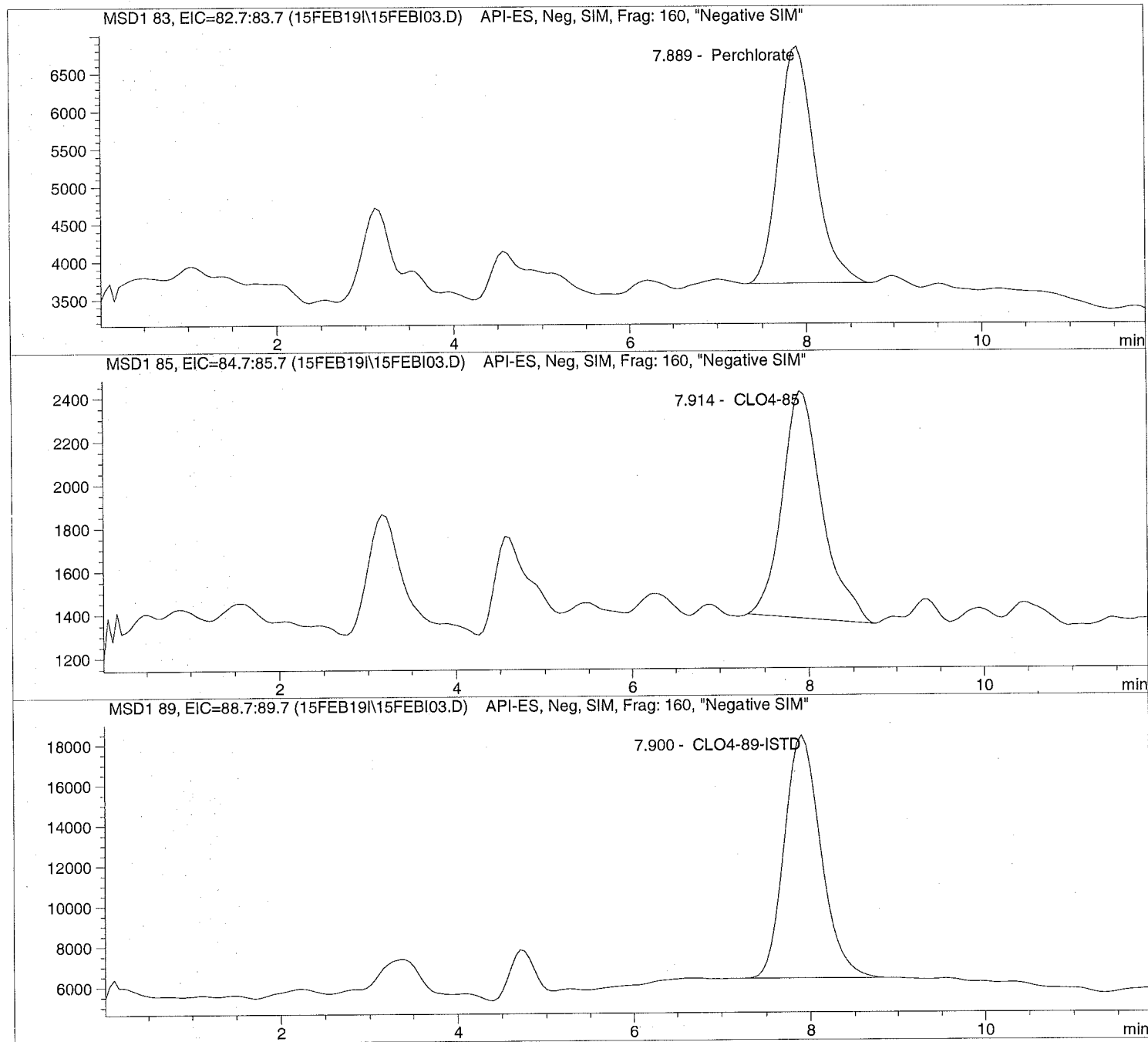
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ .20ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Injection Date: 2/15/2019 09:51:42
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====
Injection Date: 2/15/2019 09:51:42      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L           Location:  Vial 73
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.889	PBA	89400.6	0.9899	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.914	BBA	32612.1	0.9988	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	BBA	341443.2	5.0000	CLO4-89-ISTD

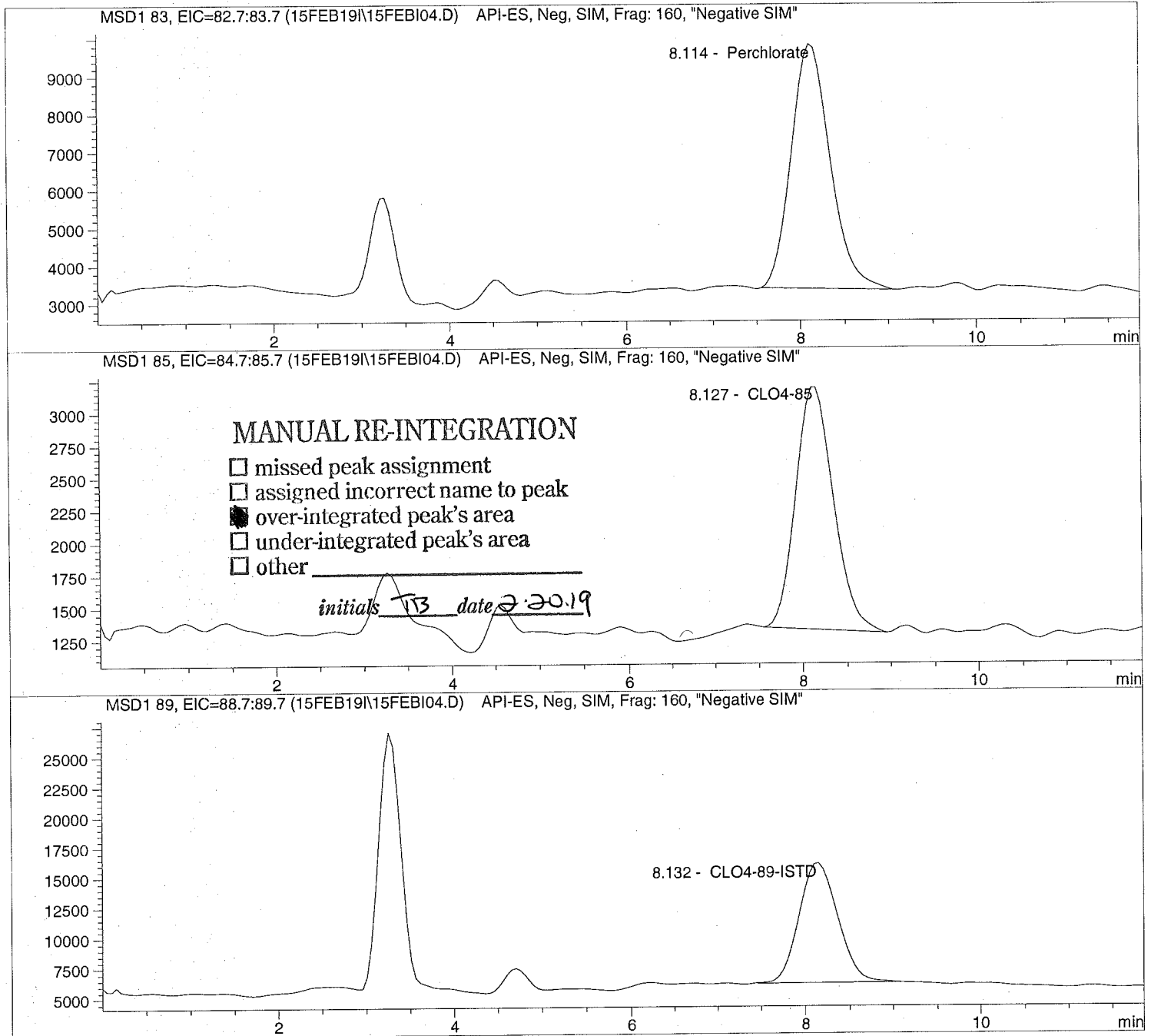
=====
*** End of Report ***

Injection Date: 2/15/2019 10:05:24
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Injection Date: 2/15/2019 10:05:24 Seq Line: 4
Sample Name: CLO4@ 2.0ug/L Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 2.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	MM	55313.4	2.1136	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

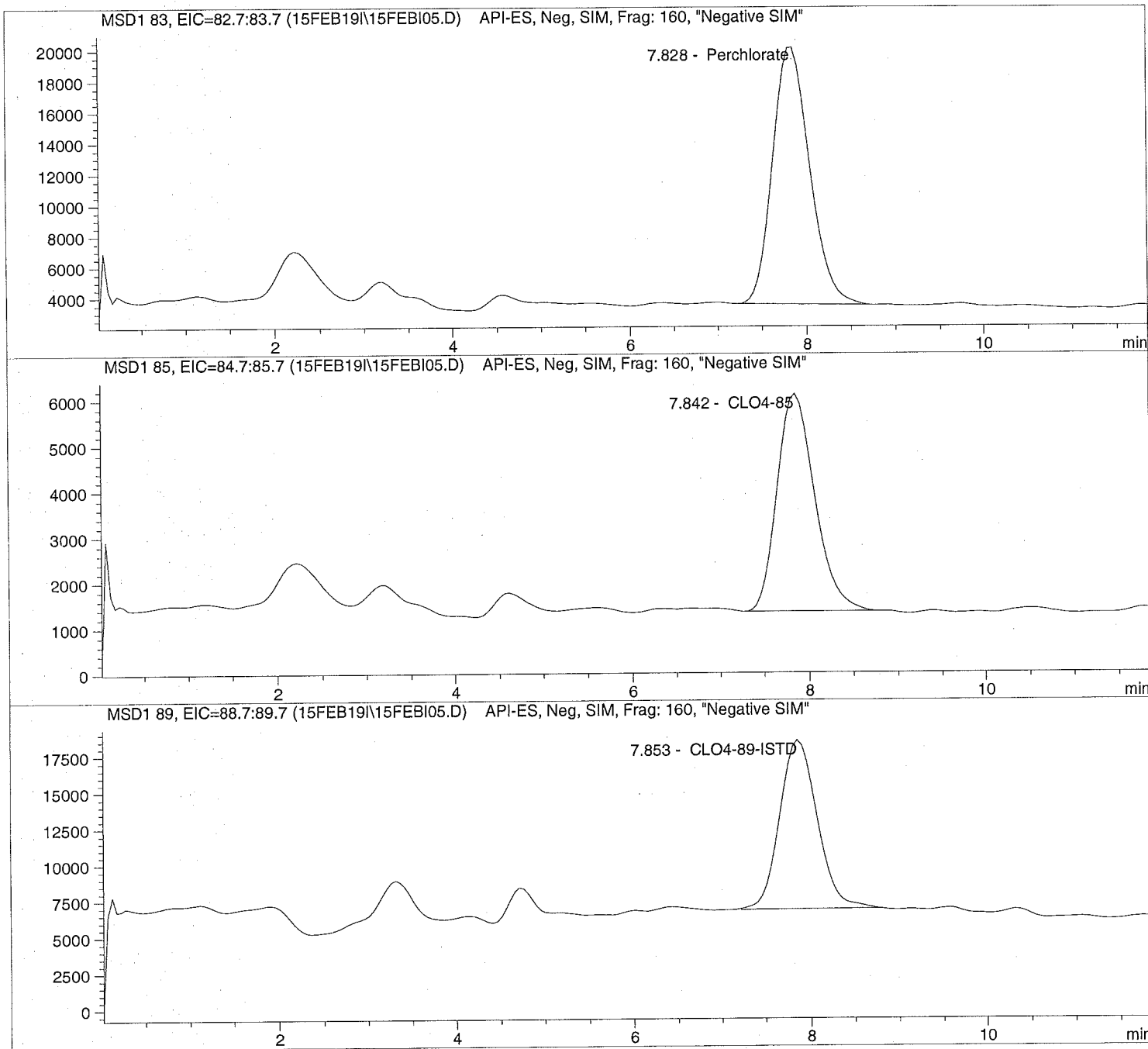
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

*** End of Report ***

=====
Injection Date: 2/15/2019 11:42:56 Seq Line: 5
Sample Name: CLO4@ 5.0ug/L Location: Vial 75
Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis
=====



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI05.D Sample Name: CLO4@ 5.0ug/L

```
=====
Injection Date: 2/15/2019 11:42:56      Seq Line:          5
Sample Name:    CLO4@ 5.0ug/L           Location:          Vial 75
Acq Operator:  TNB                      Inj. No.:         1
                                           Inj. Vol.:        25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.828	PBA	479370.4	4.6569	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.842	PBA	139246.9	4.9126	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.853	PBA	338646.3	5.0000	CLO4-89-ISTD

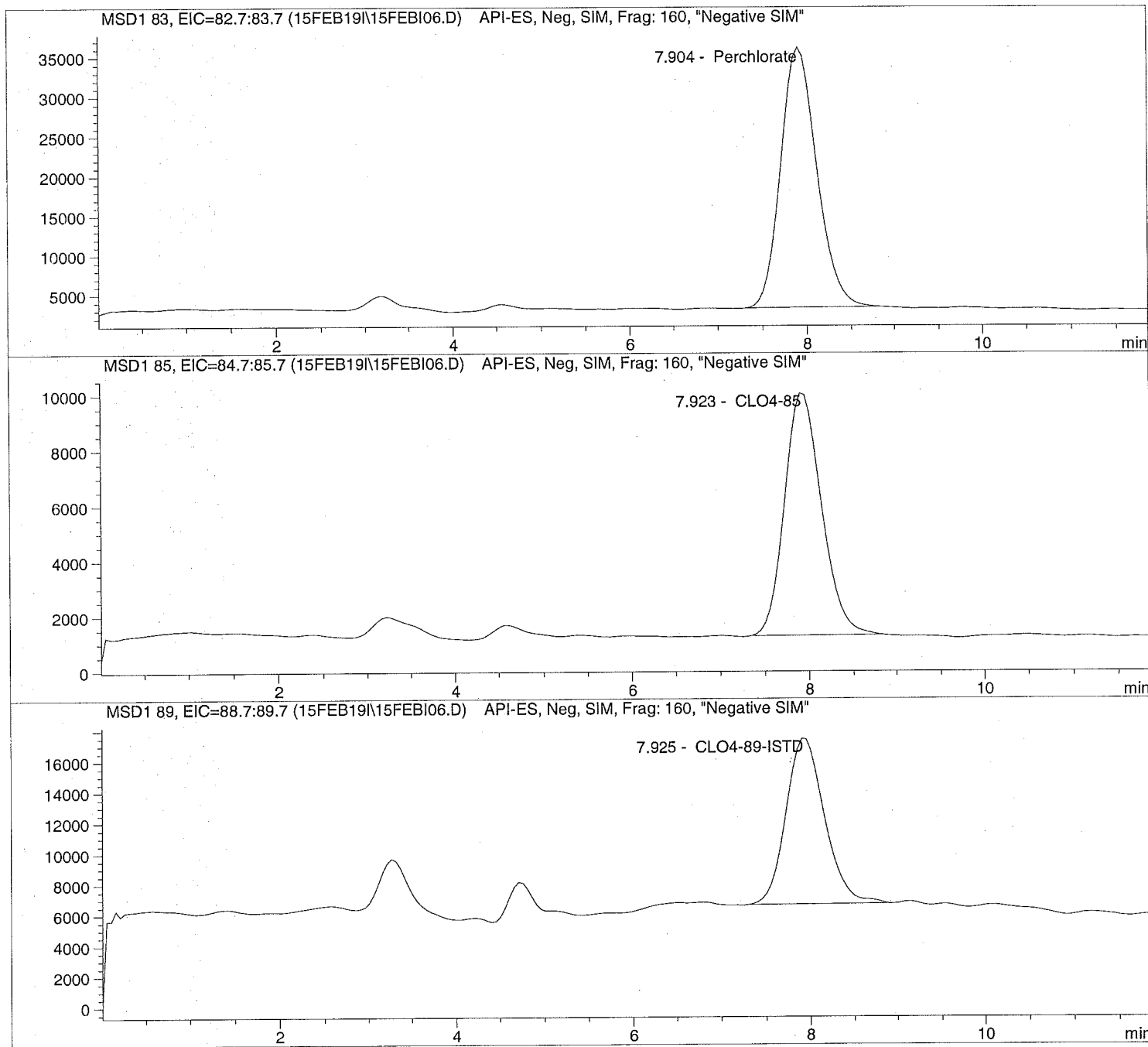
=====
*** End of Report ***

=====
Injection Date: 2/15/2019 11:56:38
Sample Name: CLO4@ 10.ug/L
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis
=====



```
=====
Injection Date: 2/15/2019 11:56:38      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.904	PBA	930135.8	9.1500	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.923	BBA	254395.6	9.3903	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.925	PBA	325154.4	5.0000	CLO4-89-ISTD

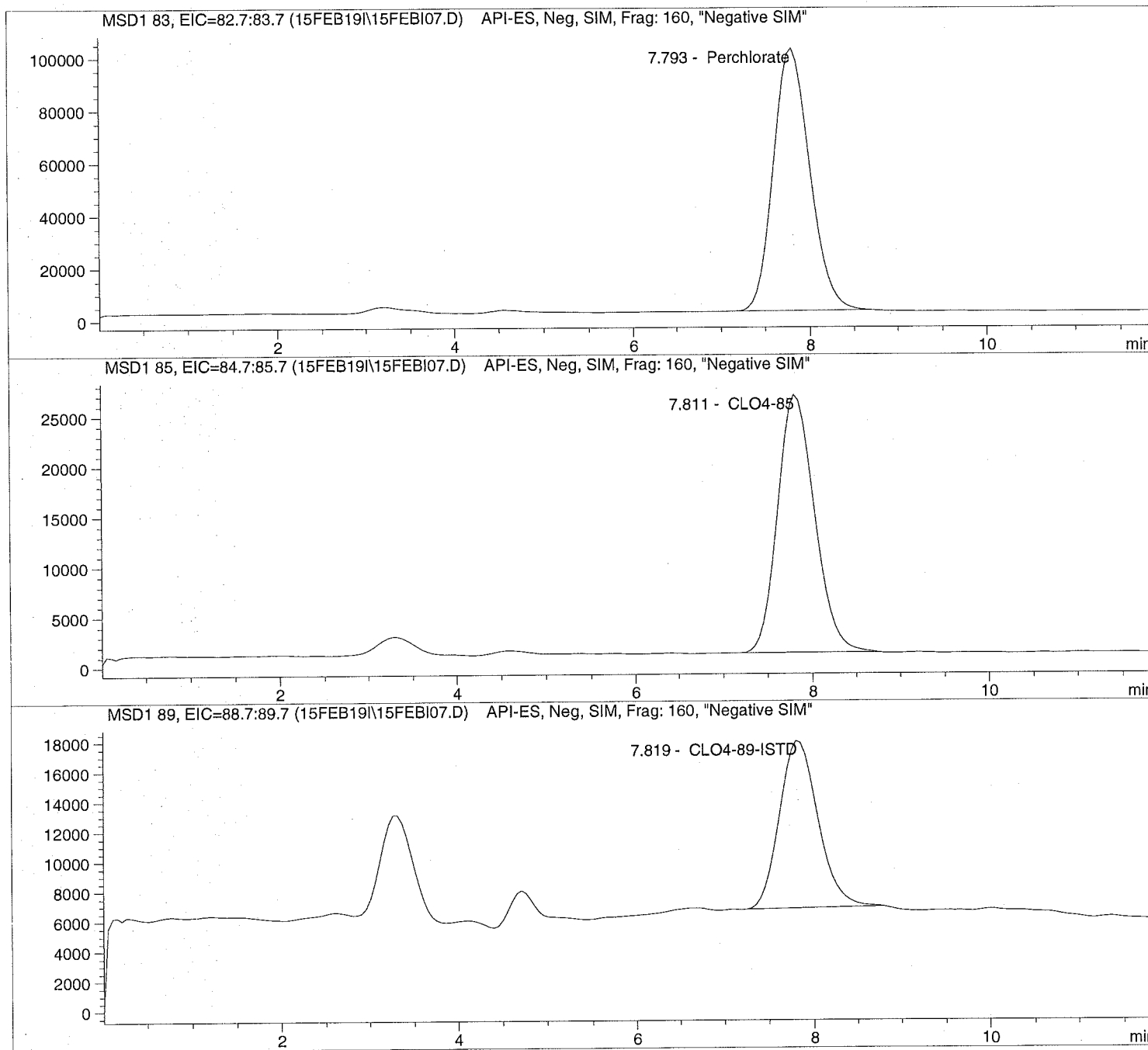
=====
*** End of Report ***

Injection Date: 2/15/2019 12:10:22
Sample Name: CLO4@ 25.ug/L
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 12:10:22      Seq Line: 7  
Sample Name:    CLO4@ 25.ug/L           Location:  Vial 77  
Acq Operator:   TNB                     Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:09:20  
=====
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 25.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.793	PBA	2810669.2	25.5264	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.811	BBA	735968.9	25.4827	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.819	PBA	333799.0	5.0000	CLO4-89-ISTD

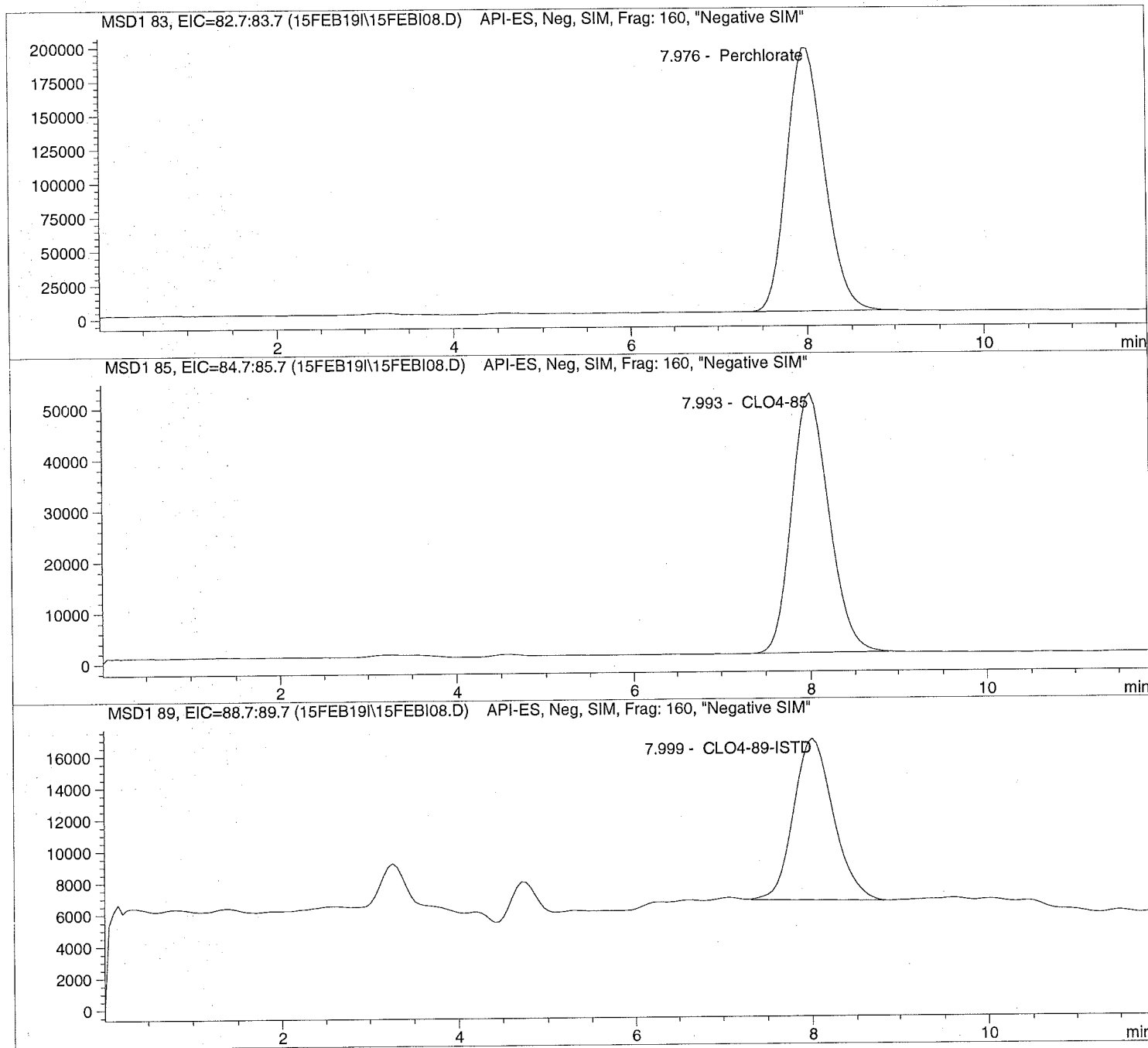
=====
*** End of Report ***
=====

Injection Date: 2/15/2019 12:24:06
Sample Name: CLO4@ 50.ug/L
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```

=====
Injection Date:  2/15/2019  12:24:06      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                      Inj. No.:     1
                                           Inj. Vol.:    25 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  09:09:20

```

Perchlorate analysis

Sample Information

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount:  50.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.976	PBA	5668301.5	51.0744	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.993	PBA	1471522.9	50.3577	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.999	BBA	314711.8	5.0000	CLO4-89-ISTD

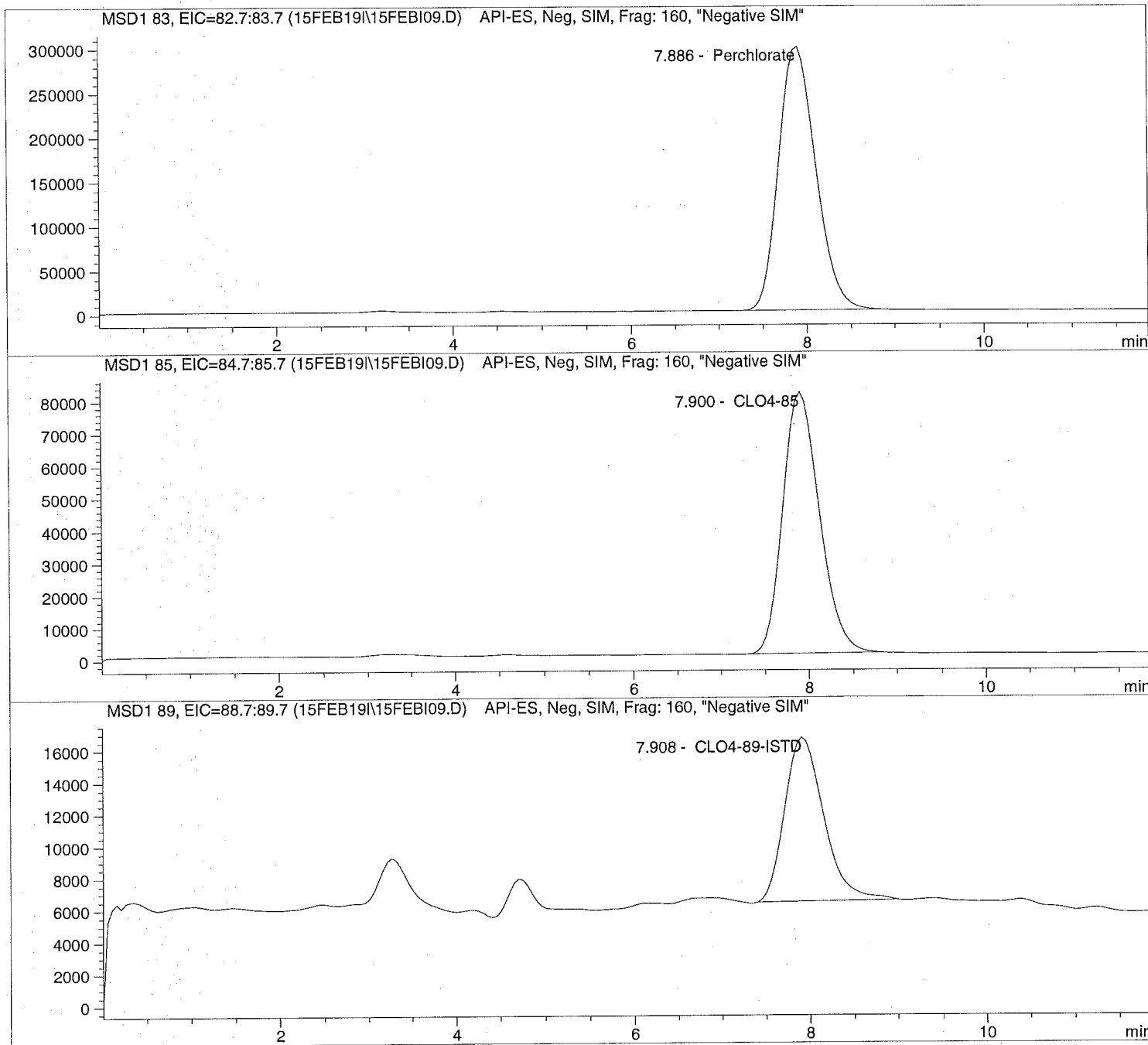
*** End of Report ***

Injection Date: 2/15/2019 12:37:48
Sample Name: CLO4@ 75.ug/L
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 12:37:48      Seq Line: 9  
Sample Name:    CLO4@ 75.ug/L           Location:  Vial 79  
Acq Operator:   TNB                     Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:09:20  
=====
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 75.000  
=====
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.886	PBA	8696239.0	74.3060	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	PBA	2328089.5	74.7223	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.908	PBA	313908.9	5.0000	CLO4-89-ISTD

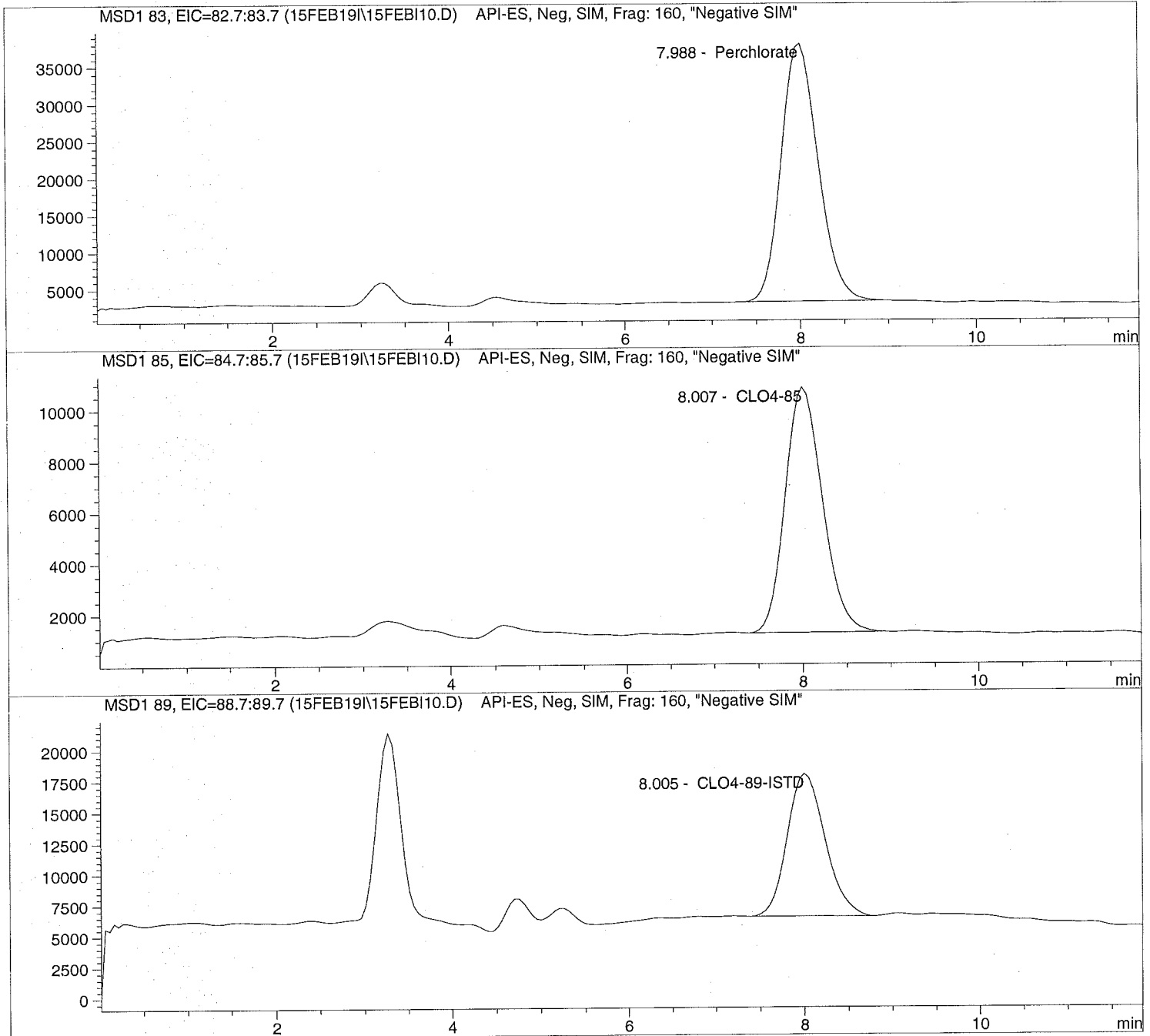
=====
*** End of Report ***
=====

Injection Date: 2/15/2019 12:51:29
Sample Name: ICAL Verf@10ug/L
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 12:51:29      Seq Line: 10  
Sample Name:    ICAL Verf@10ug/L        Location:  Vial 80  
Acq Operator:  TNB                      Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:09:20  
=====
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 10.000  
=====
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.988	BBA	1011409.8	9.4602	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.007	BBA	281229.9	9.8786	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.005	BBA	341503.2	5.0000	CLO4-89-ISTD

=====
*** End of Report ***
=====



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

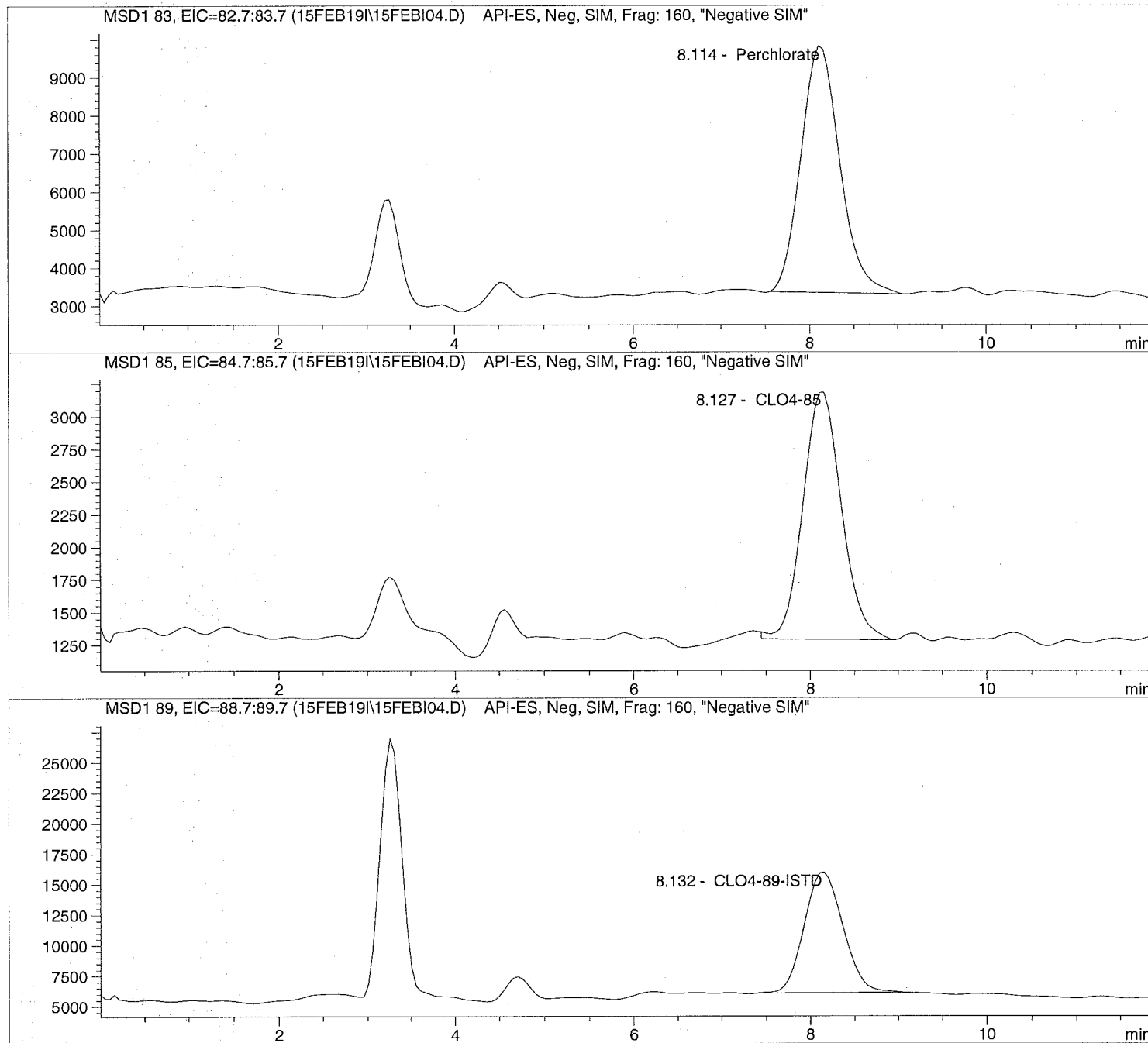
Unmodified

Injection Date: 2/15/2019 10:05:24
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:12:36

Perchlorate analysis




```
=====
Injection Date:  2/15/2019  10:05:24      Seq Line:      4
Sample Name:    CLO4@ 2.0ug/L            Location:      Vial 74
Acq Operator:   TNB                      Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  09:12:36
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified:  Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	BBA	57206.1	2.1923	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

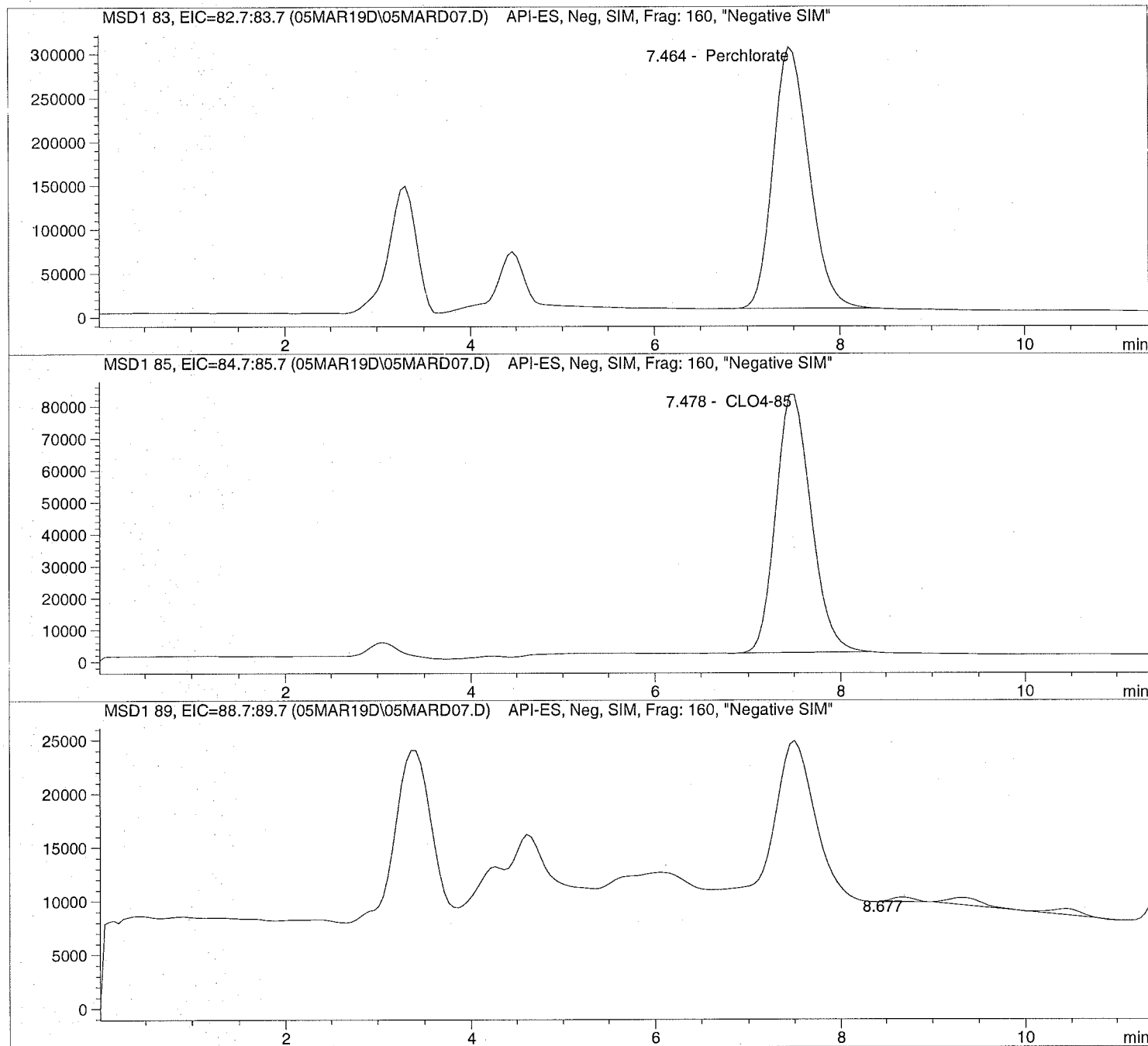
=====
*** End of Report ***

Injection Date: 3/05/2019 10:07:11
Sample Name: 1906112002 MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:07:11 Seq Line: 7
Sample Name: 1906112002 MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	419.8794	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	402.6179	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

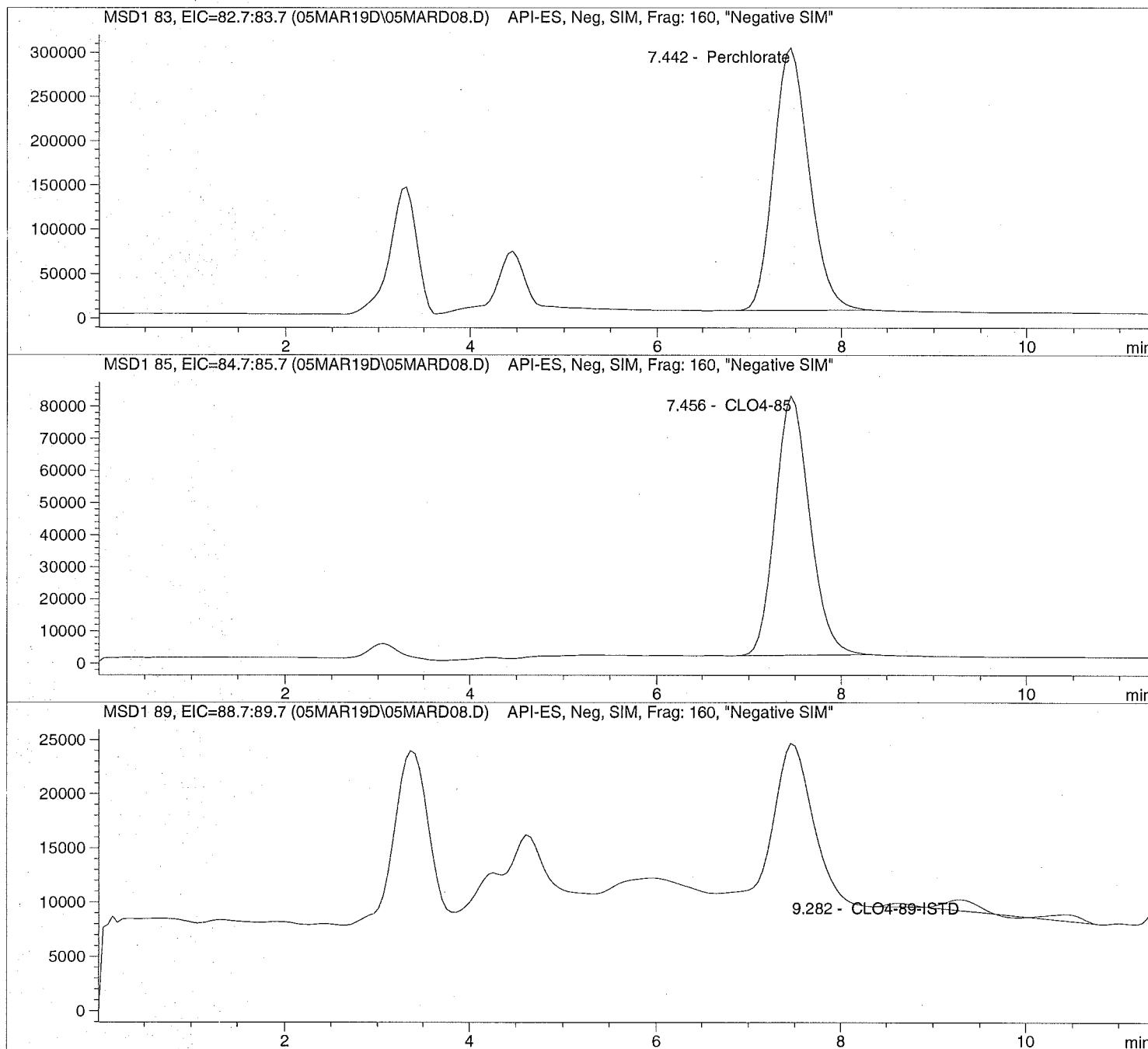
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.677	BB	7208.6	0.0000	
9.316	VBA	28561.1	5.0000	CLO4-89-ISTD

*** End of Report ***

Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 10:20:17      Seq Line:      8
Sample Name:    1906112003      MSD      Location:      Vial 78
Acq Operator:   TNB              Inj. No.:      1
                                      Inj. Vol.:     20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	359.0996	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	344.2233	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.621	VB	5769.9	0.0000	
9.282	VBA	35831.6	5.0000	CLO4-89-ISTD

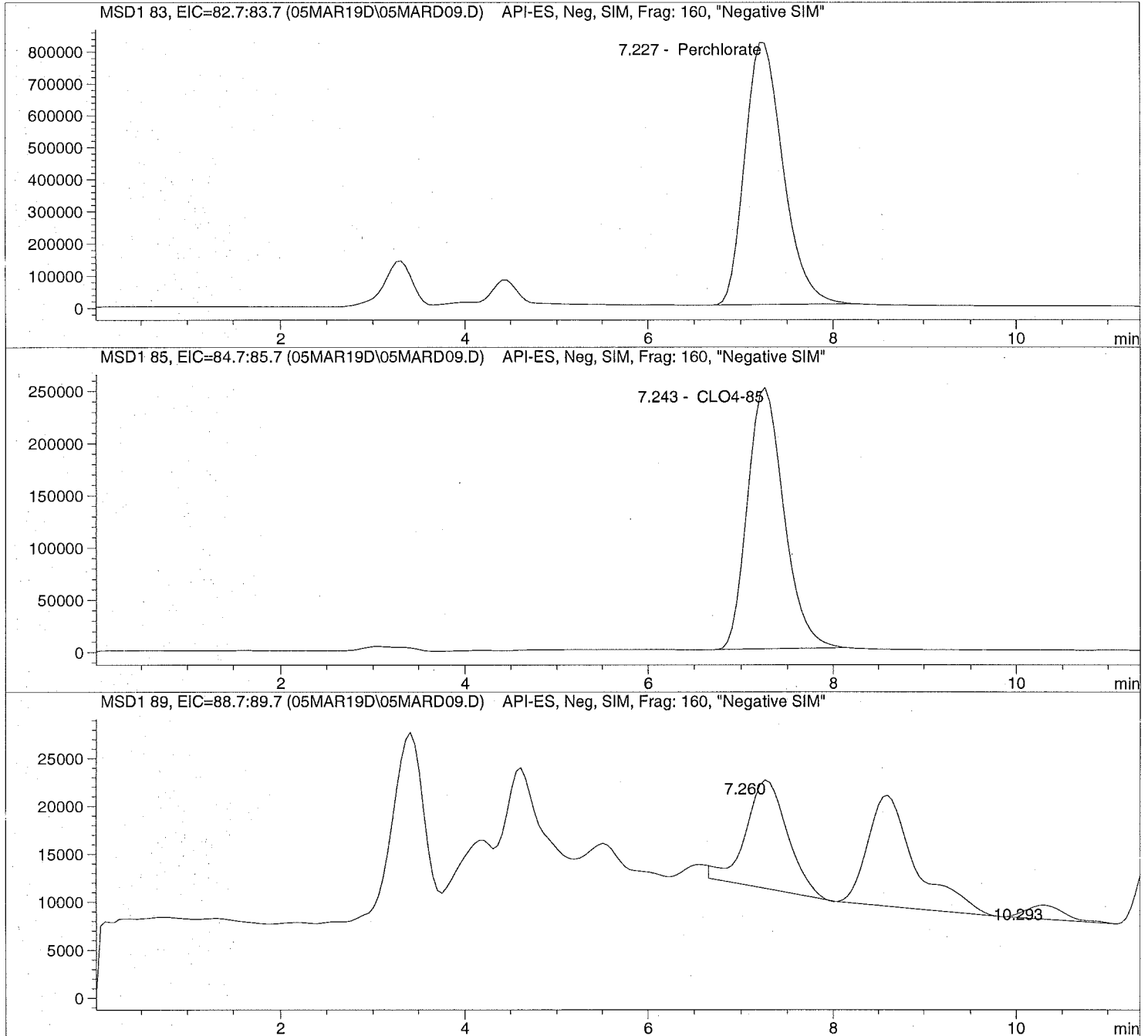
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*** End of Report ***

Injection Date: 3/05/2019 10:33:21
Sample Name: 1906112004
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 10:33:21      Seq Line:          9
Sample Name:    1906112004              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	131.1742	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	138.8050	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.260	BB	352149.7	0.0000	
8.589	VBA	421141.9	5.0000	CLO4-89-ISTD
10.293	BBA	41603.7	0.0000	

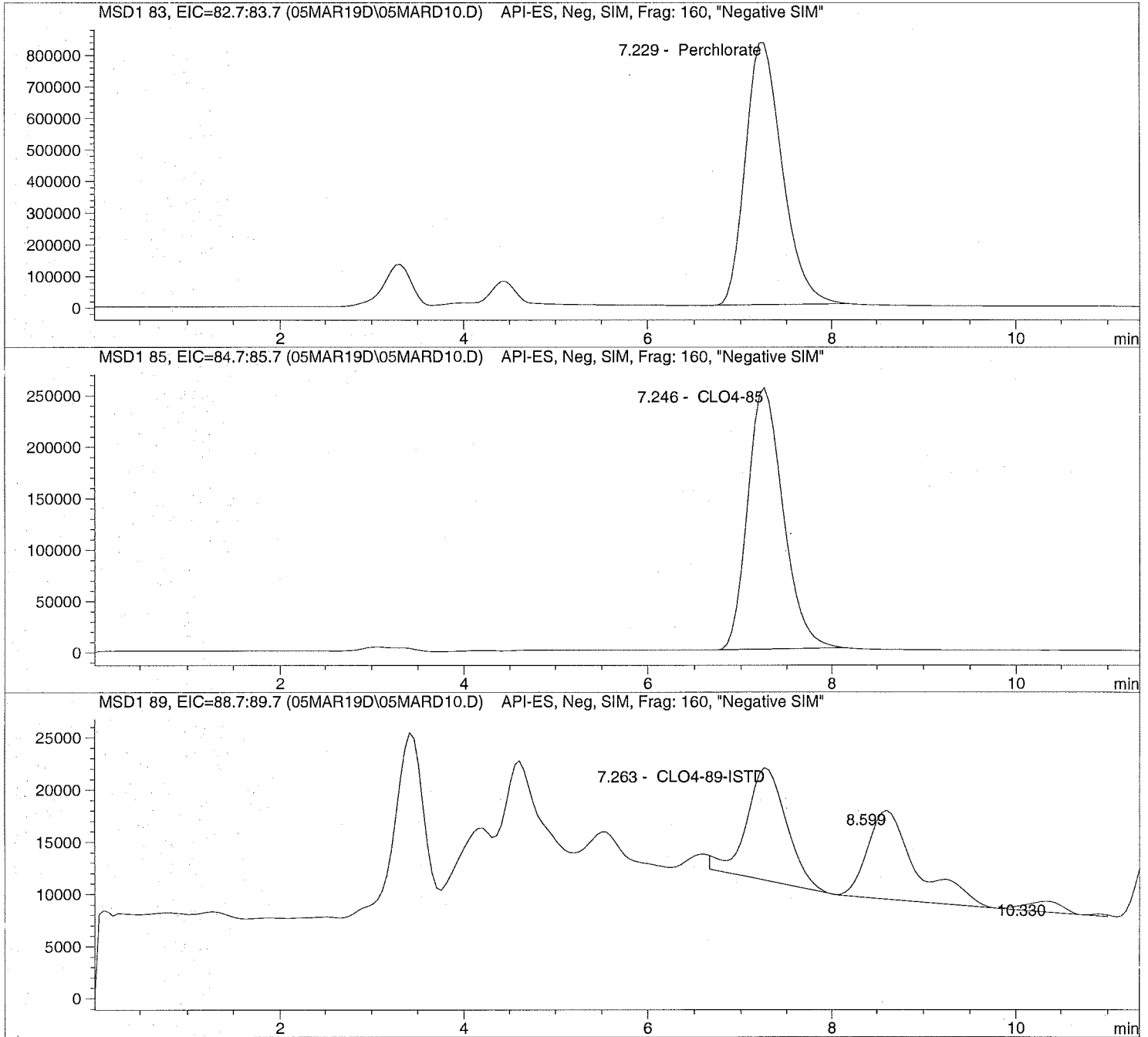
=====
*** End of Report ***

Injection Date: 3/05/2019 10:46:26
Sample Name: 1906112005
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis




```
=====
Injection Date: 3/05/2019 10:46:26      Seq Line:          10
Sample Name:    1906112005              Location:         Vial 80
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	159.0418	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	168.6882	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

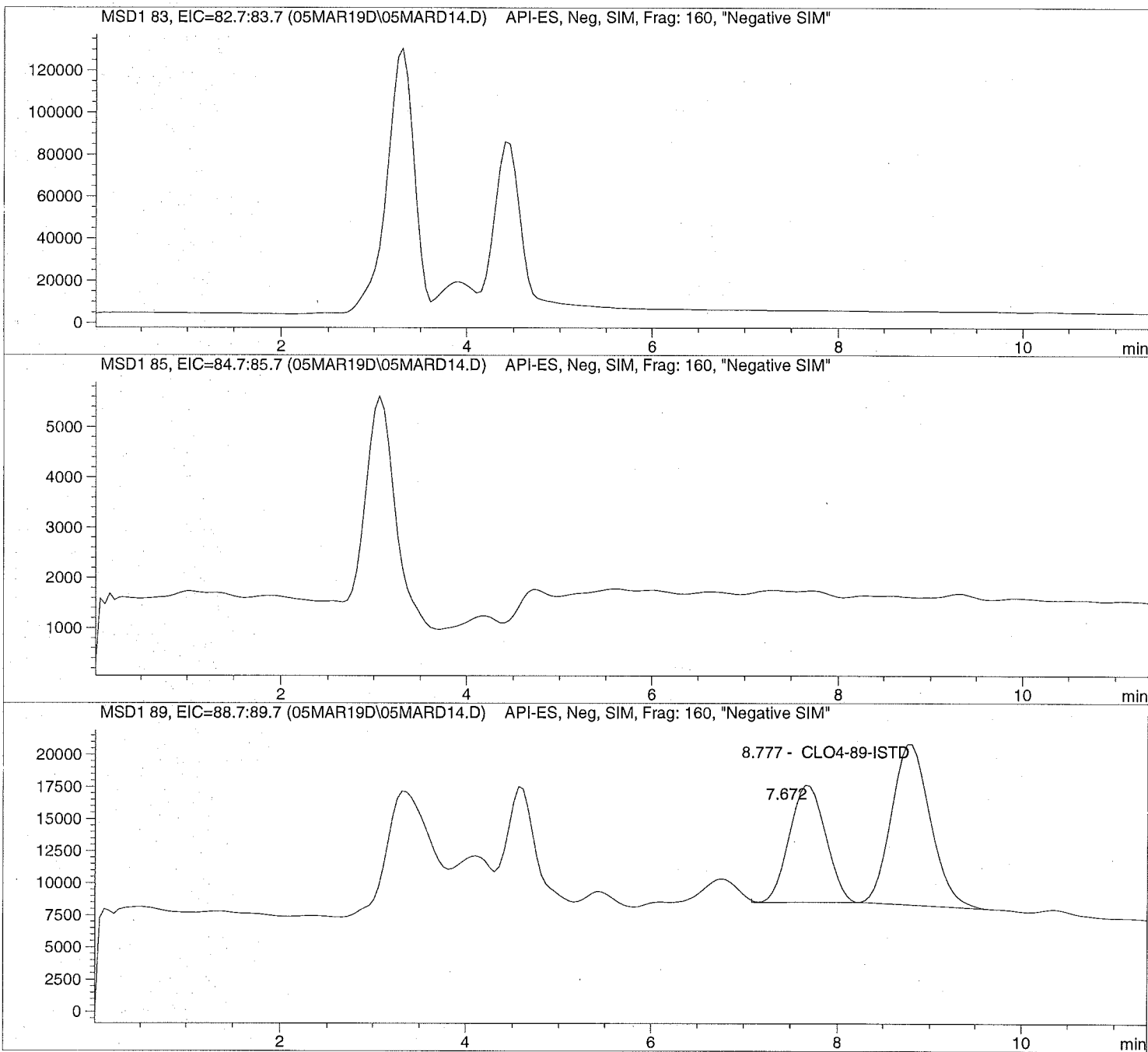
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	BB	325841.0	5.0000	CLO4-89-ISTD
8.599	VB	308921.9	0.0000	
10.330	VBA	30210.3	0.0000	

=====
*** End of Report ***

=====
 Injection Date: 3/05/2019 11:39:24 Seq Line: 14
 Sample Name: 1906112009 Location: Vial 84
 Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
 Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 11:39:24 Seq Line: 14
Sample Name: 1906112009 Location: Vial 84
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	0.0000	
8.777	VBA	362717.3	5.0000	CLO4-89-ISTD

*** End of Report ***



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 08, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19021165**

Laboratory Results for: **Longhorn GW Treatment Plant**

Dear Marcia,

ALS Environmental received 2 sample(s) on Feb 22, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: DAYNA.FISHER
RJ Modashia
Project Manager

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
Work Order: HS19021165

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19021165-01	LH18/24-SP650_022119	Water		21-Feb-2019 14:00	22-Feb-2019 09:06	<input type="checkbox"/>
HS19021165-02	Trip Blank ALS011119-60	Water		21-Feb-2019 14:00	22-Feb-2019 09:06	<input type="checkbox"/>

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
Work Order: HS19021165

CASE NARRATIVE

GCMS Volatiles by Method SW8260**Batch ID: R334099****Sample ID: HS19021185-01MS**

- MS and MSD were performed on an unrelated sample

WetChemistry by Method SW9056**Batch ID: R334211****Sample ID: HS19030163-01MSD**

- MSD is for an unrelated sample (Chloride,Sulfate)
-

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: LH18/24-SP650_022119
 Collection Date: 21-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19021165
 Lab ID:HS19021165-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
VOLATILES ORGANICS BY METHOD		Method:SW8260						
8260C								Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	05-Mar-2019 22:54
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	05-Mar-2019 22:54
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	05-Mar-2019 22:54
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	05-Mar-2019 22:54
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	05-Mar-2019 22:54
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: LH18/24-SP650_022119
 Collection Date: 21-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19021165
 Lab ID:HS19021165-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
cis-1,2-Dichloroethene	2.0		0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	05-Mar-2019 22:54	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	05-Mar-2019 22:54	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	05-Mar-2019 22:54	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Trichloroethene	0.70	J	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:54	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>82.0</i>			0	<i>81-118</i>	%REC	1	05-Mar-2019 22:54	
<i>Surr: 4-Bromofluorobenzene</i>	<i>104</i>			0	<i>85-114</i>	%REC	1	05-Mar-2019 22:54	
<i>Surr: Dibromofluoromethane</i>	<i>84.8</i>			0	<i>80-119</i>	%REC	1	05-Mar-2019 22:54	
<i>Surr: Toluene-d8</i>	<i>101</i>			0	<i>89-112</i>	%REC	1	05-Mar-2019 22:54	
ANIONS BY SW9056A		Method:SW9056						Analyst: KMU	
Chloride	320		2.00	5.00	5.00	mg/L	10	05-Mar-2019 22:00	
Sulfate	29.8		2.00	5.00	5.00	mg/L	10	05-Mar-2019 22:00	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: Trip Blank ALS011119-60
 Collection Date: 21-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19021165
 Lab ID:HS19021165-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD		Method:SW8260							Analyst: PC
8260C									
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	05-Mar-2019 22:06	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	05-Mar-2019 22:06	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	05-Mar-2019 22:06	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	05-Mar-2019 22:06	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	05-Mar-2019 22:06	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: Trip Blank ALS011119-60
 Collection Date: 21-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19021165
 Lab ID:HS19021165-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD		Method:SW8260							Analyst: PC
8260C									
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	05-Mar-2019 22:06	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	05-Mar-2019 22:06	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	05-Mar-2019 22:06	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Trichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	05-Mar-2019 22:06	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>83.7</i>			<i>0</i>	<i>81-118</i>	<i>%REC</i>	<i>1</i>	<i>05-Mar-2019 22:06</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>97.1</i>			<i>0</i>	<i>85-114</i>	<i>%REC</i>	<i>1</i>	<i>05-Mar-2019 22:06</i>	
<i>Surr: Dibromofluoromethane</i>	<i>84.9</i>			<i>0</i>	<i>80-119</i>	<i>%REC</i>	<i>1</i>	<i>05-Mar-2019 22:06</i>	
<i>Surr: Toluene-d8</i>	<i>109</i>			<i>0</i>	<i>89-112</i>	<i>%REC</i>	<i>1</i>	<i>05-Mar-2019 22:06</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R334099					Test Name : VOLATILES ORGANICS BY METHOD 8260C	Matrix: Water
HS19021165-01	LH18/24-SP650_022119	21 Feb 2019 14:00				05 Mar 2019 22:54 1
HS19021165-02	Trip Blank ALS011119-60	21 Feb 2019 14:00				05 Mar 2019 22:06 1
Batch ID R334211					Test Name : ANIONS BY SW9056A	Matrix: Water
HS19021165-01	LH18/24-SP650_022119	21 Feb 2019 14:00				05 Mar 2019 22:00 10

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334099		Instrument: VOA6		Method: SW8260						
MBLK	Sample ID: VBLKW-190305	Units: UG/L			Analysis Date: 05-Mar-2019 19:18					
Client ID:	Run ID: VOA6_334099	SeqNo: 4977030	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	40.72	1.0	50	0	81.4	81 - 118				
Surr: 4-Bromofluorobenzene	50.56	1.0	50	0	101	85 - 114				
Surr: Dibromofluoromethane	42.1	1.0	50	0	84.2	80 - 119				
Surr: Toluene-d8	53.5	1.0	50	0	107	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334099		Instrument: VOA6		Method: SW8260						
LCS	Sample ID: VLCSW-190305	Units: UG/L			Analysis Date: 05-Mar-2019 18:05					
Client ID:	Run ID: VOA6_334099	SeqNo: 4977029	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	19.19	1.0	20	0	96.0	78 - 124				
1,1,1-Trichloroethane	18.34	1.0	20	0	91.7	74 - 131				
1,1,2,2-Tetrachloroethane	19.89	1.0	20	0	99.4	71 - 121				
1,1,2-Trichloroethane	19.74	1.0	20	0	98.7	80 - 119				
1,1-Dichloroethane	18.86	1.0	20	0	94.3	77 - 125				
1,1-Dichloroethene	17.9	1.0	20	0	89.5	71 - 131				
1,1-Dichloropropene	18.25	1.0	20	0	91.3	78 - 125				
1,2,3-Trichlorobenzene	22.1	1.0	20	0	111	69 - 129				
1,2,3-Trichloropropane	19.69	1.0	20	0	98.4	73 - 122				
1,2,4-Trichlorobenzene	20.94	1.0	20	0	105	69 - 130				
1,2,4-Trimethylbenzene	18.53	1.0	20	0	92.7	76 - 124				
1,2-Dibromo-3-chloropropane	21.06	1.0	20	0	105	62 - 128				
1,2-Dibromoethane	20.36	1.0	20	0	102	77 - 121				
1,2-Dichlorobenzene	18.97	1.0	20	0	94.8	80 - 119				
1,2-Dichloroethane	19.73	1.0	20	0	98.7	73 - 128				
1,2-Dichloropropane	19.35	1.0	20	0	96.7	78 - 122				
1,3,5-Trimethylbenzene	18.35	1.0	20	0	91.8	75 - 124				
1,3-Dichlorobenzene	18.87	1.0	20	0	94.3	80 - 119				
1,3-Dichloropropane	19.67	1.0	20	0	98.4	80 - 119				
1,4-Dichlorobenzene	18.61	1.0	20	0	93.0	79 - 118				
2,2-Dichloropropane	18.33	1.0	20	0	91.6	60 - 139				
2-Butanone	43.52	2.0	40	0	109	56 - 143				
2-Chlorotoluene	18.21	1.0	20	0	91.1	79 - 122				
2-Hexanone	40.29	2.0	40	0	101	57 - 139				
4-Chlorotoluene	18.65	1.0	20	0	93.3	78 - 122				
4-Isopropyltoluene	17.75	1.0	20	0	88.7	77 - 127				
4-Methyl-2-pentanone	40.53	2.0	40	0	101	67 - 130				
Acetone	42.24	2.0	40	0	106	39 - 160				
Benzene	18.9	1.0	20	0	94.5	79 - 120				
Bromobenzene	19.12	1.0	20	0	95.6	80 - 120				
Bromochloromethane	19.26	1.0	20	0	96.3	78 - 123				
Bromodichloromethane	19.32	1.0	20	0	96.6	79 - 125				
Bromoform	20.19	1.0	20	0	101	66 - 130				
Bromomethane	23.44	1.0	20	0	117	53 - 141				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334099		Instrument: VOA6		Method: SW8260						
LCS	Sample ID: VLCSW-190305	Units: UG/L			Analysis Date: 05-Mar-2019 18:05					
Client ID:	Run ID: VOA6_334099	SeqNo: 4977029		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	36.19	2.0	40	0	90.5	64 - 133				
Carbon tetrachloride	18.61	1.0	20	0	93.0	72 - 136				
Chlorobenzene	19.07	1.0	20	0	95.4	82 - 118				
Chloroethane	18.68	1.0	20	0	93.4	60 - 138				
Chloroform	19.06	1.0	20	0	95.3	79 - 124				
Chloromethane	18.78	1.0	20	0	93.9	50 - 139				
cis-1,2-Dichloroethene	18.62	1.0	20	0	93.1	78 - 123				
cis-1,3-Dichloropropene	19.79	1.0	20	0	99.0	75 - 124				
Dibromochloromethane	20.2	1.0	20	0	101	74 - 126				
Dibromomethane	19.92	1.0	20	0	99.6	79 - 123				
Dichlorodifluoromethane	17.63	1.0	20	0	88.2	32 - 152				
Ethylbenzene	18.79	1.0	20	0	93.9	79 - 121				
Hexachlorobutadiene	19.81	1.0	20	0	99.1	66 - 134				
Isopropylbenzene	18.42	1.0	20	0	92.1	72 - 131				
m,p-Xylene	36.96	2.0	40	0	92.4	80 - 121				
Methylene chloride	19.77	2.0	20	0	98.9	74 - 124				
Naphthalene	20.69	1.0	20	0	103	61 - 128				
n-Butylbenzene	17.4	1.0	20	0	87.0	75 - 128				
n-Propylbenzene	17.93	1.0	20	0	89.7	76 - 126				
o-Xylene	19.4	1.0	20	0	97.0	78 - 122				
sec-Butylbenzene	17.21	1.0	20	0	86.1	77 - 126				
Styrene	19.77	1.0	20	0	98.8	78 - 123				
tert-Butylbenzene	17.51	1.0	20	0	87.5	78 - 124				
Tetrachloroethene	18.09	1.0	20	0	90.4	74 - 129				
Toluene	19.07	1.0	20	0	95.3	80 - 121				
trans-1,2-Dichloroethene	19.09	1.0	20	0	95.4	75 - 124				
trans-1,3-Dichloropropene	19.92	1.0	20	0	99.6	73 - 127				
Trichloroethene	19.04	1.0	20	0	95.2	79 - 123				
Trichlorofluoromethane	17.17	1.0	20	0	85.8	65 - 141				
Vinyl chloride	17.04	1.0	20	0	85.2	58 - 137				
Surr: 1,2-Dichloroethane-d4	52.42	1.0	50	0	105	81 - 118				
Surr: 4-Bromofluorobenzene	52.66	1.0	50	0	105	85 - 114				
Surr: Dibromofluoromethane	51.95	1.0	50	0	104	80 - 119				
Surr: Toluene-d8	49.26	1.0	50	0	98.5	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334099		Instrument: VOA6		Method: SW8260						
MS	Sample ID: HS19021185-01MS	Units: UG/L			Analysis Date: 06-Mar-2019 00:06					
Client ID:	Run ID: VOA6_334099	SeqNo: 4977037	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	17.93	1.0	20	0	89.7	78 - 124				
1,1,1-Trichloroethane	16.57	1.0	20	0	82.9	74 - 131				
1,1,2,2-Tetrachloroethane	19.71	1.0	20	0	98.5	71 - 121				
1,1,2-Trichloroethane	18.65	1.0	20	0	93.2	80 - 119				
1,1-Dichloroethane	15.53	1.0	20	0	77.7	77 - 125				
1,1-Dichloroethene	17.08	1.0	20	0	85.4	71 - 131				
1,1-Dichloropropene	18.65	1.0	20	0	93.3	78 - 125				
1,2,3-Trichlorobenzene	21.52	1.0	20	0	108	69 - 129				
1,2,3-Trichloropropane	19.51	1.0	20	0	97.6	73 - 122				
1,2,4-Trichlorobenzene	21.02	1.0	20	0	105	69 - 130				
1,2,4-Trimethylbenzene	18.65	1.0	20	0	93.3	76 - 124				
1,2-Dibromo-3-chloropropane	21.04	1.0	20	0	105	62 - 128				
1,2-Dibromoethane	18.83	1.0	20	0	94.2	77 - 121				
1,2-Dichlorobenzene	18.46	1.0	20	0	92.3	80 - 119				
1,2-Dichloroethane	16.99	1.0	20	0	85.0	73 - 128				
1,2-Dichloropropane	17.07	1.0	20	0	85.4	78 - 122				
1,3,5-Trimethylbenzene	19.18	1.0	20	0	95.9	75 - 124				
1,3-Dichlorobenzene	18.39	1.0	20	0	91.9	80 - 119				
1,3-Dichloropropane	18.51	1.0	20	0	92.5	80 - 119				
1,4-Dichlorobenzene	18.05	1.0	20	0	90.3	79 - 118				
2,2-Dichloropropane	14.97	1.0	20	0	74.9	60 - 139				
2-Butanone	36.36	2.0	40	0	90.9	56 - 143				
2-Chlorotoluene	18.19	1.0	20	0	91.0	79 - 122				
2-Hexanone	41.7	2.0	40	0	104	57 - 139				
4-Chlorotoluene	18.42	1.0	20	0	92.1	78 - 122				
4-Isopropyltoluene	19.79	1.0	20	0	98.9	77 - 127				
4-Methyl-2-pentanone	40.54	2.0	40	0	101	67 - 130				
Acetone	35.14	2.0	40	0	87.8	39 - 160				
Benzene	17.3	1.0	20	0	86.5	79 - 120				
Bromobenzene	18.25	1.0	20	0	91.3	80 - 120				
Bromochloromethane	14.52	1.0	20	0	72.6	78 - 123				S
Bromodichloromethane	16.54	1.0	20	0	82.7	79 - 125				
Bromoform	18.9	1.0	20	0	94.5	66 - 130				
Bromomethane	17.83	1.0	20	0	89.1	53 - 141				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334099		Instrument: VOA6		Method: SW8260						
MS	Sample ID: HS19021185-01MS	Units: UG/L			Analysis Date: 06-Mar-2019 00:06					
Client ID:	Run ID: VOA6_334099	SeqNo: 4977037	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	32.12	2.0	40	0	80.3	64 - 133				
Carbon tetrachloride	19.97	1.0	20	0	99.9	72 - 136				
Chlorobenzene	17.86	1.0	20	0	89.3	82 - 118				
Chloroethane	18.44	1.0	20	0	92.2	60 - 138				
Chloroform	15.26	1.0	20	0	76.3	79 - 124				S
Chloromethane	13.37	1.0	20	0	66.8	50 - 139				
cis-1,2-Dichloroethene	15.51	1.0	20	0	77.6	78 - 123				S
cis-1,3-Dichloropropene	16.22	1.0	20	0	81.1	75 - 124				
Dibromochloromethane	18.81	1.0	20	0	94.0	74 - 126				
Dibromomethane	17.29	1.0	20	0	86.4	79 - 123				
Dichlorodifluoromethane	12.17	1.0	20	0	60.9	32 - 152				
Ethylbenzene	18.62	1.0	20	0	93.1	79 - 121				
Hexachlorobutadiene	20.45	1.0	20	0	102	66 - 134				
Isopropylbenzene	19.69	1.0	20	0	98.4	72 - 131				
m,p-Xylene	37	2.0	40	0	92.5	80 - 121				
Methylene chloride	14.99	2.0	20	0	75.0	74 - 124				
Naphthalene	20.55	1.0	20	0	103	61 - 128				
n-Butylbenzene	19.41	1.0	20	0	97.0	75 - 128				
n-Propylbenzene	19.38	1.0	20	0	96.9	76 - 126				
o-Xylene	18.61	1.0	20	0	93.0	78 - 122				
sec-Butylbenzene	19.94	1.0	20	0	99.7	77 - 126				
Styrene	18.38	1.0	20	0	91.9	78 - 123				
tert-Butylbenzene	19.72	1.0	20	0	98.6	78 - 124				
Tetrachloroethene	20.08	1.0	20	0	100	74 - 129				
Toluene	18.67	1.0	20	0	93.3	80 - 121				
trans-1,2-Dichloroethene	16.03	1.0	20	0	80.2	75 - 124				
trans-1,3-Dichloropropene	16.42	1.0	20	0	82.1	73 - 127				
Trichloroethene	31.05	1.0	20	13.06	90.0	79 - 123				
Trichlorofluoromethane	17.41	1.0	20	0	87.1	65 - 141				
Vinyl chloride	14.49	1.0	20	0	72.4	58 - 137				
Surr: 1,2-Dichloroethane-d4	42.2	1.0	50	0	84.4	81 - 118				
Surr: 4-Bromofluorobenzene	50.35	1.0	50	0	101	85 - 114				
Surr: Dibromofluoromethane	42.74	1.0	50	0	85.5	80 - 119				
Surr: Toluene-d8	52.15	1.0	50	0	104	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334099		Instrument: VOA6		Method: SW8260						
MSD	Sample ID: HS19021185-01MSD	Units: UG/L			Analysis Date: 06-Mar-2019 00:30					
Client ID:	Run ID: VOA6_334099	SeqNo: 4977038	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	20.63	1.0	20	0	103	78 - 124	17.93	14	20	
1,1,1-Trichloroethane	19	1.0	20	0	95.0	74 - 131	16.57	13.6	20	
1,1,2,2-Tetrachloroethane	20.17	1.0	20	0	101	71 - 121	19.71	2.32	20	
1,1,2-Trichloroethane	20.67	1.0	20	0	103	80 - 119	18.65	10.3	20	
1,1-Dichloroethane	18.29	1.0	20	0	91.4	77 - 125	15.53	16.3	20	
1,1-Dichloroethene	19.12	1.0	20	0	95.6	71 - 131	17.08	11.3	20	
1,1-Dichloropropene	21.28	1.0	20	0	106	78 - 125	18.65	13.2	20	
1,2,3-Trichlorobenzene	26.24	1.0	20	0	131	69 - 129	21.52	19.8	20	S
1,2,3-Trichloropropane	19.76	1.0	20	0	98.8	73 - 122	19.51	1.25	20	
1,2,4-Trichlorobenzene	24.54	1.0	20	0	123	69 - 130	21.02	15.5	20	
1,2,4-Trimethylbenzene	21.44	1.0	20	0	107	76 - 124	18.65	13.9	20	
1,2-Dibromo-3-chloropropane	21.31	1.0	20	0	107	62 - 128	21.04	1.26	20	
1,2-Dibromoethane	20.46	1.0	20	0	102	77 - 121	18.83	8.3	20	
1,2-Dichlorobenzene	20.89	1.0	20	0	104	80 - 119	18.46	12.3	20	
1,2-Dichloroethane	19.77	1.0	20	0	98.9	73 - 128	16.99	15.1	20	
1,2-Dichloropropane	19.99	1.0	20	0	100.0	78 - 122	17.07	15.8	20	
1,3,5-Trimethylbenzene	21.94	1.0	20	0	110	75 - 124	19.18	13.4	20	
1,3-Dichlorobenzene	20.85	1.0	20	0	104	80 - 119	18.39	12.6	20	
1,3-Dichloropropane	20.52	1.0	20	0	103	80 - 119	18.51	10.3	20	
1,4-Dichlorobenzene	20.5	1.0	20	0	103	79 - 118	18.05	12.7	20	
2,2-Dichloropropane	17.36	1.0	20	0	86.8	60 - 139	14.97	14.8	20	
2-Butanone	38.53	2.0	40	0	96.3	56 - 143	36.36	5.79	20	
2-Chlorotoluene	20.59	1.0	20	0	103	79 - 122	18.19	12.4	20	
2-Hexanone	42.3	2.0	40	0	106	57 - 139	41.7	1.43	20	
4-Chlorotoluene	20.9	1.0	20	0	105	78 - 122	18.42	12.6	20	
4-Isopropyltoluene	23.18	1.0	20	0	116	77 - 127	19.79	15.8	20	
4-Methyl-2-pentanone	41.93	2.0	40	0	105	67 - 130	40.54	3.38	20	
Acetone	37.85	2.0	40	0	94.6	39 - 160	35.14	7.44	20	
Benzene	20.02	1.0	20	0	100	79 - 120	17.3	14.6	20	
Bromobenzene	20.38	1.0	20	0	102	80 - 120	18.25	11	20	
Bromochloromethane	17.38	1.0	20	0	86.9	78 - 123	14.52	17.9	20	
Bromodichloromethane	19.99	1.0	20	0	99.9	79 - 125	16.54	18.9	20	
Bromoform	20.79	1.0	20	0	104	66 - 130	18.9	9.53	20	
Bromomethane	22	1.0	20	0	110	53 - 141	17.83	21	20	R

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334099		Instrument: VOA6		Method: SW8260						
MSD	Sample ID: HS19021185-01MSD	Units: UG/L			Analysis Date: 06-Mar-2019 00:30					
Client ID:	Run ID: VOA6_334099	SeqNo: 4977038		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	37.14	2.0	40	0	92.8	64 - 133	32.12	14.5	20	
Carbon tetrachloride	22.02	1.0	20	0	110	72 - 136	19.97	9.75	20	
Chlorobenzene	20.93	1.0	20	0	105	82 - 118	17.86	15.8	20	
Chloroethane	13.33	1.0	20	0	66.7	60 - 138	18.44	32.1	20	R
Chloroform	18.11	1.0	20	0	90.6	79 - 124	15.26	17.1	20	
Chloromethane	16.37	1.0	20	0	81.8	50 - 139	13.37	20.2	20	R
cis-1,2-Dichloroethene	18.3	1.0	20	0	91.5	78 - 123	15.51	16.5	20	
cis-1,3-Dichloropropene	19.4	1.0	20	0	97.0	75 - 124	16.22	17.9	20	
Dibromochloromethane	21.17	1.0	20	0	106	74 - 126	18.81	11.8	20	
Dibromomethane	19.86	1.0	20	0	99.3	79 - 123	17.29	13.9	20	
Dichlorodifluoromethane	12.96	1.0	20	0	64.8	32 - 152	12.17	6.28	20	
Ethylbenzene	21.52	1.0	20	0	108	79 - 121	18.62	14.4	20	
Hexachlorobutadiene	26.55	1.0	20	0	133	66 - 134	20.45	26	20	R
Isopropylbenzene	22.98	1.0	20	0	115	72 - 131	19.69	15.4	20	
m,p-Xylene	42.92	2.0	40	0	107	80 - 121	37	14.8	20	
Methylene chloride	18.35	2.0	20	0	91.7	74 - 124	14.99	20.1	20	R
Naphthalene	23.3	1.0	20	0	116	61 - 128	20.55	12.5	20	
n-Butylbenzene	23.15	1.0	20	0	116	75 - 128	19.41	17.6	20	
n-Propylbenzene	22.25	1.0	20	0	111	76 - 126	19.38	13.8	20	
o-Xylene	21.83	1.0	20	0	109	78 - 122	18.61	15.9	20	
sec-Butylbenzene	23.19	1.0	20	0	116	77 - 126	19.94	15	20	
Styrene	21.46	1.0	20	0	107	78 - 123	18.38	15.5	20	
tert-Butylbenzene	22.67	1.0	20	0	113	78 - 124	19.72	13.9	20	
Tetrachloroethene	22.98	1.0	20	0	115	74 - 129	20.08	13.5	20	
Toluene	21.26	1.0	20	0	106	80 - 121	18.67	13	20	
trans-1,2-Dichloroethene	18.95	1.0	20	0	94.8	75 - 124	16.03	16.7	20	
trans-1,3-Dichloropropene	19.37	1.0	20	0	96.9	73 - 127	16.42	16.5	20	
Trichloroethene	36.16	1.0	20	13.06	115	79 - 123	31.05	15.2	20	
Trichlorofluoromethane	18.94	1.0	20	0	94.7	65 - 141	17.41	8.37	20	
Vinyl chloride	16.23	1.0	20	0	81.1	58 - 137	14.49	11.3	20	
Surr: 1,2-Dichloroethane-d4	41.82	1.0	50	0	83.6	81 - 118	42.2	0.909	20	
Surr: 4-Bromofluorobenzene	51.21	1.0	50	0	102	85 - 114	50.35	1.7	20	
Surr: Dibromofluoromethane	42.43	1.0	50	0	84.9	80 - 119	42.74	0.725	20	
Surr: Toluene-d8	51.03	1.0	50	0	102	89 - 112	52.15	2.17	20	

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT**Batch ID:** R334099**Instrument:** VOA6**Method:** SW8260

The following samples were analyzed in this batch: HS19021165-01 HS19021165-02

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334211		Instrument: ICS2100		Method: SW9056						
MBLK	Sample ID: WBLKW1-030519	Units: mg/L			Analysis Date: 05-Mar-2019 18:30					
Client ID:	Run ID: ICS2100_334211	SeqNo: 4979542		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	0.500	0.500							U	
Sulfate	0.500	0.500							U	
LCS	Sample ID: WLCSW1-030519	Units: mg/L			Analysis Date: 05-Mar-2019 18:44					
Client ID:	Run ID: ICS2100_334211	SeqNo: 4979543		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	20.24	0.500	20	0	101	80 - 120				
Sulfate	19.49	0.500	20	0	97.4	80 - 120				
LCSD	Sample ID: WLCSDW1-030519	Units: mg/L			Analysis Date: 05-Mar-2019 18:59					
Client ID:	Run ID: ICS2100_334211	SeqNo: 4979544		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	20.35	0.500	20	0	102	80 - 120	20.24	0.567	20	
Sulfate	19.55	0.500	20	0	97.7	80 - 120	19.49	0.292	20	
MS	Sample ID: HS19030163-01MS	Units: mg/L			Analysis Date: 05-Mar-2019 23:42					
Client ID:	Run ID: ICS2100_334211	SeqNo: 4979557		PrepDate:			DF: 10			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	292.6	5.00	100	187.6	105	80 - 120				
Sulfate	199.7	5.00	100	98.43	101	80 - 120				
MS	Sample ID: HS19030163-01MS	Units: mg/L			Analysis Date: 05-Mar-2019 22:58					
Client ID:	Run ID: ICS2100_334211	SeqNo: 4979554		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	202.7	0.500	10	192.4	104	80 - 120			EO	
Sulfate	117.6	0.500	10	107.3	102	80 - 120			EO	

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

QC BATCH REPORT

Batch ID: R334211		Instrument: ICS2100		Method: SW9056						
MSD	Sample ID: HS19030163-01MSD	Units: mg/L			Analysis Date: 05-Mar-2019 23:57					
Client ID:	Run ID: ICS2100_334211	SeqNo: 4979558		PrepDate:			DF: 10			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	278.4	5.00	100	187.6	90.8	80 - 120	292.6	4.96	20	
Sulfate	188.6	5.00	100	98.43	90.2	80 - 120	199.7	5.7	20	
MSD	Sample ID: HS19030163-01MSD	Units: mg/L			Analysis Date: 05-Mar-2019 23:13					
Client ID:	Run ID: ICS2100_334211	SeqNo: 4979555		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	209.7	0.500	10	192.4	173	80 - 120	202.7	3.39	20 SEO	
Sulfate	120.8	0.500	10	107.3	135	80 - 120	117.6	2.7	20 SEO	

The following samples were analyzed in this batch: HS19021165-01

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19021165

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

Unit Reported	Description
mg/L	Milligrams per Liter

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19021165

Date/Time Received: **22-Feb-2019 09:06**
 Received by: **RPG**

Checklist completed by: Raegen Giga 22-Feb-2019
 eSignature Date

Reviewed by: RJ Modashia 22-Feb-2019
 eSignature Date

Matrices: **water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.6c/1.6c uc/c IR 11
 Cooler(s)/Kit(s): 24406
 Date/Time sample(s) sent to storage: 02/22/2019 11:59

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:


Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd. Suite 210, Houston, Tx. 77099 ATTN: R.J. Modashig

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS				Project No. NWO1312.0150.0 16.0001				Analyses <div style="text-align: center;"> HS19021165 Bhate Environmental Associates, Inc. Longhorn GW Treatment Plant </div> 																																																																																																																																																																																																																																																								
Job: GROUNDWATER TREATMENT PLANT BI-WEEKLY SAMPLES						MS / MSD No. OF CONTAINERS VOC CHLORIDE, SULFATE																																																																																																																																																																																																																																																										
Prepared By: Scott Beesinger				P.O Number																																																																																																																																																																																																																																																												
<table border="1" style="width:100%; border-collapse: collapse;"> <thead> <tr> <th>Field Sample I.D.</th> <th>Sample Matrix</th> <th>Date / Time</th> <th>MS / MSD</th> <th>No. OF CONTAINERS</th> <th>VOC</th> <th>CHLORIDE, SULFATE</th> <th colspan="10">Analyses</th> <th>Remarks (Preservatives, etc.)</th> <th>Lab I.D.#</th> </tr> </thead> <tbody> <tr> <td>LH18/24-SP650_022119</td> <td>Water</td> <td>02/21/19 / 14:00</td> <td></td> <td>3</td> <td>3</td> <td></td> <td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td> <td>HCL</td> <td></td> </tr> <tr> <td>LH18/24-SP650_022119</td> <td>Water</td> <td>02/21/19 / 14:00</td> <td></td> <td>1</td> <td></td> <td>1</td> <td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td> <td>NONE</td> <td></td> </tr> <tr> <td>Trip Blank</td> <td>Water</td> <td>02/21/19</td> <td></td> <td>2</td> <td>2</td> <td></td> <td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td> <td>HCL</td> <td></td> </tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> <tr><td> </td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td><td></td></tr> </tbody> </table>																		Field Sample I.D.	Sample Matrix	Date / Time	MS / MSD	No. OF CONTAINERS	VOC	CHLORIDE, SULFATE	Analyses										Remarks (Preservatives, etc.)	Lab I.D.#	LH18/24-SP650_022119	Water	02/21/19 / 14:00		3	3													HCL		LH18/24-SP650_022119	Water	02/21/19 / 14:00		1		1												NONE		Trip Blank	Water	02/21/19		2	2													HCL																																																																																																																																																																	
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Trip Blank	Water	02/21/19		2	2													HCL																																																																																																																																																																																																																																														
Additional Remarks: STANDARD TAT ON ALL PARAMETERS.																																																																																																																																																																																																																																																																
Relinquished By: <i>Scott Beesinger</i>		Date: 02/21/19	Time: 14:30	Received By:		Date:	Time:	Relinquished By: <i>R. C. Cages</i>		Date:	Time:	Received By:		Date:	Time:	(Signature)																																																																																																																																																																																																																																																
For Lab Use Only <table border="1" style="width:100%; border-collapse: collapse;"> <tr> <td colspan="2"> Received At Lab By: </td> <td> Date </td> <td> Time </td> <td> Airbill No. </td> <td> Opened By: </td> <td> Date </td> <td> Time </td> <td> Temp of Container </td> <td> Seal No. </td> <td> Condition </td> </tr> <tr> <td colspan="11"> Remarks: </td> </tr> </table>																		Received At Lab By:		Date	Time	Airbill No.	Opened By:	Date	Time	Temp of Container	Seal No.	Condition	Remarks:																																																																																																																																																																																																																																			
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Remarks:																																																																																																																																																																																																																																																																
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ALS
 10450 Stancliff Rd., Suite 210
 Houston, Texas 77099
 Tel. +1 281 530 5856
 Fax. +1 281 530 5887

CUSTODY SEAL
 Date: 2/21/99 Time: 14:30
 Name: Susan E. Brown
 Company: ZHE-4

Seal Broken By:
 2/21/99

TRK# 4380 9530 9467
 0221

FedEx
 TRK# 4380 9530 9467
 0221

AB SGRA

RETURNS MON-SAT
PRIORITY OVERNIGHT
 FRI - 22 FEB 10:30A
PRIORITY OVERNIGHT

77099
 TX-US
 IAH



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 12, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030011**

Laboratory Results for: **LH18/24 GW Treatment Plant Monthly Influent Samples**

Dear Marcia,

ALS Environmental received 1 sample(s) on Mar 01, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: DAYNA.FISHER
RJ Modashia
Project Manager

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
Work Order: HS19030011

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030011-01	LH18/24-SP140_022819	Water		28-Feb-2019 14:00	01-Mar-2019 08:50	<input type="checkbox"/>

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
Work Order: HS19030011

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
-

Metals by Method SW6020**Batch ID: 138281**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

WetChemistry by Method SW7196**Batch ID: R334254**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 GW Treatment Plant Monthly Influent Samples
 Sample ID: LH18/24-SP140_022819
 Collection Date: 28-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030011
 Lab ID:HS19030011-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
ICP-MS METALS BY SW6020A		Method:SW6020				Prep:SW3010A / 01-Mar-2019		Analyst: JHD
Selenium	0.00250	U	0.00110	0.00250	0.00200	mg/L	1	04-Mar-2019 22:47
Silver	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	04-Mar-2019 22:47
HEXAVALENT CHROMIUM BY SW7196A		Method:SW7196						Analyst: MZD
Chromium, Hexavalent	0.0100	U	0.00600	0.0100	0.0100	mg/L	1	01-Mar-2019 11:42
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47

WEIGHT LOG

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030011

Batch ID: 138281 **Method:** ICP-MS METALS BY SW6020A **Prep:** 3010A

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19030011-01	1	10	10 (mL)	1

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030011

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID 138281		Test Name : ICP-MS METALS BY SW6020A			Matrix: Water	
HS19030011-01	LH18/24-SP140_022819	28 Feb 2019 14:00		01 Mar 2019 10:30	04 Mar 2019 22:47	1
Batch ID R334176		Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)			Matrix: Water	
HS19030011-01	LH18/24-SP140_022819	28 Feb 2019 14:00			07 Mar 2019 17:47	1
Batch ID R334254		Test Name : HEXAVALENT CHROMIUM BY SW7196A			Matrix: Water	
HS19030011-01	LH18/24-SP140_022819	28 Feb 2019 14:00			01 Mar 2019 11:42	1

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030011

QC BATCH REPORT

Batch ID: 138281		Instrument: ICPMS05		Method: SW6020						
MBLK	Sample ID: MBLK-138281	Units: mg/L		Analysis Date: 04-Mar-2019 21:51						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974208	PrepDate: 01-Mar-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.00250	0.00200							U	
Silver	0.000500	0.00200							U	
LCS	Sample ID: LCS-138281	Units: mg/L		Analysis Date: 04-Mar-2019 21:53						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974209	PrepDate: 01-Mar-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.05168	0.00200	0.05	0	103	80 - 120				
Silver	0.05504	0.00200	0.05	0	110	80 - 120				
MS	Sample ID: HS19021441-01MS	Units: mg/L		Analysis Date: 04-Mar-2019 22:00						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974212	PrepDate: 01-Mar-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.04948	0.00200	0.05	0	99.0	80 - 120				
Silver	0.0515	0.00200	0.05	0	103	80 - 120				
MSD	Sample ID: HS19021441-01MSD	Units: mg/L		Analysis Date: 04-Mar-2019 22:02						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974213	PrepDate: 01-Mar-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.04959	0.00200	0.05	0	99.2	80 - 120	0.04948	0.218	20	
Silver	0.05174	0.00200	0.05	0	103	80 - 120	0.0515	0.473	20	
PDS	Sample ID: HS19021441-01PDS	Units: mg/L		Analysis Date: 04-Mar-2019 22:04						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974214	PrepDate: 01-Mar-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.09793	0.00200	0.1	0	97.9	75 - 125				
Silver	0.09993	0.00200	0.1	0	99.9	75 - 125				

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030011

QC BATCH REPORT

Batch ID: 138281		Instrument: ICPMS05		Method: SW6020						
SD	Sample ID: HS19021441-01SD	Units: mg/L			Analysis Date: 04-Mar-2019 21:57					
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974211	PrepDate: 01-Mar-2019	DF: 5						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	%D Limit	Qual
Selenium	0.0125	0.0100					0.000452	0	10	U
Silver	0.00250	0.0100					0.00002	0	10	U

The following samples were analyzed in this batch: HS19030011-01

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030011

QC BATCH REPORT

Batch ID: R334254		Instrument: UV-2450		Method: SW7196					
MBLK	Sample ID: MBLK-334254	Units: mg/L		Analysis Date: 01-Mar-2019 11:42					
Client ID:	Run ID: UV-2450_334254	SeqNo: 4986763		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Chromium, Hexavalent	0.0100	0.0100							U
LCS	Sample ID: LCS-334254	Units: mg/L		Analysis Date: 01-Mar-2019 11:42					
Client ID:	Run ID: UV-2450_334254	SeqNo: 4986764		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Chromium, Hexavalent	0.249	0.0100	0.25	0	99.6	90 - 111			
LCSD	Sample ID: LCSD-334254	Units: mg/L		Analysis Date: 01-Mar-2019 11:42					
Client ID:	Run ID: UV-2450_334254	SeqNo: 4986765		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Chromium, Hexavalent	0.241	0.0100	0.25	0	96.4	90 - 111	0.249	3.27	20

The following samples were analyzed in this batch: HS19030011-01

ALS Houston, US

Date: 12-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030011

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030011

Date/Time Received: **01-Mar-2019 08:50**
 Received by: **JRM**

Checklist completed by: Jared R. Makan 1-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 1-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **ALS Courier**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.8c/1.8c UC/C IR25
 Cooler(s)/Kit(s): 43795
 Date/Time sample(s) sent to storage: 03/01/2019 10:30

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd., Suite 210 Houston, TX. 77099 (281) 530-5656 ATTN: RJ Modshia

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001		Analyses																			
Job: GROUNDWATER TREATMENT PLANT MONTHLY INFLUENT SAMPLES					MS / MSD	No. OF CONTAINERS	SILVER & SELENIUM	HEXAVALENT CHROMIUM	PERCHLORATE															
Prepared By: Scott Beesinger			P.O. Number																					
Field Sample I.D.		Sample Matrix	Date / Time																			Remarks (Preservatives, etc.)	Lab I.D.#	
LH18/24-SP140_022819		Water	02/28/19 / 14:00		1	X																HNO3		
LH18/24-SP140_022819		Water	02/28/19 / 14:00		2		X	X															NONE	
<p style="font-size: 1.2em; margin: 0;">HS19030011</p> <p style="margin: 0;">Bhate Environmental Associates, Inc. 18/24 GW Treatment Plant Monthly Influent Samples</p>																								
Additional Remarks: <u>STANDARD TURN AROUND TIME</u>																								
Relinquished By:		Date	Time	Received By:		Date	Time	Relinquished By:		Date	Time	Received By:		Date	Time									
<i>Scott Beesinger</i>		02/28/19	14:30	<i>J. MUMFORD</i>		3/1/19	08:50																	
For Lab Use Only																								
Received At Lab By:		Date	Time	Airbill No.		Opened By:		Date	Time	Temp of Container	Seal No.	Condition												
Remarks: Cooler 43795 Temp 1.8 1R25 CFO.0																								

(Word) S:\1-ees\Forms\Chain of Custody - BiWeekly

TRK# 4380 9530 9423
0221


FRI - 01 MAR 10:30A
PRIORITY OVERNIGHT

AB SGRA

77099
TX-US
IAH



364688 01Mar 00:38 APWH 547C210F30A17C

 <p>ALS Environmental 10450 Stancif. Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 6656 Fax. +1 281 530 6887</p>	<p>CUSTOMER</p> <p>Date: 2/27/19 Name: S. G. B. Company: S. G. B.</p>

BODY SEAL	
Type: 1430	Seal Broken By: JM
2151846	Date: 3/01/19
29	



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1905651; 1906112; 1906330;
1906332; 1906334

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2223 (233911)

General Set Information: There were thirteen field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Method QC data: The method blank (LMB 642099) was less than 1/2 the CRDL. The recovery for the LCS (642100) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0 μ l of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.0 μ g/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in μ g/L. Results were calculated in μ g/L by the equation $(A) \times (B)$,

where: A = Analyte concentration from the standard curve (μ g/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level of 4.0 μ g/L. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported. Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEBI04) along with datafiles 05MARD07-10.

Thomas Bosch March 06, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 07, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1906330**

Project ID: HS19030011

Purchase Order: HS19030011

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP140_022819	1906330001	02/28/19	03/02/19	



ANALYTICAL REPORT

Workorder: **34-1906330**Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP140_022819	Sampling Site: NA	Collected: 02/28/2019				
Lab ID: 1906330001	Media: 125 mL Nalgene	Received: 03/02/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 13:54	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	3700	100	200	400	100	

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 233911)

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/06/2019 09:17	/S/ Stephen Brose 03/07/2019 11:04

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1906330

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00934538

Analysis Information

Workorder: 1906330

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2223 (HBN: 233911)
Analyzed By: Thomas Bosch

Blank

LMB: 642099 Analyzed: 03/05/2019 09:26 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 642100 Analyzed: 03/05/2019 09:00 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.11	4.00	103	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1906112001 Analyzed: 03/05/2019 09:54 Dilution: 1 Units: ug/L			MS: 1906112002 Analyzed: 03/05/2019 10:07 Dilution: 1 Units: ug/L			MSD: 1906112003 Analyzed: 03/05/2019 10:20 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	51.0	53.9	4	▲ 65.7	78.8 123.8	53.7	▲ 61.2	0.337	0.0 20.0

Continuing Calibration Verification

CCV: 642096 Analyzed: 03/05/2019 08:44 Units: ug/L Criteria: ± 15%			CCV: 642101 Analyzed: 03/05/2019 12:18 Units: ug/L Criteria: ± 15%			CCV: 642102 Analyzed: 03/05/2019 14:07 Units: ug/L Criteria: ± 15%			
Analyte	Result	Target	% Rec.	Result	Target	% Rec.	Result	Target	% Rec.
Perchlorate	25.0	25.0	100	25.0	25.0	100	24.5	25.0	97.8

Interference Check Sample

ICSA: 642098 Analyzed: 03/05/2019 09:13 Units: ug/L Criteria: ± 30%			
Analyte	Result	Target	% Rec.
Perchlorate	3.93	4.00	98.2

Comments

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.



Quality Control Sample Batch Report

00934539

Analysis Information

Workorder: 1906330

Limits: Client SOW/Contract Specified

Preparation: NA

Analysis: EPA 6850, DoD QSM

Basis: DoD QSM

Batch: NA

Batch: ELMS/2223 (HBN: 233911)

Prepared By: NA

Analyzed By: Thomas Bosch

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/06/2019 13:45	/S/ Stephen Brose 03/07/2019 11:04

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



18698/#2

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Subcontract Chain of Custody

COC ID: 10853

1906330

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030011
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030011-01	LH18/24-SP140_022819	Water	28 Feb 2019 14:00
SUB_Perch-6850			15 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: *F*
Received By: *Meredyl Jewell*
Cooler ID(s): 7283

Date/Time: 3/1/19 1800
Date/Time: 3/2/19 1903
Temperature(s): 3

ALS Environmental
CHAIN-OF-CUSTODY



Project / Job / Task: HS19030011		Split:	Workorder ID: 1906330	Level: ENV_LVL4		Requested Analysis				
Client: ALS Environmental (Houston)			Account: 8101	Type: 125Poly						
Comments:						EPA 6850, DOD QSM				
Item	Collect Date/Time	Sample ID	Lab ID	QC	Matrix	ID(s)	Count			
1	02/28/2019 14:00	LH18/24-SP140_022819	1906330001		Water	A	1			
2										
3										
4										
5										
6										
7										
8										
9										
10										

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY					SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY				
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location	Sample Prep / Analysis for:	Prepared / Analyzed by:	Reinquired By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location
Wraith, Julie	03/02/2019 09:03	ALS Sample Receiving	Sample Login	Lab Notebook No.:	Date / Time:				
<i>Julie Wraith</i>	03-01-19 1400	13B	Storage						
B.33.1	03-01-19 16:10	T. Board	6850						

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: ALS Houston Project/Task/Site: 19060330
 Date/Time of Receipt: 3/2/19 903 Number of Coolers Received: 1

Condition of Coolers:	Acceptable/Unacceptable	Temperature Control:	Present/Not Included
Cooler Custody Seals:	Present/Absent/NA	Location Temp Taken:	Control/Between Samples
Container Custody Seals:	Present/Absent/NA	Are all temperatures within project specific guidelines?	Yes/No/NA
Ice Present:	Yes/No/NA	VOA Headspace Present?	Yes/No/NA
	Frozen/Melted/NA		

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 9203	3 °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: [Signature] [Signature] 3/2/19
Signature Printed Name Date

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Must Deliver Next Business Day
Time and Tempature Sensitive!

Part # 150488-434 RT2 Exp 11/19

ORIGIN ID:SGRA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

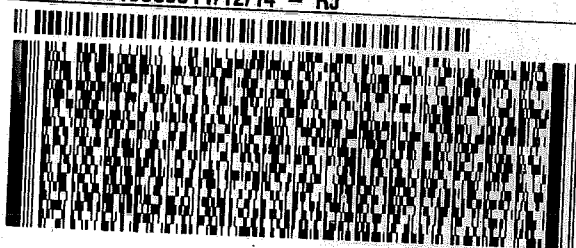
SHIP DATE: 01MAR19
ACTWGT: 9.55 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(801) 288-7700

REF: HS19030011/12/14 - RJ



FedEx
Express



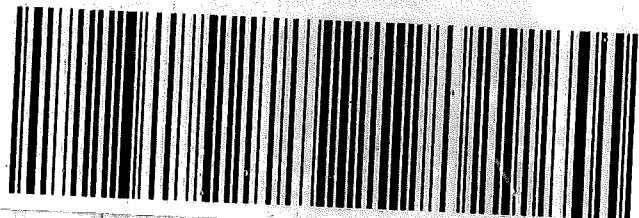
ATL10509081181JF

TRK# 4809 7831 2469
0201

SATURDAY 12:00P
PRIORITY OVERNIGHT

XO BTFA

84123
UT-US SLC



FedEx® Saturday Delivery

151956 1004 MW1

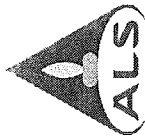


SDR

RT **B639** 1
ST **F1** 12:00



ALS
10450
Houston
Tel: +1
Fax: +1



Batch Worklist

HBN: 233911

Instrument:



Status: WP

Created: 3/5/2019 08:21

Analyst: T. Bosch

Batch: ELMS/ 2223
 Rule: EPA 6850, DoD QSM Water

- Workorder: 1905651 [ENV_LVL4]
- Workorder: 1906112 [ENV_LVL4]
- Workorder: 1906330 [ENV_LVL4]
- Workorder: 1906332 [ENV_LVL4]
- Workorder: 1906334 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mix	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	642096	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
2	642097	RLVS for HBN 233911 [ELMS/2223]				RLVS	3		E685041C3Q	5311		3/7/2019	
3	642098	ICS for HBN 233911 [ELMS/2223]				ICS	3		E6850.D3Q	5311		3/7/2019	
4	642099	LMB for HBN 233911 [ELMS/2223]				LMB	3		E6850Q413Q	5311		3/7/2019	
5	642100	LCS for HBN 233911 [ELMS/2223]				LCS	3		E6850Q413Q	5311		3/7/2019	
6	1905651001	HS19021158-02/LH18/24-SP650_02				SAMPLE	3	1905651001-A	E6850Q41.3	5480	3/21/2019	3/7/2019	
7	1906112001	EW01_022619				SAMPLE	3	1906112001-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
8	1906112002	EW01_022619MS				MS	3	1906112002-A	E6850Q413Q	5480		3/7/2019	
9	1906112003	EW01_022619MSD				MSD	3	1906112003-A	E6850Q413Q	5480		3/7/2019	
10	1906112004	EW05_022619				SAMPLE	3	1906112004-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
11	1906112005	EW05_022619_FD				FLDDUP	3	1906112005-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
12	1906112006	EW02_022619				SAMPLE	3	1906112006-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
13	1906112007	EW06_022619				SAMPLE	3	1906112007-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
14	1906112008	EW03_022619				SAMPLE	3	1906112008-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
15	1906112009	EW07_022619				SAMPLE	3	1906112009-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
16	642101	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
17	1906112010	EW04_022619				SAMPLE	3	1906112010-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
18	1906112011	EW08_022619				SAMPLE	3	1906112011-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
19	1906330001	LH18/24-SP140_022819				SAMPLE	3	1906330001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
20	1906332001	LH18/24-SP650_022819_BIX				SAMPLE	3	1906332001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
21	1906334001	LH18-24-SP650_022819_BIX				SAMPLE	3	1906334001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
22	642102	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1905651 (001); 1906112 (001-11); 1906330 (001); 1906332 (001); 1906334 (001)
 ELMS Batch/HBN ID: 2223 (233911)
 Prep Date: 03/04/2019 Analysis Date: 03/05/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\05MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by TNB. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 02/15/2019, sequence 15FEB19D.s Offline Quantitation Method: CLO4-DP1.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 3 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 642100; Target = 4.0µg/L. ASTM type II water was used for LMB 642099.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\23911-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEB104) along with datafiles 05MARD07-10.

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
Batch(es)/SDG: E LMS: 2223 HBN: 233911		
Sample Set IDs if Applicable: 1905651 / 190912 / 1906330 1906332 / 1906334		
Calibration standards analyzed and meets criteria	TB	SB
Standards traceability checked and meets criteria	TB	SB
Standard curve coefficients evaluated and meet criteria	TB	SB
ICVs analyzed and meet acceptance criteria	TB	SB
CCVs analyzed and meet acceptance criteria	TB	SB
Method Blanks analyzed and meet acceptance criteria	TB	SB
Retention Time Windows checked	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
Surrogate recoveries checked and appropriately addressed	—	—
Method Preparation Blanks analyzed and meet acceptance criteria	TB	SB
MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed	TB	SB
RLVS analyzed	TB	SB
Preparation and analysis hold times met	TB	SB
Preparation deviations and re-preparations noted when performed	TB	SB
Analysis deviations and re-analyses noted when performed	TB	SB
Sample dilution factors noted on reports	TB	SB
Electronic records in HBN transcription accuracy and completeness checked	TB	SB
Preparation and analysis calculations checked	TB	SB
NCRs are completed as necessary NC/CAR# _____	—	—
Report forms are complete and accurate	TB	SB
Manual integrations checked	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850.WKG Std 100.ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK			Description - 6850 Stock AccStd 1,000ug/mL
Standard: 43659 MFG: AccuStandard MFG Lot: 218065075 Part ID: IC-PER-10X-1		Created By: Thomas Bosch Create Date: 09/17/2018 09:09AM Amount: 100 mL Expires: 07/25/2020 Usable: No Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT			Description - 6850 Intermdt AccStd 10.ug/mL		
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK			Description: 6850 QC WKG STD 100ug/L		
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdf Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK			Description: -6850 QC Stock STD 1,000ug/mL
Standard: 36748		Created By: Thomas Bosch	Amount: 100 mL
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	Expires: 03/31/2020
MFG Lot: CP-0860			Usable: Yes
Part ID: ICC-013			Lab Lot: CLO4 QC STOCK
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730		Created By: Thomas Bosch		Amount: 25 mL	
MFG: ALS/SLC		Create Date: 09/20/2018 09:09AM		Expires: 09/20/2019	
MFG Lot: TNB: 05/09/2018		Verified By: Thomas Bosch		Usable: Yes	
Pipette ID: Not Provided		Verify Date:		Lab Lot: CLO4ISTDWRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	.CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

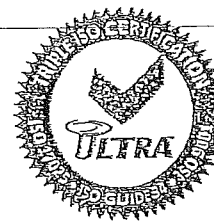
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:
This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:
Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:
The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:
This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:
This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:
Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

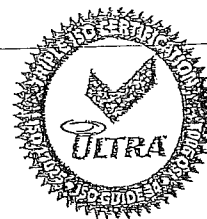
Hazards:
Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:
The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis

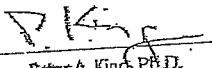


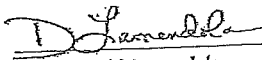
ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:
The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QAVRA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleared Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

For use in routine laboratory analysis.



Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:

ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: NaCl*O4

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 µg/mL
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 ± 2.8 µg/mL (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount	
*	642096	CCV@25	Vial 71	1	Control	1	2.57589e6	8.017	25.04585
*	642100	QC@4.0	Vial 72	1	Control	2	4.23307e5	8.139	4.10749
*	642098	ICS@4.0	Vial 73	1	Control	3	3.17619e5	7.883	3.92708
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	7.71127e6	7.438	51.25151
*	1906112002	MS	Vial 77	1	Sample	7	8.02696e6	7.464	53.87972
*	1906112003	MSD	Vial 78	1	Sample	8	7.94242e6	7.442	53.69940
*	1906112004		Vial 79	1	Sample	9	2.33017e7	7.227	172.36522
*	1906112005		Vial 80	1	Sample	10	2.30991e7	7.229	180.38568
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.33611e6	8.114	25.01987
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	2.53126e6	7.736	226.48155
*	1906112005	10X	Vial 91	1	Sample	22	2.68237e6	7.739	241.03710
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	3.77363e6	8.090	3708.65666
*	642102	CCV@25	Vial 71	1	Control	25	2.15787e6	8.144	24.45533

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount	
*	642096	CCV@25	Vial 71	1	Control	1	6.79028e5	8.036	25.15984
*	642100	QC@4.0	Vial 72	1	Control	2	1.27412e5	8.156	4.45187
*	642098	ICS@4.0	Vial 73	1	Control	3	9.68721e4	7.906	4.29888
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	2.02786e6	7.452	51.10329
*	1906112002	MS	Vial 77	1	Sample	7	2.14637e6	7.478	54.46797
*	1906112003	MSD	Vial 78	1	Sample	8	2.10991e6	7.456	53.98100
*	1906112004		Vial 79	1	Sample	9	6.79668e6	7.243	180.45962
*	1906112005		Vial 80	1	Sample	10	6.82834e6	7.246	190.34382
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	6.28961e5	8.128	25.63410
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	6.46870e5	7.754	220.91905
*	1906112005	10X	Vial 91	1	Sample	22	6.95643e5	7.756	238.44674
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	1.01021e6	8.102	3771.67475
*	642102	CCV@25	Vial 71	1	Control	25	5.83711e5	8.160	25.16652

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount	
*	642096	CCV@25	Vial 71	1	Control	1	3.12207e5	8.043	5.00000
*	642100	QC@4.0	Vial 72	1	Control	2	3.41038e5	8.157	5.00000
*	642098	ICS@4.0	Vial 73	1	Control	3	2.68237e5	7.902	5.00000
*	642099	LMB	Vial 74	1	Control	4	3.53313e5	8.102	5.00000
*	1905651001		Vial 75	1	Sample	5	3.26356e5	7.780	5.00000
*	1906112001		Vial 76	1	Sample	6	4.26473e5	7.464	5.00000
*	1906112002	MS	Vial 77	1	Sample	7	4.19549e5	7.491	5.00000
*	1906112003	MSD	Vial 78	1	Sample	8	4.16709e5	7.467	5.00000
*	1906112004		Vial 79	1	Sample	9	2.95705e5	7.253	5.00000
*	1906112005		Vial 80	1	Sample	10	2.75946e5	7.263	5.00000
*	1906112006		Vial 81	1	Sample	11	3.73575e5	7.480	5.00000
*	1906112007		Vial 82	1	Sample	12	5.00533e5	7.256	5.00000
*	1906112008		Vial 83	1	Sample	13	4.65121e5	7.237	5.00000
*	1906112009		Vial 84	1	Sample	14	2.43675e5	7.672	5.00000
*	1906112010		Vial 85	1	Sample	15	2.51865e5	7.710	5.00000
*	1906112011		Vial 86	1	Sample	16	2.80792e5	7.876	5.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.83460e5	8.140	5.00000
*	1906332001		Vial 88	1	Sample	19	2.64674e5	7.827	5.00000
*	1906334001		Vial 89	1	Sample	20	2.65662e5	7.808	5.00000
*	1906112004	10X	Vial 90	1	Sample	21	3.41608e5	7.756	50.00000
*	1906112005	10X	Vial 91	1	Sample	22	3.38724e5	7.764	50.00000
*	1906112007	RE	Vial 82	1	Sample	23	4.31135e5	7.332	5.00000
*	1906330001	100	Vial 92	1	Sample	24	2.98985e5	8.111	500.00000
*	642102	CCV@25	Vial 71	1	Control	25	2.68305e5	8.165	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

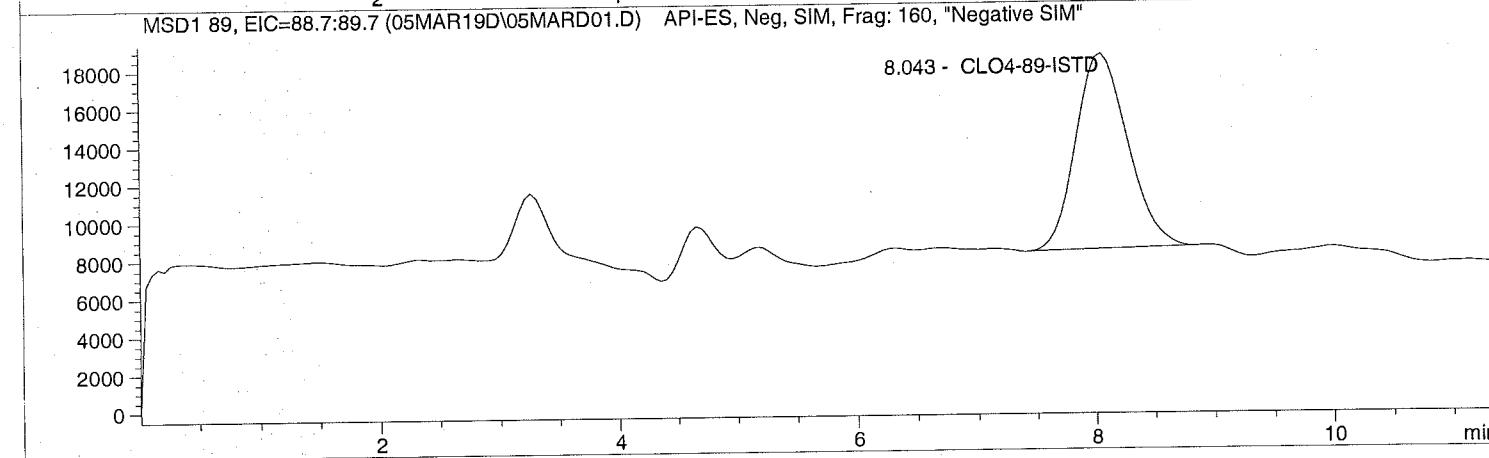
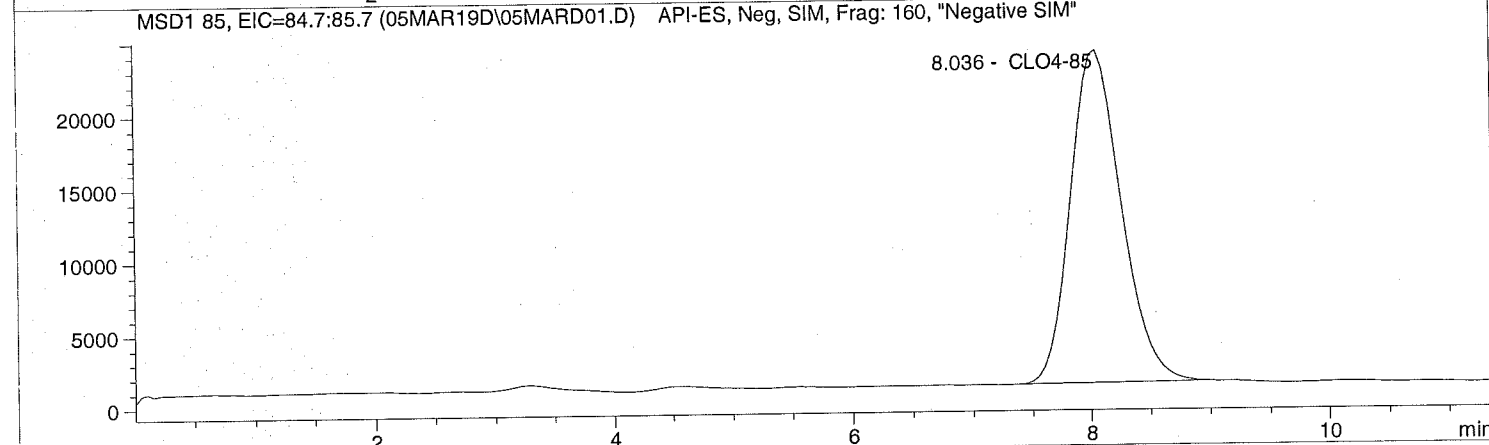
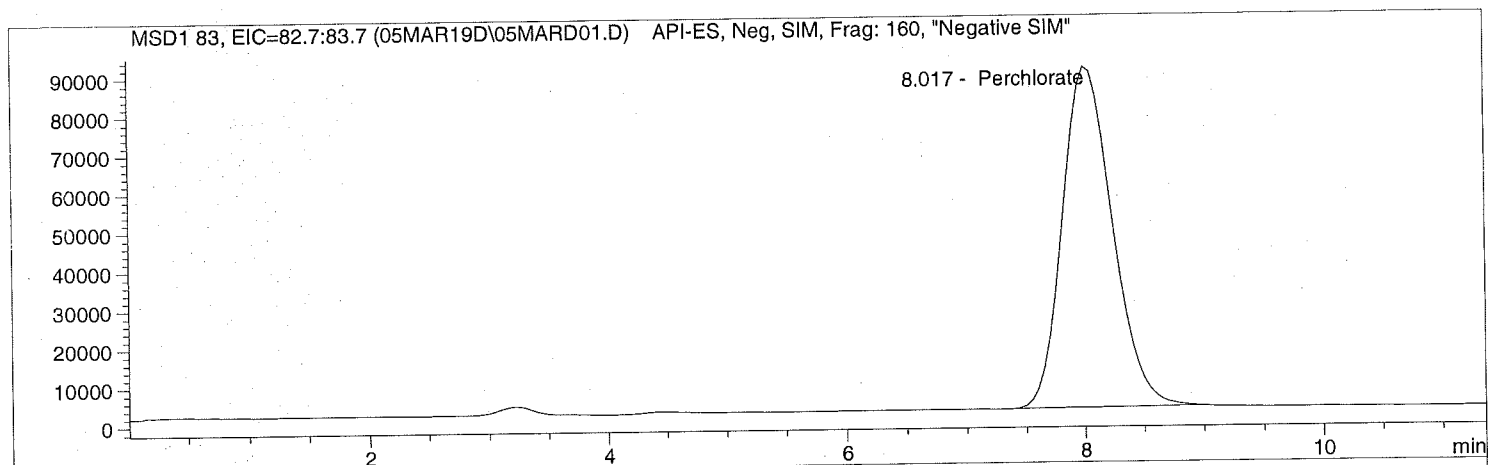
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	642096	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	642100	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	642098	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	642099	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1905651001		CLO4-AQN	1	Sample	
6	Vial 76	1906112001		CLO4-AQN	1	Sample	
7	Vial 77	1906112002	MS	CLO4-AQN	1	Sample	
8	Vial 78	1906112003	MSD	CLO4-AQN	1	Sample	
9	Vial 79	1906112004		CLO4-AQN	1	Sample	
10	Vial 80	1906112005		CLO4-AQN	1	Sample	
11	Vial 81	1906112006		CLO4-AQN	1	Sample	
12	Vial 82	1906112007		CLO4-AQN	1	Sample	
13	Vial 83	1906112008		CLO4-AQN	1	Sample	
14	Vial 84	1906112009		CLO4-AQN	1	Sample	
15	Vial 85	1906112010		CLO4-AQN	1	Sample	
16	Vial 86	1906112011		CLO4-AQN	1	Sample	
17	Vial 71	642101	CCV@25	CLO4-AQN	1	Ctrl Samp	
18	Vial 87	1906330001	1K	CLO4-AQN	1	Sample	
19	Vial 88	1906332001		CLO4-AQN	1	Sample	
20	Vial 89	1906334001		CLO4-AQN	1	Sample	
21	Vial 90	1906112004	10X	CLO4-AQN	1	Sample	
22	Vial 91	1906112005	10X	CLO4-AQN	1	Sample	
23	Vial 82	1906112007	RE	CLO4-AQN	1	Sample	
24	Vial 92	1906330001	100	CLO4-AQN	1	Sample	
25	Vial 71	642102	CCV@25	CLO4-AQN	1	Ctrl Samp	

Injection Date: 3/05/2019 08:44:45
Sample Name: 642096 CCV@25
Acq Operator: TNB

Seq Line: 1
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 08:44:45 Seq Line: 1
Sample Name: 642096 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.017	PBA	2575886.3	25.0459	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.036	PBA	679028.4	25.1598	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.043	PBA	312206.9	5.0000	CLO4-89-ISTD

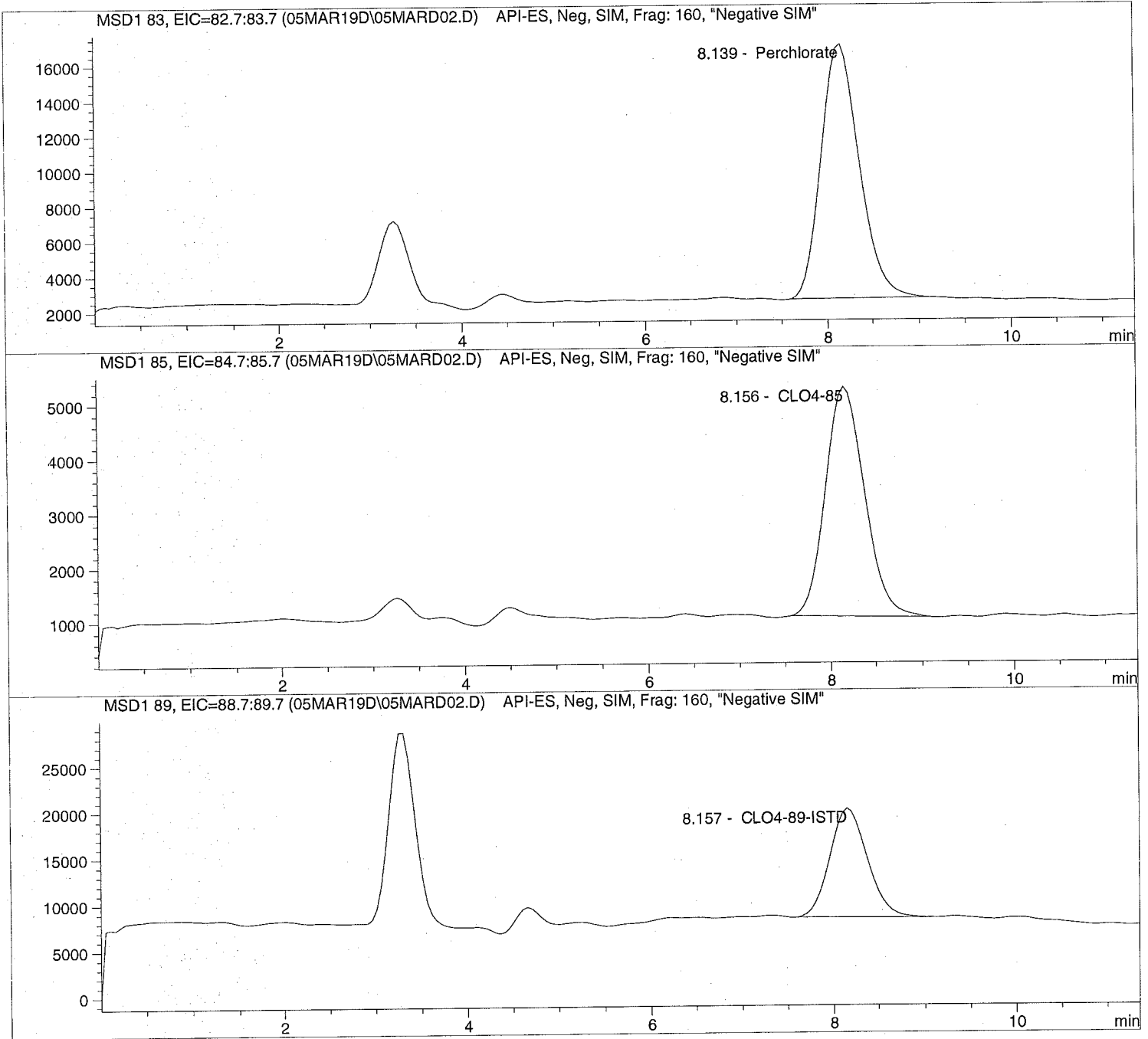
*** End of Report ***

Injection Date: 3/05/2019 09:00:30
Sample Name: 642100 QC@4.0
Acq Operator: TNB

Seq Line: 2
Location: Vial 72
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 09:00:30 Seq Line: 2
Sample Name: 642100 QC@4.0 Location: Vial 72
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.139	PBA	423307.1	4.1075	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.156	PBA	127412.0	4.4519	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.157	PBA	341038.3	5.0000	CLO4-89-ISTD

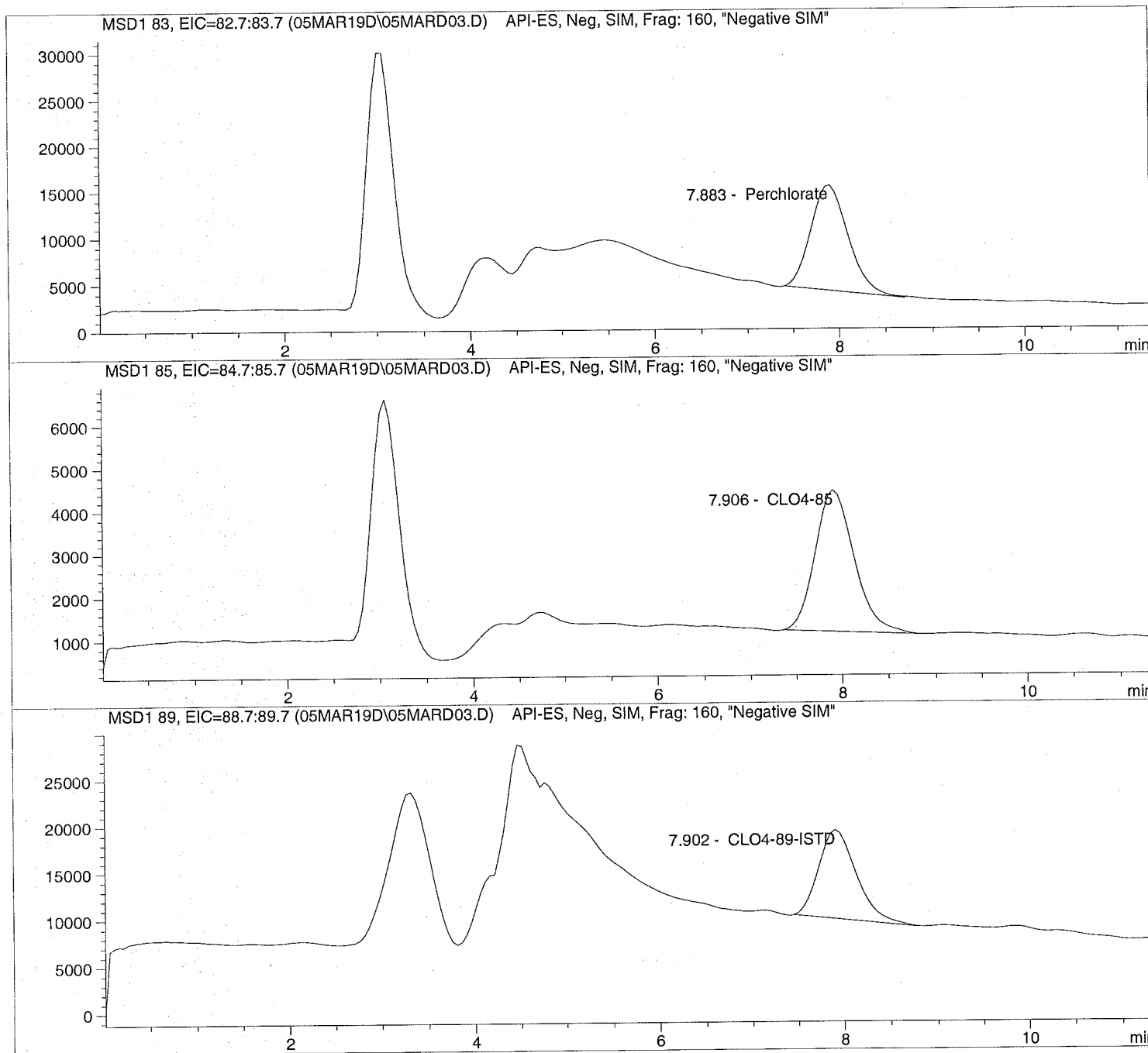
*** End of Report ***

Injection Date: 3/05/2019 09:13:34
Sample Name: 642098 ICS@4.0
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 09:13:34      Seq Line: 3  
Sample Name: 642098 ICS@4.0           Location: Vial 73  
Acq Operator: TNB                      Inj. No.: 1  
                                         Inj. Vol.: 20 µl  
=====
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 4.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.883	PBA	317618.9	3.9271	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.906	PBA	96872.1	4.2989	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.902	PBA	268236.7	5.0000	CLO4-89-ISTD

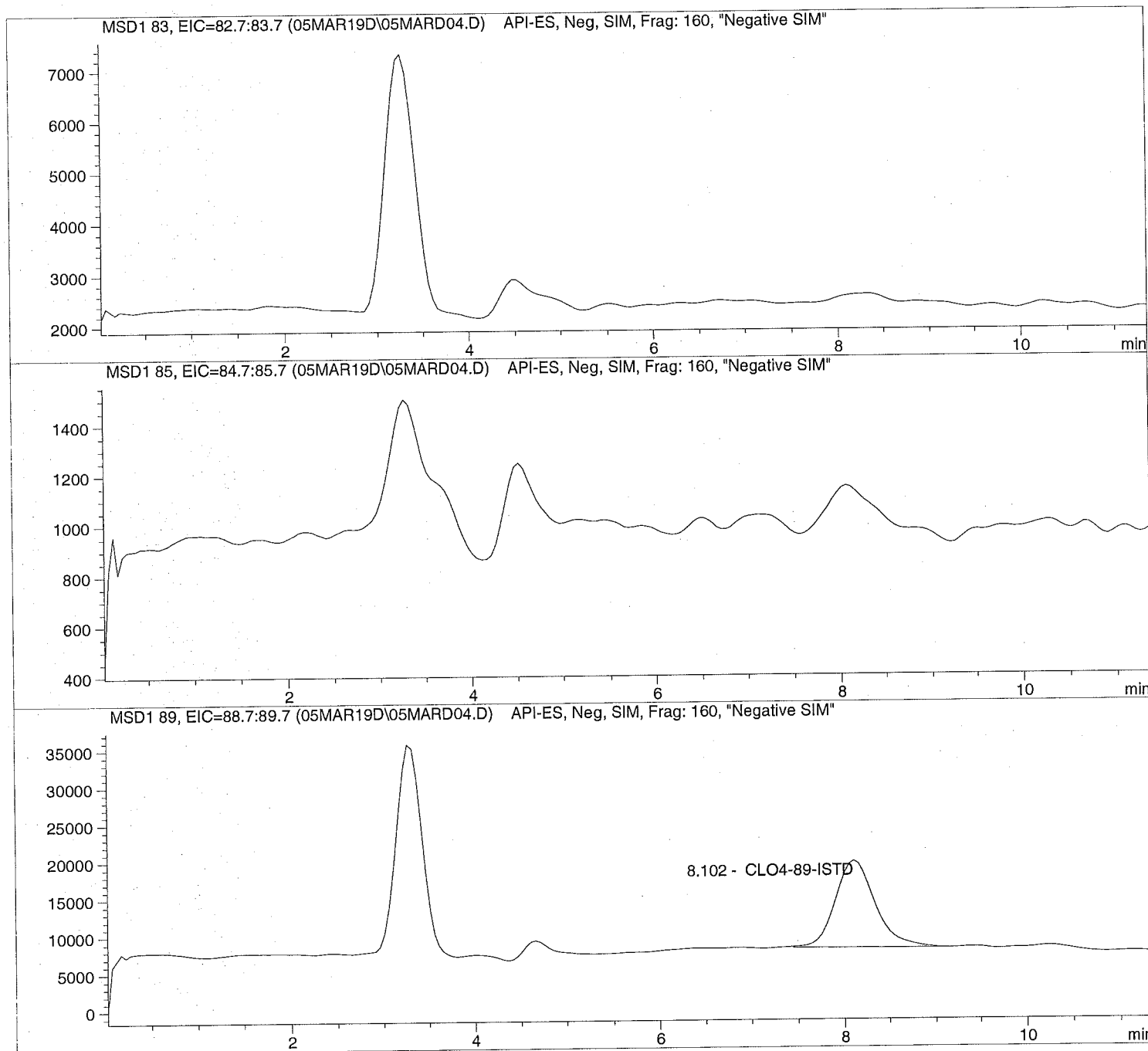
=====
*** End of Report ***
=====

Injection Date: 3/05/2019 09:26:40
Sample Name: 642099 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 09:26:40 Seq Line: 4
Sample Name: 642099 LMB Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	BBA	353313.1	5.0000	CLO4-89-ISTD

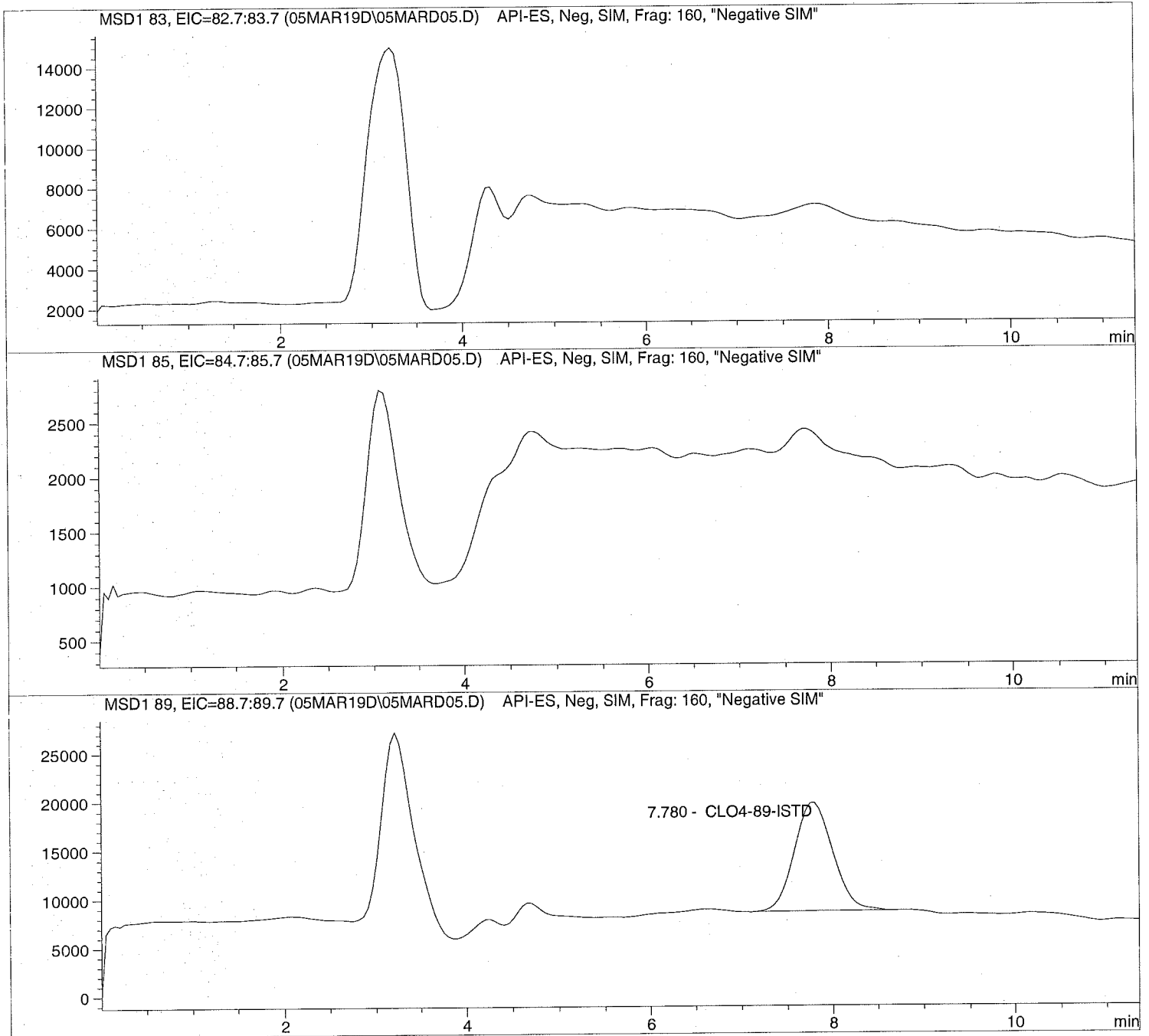
*** End of Report ***

Injection Date: 3/05/2019 09:40:58
Sample Name: 1905651001
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 09:40:58      Seq Line: 5
Sample Name: 1905651001                Location: Vial 75
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.780	PBA	326356.2	5.0000	CLO4-89-ISTD

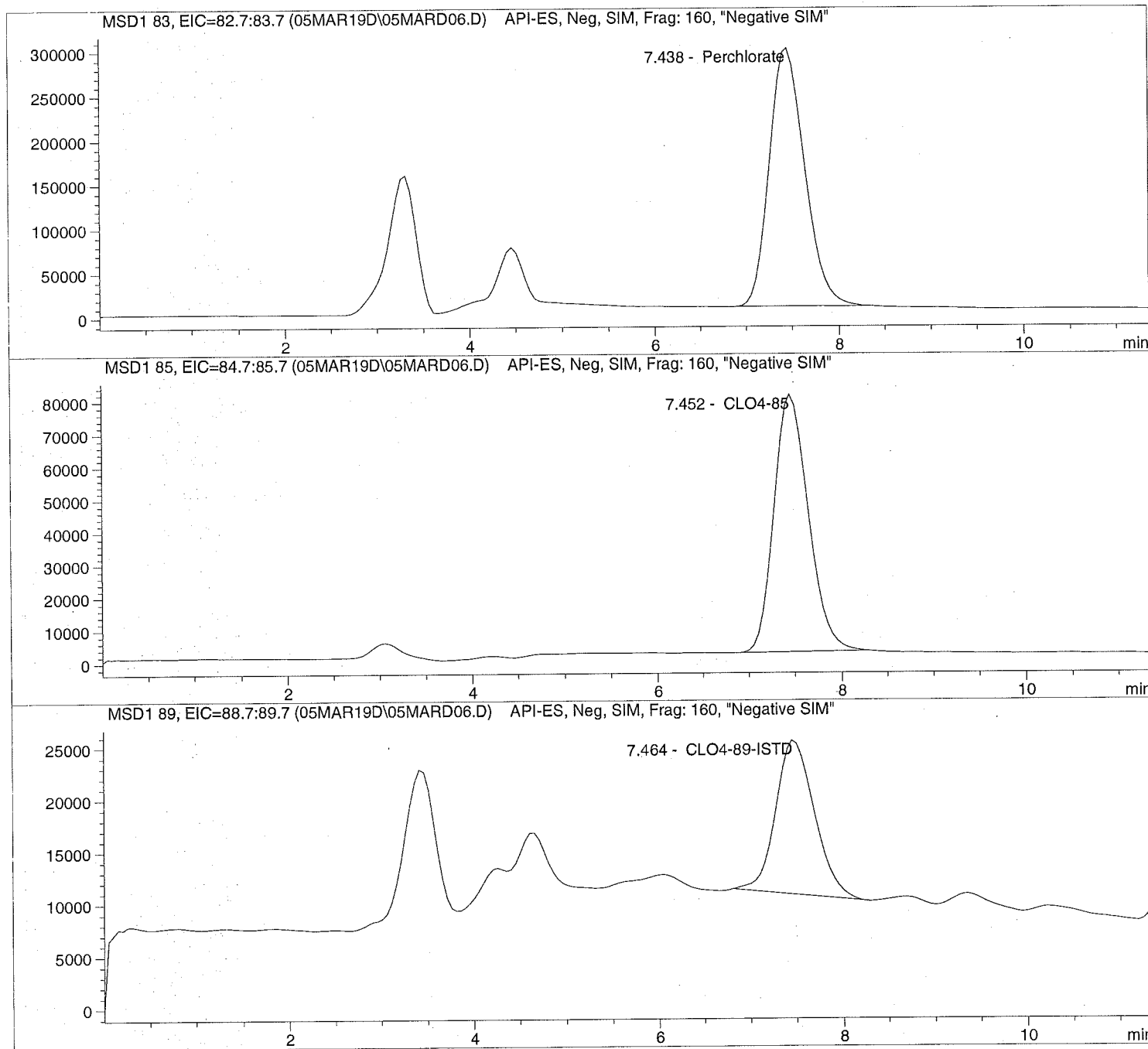
=====
*** End of Report ***

Injection Date: 3/05/2019 09:54:04
Sample Name: 1906112001
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 09:54:04      Seg Line: 6
Sample Name: 1906112001                 Location: Vial 76
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.438	PBA	7711270.5	51.2515	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.452	PBA	2027855.1	51.1033	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PB	426473.5	5.0000	CLO4-89-ISTD

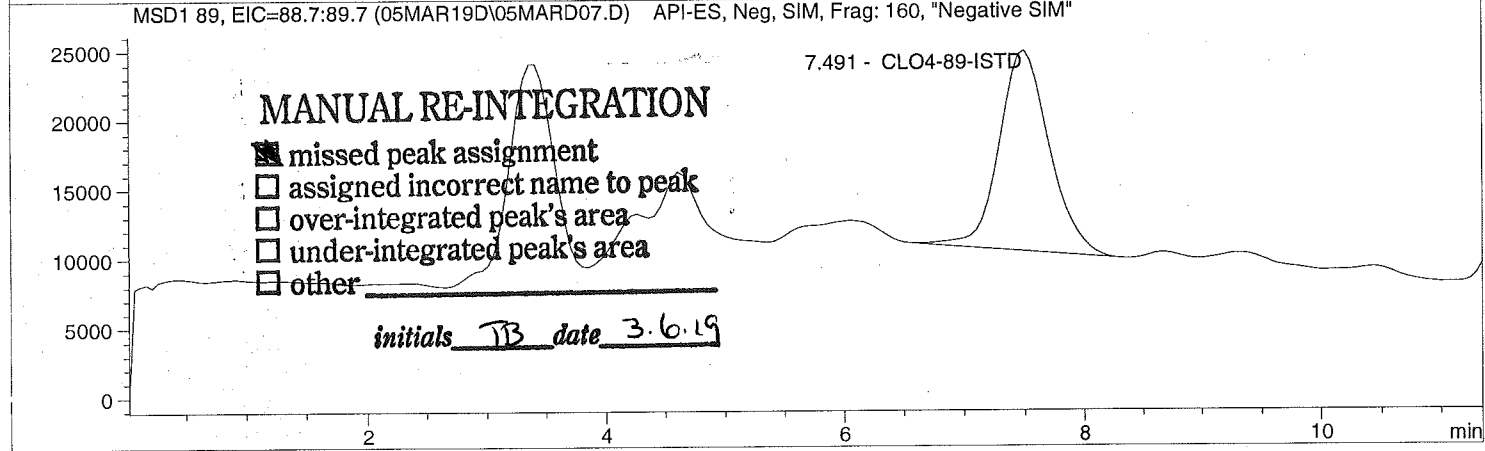
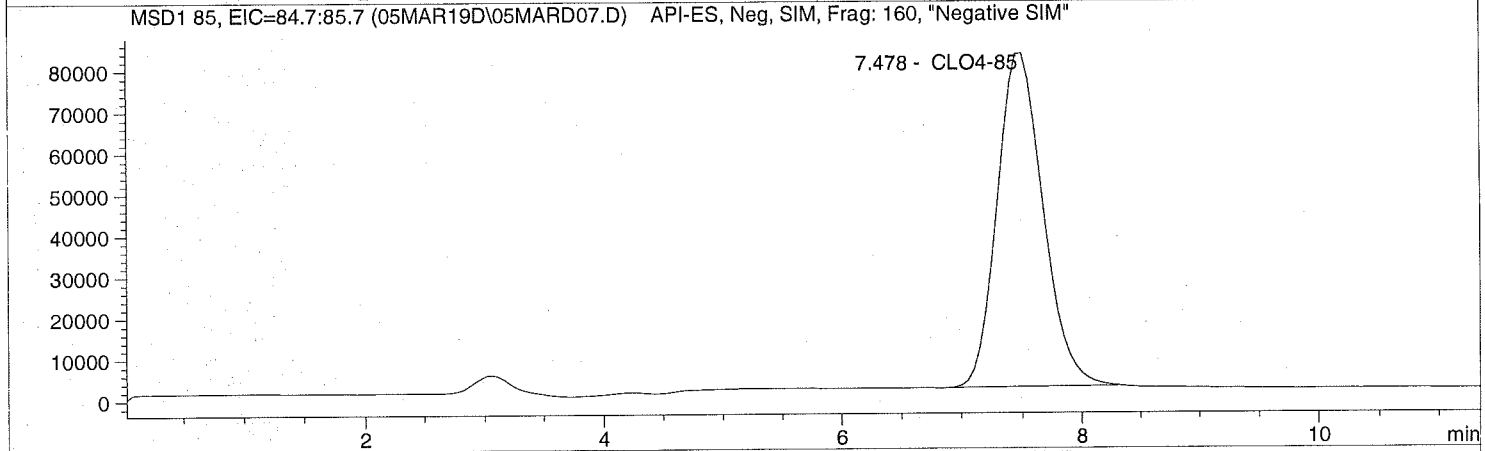
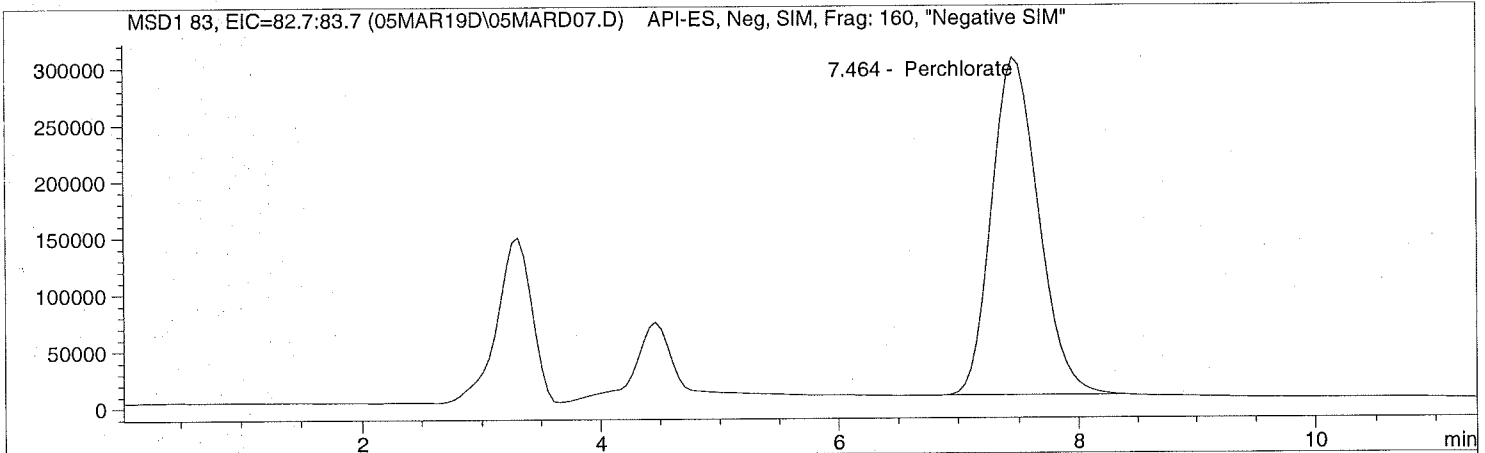
=====
*** End of Report ***

Injection Date: 3/05/2019 10:07:11
Sample Name: 1906112002 MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:07:11 Seq Line: 7
Sample Name: 1906112002 MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	53.8797	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	54.4680	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.491	MM	419549.0	5.0000	CLO4-89-ISTD

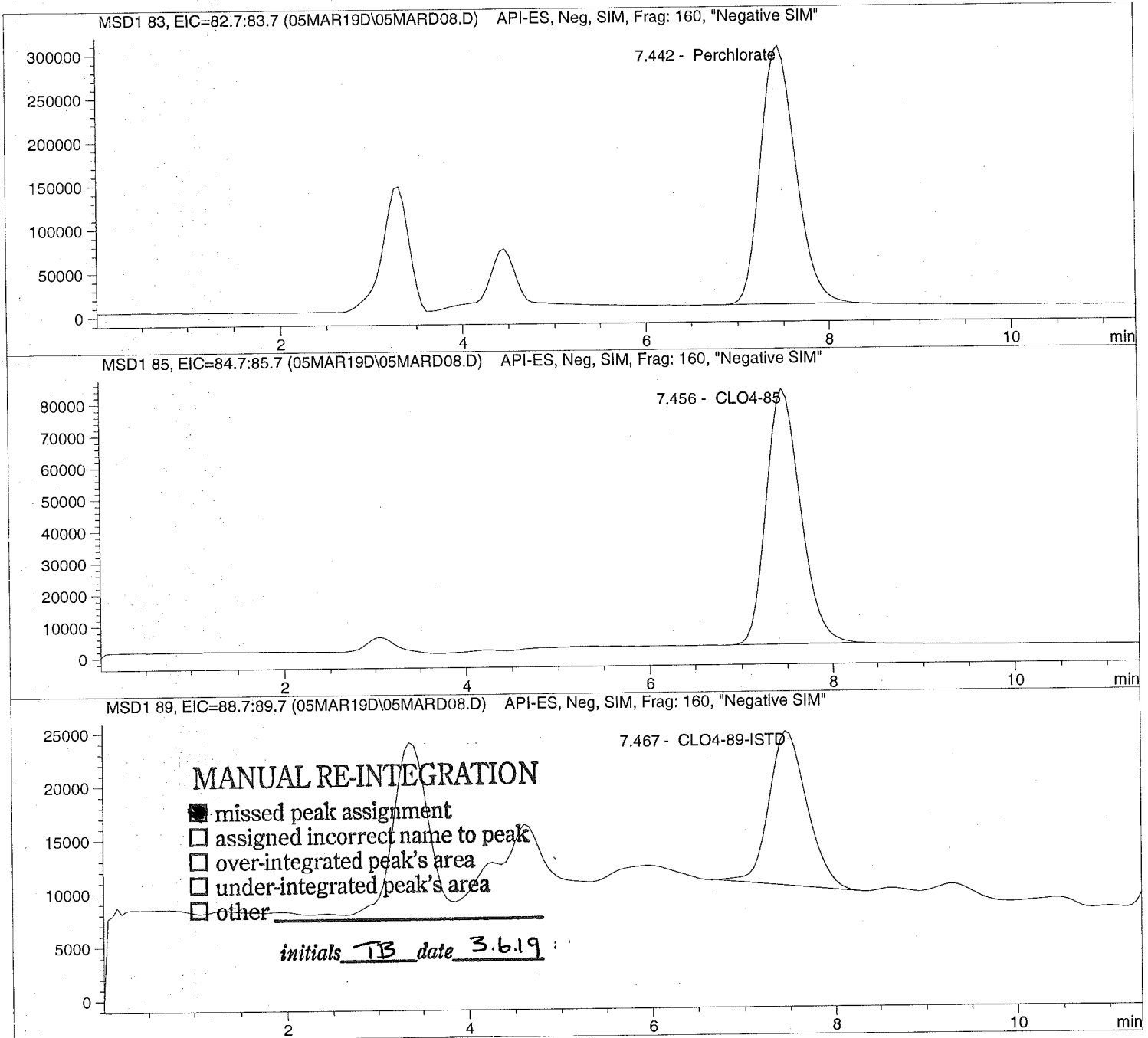
*** End of Report ***

Injection Date: 3/05/2019 10:20:17
Sample Name: 1906112003 MSD
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	53.6994	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	53.9810	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.467	MM	416709.1	5.0000	CLO4-89-ISTD

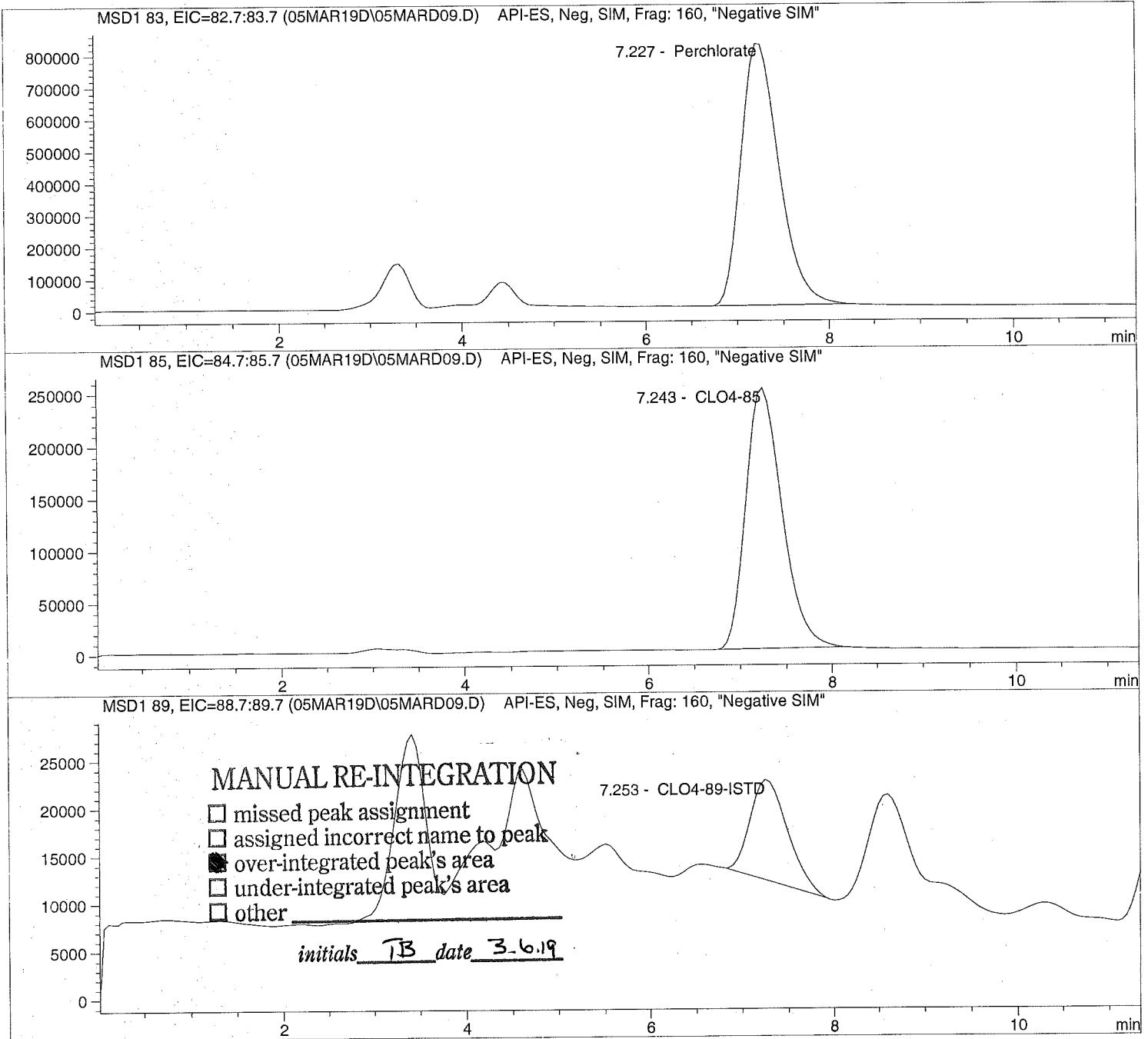
*** End of Report ***

Injection Date: 3/05/2019 10:33:21
Sample Name: 1906112004
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 10:33:21      Seq Line:          9  
Sample Name:    1906112004              Location:         Vial 79  
Acq Operator:   TNB                     Inj. No.:        1  
                                           Inj. Vol.:       20 µl
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	172.3652	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	180.4596	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.253	MM	295705.2	5.0000	CLO4-89-ISTD

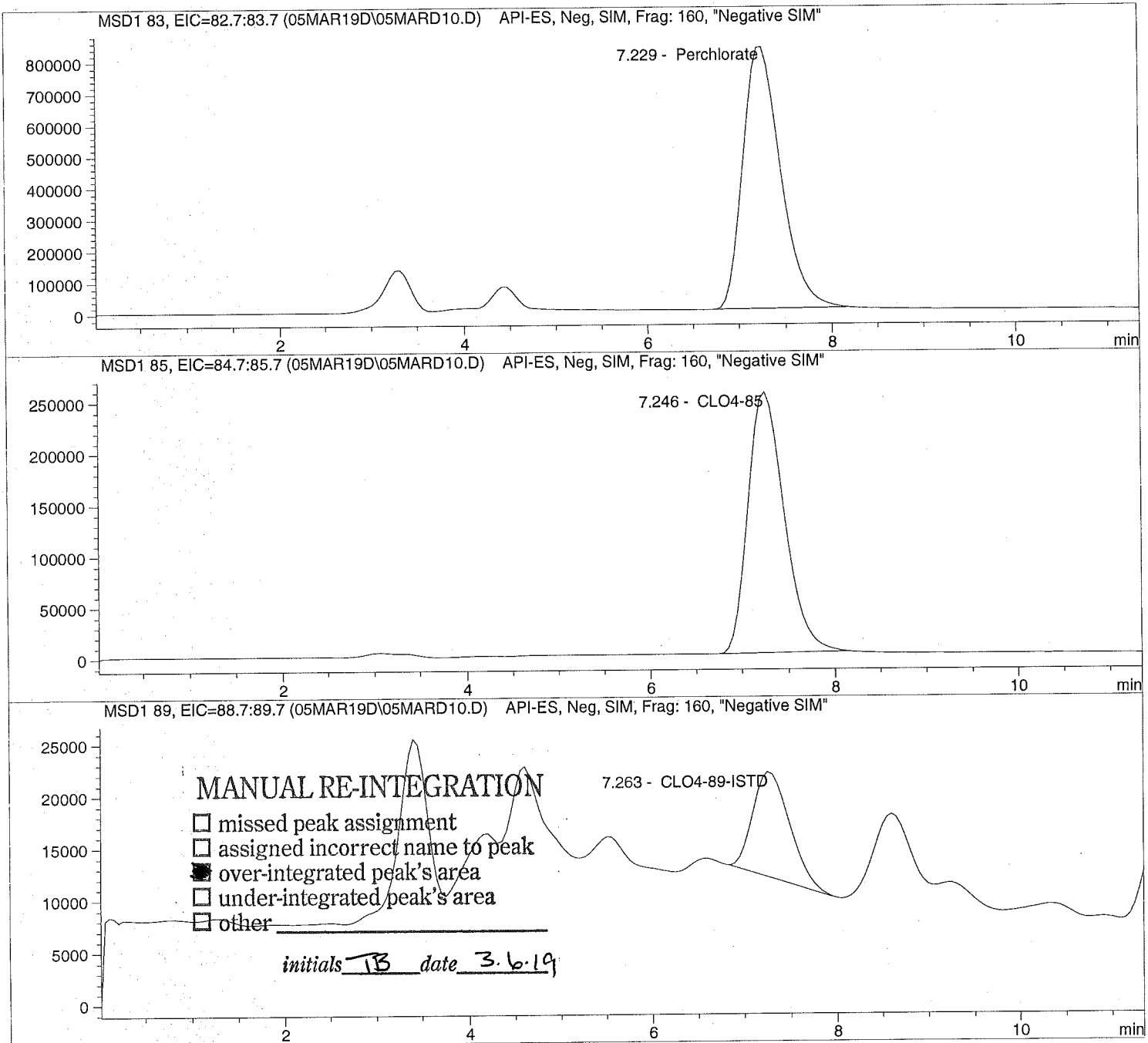
=====
*** End of Report ***

Injection Date: 3/05/2019 10:46:26
Sample Name: 1906112005
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

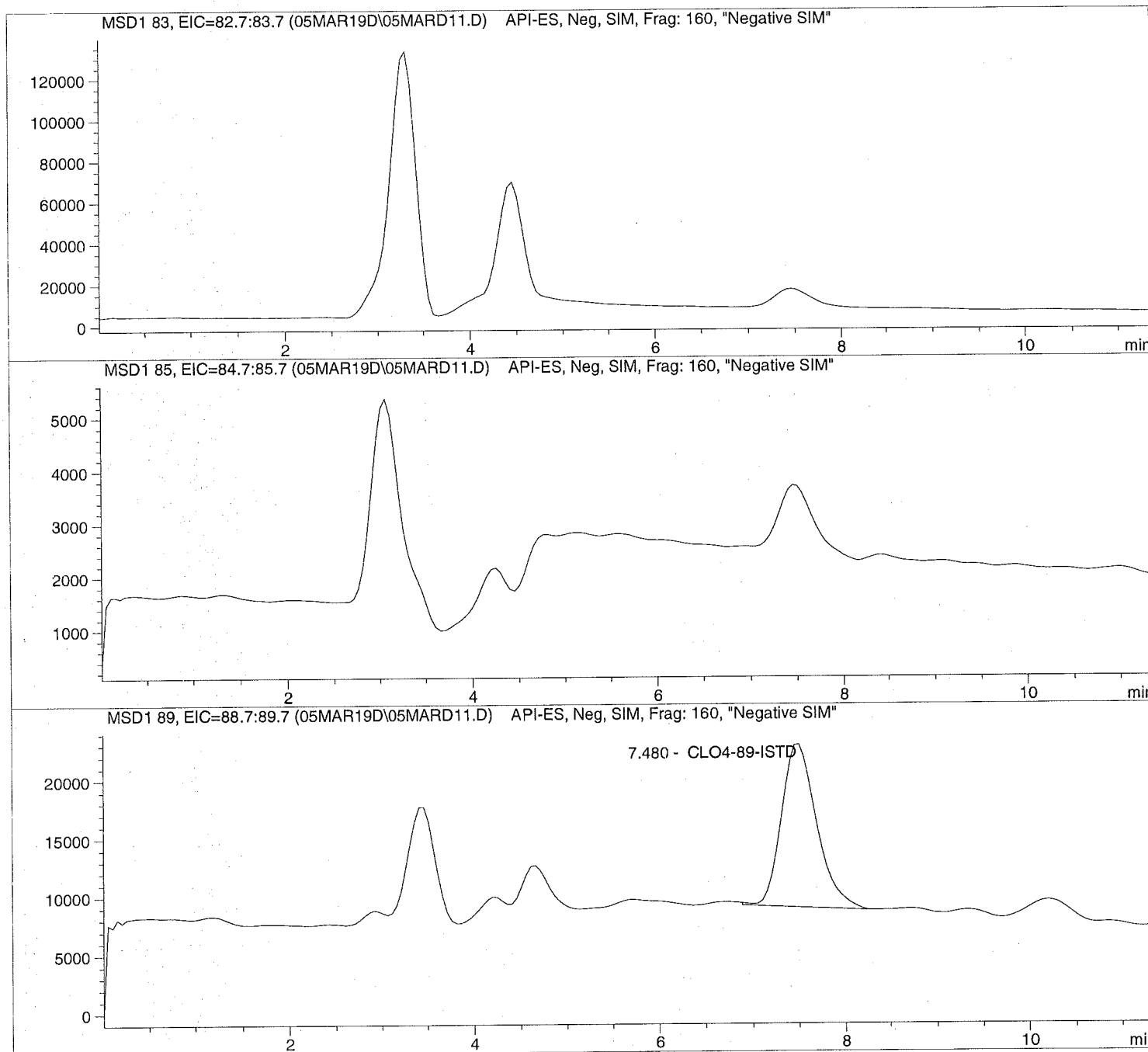


Injection Date: 3/05/2019 10:59:36
Sample Name: 1906112006
Acq Operator: TNB

Seq Line: 11
Location: Vial 81
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:59:36 Seq Line: 11
Sample Name: 1906112006 Location: Vial 81
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.480	BBA	373575.4	5.0000	CLO4-89-ISTD

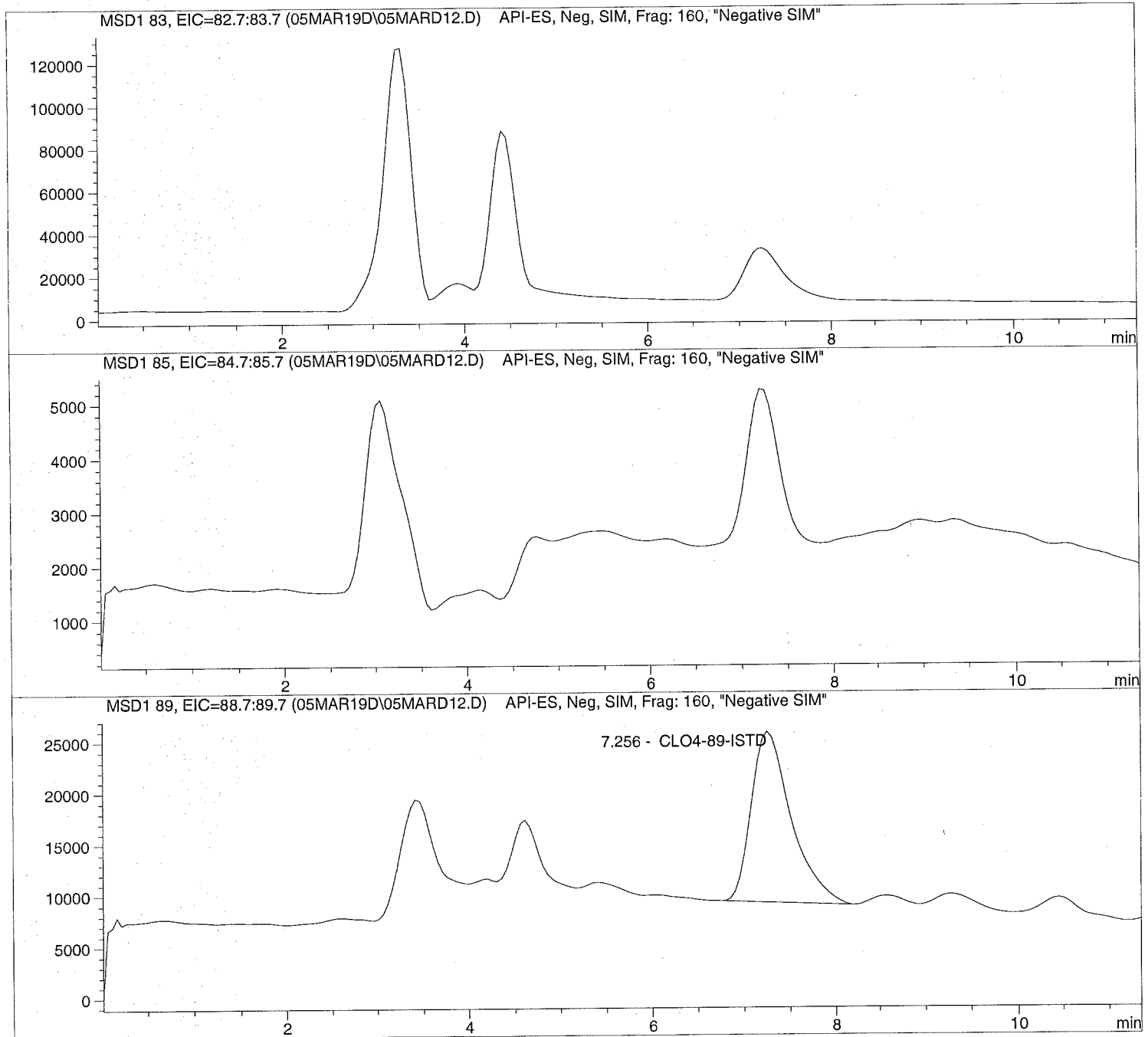
*** End of Report ***

Injection Date: 3/05/2019 11:12:28
Sample Name: 1906112007
Acq Operator: TNB

Seq Line: 12
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 11:12:28      Seq Line: 12  
Sample Name: 1906112007                Location: Vial 82  
Acq Operator: TNB                       Inj. No.: 1  
                                           Inj. Vol.: 20 µl
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.256	PB	500532.9	5.0000	CLO4-89-ISTD

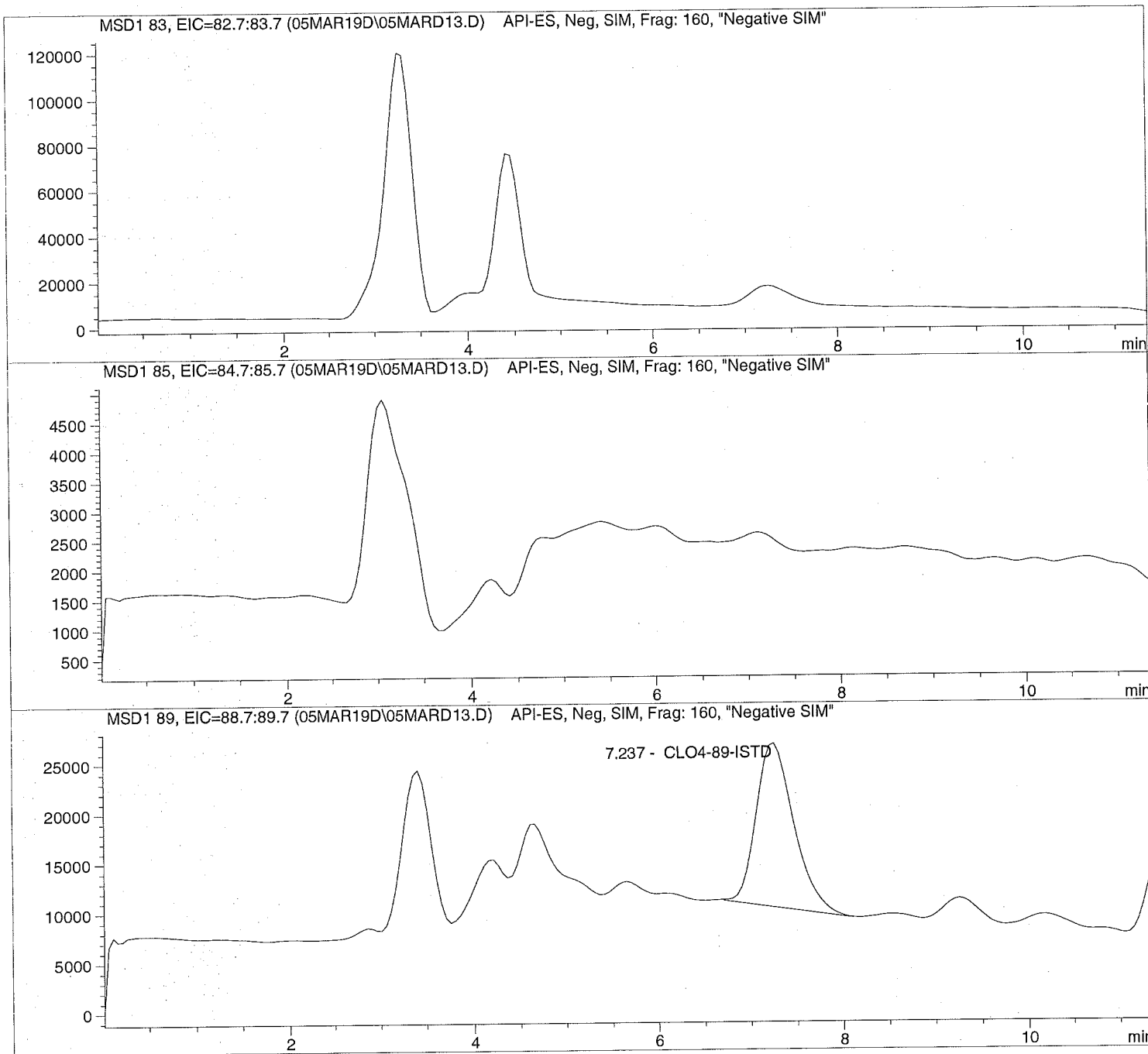
=====
*** End of Report ***

Injection Date: 3/05/2019 11:26:23
Sample Name: 1906112008
Acq Operator: TNB

Seq Line: 13
Location: Vial 83
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



=====
Injection Date: 3/05/2019 11:26:23 Seq Line: 13
Sample Name: 1906112008 Location: Vial 83
Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 20 µl
=====

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

=====
Sample Information
=====

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.237	BBA	465121.2	5.0000	CLO4-89-ISTD

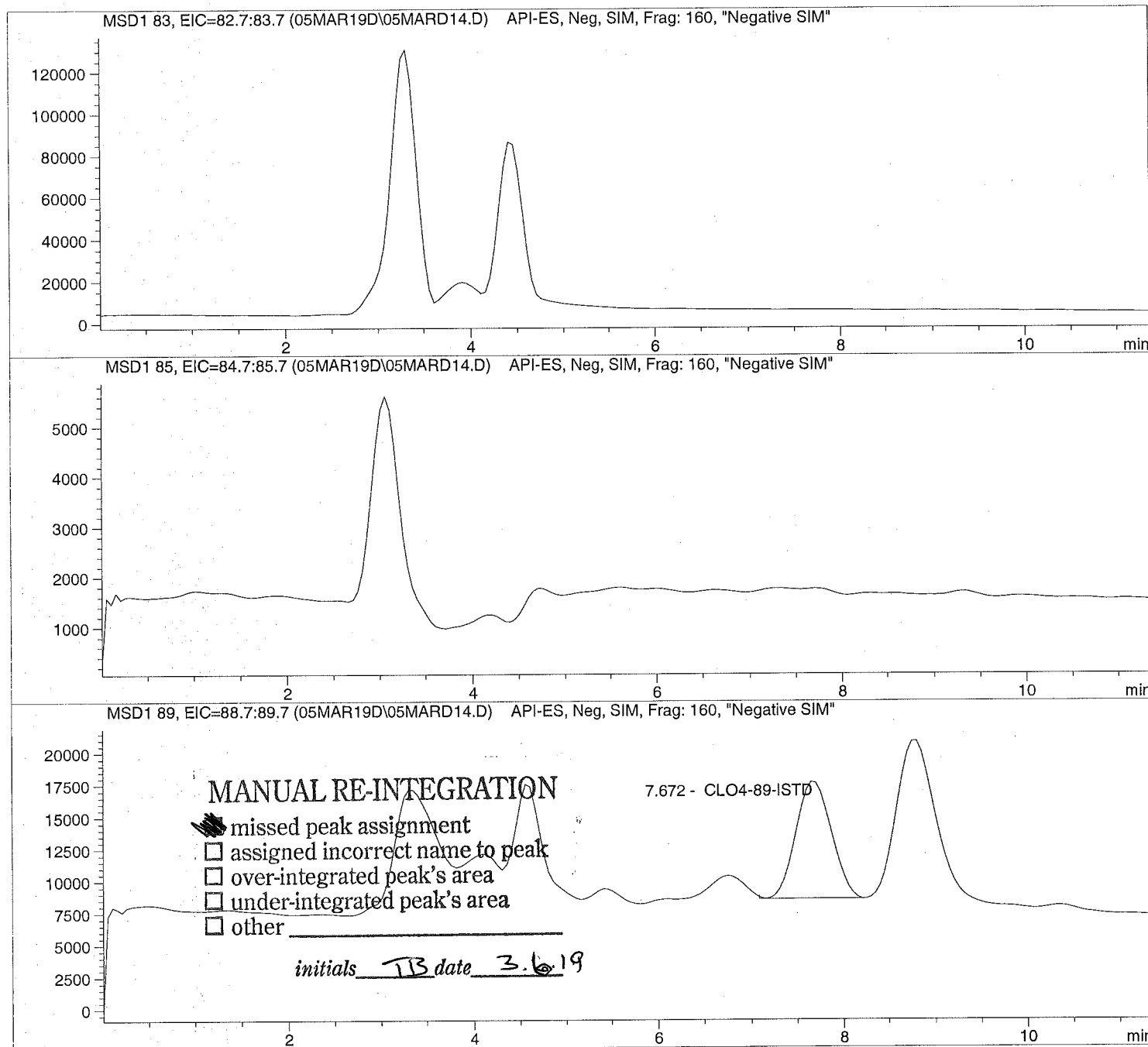
=====
*** End of Report ***
=====

Injection Date: 3/05/2019 11:39:24
Sample Name: 1906112009
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 11:39:24 Seq Line: 14
Sample Name: 1906112009 Location: Vial 84
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	5.0000	CLO4-89-ISTD

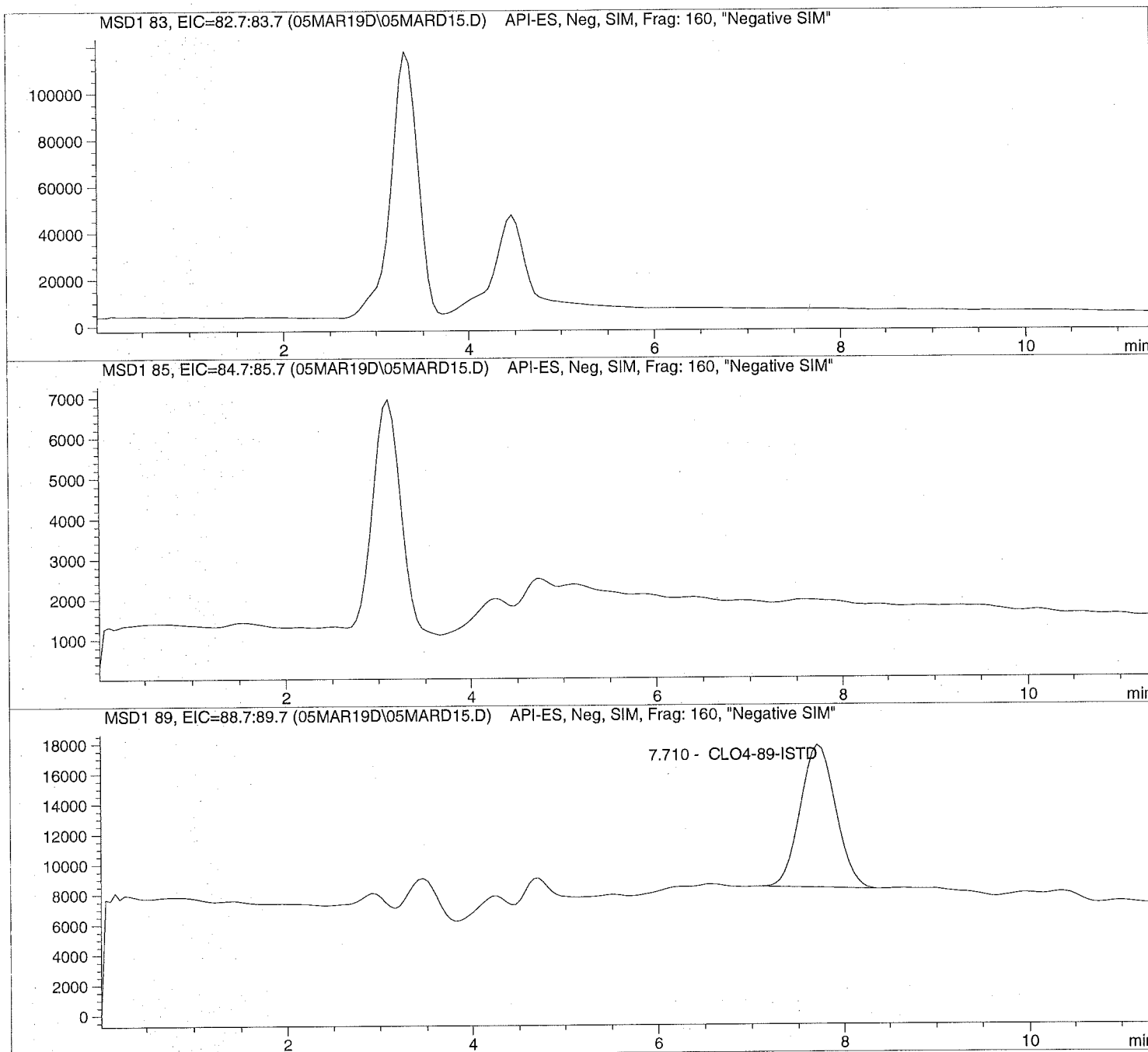
*** End of Report ***

Injection Date: 3/05/2019 11:52:27
Sample Name: 1906112010
Acq Operator: TNB

Seq Line: 15
Location: Vial 85
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date:  3/05/2019  11:52:27      Seq Line:      15
Sample Name:    1906112010                Location:      Vial 85
Acq Operator:  TNB                       Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019  12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.710	BBA	251865.4	5.0000	CLO4-89-ISTD

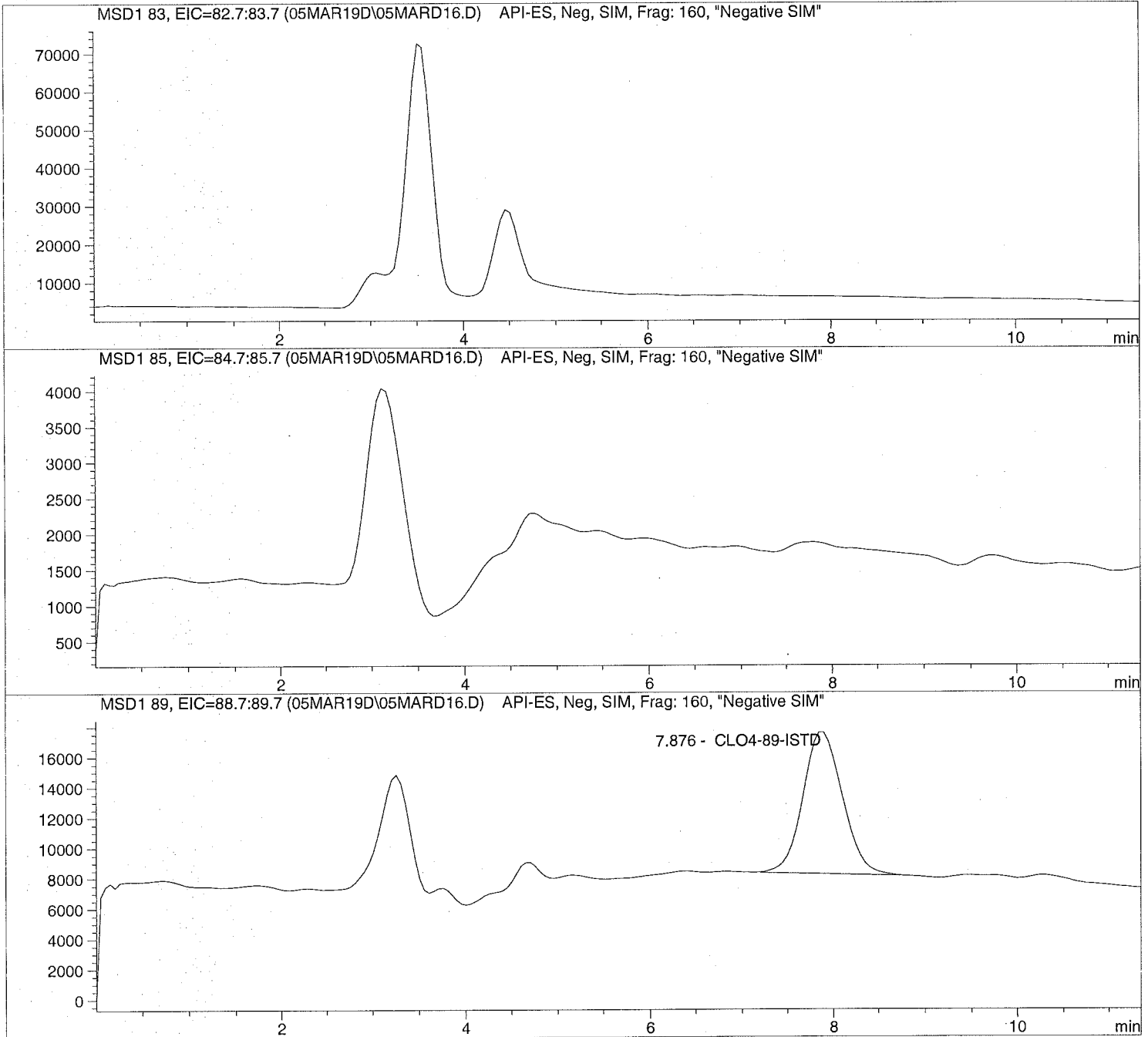
=====
*** End of Report ***

Injection Date: 3/05/2019 12:05:39
Sample Name: 1906112011
Acq Operator: TNB

Seq Line: 16
Location: Vial 86
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 12:05:39 Seq Line: 16
Sample Name: 1906112011 Location: Vial 86
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

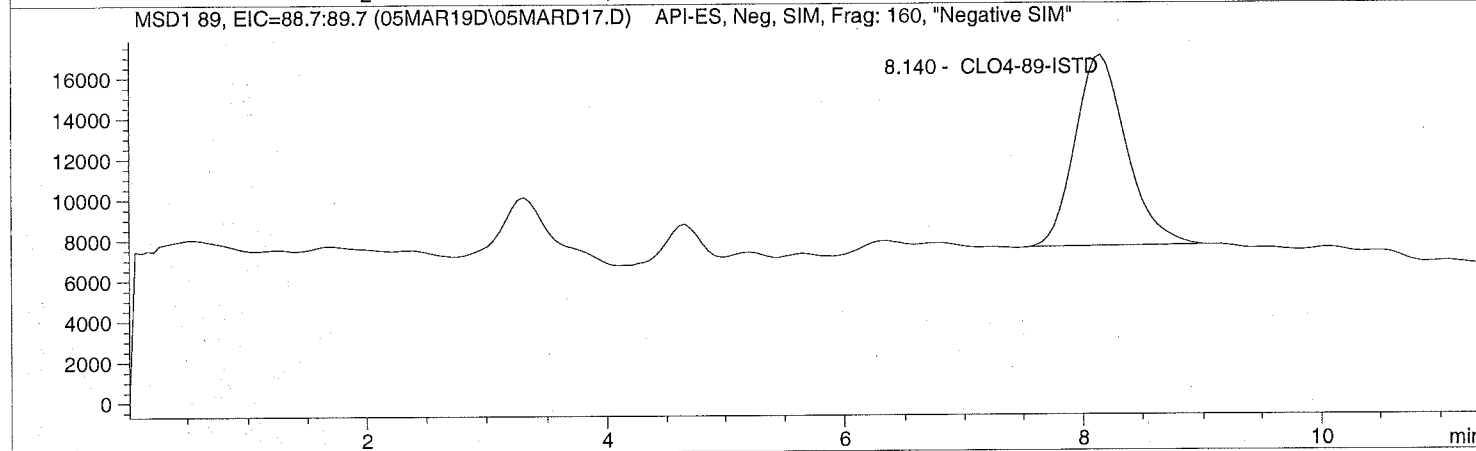
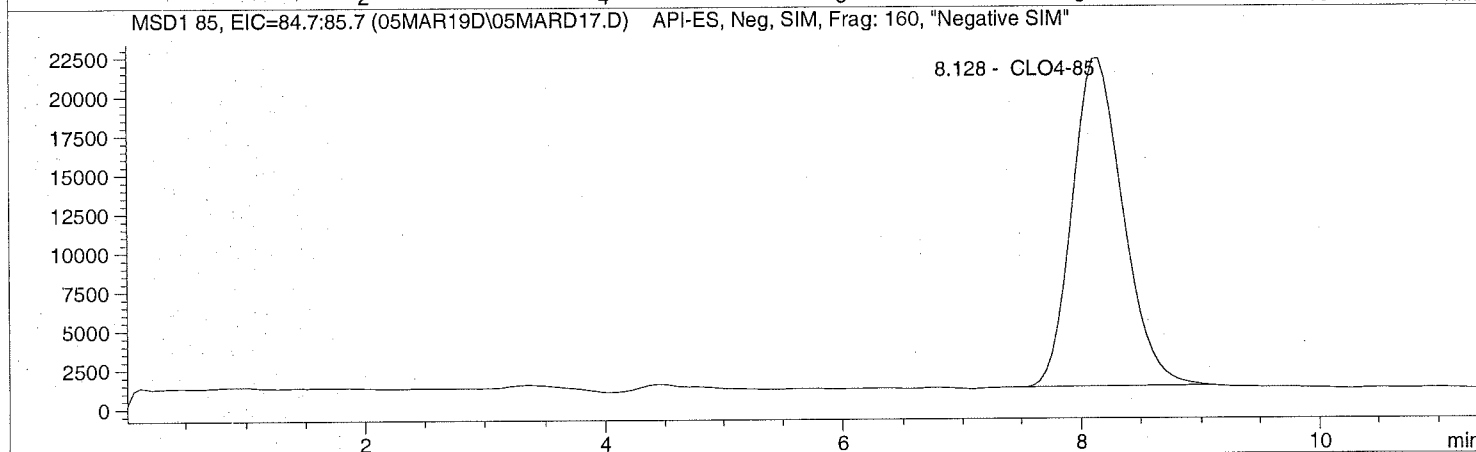
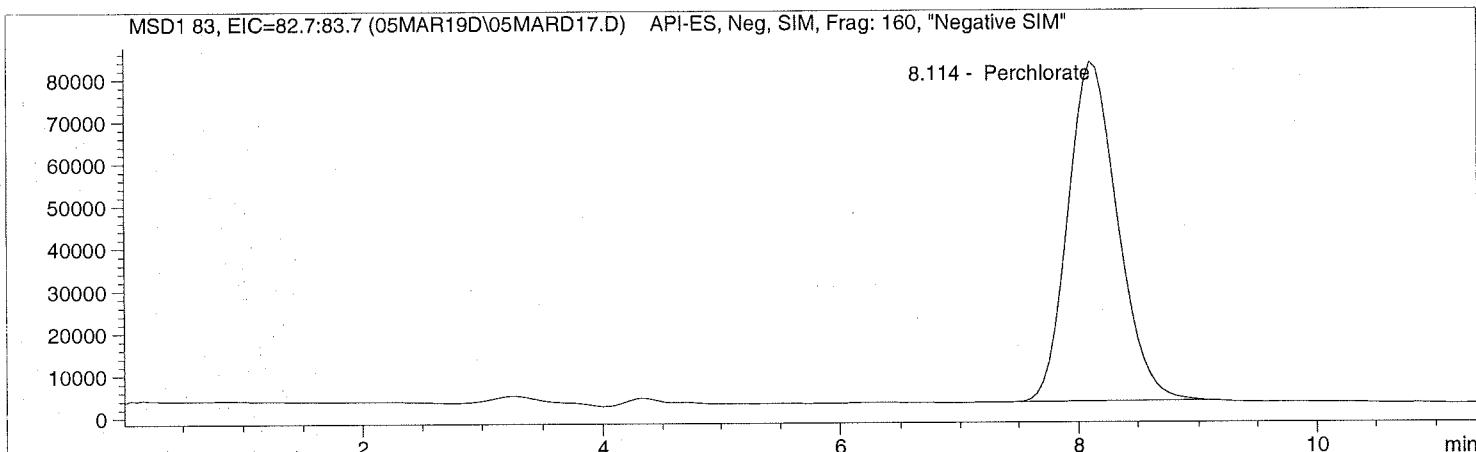
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.876	BBA	280792.2	5.0000	CLO4-89-ISTD

*** End of Report ***

Injection Date: 3/05/2019 12:18:41 Seq Line: 17
Sample Name: 642101 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 12:18:41 Seq Line: 17
Sample Name: 642101 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	2336112.7	25.0199	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.128	PBA	628961.2	25.6341	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.140	PBA	283460.4	5.0000	CLO4-89-ISTD

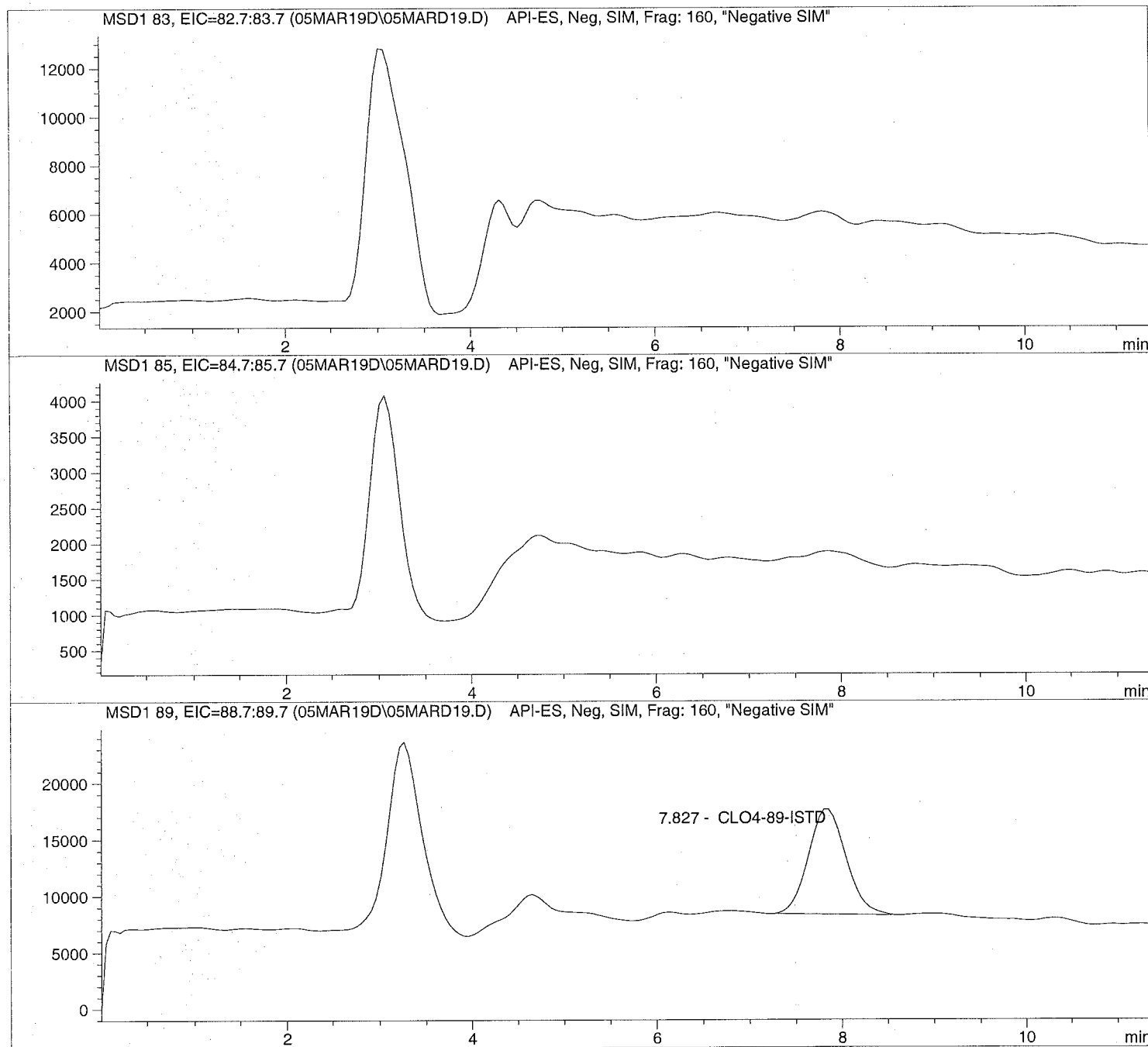
*** End of Report ***

Injection Date: 3/05/2019 12:49:07
Sample Name: 1906332001
Acq Operator: TNB

Seq Line: 19
Location: Vial 88
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 12:49:07      Seq Line: 19
Sample Name: 1906332001                 Location: Vial 88
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.827	PBA	264674.3	5.0000	CLO4-89-ISTD

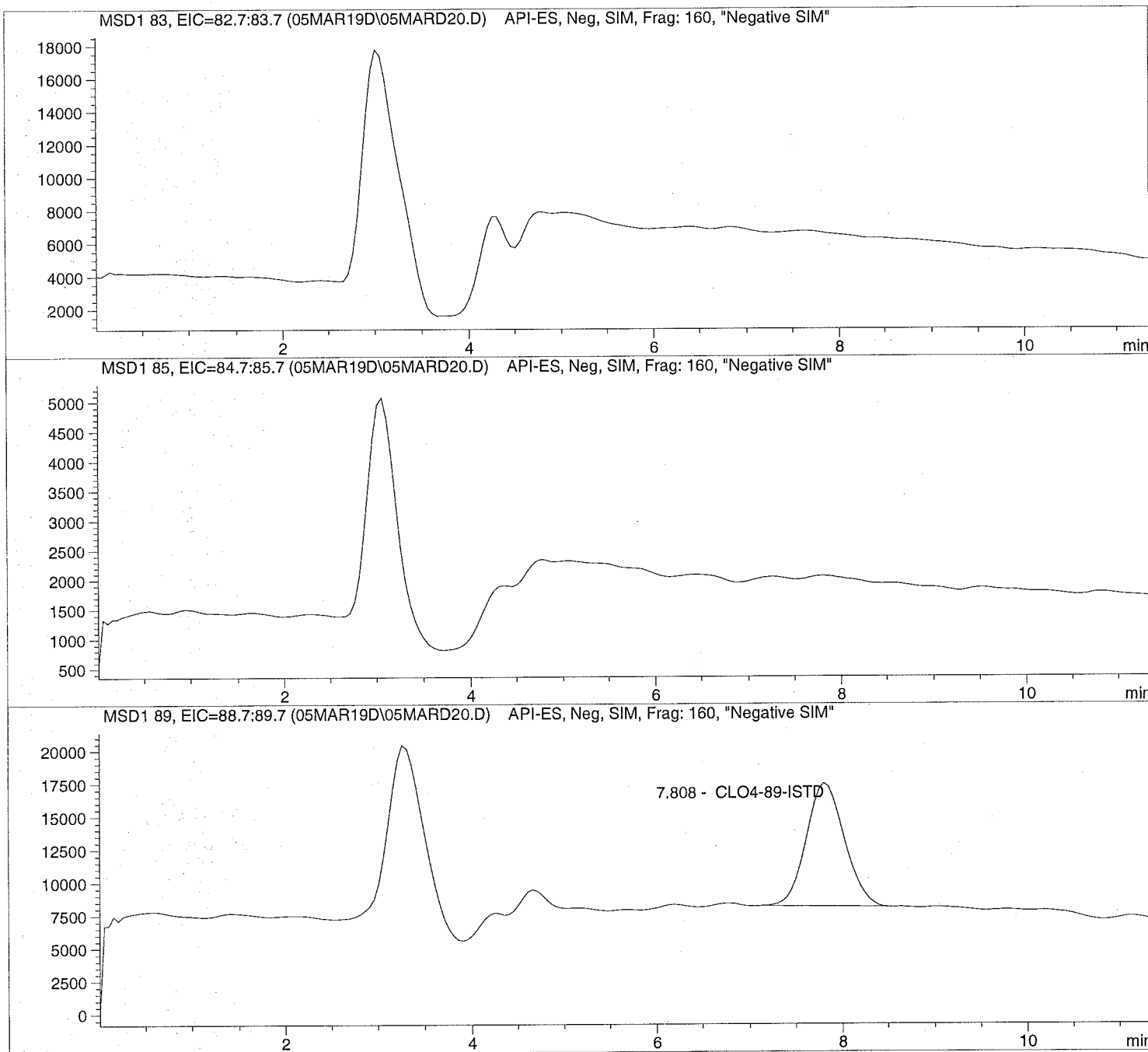
=====
*** End of Report ***

Injection Date: 3/05/2019 13:02:15
Sample Name: 1906334001
Acq Operator: TNB

Seq Line: 20
Location: Vial 89
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 13:02:15      Seq Line: 20
Sample Name: 1906334001                 Location: Vial 89
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

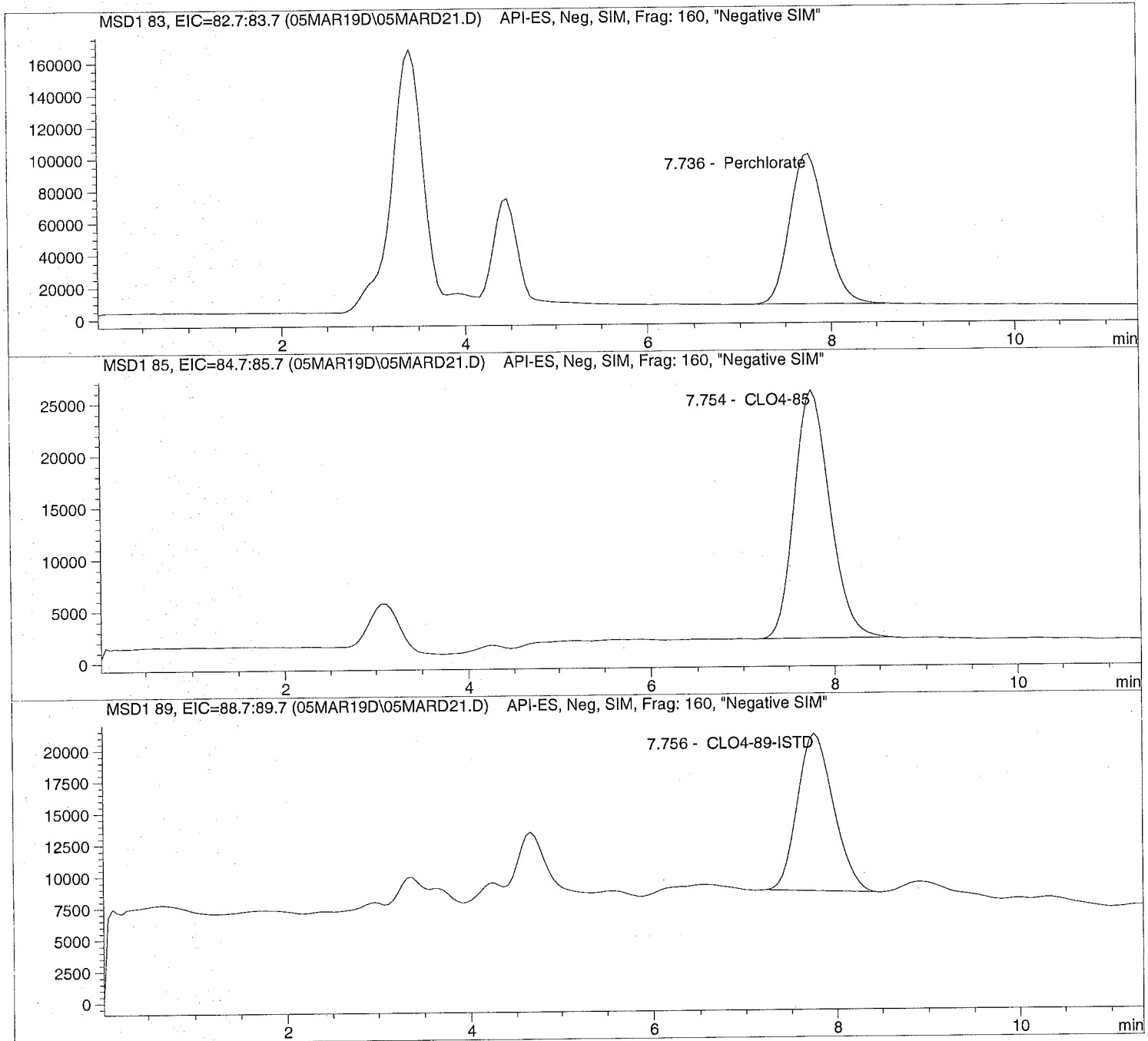
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.808	PBA	265662.0	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

Injection Date: 3/05/2019 13:15:18 Seq Line: 21
Sample Name: 1906112004 10X Location: Vial 90
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 13:15:18      Seq Line: 21  
Sample Name: 1906112004 10X            Location: Vial 90  
Acq Operator: TNB                      Inj. No.: 1  
                                         Inj. Vol.: 20 µl
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier: 1.000000  
Dilution: 10.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.736	PBA	2531259.3	226.4815	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.754	PBA	646870.4	220.9191	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	PB	341607.9	50.0000	CLO4-89-ISTD

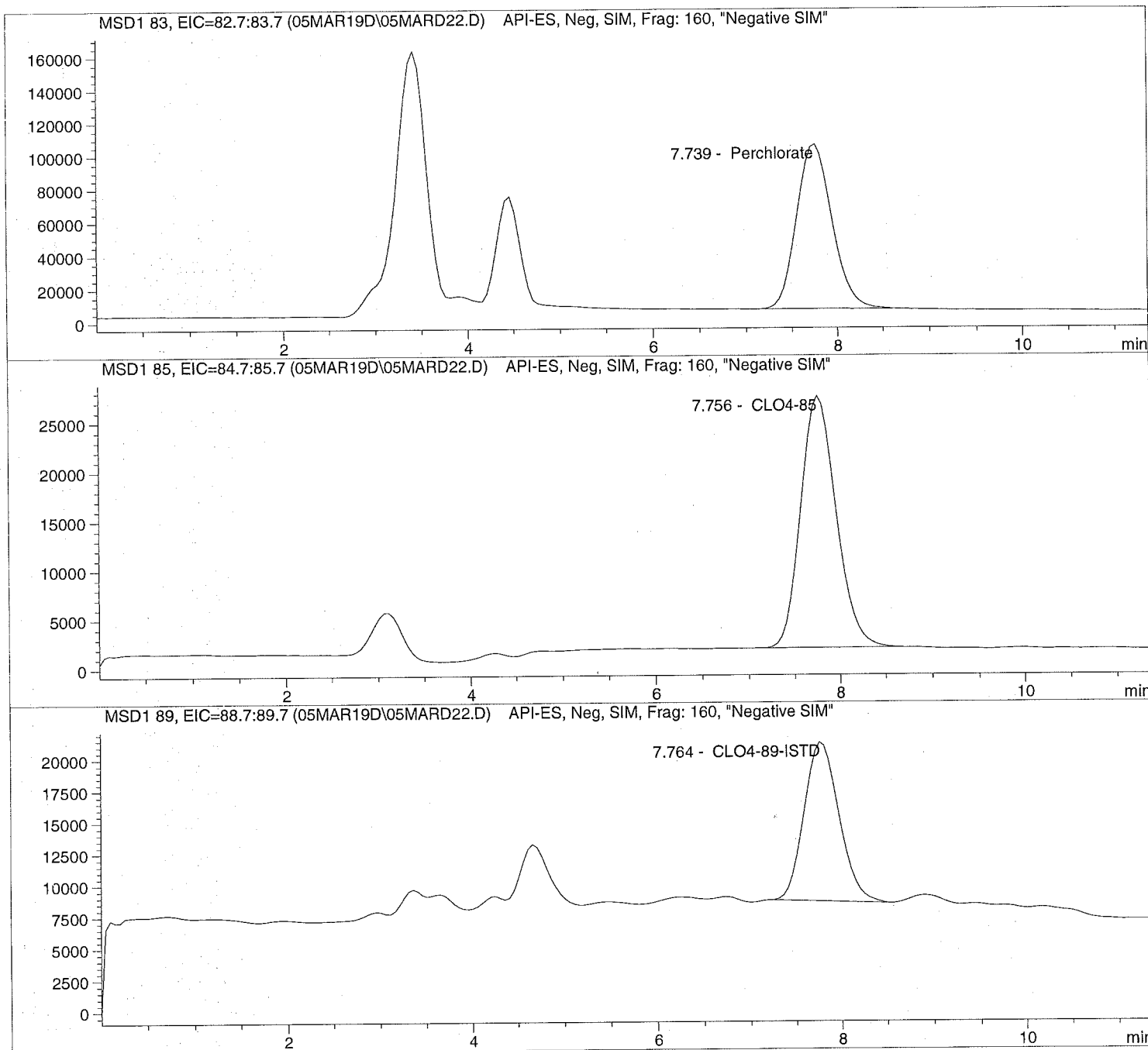
=====
*** End of Report ***
=====

Injection Date: 3/05/2019 13:28:32
Sample Name: 1906112005 10X
Acq Operator: TNB

Seq Line: 22
Location: Vial 91
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 13:28:32      Seq Line:      22
Sample Name:    1906112005 10X          Location:      Vial 91
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       10.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.739	PBA	2682370.5	241.0371	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	BBA	695643.1	238.4467	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.764	PB	338724.1	50.0000	CLO4-89-ISTD

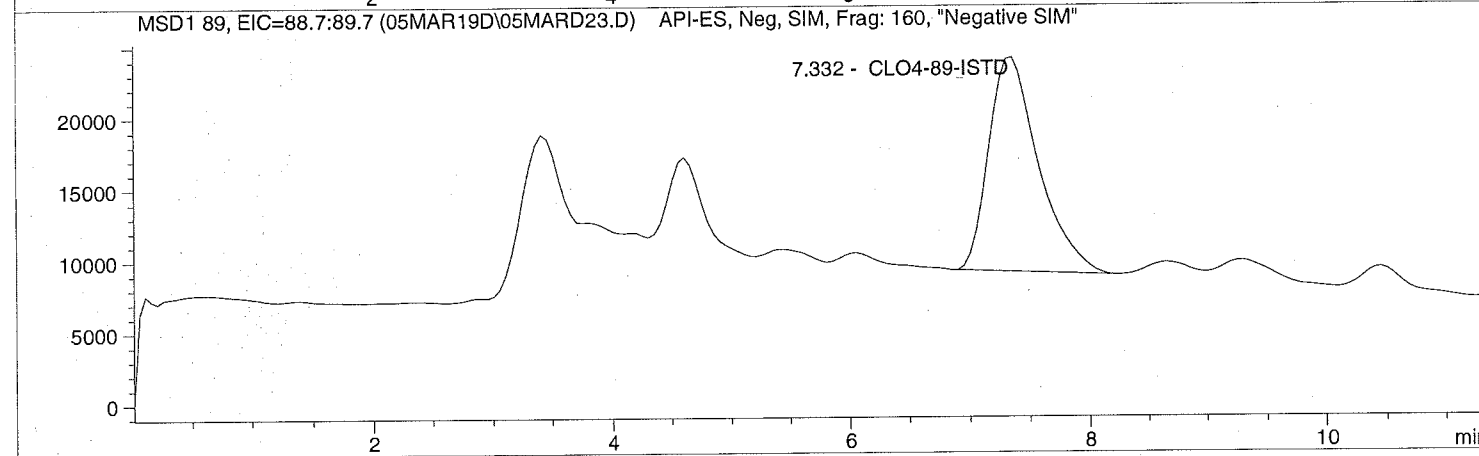
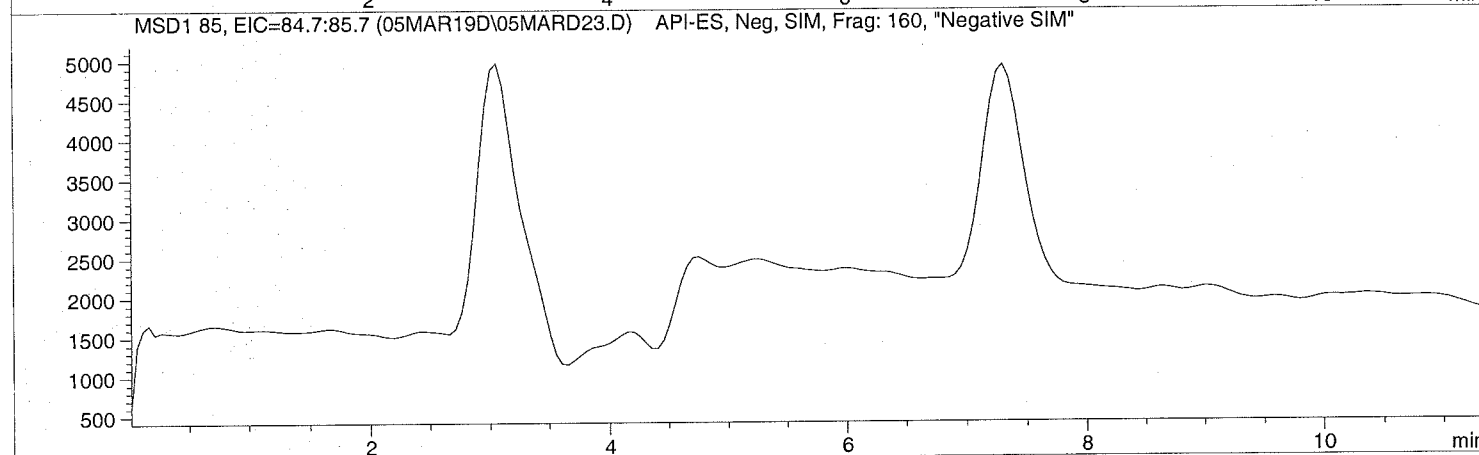
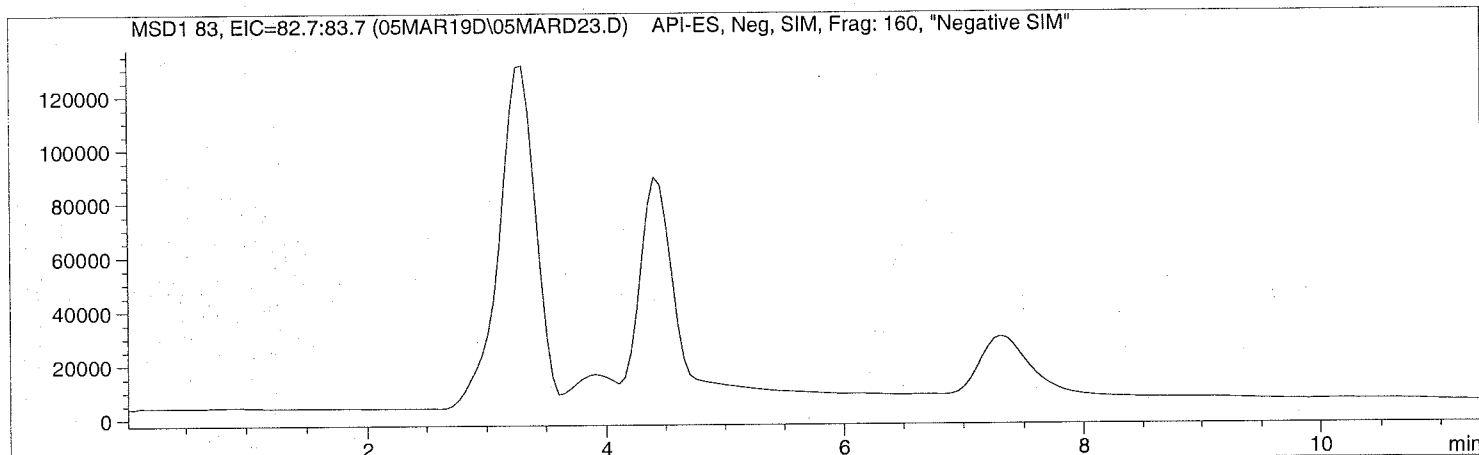
=====
*** End of Report ***

Injection Date: 3/05/2019 13:41:34
Sample Name: 1906112007 RE
Acq Operator: TNB

Seq Line: 23
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 13:41:34 Seq Line: 23
Sample Name: 1906112007 RE Location: Vial 82
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.332	PB	431134.6	5.0000	CLO4-89-ISTD

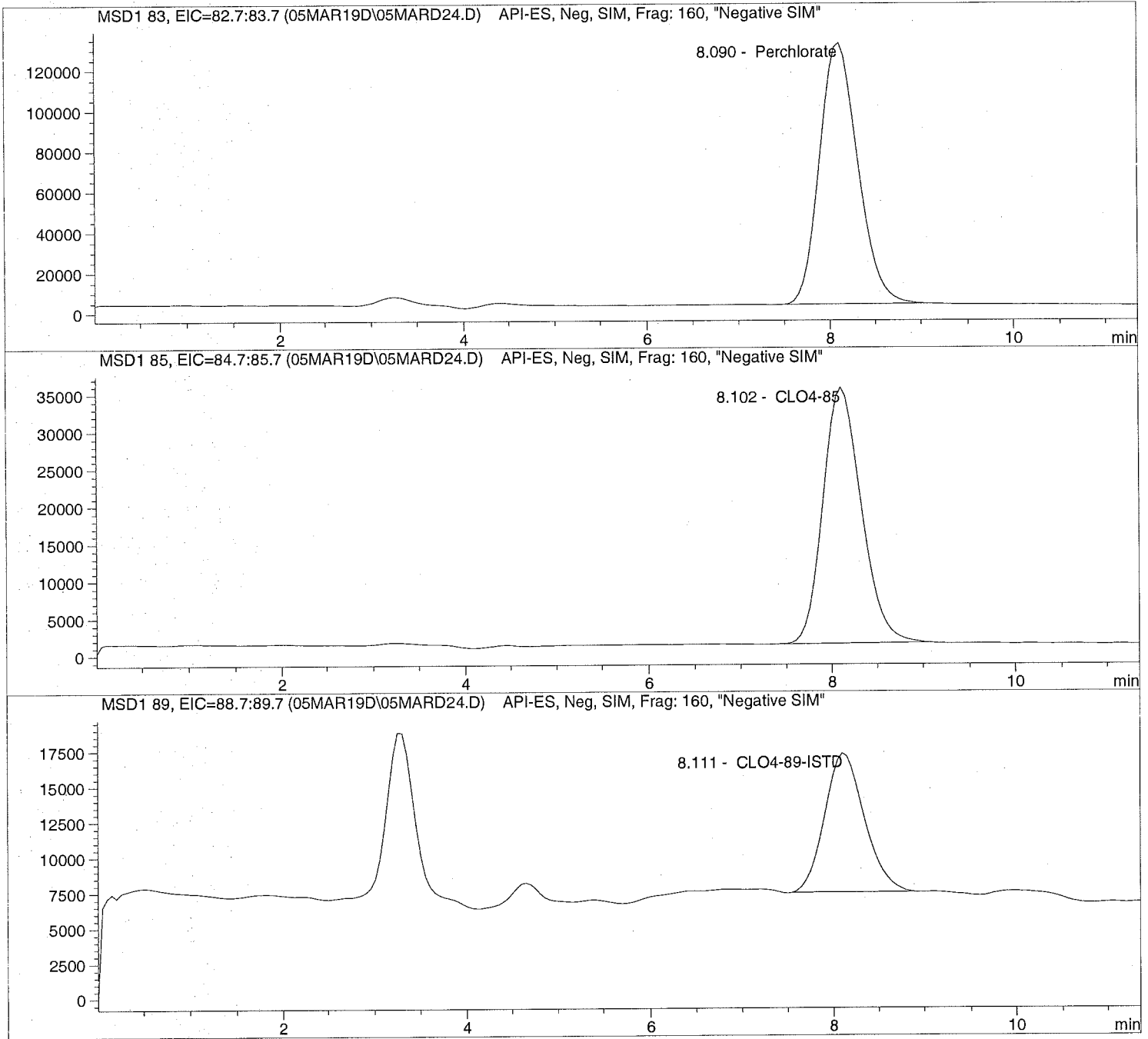
*** End of Report ***

Injection Date: 3/05/2019 13:54:34
Sample Name: 1906330001 100
Acq Operator: TNB

Seq Line: 24
Location: Vial 92
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 13:54:34      Seq Line:      24
Sample Name:    1906330001 100          Location:      Vial 92
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       100.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.090	PBA	3773628.7	3708.6567	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	PBA	1010205.2	3771.6748	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.111	PBA	298984.6	500.0000	CLO4-89-ISTD

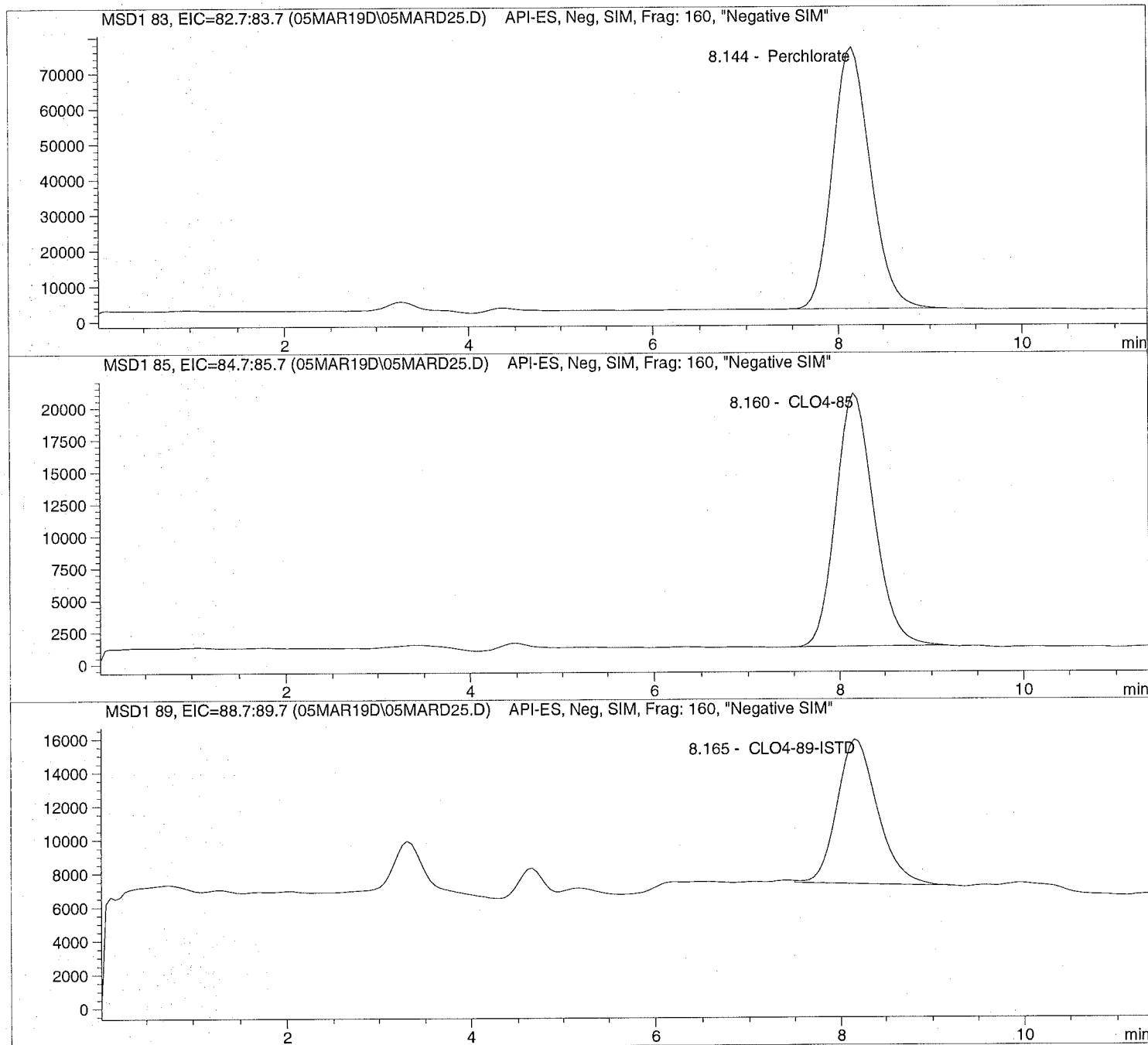
=====
*** End of Report ***

Injection Date: 3/05/2019 14:07:36
Sample Name: 642102 CCV@25
Acq Operator: TNB

Seq Line: 25
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 14:07:36      Seq Line:          25  
Sample Name:    642102  CCV@25          Location:          Vial 71  
Acq Operator:  TNB                      Inj. No.:         1  
                                           Inj. Vol.:        20 µl
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 25.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.144	PBA	2157871.5	24.4553	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.160	PBA	583711.5	25.1665	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.165	BBA	268305.4	5.0000	CLO4-89-ISTD

=====
*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

**Initial
Calibration**

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	8.94006e4	7.889	9.89924e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.97443e5	8.114	2.26028
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	4.79370e5	7.828	4.65688
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	9.30136e5	7.904	9.14998
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.81067e6	7.793	25.52636
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	5.66830e6	7.976	51.07439
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	8.69624e6	7.886	74.30603
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	1.01141e6	7.988	9.46019

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.26121e4	7.914	9.98836e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	5.53134e4	8.127	2.11360
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.39247e5	7.842	4.91261
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.54396e5	7.923	9.39034
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	7.35969e5	7.811	25.48268
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.47152e6	7.993	50.35774
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.32809e6	7.900	74.72233
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.81230e5	8.007	9.87858

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.41443e5	7.900	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.99651e5	8.132	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.38646e5	7.853	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	3.25154e5	7.925	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	3.33799e5	7.819	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	3.14712e5	7.999	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	3.13909e5	7.908	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	3.41503e5	8.005	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

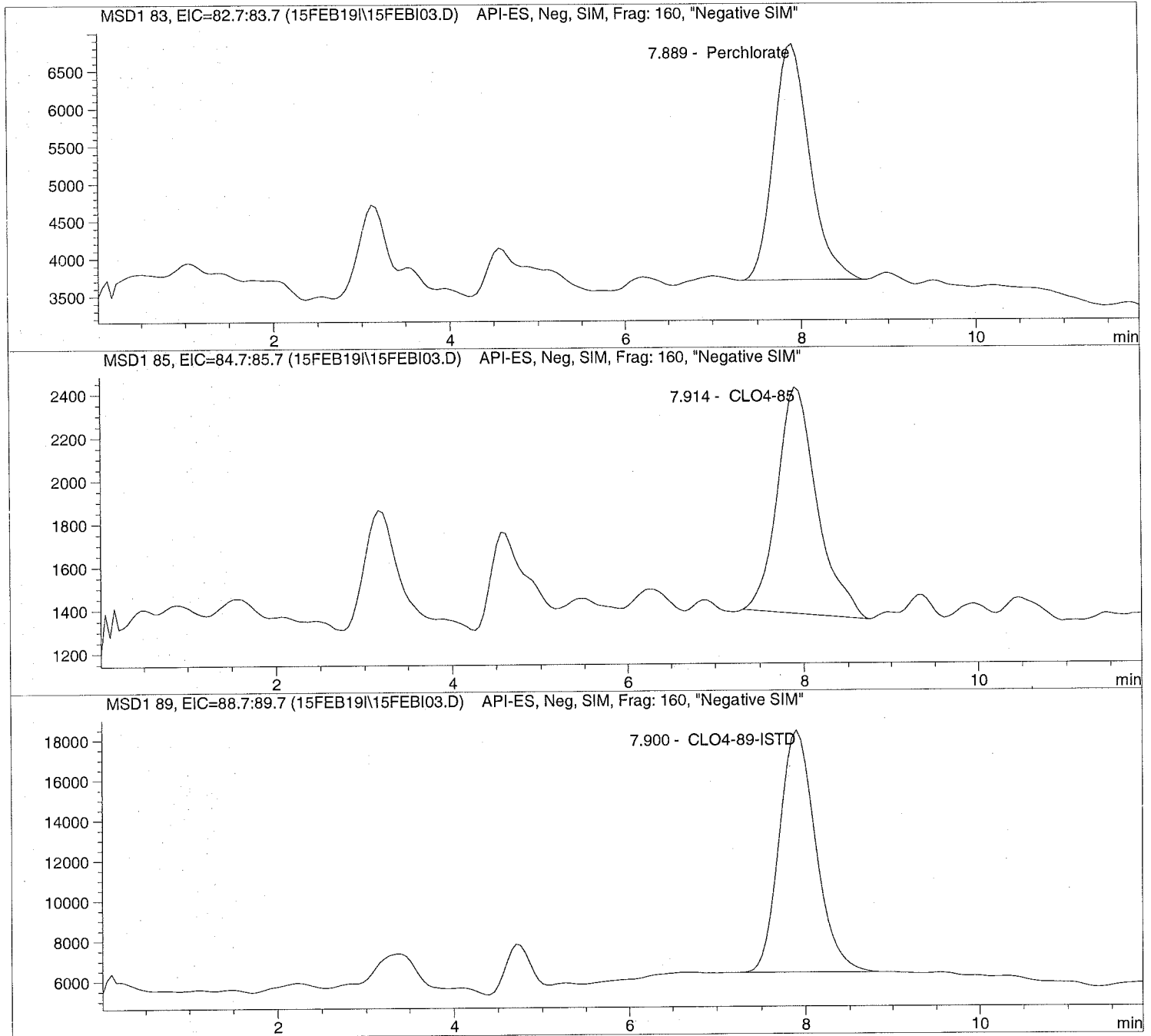
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ .20ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Injection Date: 2/15/2019 09:51:42
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====
Injection Date: 2/15/2019 09:51:42      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L          Location:  Vial 73
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.889	PBA	89400.6	0.9899	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.914	BBA	32612.1	0.9988	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	BBA	341443.2	5.0000	CLO4-89-ISTD

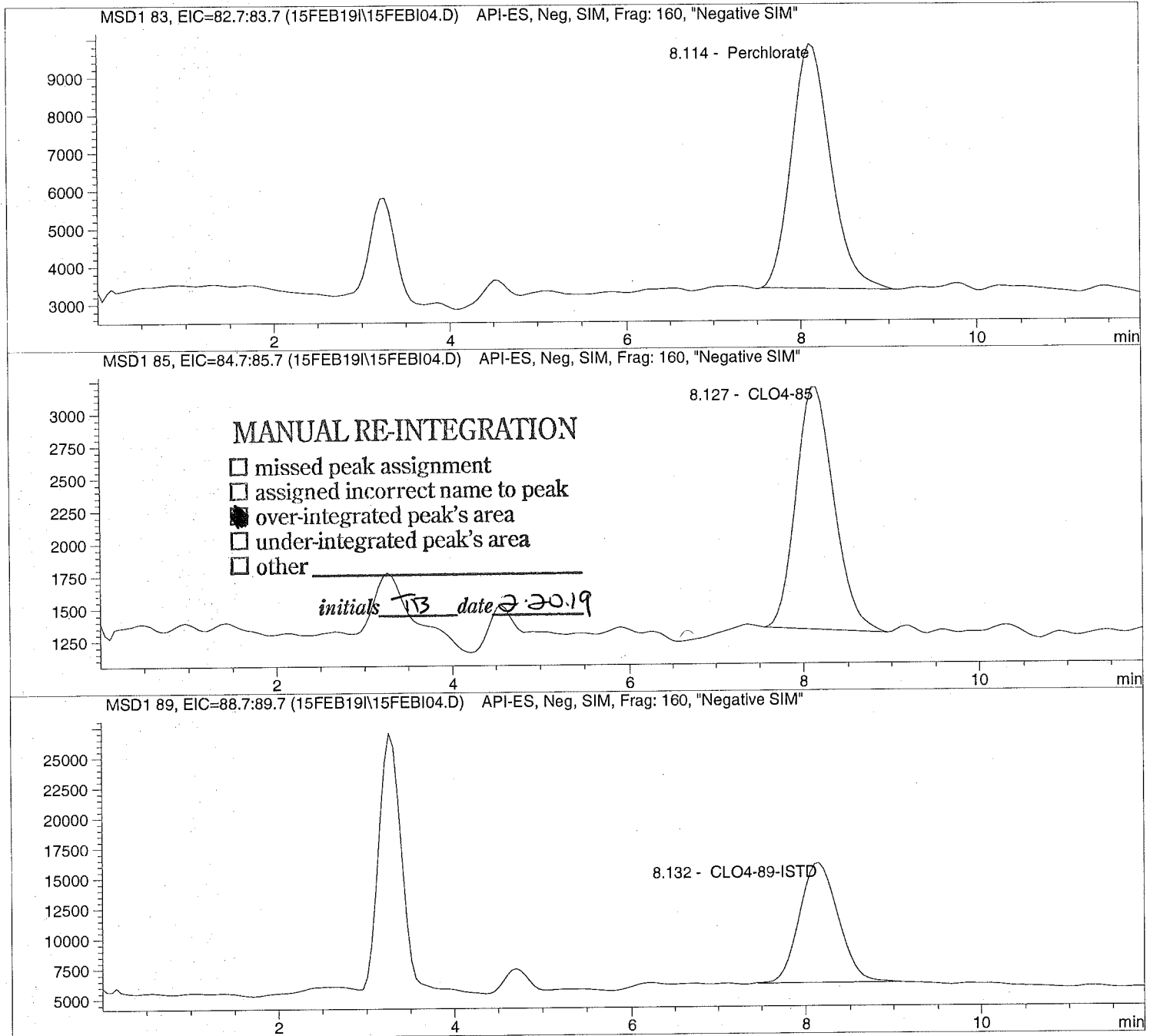
=====
*** End of Report ***

Injection Date: 2/15/2019 10:05:24
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Injection Date: 2/15/2019 10:05:24 Seq Line: 4
Sample Name: CLO4@ 2.0ug/L Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 2.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	MM	55313.4	2.1136	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

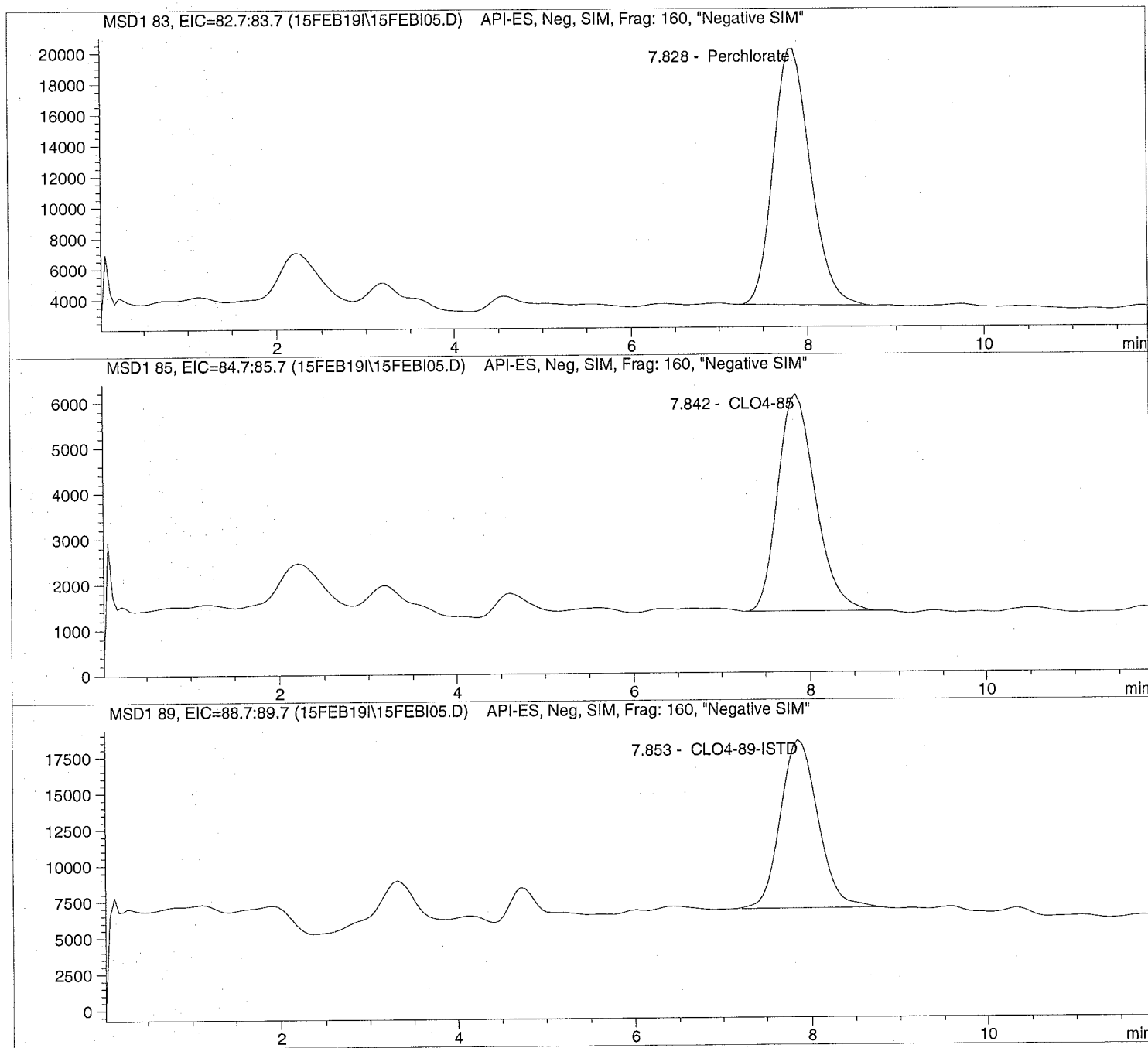
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

*** End of Report ***

=====
Injection Date: 2/15/2019 11:42:56 Seq Line: 5
Sample Name: CLO4@ 5.0ug/L Location: Vial 75
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis
=====



```
=====
Injection Date: 2/15/2019 11:42:56      Seq Line: 5
Sample Name:    CLO4@ 5.0ug/L           Location:  Vial 75
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

Sample Information

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.828	PBA	479370.4	4.6569	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.842	PBA	139246.9	4.9126	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.853	PBA	338646.3	5.0000	CLO4-89-ISTD

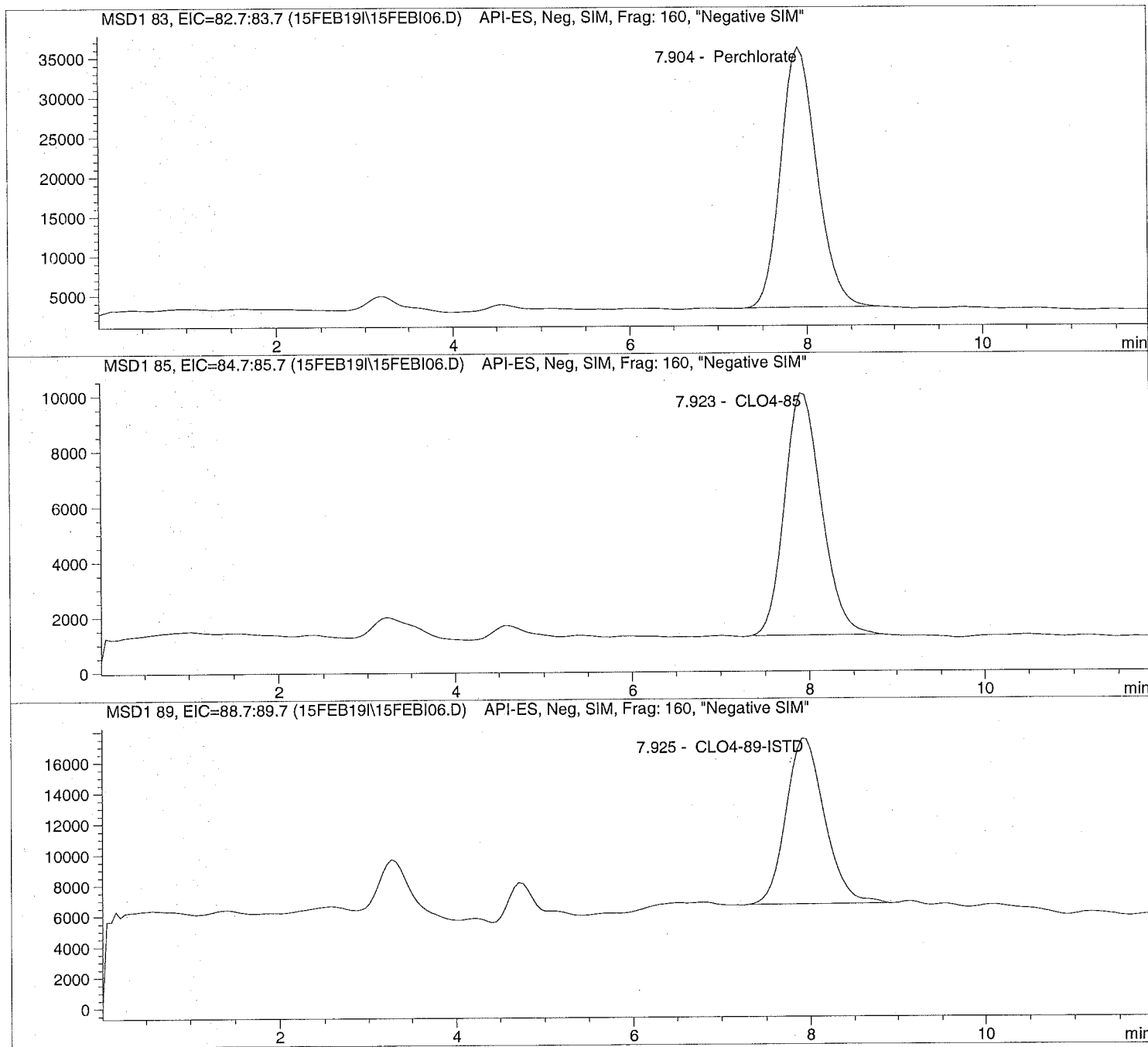
*** End of Report ***

=====
Injection Date: 2/15/2019 11:56:38
Sample Name: CLO4@ 10.ug/L
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis
=====



```
=====
Injection Date: 2/15/2019 11:56:38      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.904	PBA	930135.8	9.1500	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.923	BBA	254395.6	9.3903	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.925	PBA	325154.4	5.0000	CLO4-89-ISTD

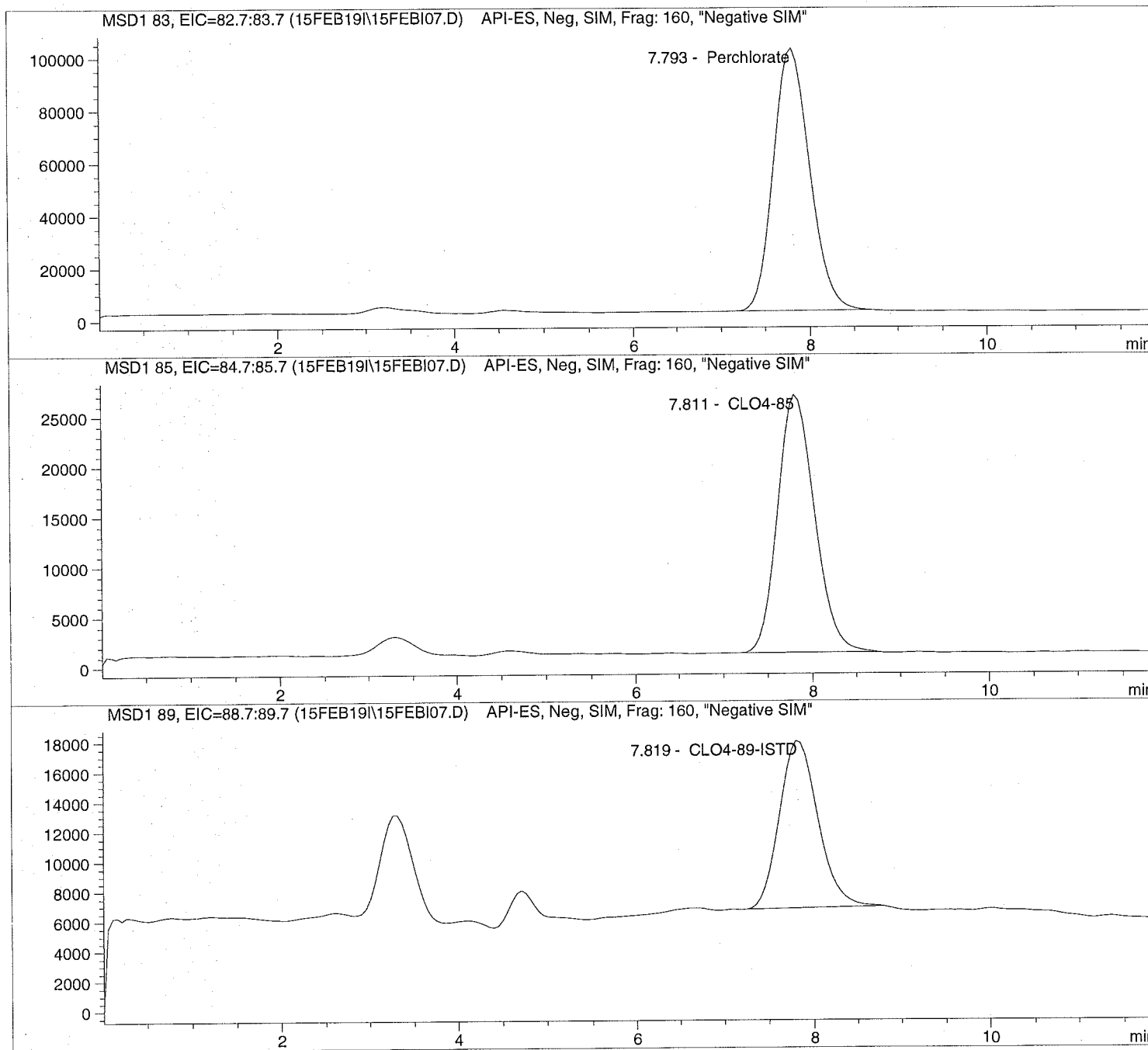
=====
*** End of Report ***

Injection Date: 2/15/2019 12:10:22
Sample Name: CLO4@ 25.ug/L
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 12:10:22      Seq Line: 7  
Sample Name:    CLO4@ 25.ug/L           Location:  Vial 77  
Acq Operator:  TNB                     Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:09:20  
=====
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 25.000  
=====
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.793	PBA	2810669.2	25.5264	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.811	BBA	735968.9	25.4827	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.819	PBA	333799.0	5.0000	CLO4-89-ISTD

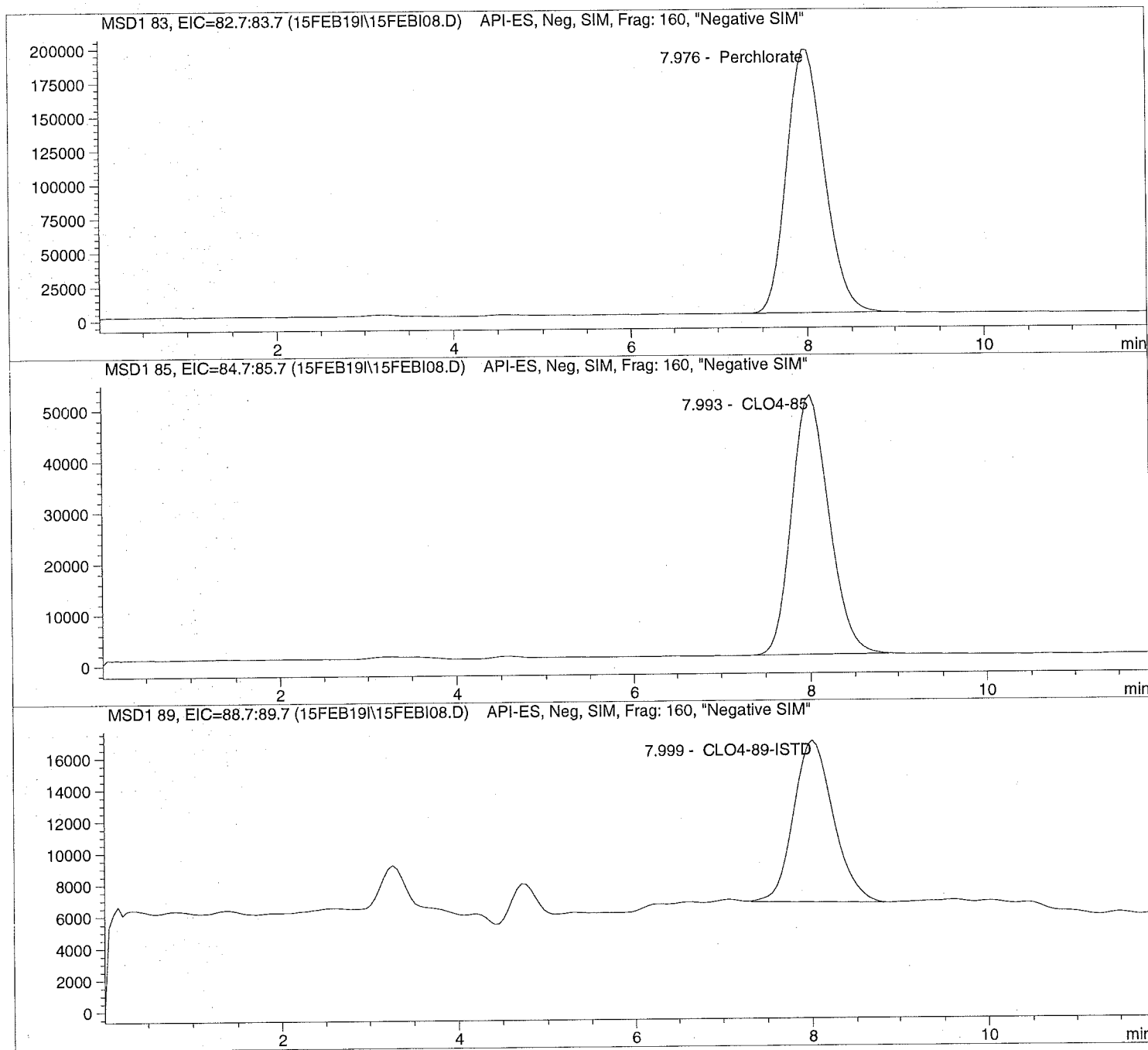
=====
*** End of Report ***
=====

Injection Date: 2/15/2019 12:24:06
Sample Name: CLO4@ 50.ug/L
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis




```
=====
Injection Date: 2/15/2019 12:24:06      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  50.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.976	PBA	5668301.5	51.0744	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.993	PBA	1471522.9	50.3577	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.999	BBA	314711.8	5.0000	CLO4-89-ISTD

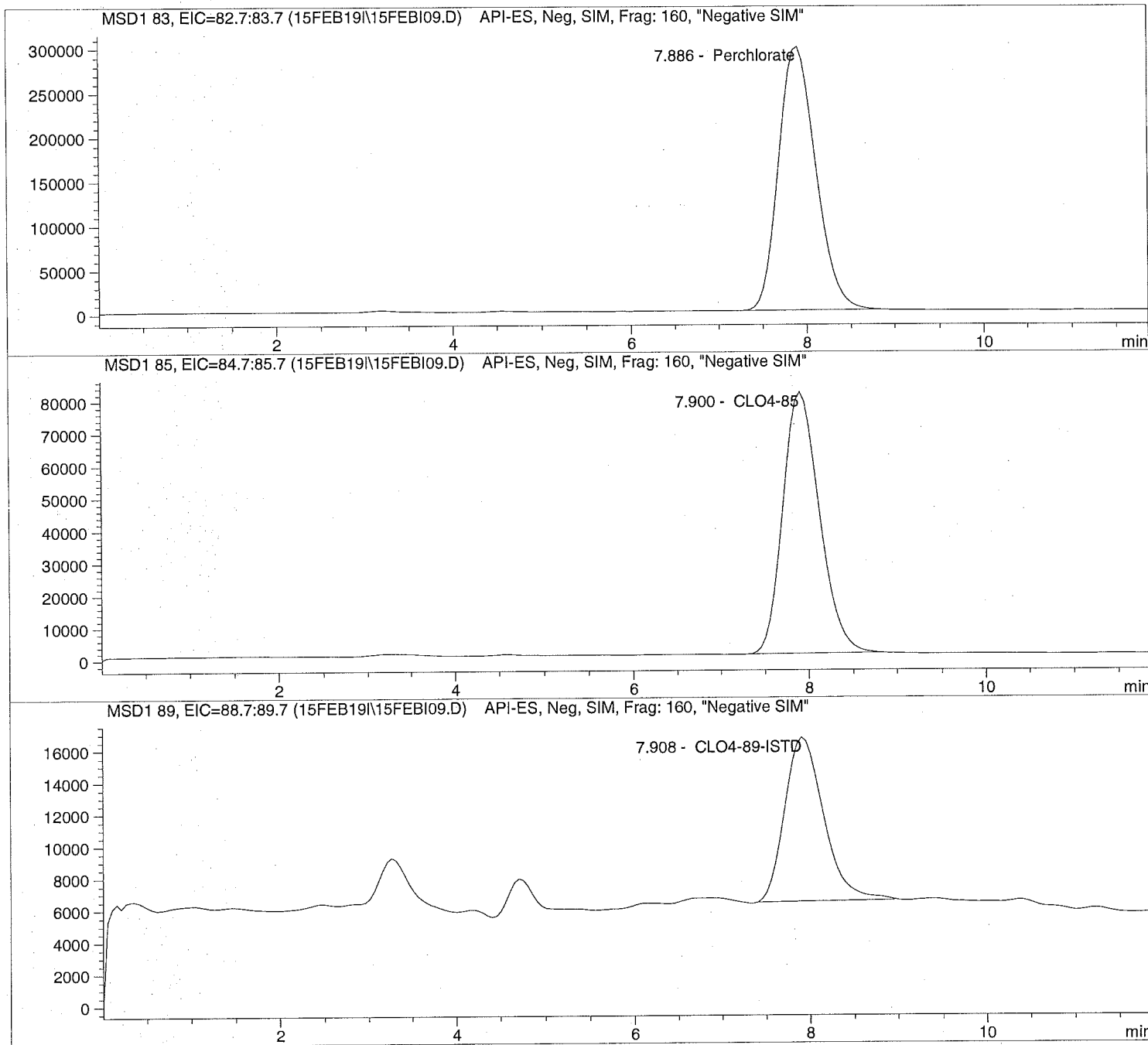
=====
*** End of Report ***

Injection Date: 2/15/2019 12:37:48
Sample Name: CLO4@ 75.ug/L
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 12:37:48      Seq Line: 9  
Sample Name:    CLO4@ 75.ug/L          Location:  Vial 79  
Acq Operator:   TNB                    Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:09:20  
=====
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 75.000  
=====
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.886	PBA	8696239.0	74.3060	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	PBA	2328089.5	74.7223	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.908	PBA	313908.9	5.0000	CLO4-89-ISTD

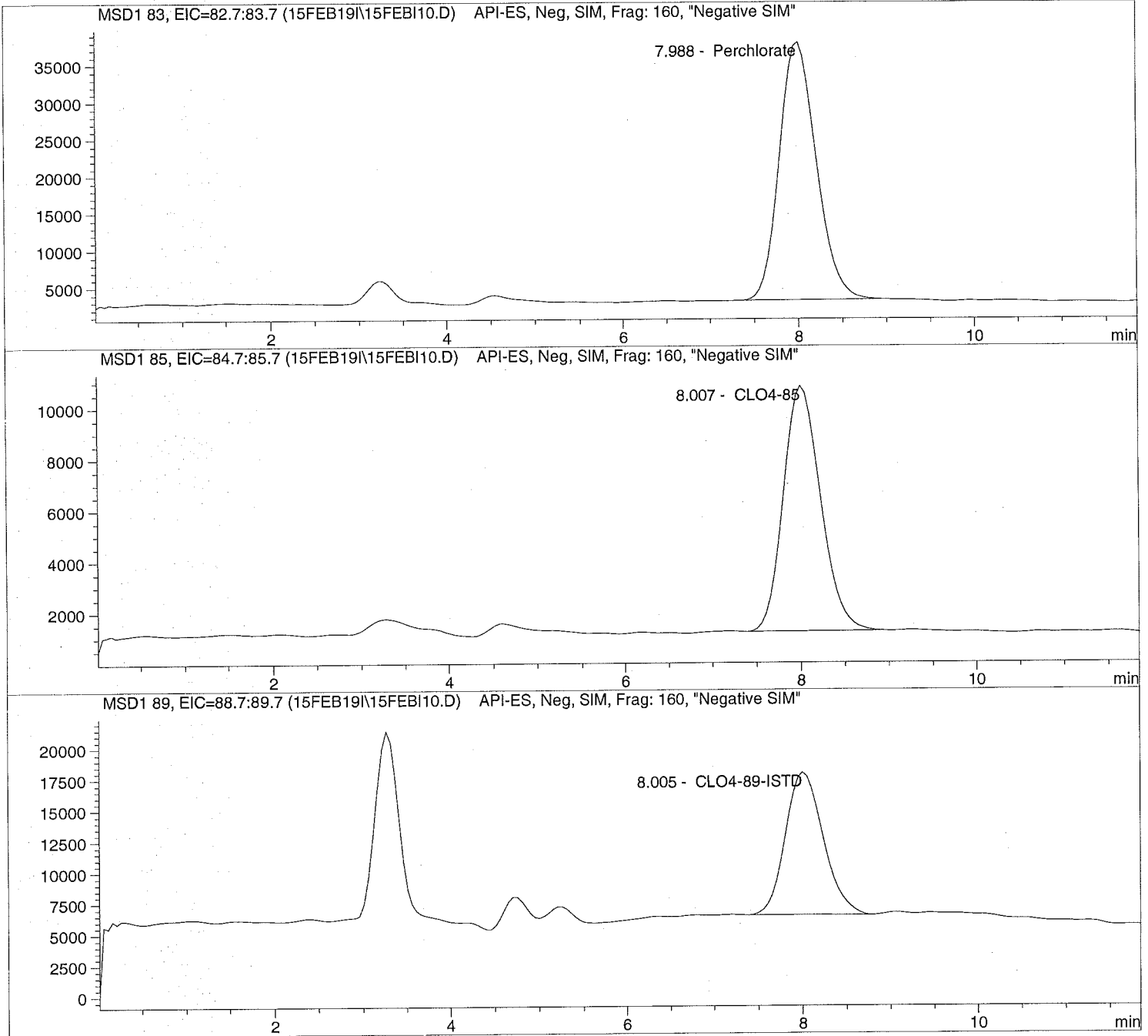
=====
*** End of Report ***
=====

Injection Date: 2/15/2019 12:51:29
Sample Name: ICAL Verf@10ug/L
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====
Injection Date: 2/15/2019 12:51:29      Seq Line: 10
Sample Name:    ICAL Verf@10ug/L        Location:  Vial 80
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.988	BBA	1011409.8	9.4602	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.007	BBA	281229.9	9.8786	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.005	BBA	341503.2	5.0000	CLO4-89-ISTD

=====
*** End of Report ***
=====



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

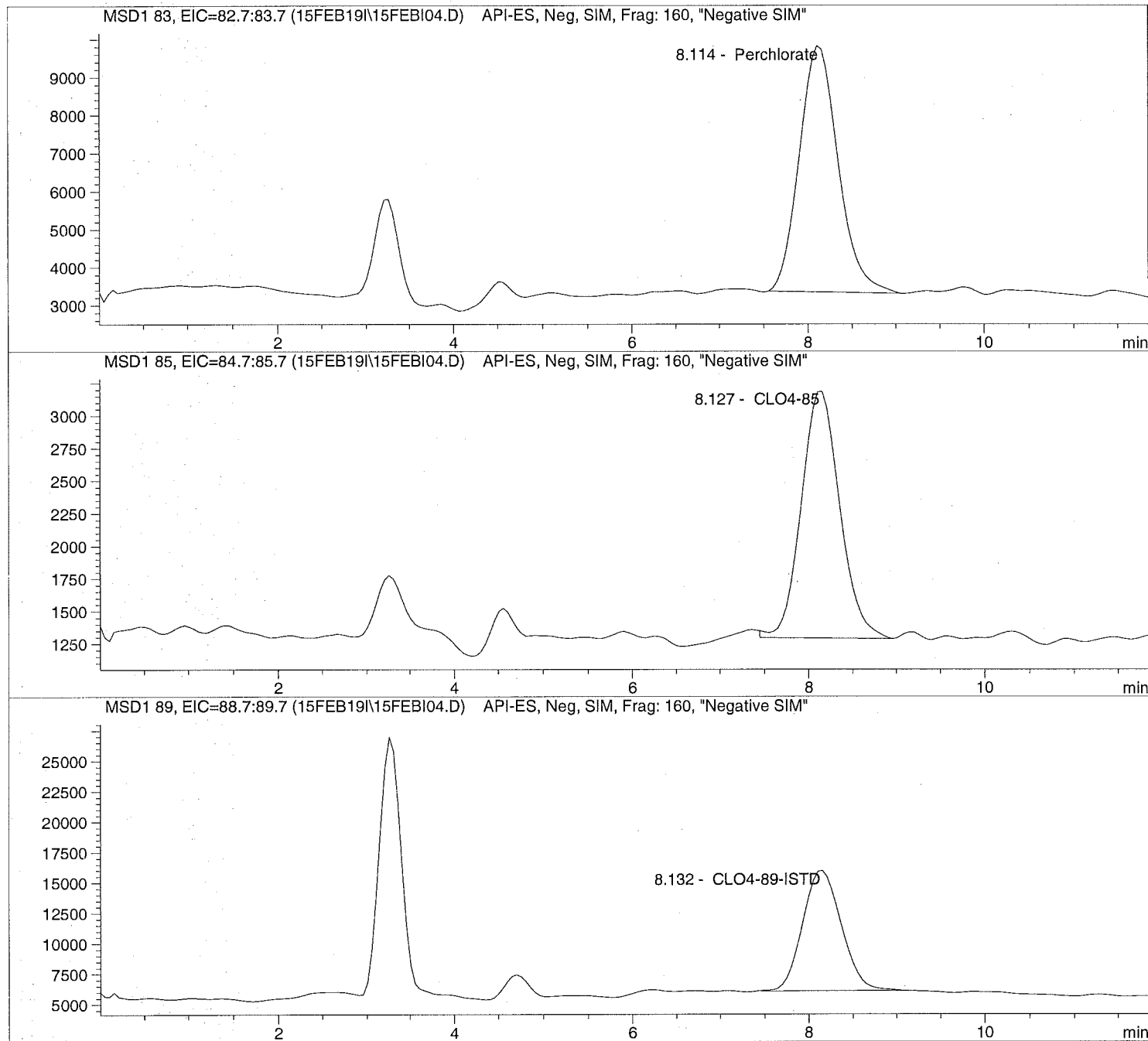
Unmodified

Injection Date: 2/15/2019 10:05:24
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:12:36

Perchlorate analysis



=====
Injection Date: 2/15/2019 10:05:24 Seq Line: 4
Sample Name: CLO4@ 2.0ug/L Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:12:36

Perchlorate analysis

=====
Sample Information
=====

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 2.000

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	BBA	57206.1	2.1923	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

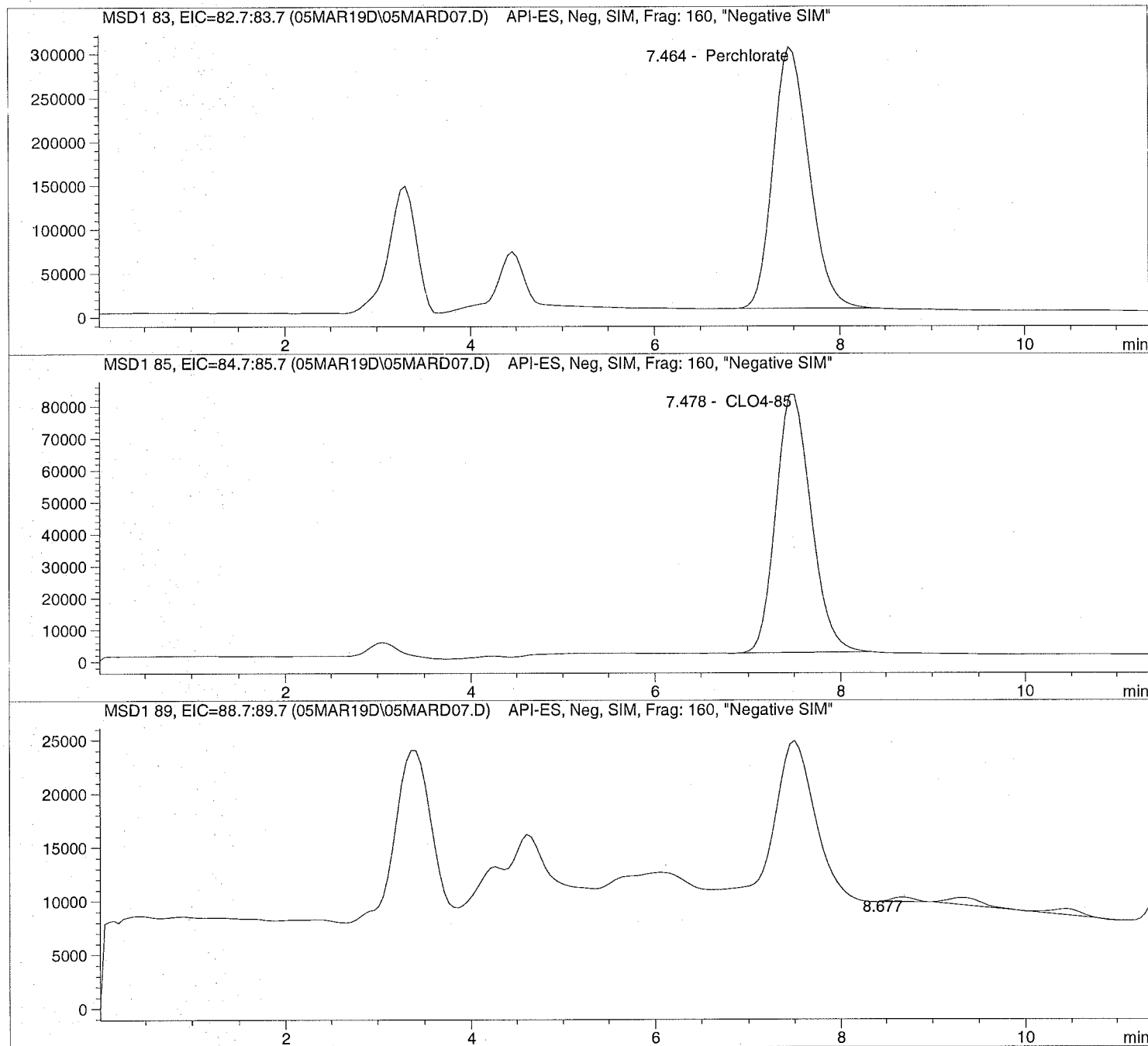
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*** End of Report ***

Injection Date: 3/05/2019 10:07:11
Sample Name: 1906112002 MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:07:11 Seq Line: 7
Sample Name: 1906112002 MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	419.8794	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	402.6179	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

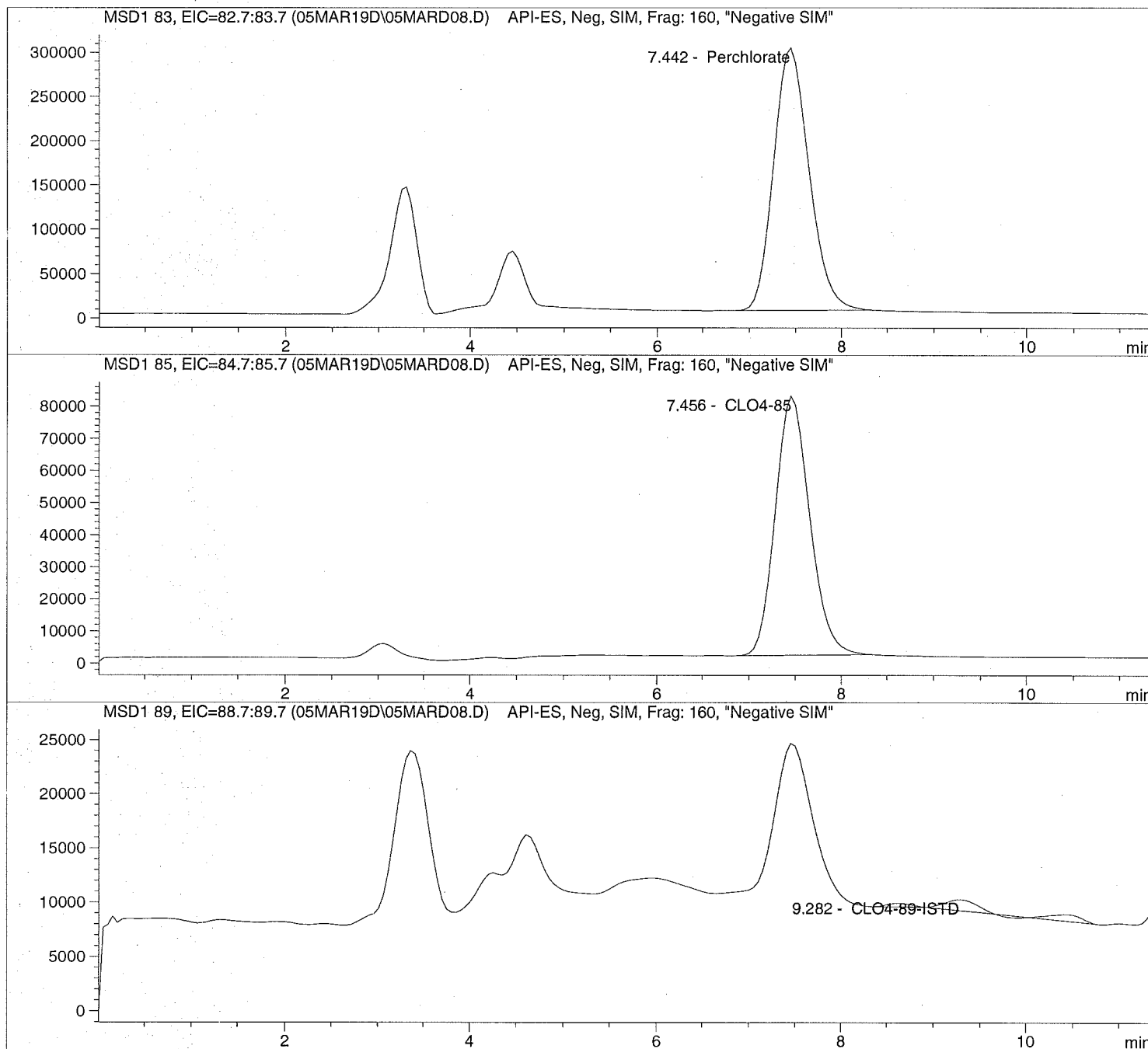
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.677	BB	7208.6	0.0000	
9.316	VBA	28561.1	5.0000	CLO4-89-ISTD

*** End of Report ***

Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	359.0996	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	344.2233	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.621	VB	5769.9	0.0000	
9.282	VBA	35831.6	5.0000	CLO4-89-ISTD

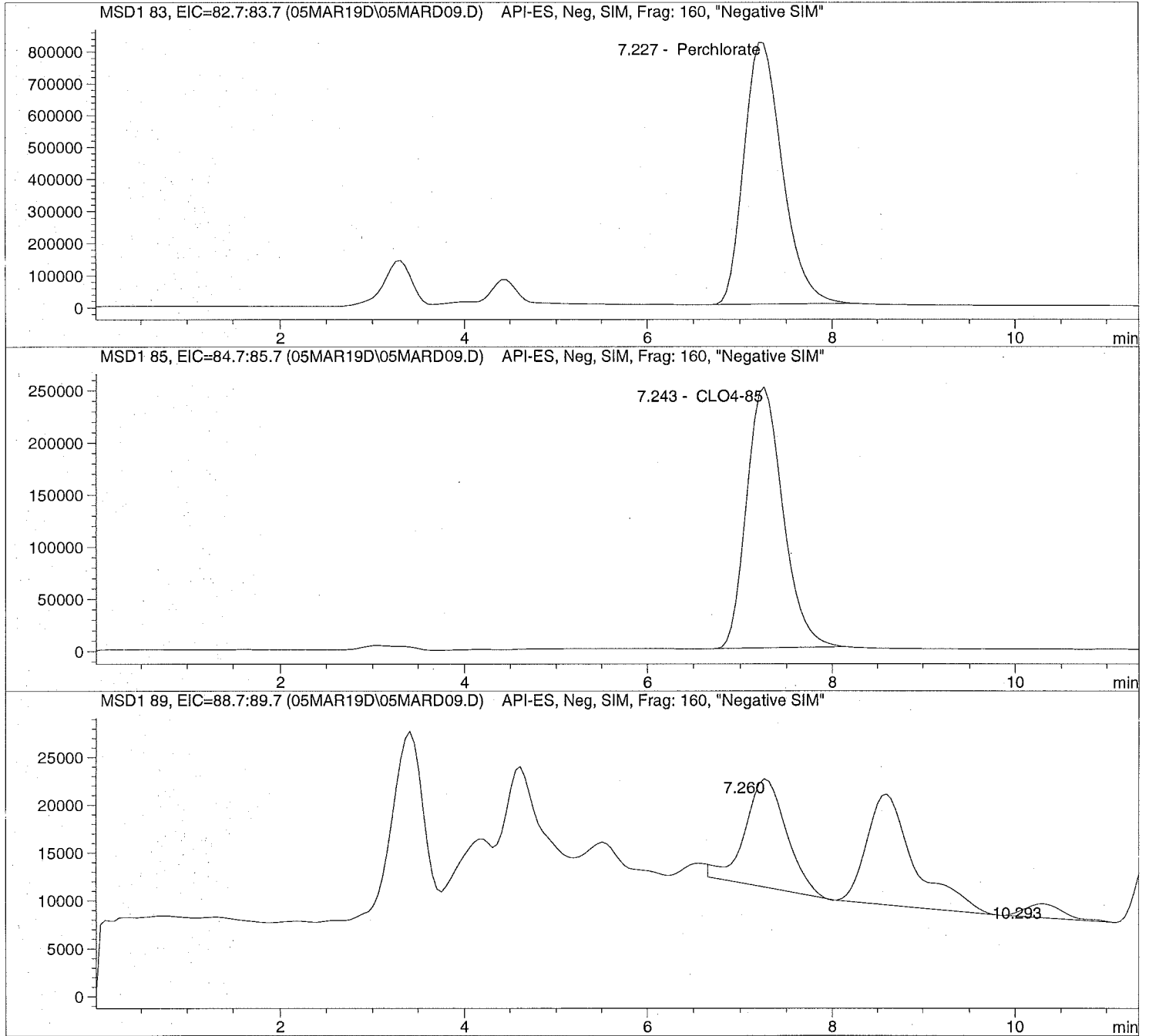
*** End of Report ***

Injection Date: 3/05/2019 10:33:21
Sample Name: 1906112004
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:33:21 Seq Line: 9
Sample Name: 1906112004 Location: Vial 79
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	131.1742	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	138.8050	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.260	BB	352149.7	0.0000	
8.589	VBA	421141.9	5.0000	CLO4-89-ISTD
10.293	BBA	41603.7	0.0000	

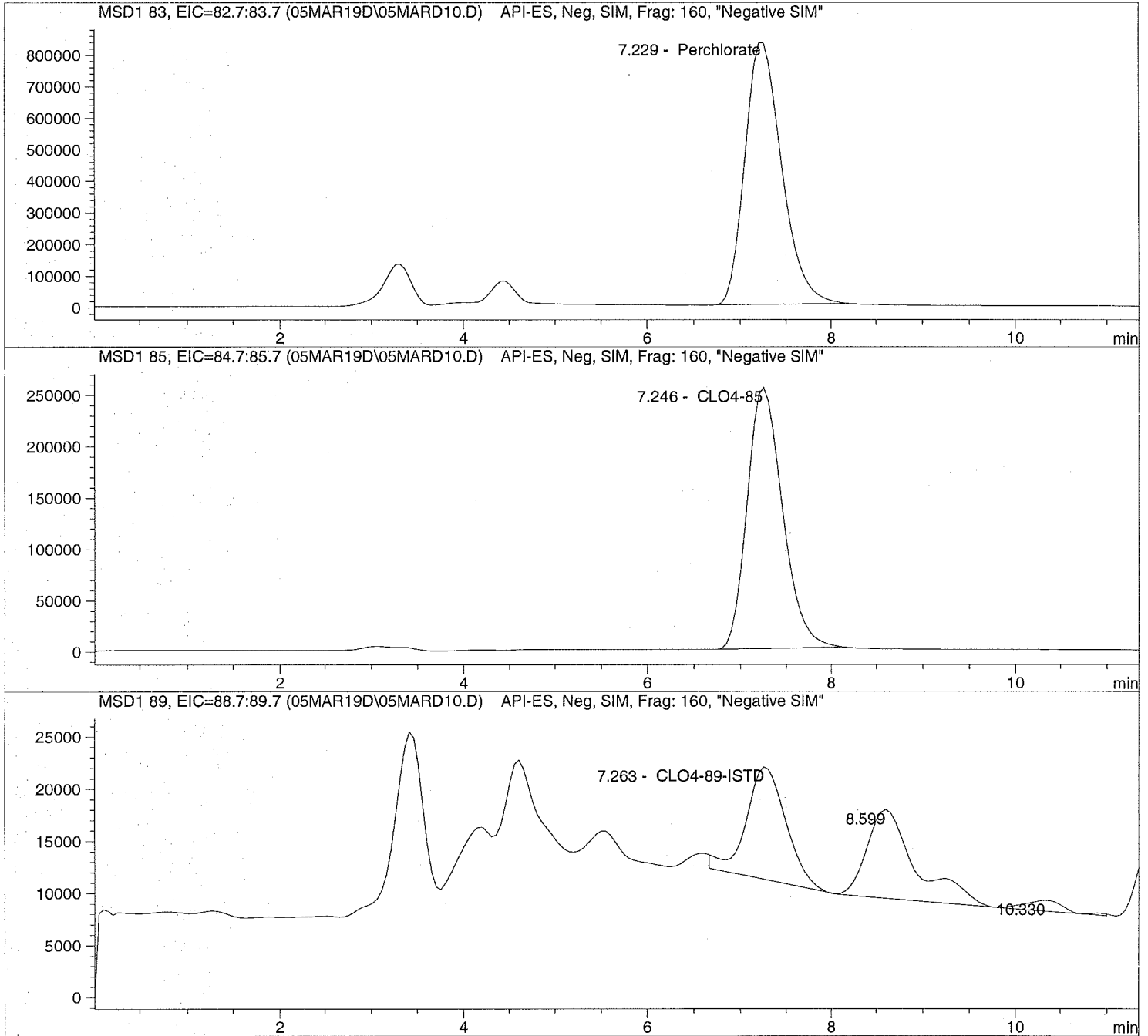
*** End of Report ***

Injection Date: 3/05/2019 10:46:26
Sample Name: 1906112005
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 10:46:26      Seq Line:          10
Sample Name:    1906112005              Location:         Vial 80
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	159.0418	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	168.6882	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	BB	325841.0	5.0000	CLO4-89-ISTD
8.599	VB	308921.9	0.0000	
10.330	VBA	30210.3	0.0000	

=====
*** End of Report ***



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
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March 18, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030012**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples**

Dear Marcia,

ALS Environmental received 3 sample(s) on Mar 01, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read "Raj. P. Modashia", enclosed in a simple oval scribble.

Generated By: DAYNA.FISHER

RJ Modashia
Project Manager

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
Work Order: HS19030012

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030012-01	LH18/24-SP650_022819	Water		28-Feb-2019 14:00	01-Mar-2019 08:50	<input type="checkbox"/>
HS19030012-02	LH18/24-SP650_022819_BIX	Water		28-Feb-2019 14:00	01-Mar-2019 08:50	<input type="checkbox"/>
HS19030012-03	Trip Blank	Water	ALS 112818-41	28-Feb-2019 00:00	01-Mar-2019 08:50	<input type="checkbox"/>

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
-

GCMS Semivolatiles by Method SW8270SIM**Batch ID: 138390****Sample ID: LCSD-138390**

- The RPD between the LCS and LCSD was outside of the control limit for surrogate Nitrobenzene-d5

Sample ID: MBLK-138390/LCS-138390/LCSD-138390

- Surrogates double spiked, however calculations were adjusted accordingly and the recoveries were within control limits.
-

GCMS Volatiles by Method SW8260**Batch ID: R334128****Sample ID: HS19021428-01MS**

- MS and MSD were performed on unrelated sample
-

Metals by Method SW6020**Batch ID: 138281****Sample ID: HS19021441-01MS**

- MS and MSD are for an unrelated sample
-

Wet Chemistry by Method SW7196**Batch ID: R334254**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: LH18/24-SP650_022819
 Collection Date: 28-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030012
 Lab ID:HS19030012-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 12:39	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	06-Mar-2019 12:39	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	06-Mar-2019 12:39	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	06-Mar-2019 12:39	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	06-Mar-2019 12:39	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: LH18/24-SP650_022819
 Collection Date: 28-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030012
 Lab ID:HS19030012-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
cis-1,2-Dichloroethene	2.1		0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	06-Mar-2019 12:39	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 12:39	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	06-Mar-2019 12:39	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Trichloroethene	0.80	J	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:39	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>81.4</i>			0	<i>81-118</i>	%REC	<i>1</i>	<i>06-Mar-2019 12:39</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>96.6</i>			0	<i>85-114</i>	%REC	<i>1</i>	<i>06-Mar-2019 12:39</i>	
<i>Surr: Dibromofluoromethane</i>	<i>83.6</i>			0	<i>80-119</i>	%REC	<i>1</i>	<i>06-Mar-2019 12:39</i>	
<i>Surr: Toluene-d8</i>	<i>107</i>			0	<i>89-112</i>	%REC	<i>1</i>	<i>06-Mar-2019 12:39</i>	
SEMIVOLATILES SIM		Method:SW8270SIM				Prep:SW3510 / 06-Mar-2019		Analyst: QX	
1,4-Dioxane	4.3		0.51	0.51	0.51	ug/L	50	06-Mar-2019 13:23	
<i>Surr: 2-Fluorobiphenyl</i>	<i>55.4</i>			0	<i>40-140</i>	%REC	<i>50</i>	<i>06-Mar-2019 13:23</i>	
<i>Surr: 4-Terphenyl-d14</i>	<i>70.0</i>			0	<i>40-140</i>	%REC	<i>50</i>	<i>06-Mar-2019 13:23</i>	
<i>Surr: Nitrobenzene-d5</i>	<i>54.5</i>			0	<i>40-140</i>	%REC	<i>50</i>	<i>06-Mar-2019 13:23</i>	
ICP-MS METALS BY SW6020A		Method:SW6020				Prep:SW3010A / 01-Mar-2019		Analyst: JHD	
Barium	0.137		0.00190	0.00250	0.00400	mg/L	1	04-Mar-2019 22:49	
Lead	0.00100	U	0.000600	0.00100	0.00200	mg/L	1	04-Mar-2019 22:49	
Selenium	0.00250	U	0.00110	0.00250	0.00200	mg/L	1	04-Mar-2019 22:49	
Silver	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	04-Mar-2019 22:49	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Mar-19

Client:	Bhate Environmental Associates, Inc.	ANALYTICAL REPORT
Project:	LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples	WorkOrder:HS19030012
Sample ID:	LH18/24-SP650_022819	Lab ID:HS19030012-01
Collection Date:	28-Feb-2019 14:00	Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
HEXAVALENT CHROMIUM BY SW7196A		Method:SW7196		Analyst: MZD				
Chromium, Hexavalent	0.0100	U	0.00600	0.0100	0.0100	mg/L	1	01-Mar-2019 11:42

ALS Houston, US

Date: 18-Mar-19

Client:	Bhate Environmental Associates, Inc.	ANALYTICAL REPORT
Project:	LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples	WorkOrder:HS19030012
Sample ID:	LH18/24-SP650_022819_BIX	Lab ID:HS19030012-02
Collection Date:	28-Feb-2019 14:00	Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: Trip Blank
 Collection Date: 28-Feb-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19030012
 Lab ID:HS19030012-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 12:15	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	06-Mar-2019 12:15	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	06-Mar-2019 12:15	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	06-Mar-2019 12:15	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	06-Mar-2019 12:15	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: Trip Blank
 Collection Date: 28-Feb-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19030012
 Lab ID:HS19030012-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD		Method:SW8260							Analyst: PC
8260C									
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	06-Mar-2019 12:15	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 12:15	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	06-Mar-2019 12:15	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Trichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 12:15	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>82.7</i>			0	<i>81-118</i>	<i>%REC</i>	<i>1</i>	<i>06-Mar-2019 12:15</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>99.5</i>			0	<i>85-114</i>	<i>%REC</i>	<i>1</i>	<i>06-Mar-2019 12:15</i>	
<i>Surr: Dibromofluoromethane</i>	<i>86.2</i>			0	<i>80-119</i>	<i>%REC</i>	<i>1</i>	<i>06-Mar-2019 12:15</i>	
<i>Surr: Toluene-d8</i>	<i>107</i>			0	<i>89-112</i>	<i>%REC</i>	<i>1</i>	<i>06-Mar-2019 12:15</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

WEIGHT LOG

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

Batch ID: 138281 **Method:** ICP-MS METALS BY SW6020A **Prep:** 3010A

SamplID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19030012-01	1	10	10 (mL)	1

Batch ID: 138390 **Method:** SEMIVOLATILES SIM **Prep:** 3510_B_SIM

SamplID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19030012-01	1	990	1 (mL)	0.00101

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID 138281	Test Name : ICP-MS METALS BY SW6020A		Matrix: Water			
HS19030012-01	LH18/24-SP650_022819	28 Feb 2019 14:00		01 Mar 2019 10:30	04 Mar 2019 22:49	1
Batch ID 138390	Test Name : SEMIVOLATILES SIM		Matrix: Water			
HS19030012-01	LH18/24-SP650_022819	28 Feb 2019 14:00		06 Mar 2019 09:10	06 Mar 2019 13:23	50
Batch ID R334128	Test Name : VOLATILES ORGANICS BY METHOD 8260C		Matrix: Water			
HS19030012-01	LH18/24-SP650_022819	28 Feb 2019 14:00			06 Mar 2019 12:39	1
HS19030012-03	Trip Blank	28 Feb 2019 00:00			06 Mar 2019 12:15	1
Batch ID R334176	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19030012-02	LH18/24-SP650_022819_BIX	28 Feb 2019 14:00			07 Mar 2019 17:47	1
Batch ID R334254	Test Name : HEXAVALENT CHROMIUM BY SW7196A		Matrix: Water			
HS19030012-01	LH18/24-SP650_022819	28 Feb 2019 14:00			01 Mar 2019 11:42	1

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: 138281 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
MBLK	Sample ID: MBLK-138281	Units: mg/L		Analysis Date: 04-Mar-2019 21:51						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974208		PrepDate: 01-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.00250	0.00400								U
Lead	0.00100	0.00200								U
Selenium	0.00250	0.00200								U
Silver	0.000500	0.00200								U
LCS	Sample ID: LCS-138281	Units: mg/L		Analysis Date: 04-Mar-2019 21:53						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974209		PrepDate: 01-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.05028	0.00400	0.05	0	101	80 - 120				
Lead	0.05263	0.00200	0.05	0	105	80 - 120				
Selenium	0.05168	0.00200	0.05	0	103	80 - 120				
Silver	0.05504	0.00200	0.05	0	110	80 - 120				
MS	Sample ID: HS19021441-01MS	Units: mg/L		Analysis Date: 04-Mar-2019 22:00						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974212		PrepDate: 01-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.3606	0.00400	0.05	0.3093	103	80 - 120				O
Lead	0.0512	0.00200	0.05	0	102	80 - 120				
Selenium	0.04948	0.00200	0.05	0	99.0	80 - 120				
Silver	0.0515	0.00200	0.05	0	103	80 - 120				
MSD	Sample ID: HS19021441-01MSD	Units: mg/L		Analysis Date: 04-Mar-2019 22:02						
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974213		PrepDate: 01-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.3437	0.00400	0.05	0.3093	68.8	80 - 120	0.3606	4.8	20	SO
Lead	0.0515	0.00200	0.05	0	103	80 - 120	0.0512	0.582	20	
Selenium	0.04959	0.00200	0.05	0	99.2	80 - 120	0.04948	0.218	20	
Silver	0.05174	0.00200	0.05	0	103	80 - 120	0.0515	0.473	20	

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: 138281 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
PDS	Sample ID: HS19021441-01PDS	Units: mg/L			Analysis Date: 04-Mar-2019 22:04					
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974214		PrepDate: 01-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.4021	0.00400	0.1	0.3093	92.9	75 - 125				
Lead	0.1019	0.00200	0.1	0	102	75 - 125				
Selenium	0.09793	0.00200	0.1	0	97.9	75 - 125				
Silver	0.09993	0.00200	0.1	0	99.9	75 - 125				
SD	Sample ID: HS19021441-01SD	Units: mg/L			Analysis Date: 04-Mar-2019 21:57					
Client ID:	Run ID: ICPMS05_333923	SeqNo: 4974211		PrepDate: 01-Mar-2019		DF: 5				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	RPD Limit	Qual
Barium	0.3016	0.0200					0.3093	2.48	10	
Lead	0.00500	0.0100					0.000204	0	10	U
Selenium	0.0125	0.0100					0.000452	0	10	U
Silver	0.00250	0.0100					0.00002	0	10	U

The following samples were analyzed in this batch:

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: 138390 (0)		Instrument: SV-5		Method: SEMIVOLATILES SIM						
MBLK	Sample ID: MBLK-138390	Units: ug/L			Analysis Date: 06-Mar-2019 11:36					
Client ID:	Run ID: SV-5_334088	SeqNo: 4976850		PrepDate: 06-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
1,4-Dioxane	0.010	0.010							U	
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.1248</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>78.0</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.1145</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>71.6</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.1894</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>118</i>	<i>40 - 140</i>				
LCS	Sample ID: LCS-138390	Units: ug/L			Analysis Date: 06-Mar-2019 11:56					
Client ID:	Run ID: SV-5_334088	SeqNo: 4976851		PrepDate: 06-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
1,4-Dioxane	0.1105	0.010	0.08	0	138	40 - 140				
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.1196</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>74.8</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.1031</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>64.4</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.1636</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>102</i>	<i>40 - 140</i>				
LCSD	Sample ID: LCSD-138390	Units: ug/L			Analysis Date: 06-Mar-2019 12:17					
Client ID:	Run ID: SV-5_334088	SeqNo: 4976852		PrepDate: 06-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
1,4-Dioxane	0.1033	0.010	0.08	0	129	40 - 140	0.1105	6.7	20	
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.1048</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>65.5</i>	<i>40 - 140</i>	<i>0.1196</i>	<i>13.2</i>	<i>20</i>	
<i>Surr: 4-Terphenyl-d14</i>	<i>0.1045</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>65.3</i>	<i>40 - 140</i>	<i>0.1031</i>	<i>1.42</i>	<i>20</i>	
<i>Surr: Nitrobenzene-d5</i>	<i>0.1167</i>	<i>0</i>	<i>0.16</i>	<i>0</i>	<i>72.9</i>	<i>40 - 140</i>	<i>0.1636</i>	<i>33.5</i>	<i>20</i>	

The following samples were analyzed in this batch: HS19030012-01

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334128 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190306	Units: UG/L			Analysis Date: 06-Mar-2019 11:51					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977685	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	41.51	1.0	50	0	83.0	81 - 118				
Surr: 4-Bromofluorobenzene	49.32	1.0	50	0	98.6	85 - 114				
Surr: Dibromofluoromethane	43.47	1.0	50	0	86.9	80 - 119				
Surr: Toluene-d8	53.24	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334128 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190306	Units: UG/L			Analysis Date: 06-Mar-2019 11:03					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977684	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	20.28	1.0	20	0	101	78 - 124				
1,1,1-Trichloroethane	20.23	1.0	20	0	101	74 - 131				
1,1,2,2-Tetrachloroethane	19.2	1.0	20	0	96.0	71 - 121				
1,1,2-Trichloroethane	19.93	1.0	20	0	99.6	80 - 119				
1,1-Dichloroethane	19.67	1.0	20	0	98.3	77 - 125				
1,1-Dichloroethene	19.73	1.0	20	0	98.6	71 - 131				
1,1-Dichloropropene	19.78	1.0	20	0	98.9	78 - 125				
1,2,3-Trichlorobenzene	20.06	1.0	20	0	100	69 - 129				
1,2,3-Trichloropropane	18.96	1.0	20	0	94.8	73 - 122				
1,2,4-Trichlorobenzene	19.89	1.0	20	0	99.4	69 - 130				
1,2,4-Trimethylbenzene	19.37	1.0	20	0	96.8	76 - 124				
1,2-Dibromo-3-chloropropane	18.77	1.0	20	0	93.9	62 - 128				
1,2-Dibromoethane	20.52	1.0	20	0	103	77 - 121				
1,2-Dichlorobenzene	19.02	1.0	20	0	95.1	80 - 119				
1,2-Dichloroethane	20.54	1.0	20	0	103	73 - 128				
1,2-Dichloropropane	20.31	1.0	20	0	102	78 - 122				
1,3,5-Trimethylbenzene	19.43	1.0	20	0	97.1	75 - 124				
1,3-Dichlorobenzene	19.44	1.0	20	0	97.2	80 - 119				
1,3-Dichloropropane	20.12	1.0	20	0	101	80 - 119				
1,4-Dichlorobenzene	19.16	1.0	20	0	95.8	79 - 118				
2,2-Dichloropropane	19.96	1.0	20	0	99.8	60 - 139				
2-Butanone	42.82	2.0	40	0	107	56 - 143				
2-Chlorotoluene	19.13	1.0	20	0	95.7	79 - 122				
2-Hexanone	40.07	2.0	40	0	100	57 - 139				
4-Chlorotoluene	19.38	1.0	20	0	96.9	78 - 122				
4-Isopropyltoluene	19.42	1.0	20	0	97.1	77 - 127				
4-Methyl-2-pentanone	39.73	2.0	40	0	99.3	67 - 130				
Acetone	42.44	2.0	40	0	106	39 - 160				
Benzene	19.99	1.0	20	0	99.9	79 - 120				
Bromobenzene	19.33	1.0	20	0	96.6	80 - 120				
Bromochloromethane	19.66	1.0	20	0	98.3	78 - 123				
Bromodichloromethane	19.92	1.0	20	0	99.6	79 - 125				
Bromoform	20.37	1.0	20	0	102	66 - 130				
Bromomethane	24.28	1.0	20	0	121	53 - 141				

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334128 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190306	Units: UG/L			Analysis Date: 06-Mar-2019 11:03					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977684		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	40.1	2.0	40	0	100	64 - 133				
Carbon tetrachloride	20.59	1.0	20	0	103	72 - 136				
Chlorobenzene	19.9	1.0	20	0	99.5	82 - 118				
Chloroethane	19.71	1.0	20	0	98.5	60 - 138				
Chloroform	19.55	1.0	20	0	97.7	79 - 124				
Chloromethane	20.54	1.0	20	0	103	50 - 139				
cis-1,2-Dichloroethene	19.71	1.0	20	0	98.6	78 - 123				
cis-1,3-Dichloropropene	20.44	1.0	20	0	102	75 - 124				
Dibromochloromethane	20.25	1.0	20	0	101	74 - 126				
Dibromomethane	20.09	1.0	20	0	100	79 - 123				
Dichlorodifluoromethane	20.24	1.0	20	0	101	32 - 152				
Ethylbenzene	20.01	1.0	20	0	100	79 - 121				
Hexachlorobutadiene	21.69	1.0	20	0	108	66 - 134				
Isopropylbenzene	19.97	1.0	20	0	99.8	72 - 131				
m,p-Xylene	40.08	2.0	40	0	100	80 - 121				
Methylene chloride	20.41	2.0	20	0	102	74 - 124				
Naphthalene	18.14	1.0	20	0	90.7	61 - 128				
n-Butylbenzene	19.01	1.0	20	0	95.0	75 - 128				
n-Propylbenzene	19.22	1.0	20	0	96.1	76 - 126				
o-Xylene	20.19	1.0	20	0	101	78 - 122				
sec-Butylbenzene	19	1.0	20	0	95.0	77 - 126				
Styrene	20.48	1.0	20	0	102	78 - 123				
tert-Butylbenzene	18.99	1.0	20	0	95.0	78 - 124				
Tetrachloroethene	19.96	1.0	20	0	99.8	74 - 129				
Toluene	19.89	1.0	20	0	99.5	80 - 121				
trans-1,2-Dichloroethene	20.32	1.0	20	0	102	75 - 124				
trans-1,3-Dichloropropene	20.39	1.0	20	0	102	73 - 127				
Trichloroethene	20.14	1.0	20	0	101	79 - 123				
Trichlorofluoromethane	19.72	1.0	20	0	98.6	65 - 141				
Vinyl chloride	19.12	1.0	20	0	95.6	58 - 137				
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>50.81</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>102</i>	<i>81 - 118</i>				
<i>Surr: 4-Bromofluorobenzene</i>	<i>51.81</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>104</i>	<i>85 - 114</i>				
<i>Surr: Dibromofluoromethane</i>	<i>51.53</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>103</i>	<i>80 - 119</i>				
<i>Surr: Toluene-d8</i>	<i>48.19</i>	<i>1.0</i>	<i>50</i>	<i>0</i>	<i>96.4</i>	<i>89 - 112</i>				

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334128 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19021428-01MS	Units: UG/L			Analysis Date: 06-Mar-2019 14:39					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977690	PrepDate:	DF: 50						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	1131	50	1000	0	113	78 - 124				
1,1,1-Trichloroethane	987.3	50	1000	0	98.7	74 - 131				
1,1,2,2-Tetrachloroethane	1076	50	1000	0	108	71 - 121				
1,1,2-Trichloroethane	1111	50	1000	0	111	80 - 119				
1,1-Dichloroethane	940.2	50	1000	0	94.0	77 - 125				
1,1-Dichloroethene	1011	50	1000	0	101	71 - 131				
1,1-Dichloropropene	1112	50	1000	0	111	78 - 125				
1,2,3-Trichlorobenzene	1082	50	1000	0	108	69 - 129				
1,2,3-Trichloropropane	1033	50	1000	0	103	73 - 122				
1,2,4-Trichlorobenzene	1136	50	1000	0	114	69 - 130				
1,2,4-Trimethylbenzene	1203	50	1000	0	120	76 - 124				
1,2-Dibromo-3-chloropropane	1044	50	1000	0	104	62 - 128				
1,2-Dibromoethane	1119	50	1000	0	112	77 - 121				
1,2-Dichlorobenzene	1128	50	1000	0	113	80 - 119				
1,2-Dichloroethane	1026	50	1000	0	103	73 - 128				
1,2-Dichloropropane	1034	50	1000	0	103	78 - 122				
1,3,5-Trimethylbenzene	1243	50	1000	0	124	75 - 124				S
1,3-Dichlorobenzene	1180	50	1000	0	118	80 - 119				
1,3-Dichloropropane	1110	50	1000	0	111	80 - 119				
1,4-Dichlorobenzene	1153	50	1000	0	115	79 - 118				
2,2-Dichloropropane	1001	50	1000	0	100	60 - 139				
2-Butanone	1842	100	2000	0	92.1	56 - 143				
2-Chlorotoluene	1181	50	1000	0	118	79 - 122				
2-Hexanone	2034	100	2000	0	102	57 - 139				
4-Chlorotoluene	1190	50	1000	0	119	78 - 122				
4-Isopropyltoluene	1296	50	1000	0	130	77 - 127				S
4-Methyl-2-pentanone	2036	100	2000	0	102	67 - 130				
Acetone	1745	100	2000	0	87.3	39 - 160				
Benzene	1045	50	1000	0	104	79 - 120				
Bromobenzene	1145	50	1000	0	114	80 - 120				
Bromochloromethane	903.7	50	1000	0	90.4	78 - 123				
Bromodichloromethane	1027	50	1000	0	103	79 - 125				
Bromoform	1094	50	1000	0	109	66 - 130				
Bromomethane	1198	50	1000	0	120	53 - 141				

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334128 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19021428-01MS	Units: UG/L			Analysis Date: 06-Mar-2019 14:39					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977690	PrepDate:	DF: 50						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1925	100	2000	0	96.2	64 - 133				
Carbon tetrachloride	1169	50	1000	0	117	72 - 136				
Chlorobenzene	1159	50	1000	0	116	82 - 118				
Chloroethane	922.9	50	1000	0	92.3	60 - 138				
Chloroform	938.1	50	1000	0	93.8	79 - 124				
Chloromethane	804.4	50	1000	0	80.4	50 - 139				
cis-1,2-Dichloroethene	4602	50	1000	3749	85.3	78 - 123				
cis-1,3-Dichloropropene	1038	50	1000	0	104	75 - 124				
Dibromochloromethane	1148	50	1000	0	115	74 - 126				
Dibromomethane	1016	50	1000	0	102	79 - 123				
Dichlorodifluoromethane	668.3	50	1000	0	66.8	32 - 152				
Ethylbenzene	1202	50	1000	0	120	79 - 121				
Hexachlorobutadiene	1396	50	1000	0	140	66 - 134				S
Isopropylbenzene	1253	50	1000	0	125	72 - 131				
m,p-Xylene	2393	100	2000	0	120	80 - 121				
Methylene chloride	955.4	100	1000	0	95.5	74 - 124				
Naphthalene	970.7	50	1000	0	97.1	61 - 128				
n-Butylbenzene	1323	50	1000	0	132	75 - 128				S
n-Propylbenzene	1256	50	1000	0	126	76 - 126				
o-Xylene	1200	50	1000	0	120	78 - 122				
sec-Butylbenzene	1306	50	1000	0	131	77 - 126				S
Styrene	1164	50	1000	0	116	78 - 123				
tert-Butylbenzene	1270	50	1000	0	127	78 - 124				S
Tetrachloroethene	1293	50	1000	0	129	74 - 129				S
Toluene	1184	50	1000	0	118	80 - 121				
trans-1,2-Dichloroethene	997.2	50	1000	0	99.7	75 - 124				
trans-1,3-Dichloropropene	1039	50	1000	0	104	73 - 127				
Trichloroethene	4573	50	1000	3686	88.7	79 - 123				
Trichlorofluoromethane	1014	50	1000	0	101	65 - 141				
Vinyl chloride	865.2	50	1000	0	86.5	58 - 137				
Surr: 1,2-Dichloroethane-d4	2074	50	2500	0	83.0	81 - 118				
Surr: 4-Bromofluorobenzene	2472	50	2500	0	98.9	85 - 114				
Surr: Dibromofluoromethane	2130	50	2500	0	85.2	80 - 119				
Surr: Toluene-d8	2648	50	2500	0	106	89 - 112				

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334128 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19021428-01MSD	Units: UG/L			Analysis Date: 06-Mar-2019 15:03					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977691	PrepDate:	DF: 50						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	1074	50	1000	0	107	78 - 124	1131	5.23	20	
1,1,1-Trichloroethane	1004	50	1000	0	100	74 - 131	987.3	1.64	20	
1,1,2,2-Tetrachloroethane	977.6	50	1000	0	97.8	71 - 121	1076	9.59	20	
1,1,2-Trichloroethane	1026	50	1000	0	103	80 - 119	1111	7.97	20	
1,1-Dichloroethane	918.7	50	1000	0	91.9	77 - 125	940.2	2.32	20	
1,1-Dichloroethene	1003	50	1000	0	100	71 - 131	1011	0.851	20	
1,1-Dichloropropene	1155	50	1000	0	116	78 - 125	1112	3.85	20	
1,2,3-Trichlorobenzene	1116	50	1000	0	112	69 - 129	1082	3.06	20	
1,2,3-Trichloropropane	947	50	1000	0	94.7	73 - 122	1033	8.7	20	
1,2,4-Trichlorobenzene	1120	50	1000	0	112	69 - 130	1136	1.43	20	
1,2,4-Trimethylbenzene	1139	50	1000	0	114	76 - 124	1203	5.47	20	
1,2-Dibromo-3-chloropropane	993.9	50	1000	0	99.4	62 - 128	1044	4.94	20	
1,2-Dibromoethane	1034	50	1000	0	103	77 - 121	1119	7.94	20	
1,2-Dichlorobenzene	1051	50	1000	0	105	80 - 119	1128	7.08	20	
1,2-Dichloroethane	1012	50	1000	0	101	73 - 128	1026	1.39	20	
1,2-Dichloropropane	1017	50	1000	0	102	78 - 122	1034	1.63	20	
1,3,5-Trimethylbenzene	1168	50	1000	0	117	75 - 124	1243	6.24	20	
1,3-Dichlorobenzene	1093	50	1000	0	109	80 - 119	1180	7.6	20	
1,3-Dichloropropane	1025	50	1000	0	102	80 - 119	1110	7.93	20	
1,4-Dichlorobenzene	1063	50	1000	0	106	79 - 118	1153	8.16	20	
2,2-Dichloropropane	982.5	50	1000	0	98.2	60 - 139	1001	1.83	20	
2-Butanone	1713	100	2000	0	85.7	56 - 143	1842	7.25	20	
2-Chlorotoluene	1102	50	1000	0	110	79 - 122	1181	6.9	20	
2-Hexanone	1939	100	2000	0	96.9	57 - 139	2034	4.78	20	
4-Chlorotoluene	1105	50	1000	0	110	78 - 122	1190	7.4	20	
4-Isopropyltoluene	1246	50	1000	0	125	77 - 127	1296	3.91	20	
4-Methyl-2-pentanone	1953	100	2000	0	97.6	67 - 130	2036	4.18	20	
Acetone	1701	100	2000	0	85.0	39 - 160	1745	2.58	20	
Benzene	1032	50	1000	0	103	79 - 120	1045	1.18	20	
Bromobenzene	1050	50	1000	0	105	80 - 120	1145	8.62	20	
Bromochloromethane	886.4	50	1000	0	88.6	78 - 123	903.7	1.92	20	
Bromodichloromethane	1012	50	1000	0	101	79 - 125	1027	1.47	20	
Bromoform	1026	50	1000	0	103	66 - 130	1094	6.41	20	
Bromomethane	1111	50	1000	0	111	53 - 141	1198	7.53	20	

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334128 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19021428-01MSD	Units: UG/L			Analysis Date: 06-Mar-2019 15:03					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977691		PrepDate:		DF: 50				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit	Qual
Carbon disulfide	1914	100	2000	0	95.7	64 - 133	1925	0.566	20	
Carbon tetrachloride	1209	50	1000	0	121	72 - 136	1169	3.36	20	
Chlorobenzene	1092	50	1000	0	109	82 - 118	1159	5.98	20	
Chloroethane	929.6	50	1000	0	93.0	60 - 138	922.9	0.728	20	
Chloroform	908	50	1000	0	90.8	79 - 124	938.1	3.27	20	
Chloromethane	783	50	1000	0	78.3	50 - 139	804.4	2.7	20	
cis-1,2-Dichloroethene	4457	50	1000	3749	70.8	78 - 123	4602	3.18	20	S
cis-1,3-Dichloropropene	1021	50	1000	0	102	75 - 124	1038	1.73	20	
Dibromochloromethane	1059	50	1000	0	106	74 - 126	1148	8.06	20	
Dibromomethane	990.7	50	1000	0	99.1	79 - 123	1016	2.54	20	
Dichlorodifluoromethane	688.8	50	1000	0	68.9	32 - 152	668.3	3.02	20	
Ethylbenzene	1160	50	1000	0	116	79 - 121	1202	3.53	20	
Hexachlorobutadiene	1434	50	1000	0	143	66 - 134	1396	2.69	20	S
Isopropylbenzene	1231	50	1000	0	123	72 - 131	1253	1.82	20	
m,p-Xylene	2294	100	2000	0	115	80 - 121	2393	4.21	20	
Methylene chloride	914.4	100	1000	0	91.4	74 - 124	955.4	4.38	20	
Naphthalene	939.7	50	1000	0	94.0	61 - 128	970.7	3.25	20	
n-Butylbenzene	1292	50	1000	0	129	75 - 128	1323	2.38	20	S
n-Propylbenzene	1191	50	1000	0	119	76 - 126	1256	5.27	20	
o-Xylene	1141	50	1000	0	114	78 - 122	1200	5.07	20	
sec-Butylbenzene	1272	50	1000	0	127	77 - 126	1306	2.68	20	S
Styrene	1098	50	1000	0	110	78 - 123	1164	5.82	20	
tert-Butylbenzene	1210	50	1000	0	121	78 - 124	1270	4.84	20	
Tetrachloroethene	1247	50	1000	0	125	74 - 129	1293	3.64	20	
Toluene	1121	50	1000	0	112	80 - 121	1184	5.5	20	
trans-1,2-Dichloroethene	971.6	50	1000	0	97.2	75 - 124	997.2	2.59	20	
trans-1,3-Dichloropropene	1012	50	1000	0	101	73 - 127	1039	2.6	20	
Trichloroethene	4546	50	1000	3686	86.1	79 - 123	4573	0.581	20	
Trichlorofluoromethane	1013	50	1000	0	101	65 - 141	1014	0.0243	20	
Vinyl chloride	860.2	50	1000	0	86.0	58 - 137	865.2	0.586	20	
Surr: 1,2-Dichloroethane-d4	2035	50	2500	0	81.4	81 - 118	2074	1.92	20	
Surr: 4-Bromofluorobenzene	2550	50	2500	0	102	85 - 114	2472	3.09	20	
Surr: Dibromofluoromethane	2117	50	2500	0	84.7	80 - 119	2130	0.579	20	
Surr: Toluene-d8	2586	50	2500	0	103	89 - 112	2648	2.39	20	

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent
Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW**Batch ID:** R334128 (0)**Instrument:** VOA6**Method:** VOLATILES ORGANICS BY METHOD
8260C

The following samples were analyzed in this batch: HS19030012-01 HS19030012-03

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030012

QC BATCH REPORT NEW

Batch ID: R334254 (0) **Instrument:** UV-2450 **Method:** HEXAVALENT CHROMIUM BY SW7196A

MBLK		Sample ID: MBLK-334254		Units: mg/L		Analysis Date: 01-Mar-2019 11:42			
Client ID:		Run ID: UV-2450_334254		SeqNo: 4986763		PrepDate:		DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Chromium, Hexavalent	0.0100	0.0100							U

LCS		Sample ID: LCS-334254		Units: mg/L		Analysis Date: 01-Mar-2019 11:42			
Client ID:		Run ID: UV-2450_334254		SeqNo: 4986764		PrepDate:		DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Chromium, Hexavalent	0.249	0.0100	0.25	0	99.6	90 - 111			

LCSD		Sample ID: LCSD-334254		Units: mg/L		Analysis Date: 01-Mar-2019 11:42			
Client ID:		Run ID: UV-2450_334254		SeqNo: 4986765		PrepDate:		DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Chromium, Hexavalent	0.241	0.0100	0.25	0	96.4	90 - 111	0.249	3.27	20

The following samples were analyzed in this batch:

ALS Houston, US

Date: 18-Mar-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples	
WorkOrder:	HS19030012	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030012

Date/Time Received: **01-Mar-2019 08:50**
 Received by: **JRM**

Checklist completed by: Jared R. Makan 1-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 1-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.8c/1.8c UC/C IR25
 Cooler(s)/Kit(s): 43795
 Date/Time sample(s) sent to storage: 03/01/2019 10:35

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd., Suite 219 Houston, TX 77099 (281) 530 - 5656 ATTN: RJ Modashia

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001		Analyses										Remarks (Preservatives, etc.)	Lab I.D.#		
Job: GROUNDWATER TREATMENT PLANT MONTHLY EFFLUENT SAMPLES					MS / MSD	No. OF CONTAINERS	VOLATILES	SILVER, SELENIUM, LEAD, BARIUM	HEXAVALENT CHROMIUM	1, 4 - DIOXANE	PERCHLORATE							
Field Sample I.D.	Sample Matrix	Date / Time																
LH18/24-SP650_022819	Water	02/28/19 / 14:00	3	X														HCL
LH18/24-SP650_022819	Water	02/28/19 / 14:00	2			X	X											NONE
LH18/24-SP650_022819_BIX	Water	02/28/19 / 14:00	1							X								NONE
LH18/24-SP650_022819	Water	02/28/19 / 14:00	1		X													HNO3
Trip Blank	Water	02/28/19	2	X														HCL

Additional Remarks: **STANDARD TURN AROUND TIME**

Relinquished By:	Date	Time	Received By:	Date	Time	Relinquished By:	Date	Time	Received By:	Date	Time
<i>Scott Beesinger</i>	02/28/19	14:30	<i>J. W. ...</i>	3/1/19	08:50						

For Lab Use Only									
Received At Lab By:	Date	Time	Airbill No.	Opened By:	Date	Time	Temp of Container	Seal No.	Condition

Remarks: *Cooler 43795 Temp 1.8*
11/25 CFO-D

HS19030012
Bhate Environmental Associates, Inc.
18/24 Longhorn GW Treatment Plant Monthly Effluer



TRK# 4380 9530 9423
0221

FRI - 01 MAR 10:30A
PRIORITY OVERNIGHT

AB SGRA

77099
TX-US
IAH



3646088 01Mar 00:38 AFWH 547C2/0E2D/A17C

 <p>ALS Environmental 10450 Standif. Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887</p>	<p>CUSTOMER</p>
	<p>Date: 2/28/19 Name: [unclear] Company: [unclear]</p>

BODY SEAL	
<p>Weight: 1430 [unclear] [unclear]</p>	<p>Seal Broken By: JM Date: 2/01/19</p>



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1905651; 1906112; 1906330;
1906332; 1906334

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2223 (233911)

General Set Information: There were thirteen field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Method QC data: The method blank (LMB 642099) was less than 1/2 the CRDL. The recovery for the LCS (642100) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.µg/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in µg/L. Results were calculated in µg/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (µg/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level of 4.0µg/L. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported. Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEBI04) along with datafiles 05MARD07-10.

Thomas Bosch March 06, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 07, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1906332**

Project ID: HS19030012

Purchase Order: HS19030012

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP650_022819_BIX	1906332001	02/28/19	03/02/19	



ANALYTICAL REPORT

Workorder: 34-1906332

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP650_022819_BIX	Sampling Site: NA	Collected: 02/28/2019				
Lab ID: 1906332001	Media: 125 mL Nalgene	Received: 03/02/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 12:49	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 233911)

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/06/2019 09:17	/S/ Stephen Brose 03/07/2019 11:04

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als.lt.lab@ALSGlobal.com
Web: www.alssl.com



ANALYTICAL REPORT

Workorder: 34-1906332

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00934680

Analysis Information

Workorder: 1906332

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2223 (HBN: 233911)
Analyzed By: Thomas Bosch

Blank

LMB: 642099 Analyzed: 03/05/2019 09:26 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 642100 Analyzed: 03/05/2019 09:00 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.11	4.00	103	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1906112001 Analyzed: 03/05/2019 09:54 Dilution: 1 Units: ug/L		MS: 1906112002 Analyzed: 03/05/2019 10:07 Dilution: 1 Units: ug/L				MSD: 1906112003 Analyzed: 03/05/2019 10:20 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	51.0	53.9	4	▲ 65.7	78.8 123.8	53.7	▲ 61.2	0.337	0.0 20.0

Continuing Calibration Verification

CCV: 642096 Analyzed: 03/05/2019 08:44 Units: ug/L Criteria: ± 15%			CCV: 642101 Analyzed: 03/05/2019 12:18 Units: ug/L Criteria: ± 15%			CCV: 642102 Analyzed: 03/05/2019 14:07 Units: ug/L Criteria: ± 15%			
Analyte	Result	Target	% Rec.	Result	Target	% Rec.	Result	Target	% Rec.
Perchlorate	25.0	25.0	100	25.0	25.0	100	24.5	25.0	97.8

Interference Check Sample

ICSA: 642098 Analyzed: 03/05/2019 09:13 Units: ug/L Criteria: ± 30%			
Analyte	Result	Target	% Rec.
Perchlorate	3.93	4.00	98.2

Comments

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.



Quality Control Sample Batch Report

00934681

Analysis Information

Workorder: 1906332

Limits: Client SOW/Contract Specified

Preparation: NA

Analysis: EPA 6850, DoD QSM

Basis: DoD QSM

Batch: NA

Batch: ELMS/2223 (HBN: 233911)

Prepared By: NA

Analyzed By: Thomas Bosch

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/06/2019 13:45	/S/ Stephen Brose 03/07/2019 11:04

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



18698/#2

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10854

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

1906332

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com


INVOICE INFORMATION:

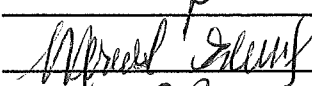
Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030012
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030012-02	LH18/24-SP650_022819_BIX	Water	28 Feb 2019 14:00
SUB_Perch-6850			15 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: 

Received By: 

Cooler ID(s): 9203

Date/Time: 3/1/19 1800

Date/Time: 3/2/19 1903

Temperature(s): 3

ALS Environmental
CHAIN-OF-CUSTODY



Project / Job / Task: HS19030012		Split:	Workorder ID: 1906332	Level: ENV_LVL4	Requested Analysis	
Client: ALS Environmental (Houston)		Account: 8101		Type: 125Poly		
Comments:						
Item	Collect Date/Time	Sample ID	Lab ID	QC	Matrix	
1	02/28/2019 14:00	LH18/24-SP650_022819_BIX	1906332001		Water	
2						
3						
4						
5						
6						
7						
8						
9						
10						

EP A 6850, D+D QSM

Containers	Count
A	1

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY				SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY			
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location	Sample Prep / Analysis for:	Lab Notebook No.:	Prepared / Analyzed by:	Date / Time:
<i>Wafath, Jale</i>	03/02/2019 09:03	ALS Sample Receiving	Sample Login				
<i>Andh W...</i>	<i>03/04/19 16:10</i>	<i>13P</i>	<i>Storage</i>				
<i>R.33.1</i>		<i>T.B. B...</i>	<i>6850</i>				

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: <u>MS Houston</u>		Project/Task/Site: <u>1906332</u>				
Date/Time of Receipt: <u>3/2/17 903</u>		Number of Coolers Received: <u>1</u>				
Condition of Coolers: <u>Acceptable/Unacceptable</u>		Temperature Control: <u>Present/Not Included</u>				
Cooler Custody Seals: <u>Present/Absent/NA</u>		Location Temp Taken: <u>Control/Between Samples</u>				
Container Custody Seals: <u>Present/Absent/NA</u>		Are all temperatures within project specific guidelines? <u>Yes/No/NA</u>				
Ice Present: <u>Yes/No/NA</u>		VOA Headspace Present? <u>Yes/No/NA</u>				
pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9203</u>	<u>3</u> °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: [Signature] [Signature] 3/2/17
Signature Printed Name Date

CLIENT-RELATED INFORMATION

- | | | | |
|---|--|---|--|
| <input type="checkbox"/> Missing Cooler
<input type="checkbox"/> Cooler Conditions
<input type="checkbox"/> Missing Paperwork
<input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Missing Samples/Bottles
<input type="checkbox"/> Broken/Leaking Samples
<input type="checkbox"/> Incorrect Bottle Type
<input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Incorrect Preservation
<input type="checkbox"/> pH Criteria Not Met
<input type="checkbox"/> Residual Chlorine Present
<input type="checkbox"/> Head Space in Bottles | <input type="checkbox"/> Insufficient Sample Volume
<input type="checkbox"/> Chain of Custody Problems
<input type="checkbox"/> Other: |
|---|--|---|--|

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Must Deliver Next Business Day
Time and Tempature Sensitive!

Part # 159469-434 RIT2 EXP 11/19

ORIGIN ID:SGRA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

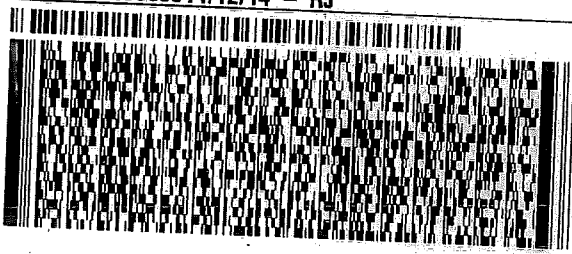
SHIP DATE: 01MAR19
ACTWGT: 8.55 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(801) 266-7700

REF: HS19030011/12/14 - RJ



FedEx
Express



AN L05090211131E

TRK# 4809 7831 2469
0201

SATURDAY 12:00P
PRIORITY OVERNIGHT

XO BTFA

84123
UT-US SLC



FedEx® Saturday Delivery

151956 10/04 MWI

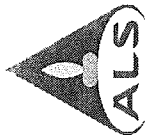


SDR

RY **B639** 1
ST **F1** 12:00



ALS
10450
Houston
Tel. +1
Fax. +1



Batch Worklist

Batch: ELMS/ 2223

Created: 3/5/2019 08:21

Rule: EPA 6850, DoD QSM Water

Instrument:



Status: WP

- Workorder: 1905651 [ENV_LVL4]
- Workorder: 1906112 [ENV_LVL4]
- Workorder: 1906330 [ENV_LVL4]
- Workorder: 1906332 [ENV_LVL4]
- Workorder: 1906334 [ENV_LVL4]

HBN: 233911

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mix	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	642096	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
2	642097	RLVS for HBN 233911 [ELMS/2223]				RLVS	3		E685041C3Q	5311		3/7/2019	
3	642098	ICS for HBN 233911 [ELMS/2223]				ICS	3		E6850.D3Q	5311		3/7/2019	
4	642099	LMB for HBN 233911 [ELMS/2223]				LMB	3		E6850Q413Q	5311		3/7/2019	
5	642100	LCS for HBN 233911 [ELMS/2223]				LCS	3		E6850Q413Q	5311		3/7/2019	
6	1905651001	HS19021158-02/LH18/24-SP650_02				SAMPLE	3	1905651001-A	E6850Q41.3	5480	3/21/2019	3/7/2019	
7	1906112001	EW01_022619				SAMPLE	3	1906112001-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
8	1906112002	EW01_022619MS				MS	3	1906112002-A	E6850Q413Q	5480		3/7/2019	
9	1906112003	EW01_022619MSD				MSD	3	1906112003-A	E6850Q413Q	5480		3/7/2019	
10	1906112004	EW05_022619				SAMPLE	3	1906112004-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
11	1906112005	EW05_022619_FD				FLDDUP	3	1906112005-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
12	1906112006	EW02_022619				SAMPLE	3	1906112006-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
13	1906112007	EW06_022619				SAMPLE	3	1906112007-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
14	1906112008	EW03_022619				SAMPLE	3	1906112008-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
15	1906112009	EW07_022619				SAMPLE	3	1906112009-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
16	642101	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
17	1906112010	EW04_022619				SAMPLE	3	1906112010-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
18	1906112011	EW08_022619				SAMPLE	3	1906112011-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
19	1906330001	LH18/24-SP140_022819				SAMPLE	3	1906330001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
20	1906332001	LH18/24-SP650_022819_BIX				SAMPLE	3	1906332001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
21	1906334001	LH18-24-SP650_022819_BIX				SAMPLE	3	1906334001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
22	642102	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1905651 (001); 1906112 (001-11); 1906330 (001); 1906332 (001); 1906334 (001)
 ELMS Batch/HBN ID: 2223 (233911)
 Prep Date: 03/04/2019 Analysis Date: 03/05/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\05MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by TNB. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 02/15/2019, sequence 15FEB19D.s Offline Quantitation Method: CLO4-DP1.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 3 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 642100; Target = 4.0µg/L. ASTM type II water was used for LMB 642099.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\23911-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEB104) along with datafiles 05MARD07-10.

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
Batch(es)/SDG: <u>E LMS: 2223 HBN: 233911</u>		
Sample Set IDs if Applicable: <u>1905651 / 190912 / 1906330</u> <u>1906332 / 1906334</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850.WKG Std 100.ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK			Description - 6850 Stock AccStd 1,000ug/mL
Standard: 43659		Created By: Thomas Bosch	
MFG: AccuStandard		Create Date: 09/17/2018 09:09AM	
MFG Lot: 218065075		Amount: 100 mL	
Part ID: IC-PER-10X-1		Expires: 07/25/2020	
		Usable: No	
		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT			Description - 6850 Intermdt AccStd 10.ug/mL		
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK			Description: 6850 QC WKG STD 100ug/L		
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdf Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type: II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK			Description: -6850 QC Stock STD 1,000ug/mL
Standard: 36748		Created By: Thomas Bosch	
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	
MFG Lot: CP-0860		Amount: 100 mL	
Part ID: ICC-013		Expires: 03/31/2020	
		Usable: Yes	
		Lab Lot: CLO4 QC STOCK	
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730		Created By: Thomas Bosch		Amount: 25 mL	
MFG: ALS/SLC		Create Date: 09/20/2018 09:09AM		Expires: 09/20/2019	
MFG Lot: TNB: 05/09/2018		Verified By: Thomas Bosch		Usable: Yes	
Pipette ID: Not Provided		Verify Date:		Lab Lot: CLO4ISTDWRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	.CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

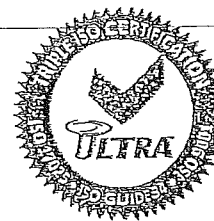
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:
This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:
Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:
The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:
This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:
This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:
Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

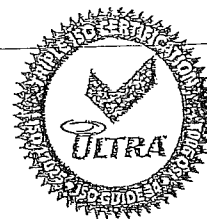
Hazards:
Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:
The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis

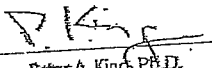


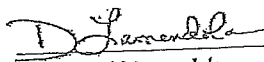
ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:
The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QAVRA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleared Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

For use in routine laboratory analysis.



Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:

ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: NaCl*O4

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 µg/mL
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 ± 2.8 µg/mL (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount	
*	642096	CCV@25	Vial 71	1	Control	1	2.57589e6	8.017	25.04585
*	642100	QC@4.0	Vial 72	1	Control	2	4.23307e5	8.139	4.10749
*	642098	ICS@4.0	Vial 73	1	Control	3	3.17619e5	7.883	3.92708
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	7.71127e6	7.438	51.25151
*	1906112002	MS	Vial 77	1	Sample	7	8.02696e6	7.464	53.87972
*	1906112003	MSD	Vial 78	1	Sample	8	7.94242e6	7.442	53.69940
*	1906112004		Vial 79	1	Sample	9	2.33017e7	7.227	172.36522
*	1906112005		Vial 80	1	Sample	10	2.30991e7	7.229	180.38568
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.33611e6	8.114	25.01987
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	2.53126e6	7.736	226.48155
*	1906112005	10X	Vial 91	1	Sample	22	2.68237e6	7.739	241.03710
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	3.77363e6	8.090	3708.65666
*	642102	CCV@25	Vial 71	1	Control	25	2.15787e6	8.144	24.45533

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount	
*	642096	CCV@25	Vial 71	1	Control	1	6.79028e5	8.036	25.15984
*	642100	QC@4.0	Vial 72	1	Control	2	1.27412e5	8.156	4.45187
*	642098	ICS@4.0	Vial 73	1	Control	3	9.68721e4	7.906	4.29888
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	2.02786e6	7.452	51.10329
*	1906112002	MS	Vial 77	1	Sample	7	2.14637e6	7.478	54.46797
*	1906112003	MSD	Vial 78	1	Sample	8	2.10991e6	7.456	53.98100
*	1906112004		Vial 79	1	Sample	9	6.79668e6	7.243	180.45962
*	1906112005		Vial 80	1	Sample	10	6.82834e6	7.246	190.34382
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	6.28961e5	8.128	25.63410
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	6.46870e5	7.754	220.91905
*	1906112005	10X	Vial 91	1	Sample	22	6.95643e5	7.756	238.44674
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	1.01021e6	8.102	3771.67475
*	642102	CCV@25	Vial 71	1	Control	25	5.83711e5	8.160	25.16652

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount	
*	642096	CCV@25	Vial 71	1	Control	1	3.12207e5	8.043	5.00000
*	642100	QC@4.0	Vial 72	1	Control	2	3.41038e5	8.157	5.00000
*	642098	ICS@4.0	Vial 73	1	Control	3	2.68237e5	7.902	5.00000
*	642099	LMB	Vial 74	1	Control	4	3.53313e5	8.102	5.00000
*	1905651001		Vial 75	1	Sample	5	3.26356e5	7.780	5.00000
*	1906112001		Vial 76	1	Sample	6	4.26473e5	7.464	5.00000
*	1906112002	MS	Vial 77	1	Sample	7	4.19549e5	7.491	5.00000
*	1906112003	MSD	Vial 78	1	Sample	8	4.16709e5	7.467	5.00000
*	1906112004		Vial 79	1	Sample	9	2.95705e5	7.253	5.00000
*	1906112005		Vial 80	1	Sample	10	2.75946e5	7.263	5.00000
*	1906112006		Vial 81	1	Sample	11	3.73575e5	7.480	5.00000
*	1906112007		Vial 82	1	Sample	12	5.00533e5	7.256	5.00000
*	1906112008		Vial 83	1	Sample	13	4.65121e5	7.237	5.00000
*	1906112009		Vial 84	1	Sample	14	2.43675e5	7.672	5.00000
*	1906112010		Vial 85	1	Sample	15	2.51865e5	7.710	5.00000
*	1906112011		Vial 86	1	Sample	16	2.80792e5	7.876	5.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.83460e5	8.140	5.00000
*	1906332001		Vial 88	1	Sample	19	2.64674e5	7.827	5.00000
*	1906334001		Vial 89	1	Sample	20	2.65662e5	7.808	5.00000
*	1906112004	10X	Vial 90	1	Sample	21	3.41608e5	7.756	50.00000
*	1906112005	10X	Vial 91	1	Sample	22	3.38724e5	7.764	50.00000
*	1906112007	RE	Vial 82	1	Sample	23	4.31135e5	7.332	5.00000
*	1906330001	100	Vial 92	1	Sample	24	2.98985e5	8.111	500.00000
*	642102	CCV@25	Vial 71	1	Control	25	2.68305e5	8.165	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

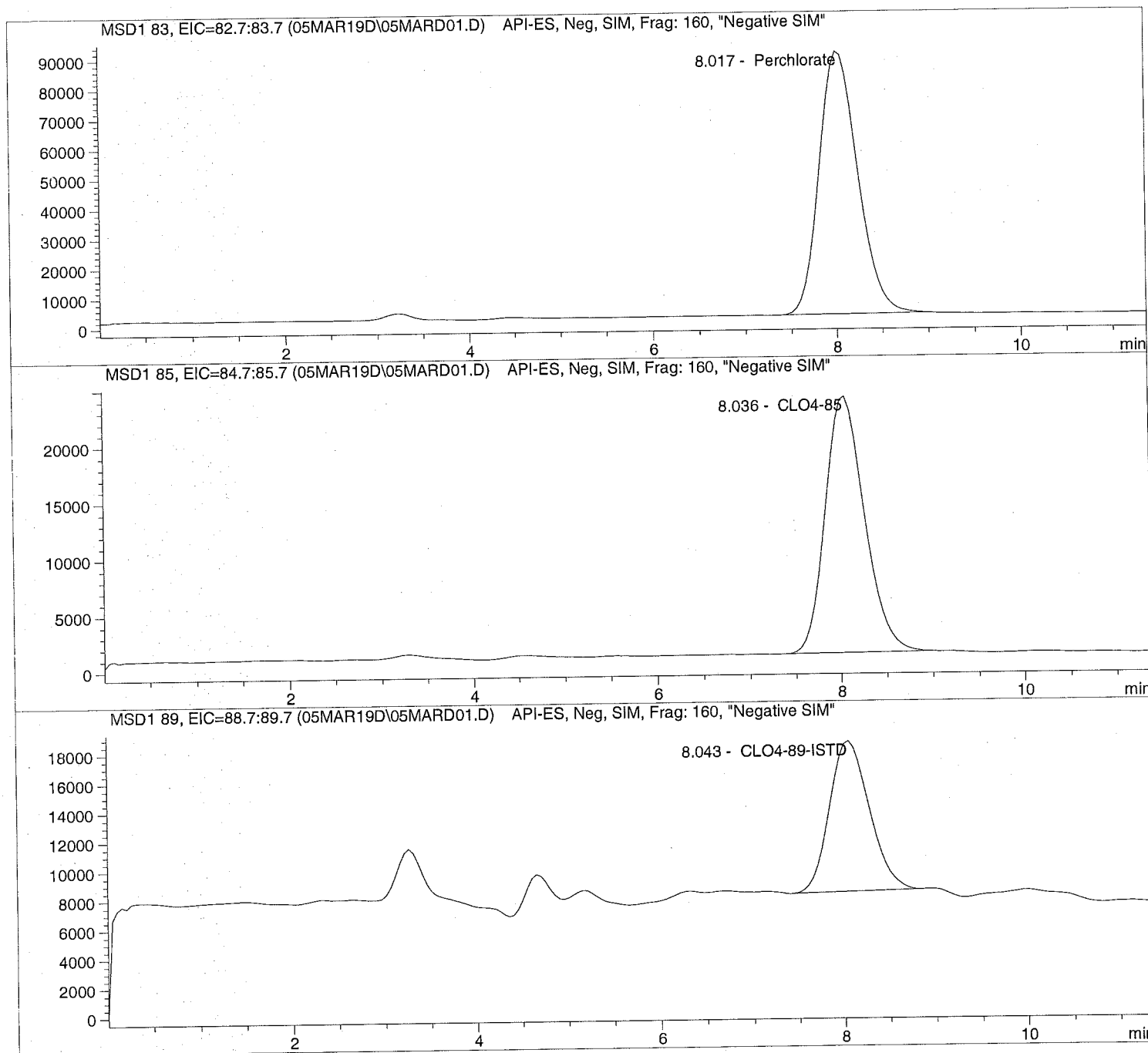
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	642096	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	642100	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	642098	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	642099	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1905651001		CLO4-AQN	1	Sample	
6	Vial 76	1906112001		CLO4-AQN	1	Sample	
7	Vial 77	1906112002	MS	CLO4-AQN	1	Sample	
8	Vial 78	1906112003	MSD	CLO4-AQN	1	Sample	
9	Vial 79	1906112004		CLO4-AQN	1	Sample	
10	Vial 80	1906112005		CLO4-AQN	1	Sample	
11	Vial 81	1906112006		CLO4-AQN	1	Sample	
12	Vial 82	1906112007		CLO4-AQN	1	Sample	
13	Vial 83	1906112008		CLO4-AQN	1	Sample	
14	Vial 84	1906112009		CLO4-AQN	1	Sample	
15	Vial 85	1906112010		CLO4-AQN	1	Sample	
16	Vial 86	1906112011		CLO4-AQN	1	Sample	
17	Vial 71	642101	CCV@25	CLO4-AQN	1	Ctrl Samp	
18	Vial 87	1906330001	1K	CLO4-AQN	1	Sample	
19	Vial 88	1906332001		CLO4-AQN	1	Sample	
20	Vial 89	1906334001		CLO4-AQN	1	Sample	
21	Vial 90	1906112004	10X	CLO4-AQN	1	Sample	
22	Vial 91	1906112005	10X	CLO4-AQN	1	Sample	
23	Vial 82	1906112007	RE	CLO4-AQN	1	Sample	
24	Vial 92	1906330001	100	CLO4-AQN	1	Sample	
25	Vial 71	642102	CCV@25	CLO4-AQN	1	Ctrl Samp	

Injection Date: 3/05/2019 08:44:45
Sample Name: 642096 CCV@25
Acq Operator: TNB

Seq Line: 1
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 08:44:45 Seq Line: 1
Sample Name: 642096 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.017	PBA	2575886.3	25.0459	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.036	PBA	679028.4	25.1598	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.043	PBA	312206.9	5.0000	CLO4-89-ISTD

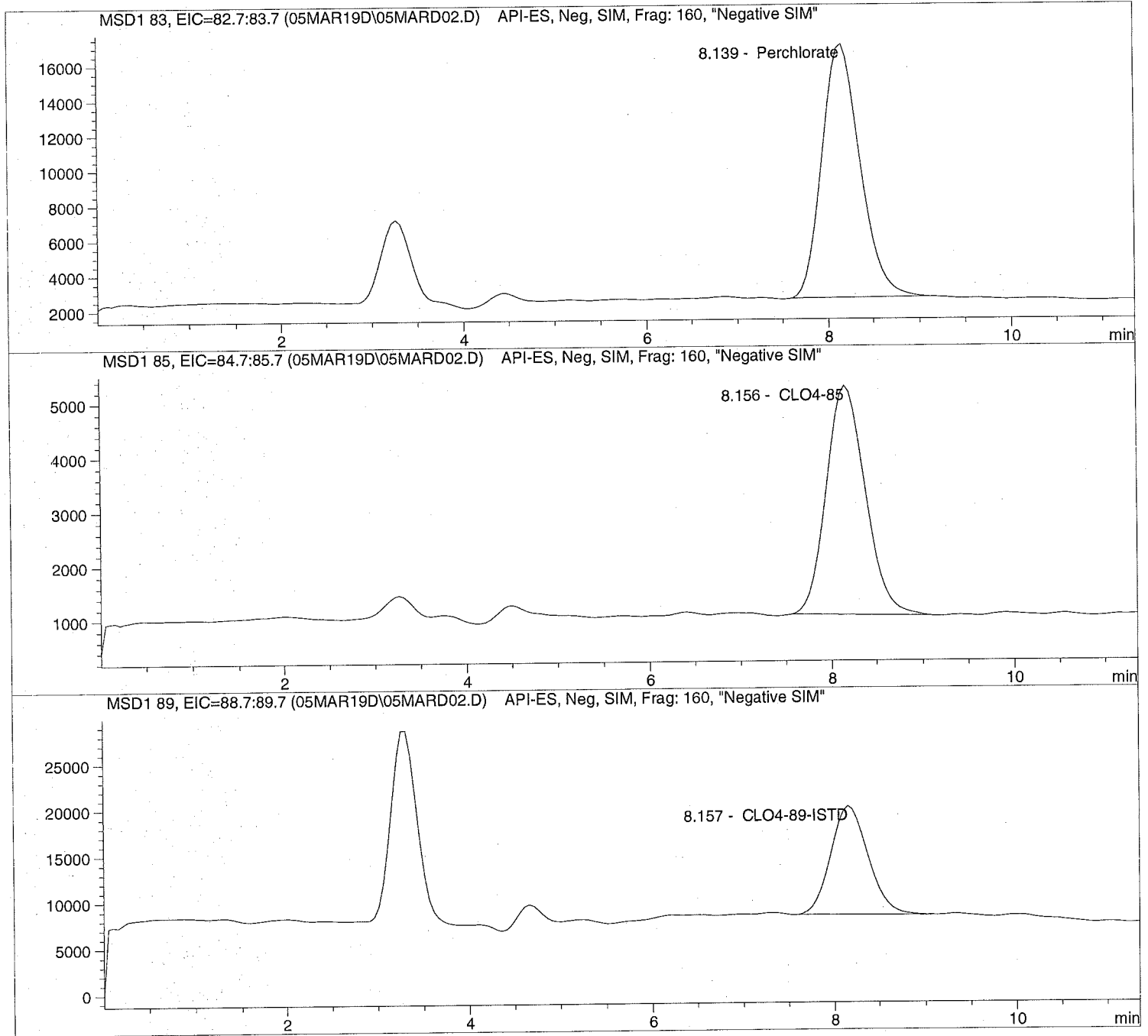
*** End of Report ***

Injection Date: 3/05/2019 09:00:30
Sample Name: 642100 QC@4.0
Acq Operator: TNB

Seq Line: 2
Location: Vial 72
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 09:00:30 Seq Line: 2
Sample Name: 642100 QC@4.0 Location: Vial 72
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.139	PBA	423307.1	4.1075	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.156	PBA	127412.0	4.4519	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.157	PBA	341038.3	5.0000	CLO4-89-ISTD

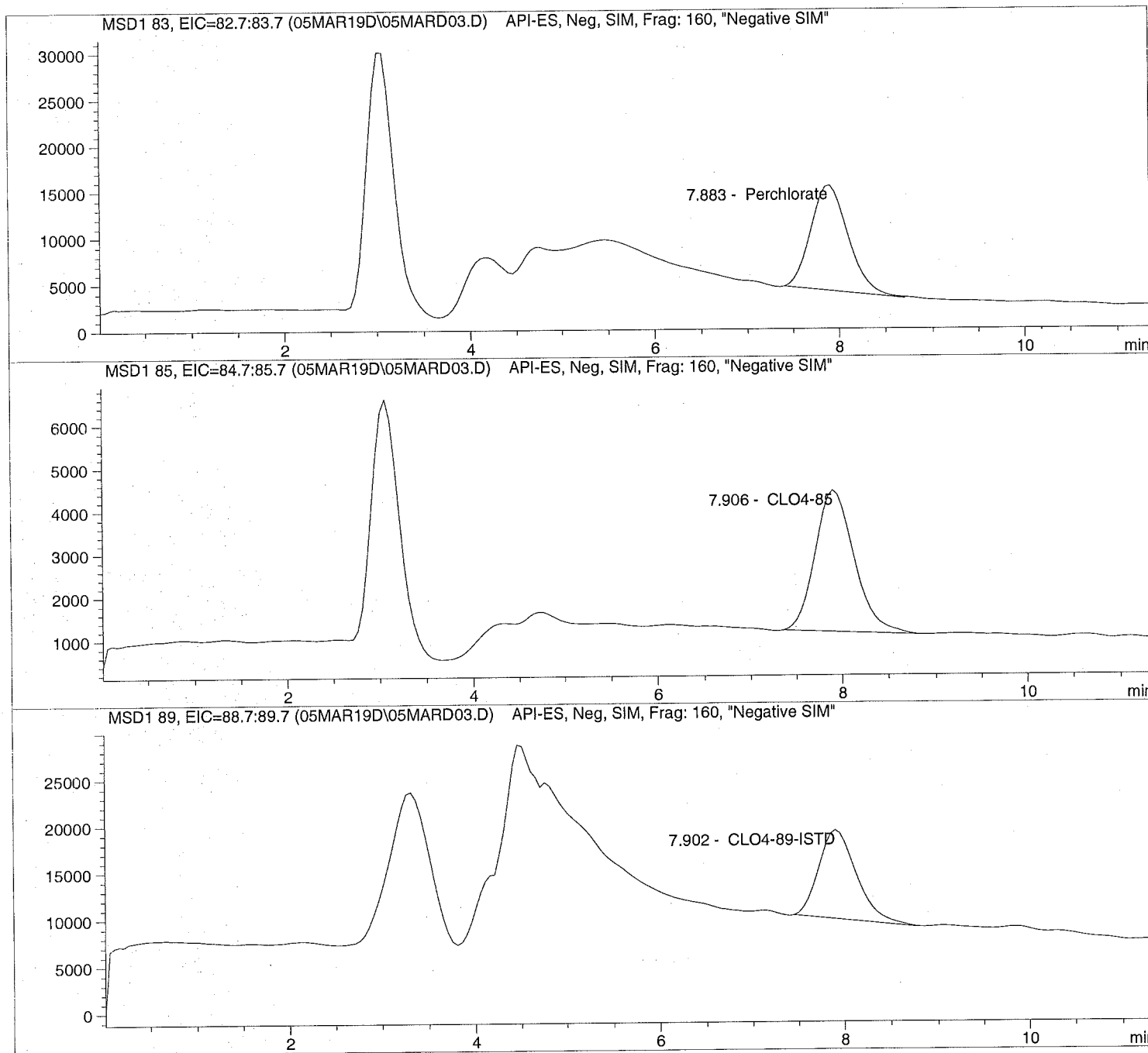
*** End of Report ***

Injection Date: 3/05/2019 09:13:34
Sample Name: 642098 ICS@4.0
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 09:13:34      Seq Line: 3  
Sample Name: 642098 ICS@4.0            Location: Vial 73  
Acq Operator: TNB                       Inj. No.: 1  
                                           Inj. Vol.: 20 µl
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 4.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.883	PBA	317618.9	3.9271	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.906	PBA	96872.1	4.2989	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.902	PBA	268236.7	5.0000	CLO4-89-ISTD

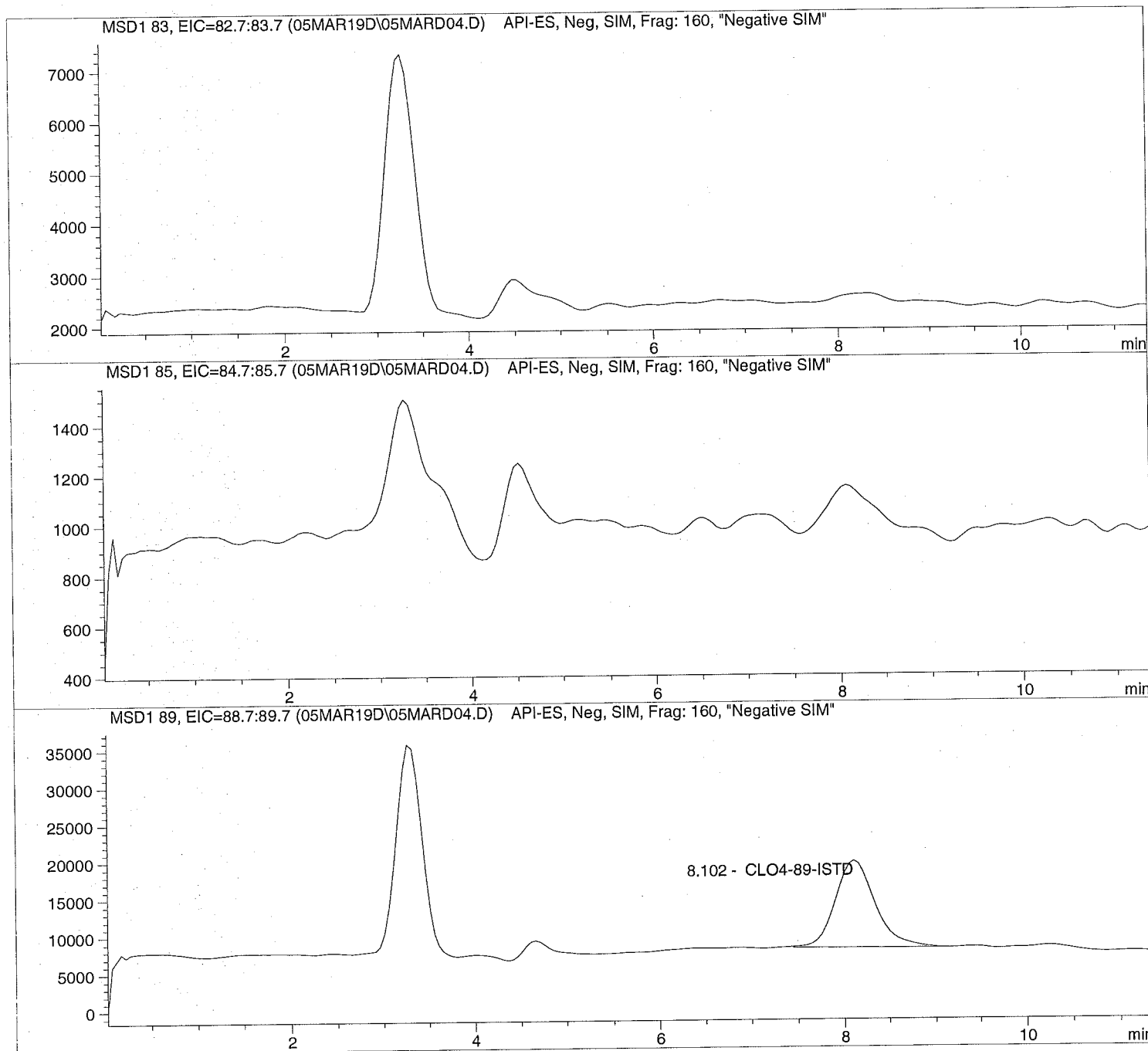
=====
*** End of Report ***
=====

Injection Date: 3/05/2019 09:26:40
Sample Name: 642099 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 09:26:40      Seq Line: 4  
Sample Name: 642099 LMB                  Location: Vial 74  
Acq Operator: TNB                        Inj. No.: 1  
                                           Inj. Vol.: 20 µl  
=====
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	BBA	353313.1	5.0000	CLO4-89-ISTD

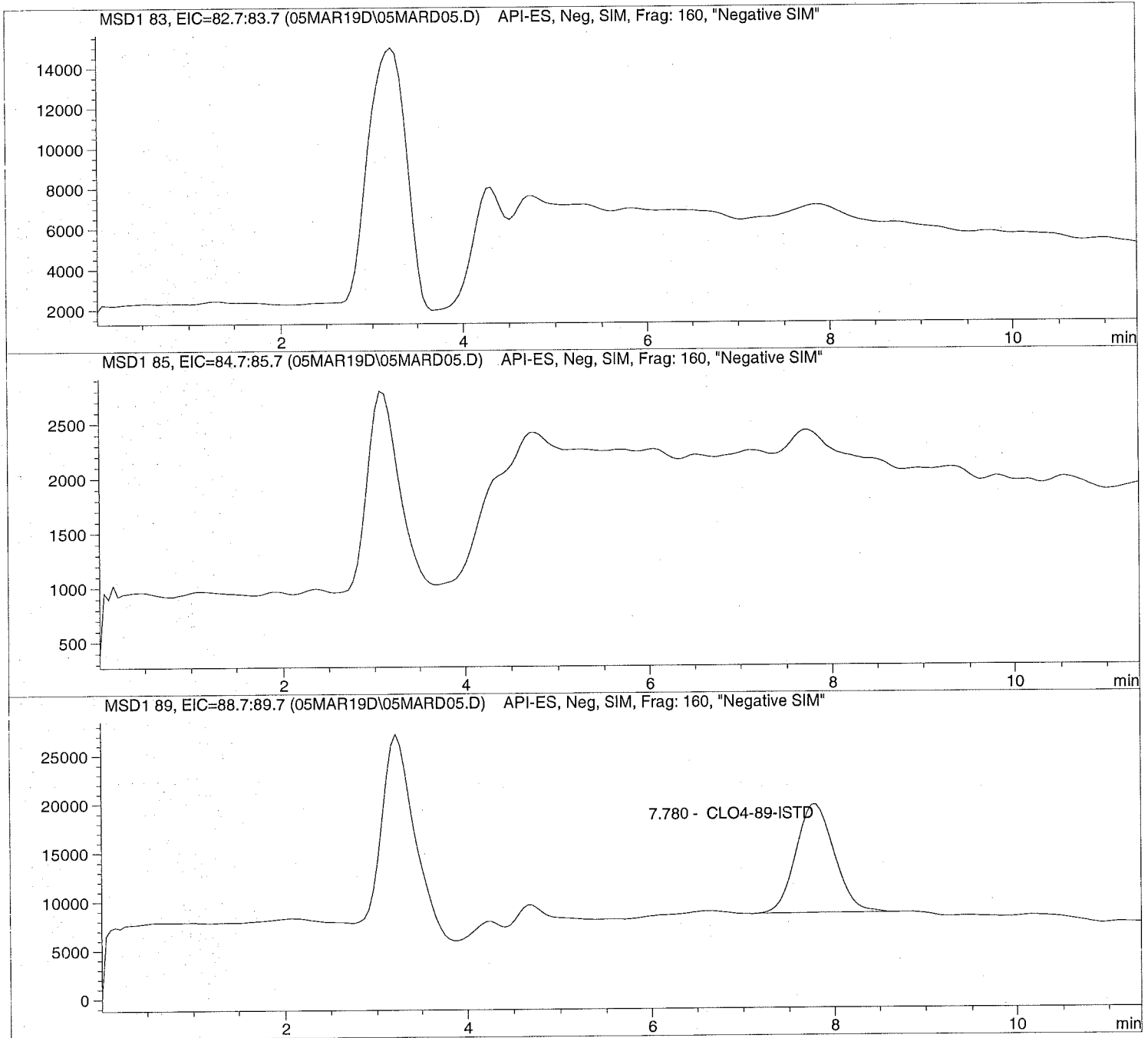
=====
*** End of Report ***

Injection Date: 3/05/2019 09:40:58
Sample Name: 1905651001
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 09:40:58      Seq Line: 5
Sample Name: 1905651001                 Location: Vial 75
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.780	PBA	326356.2	5.0000	CLO4-89-ISTD

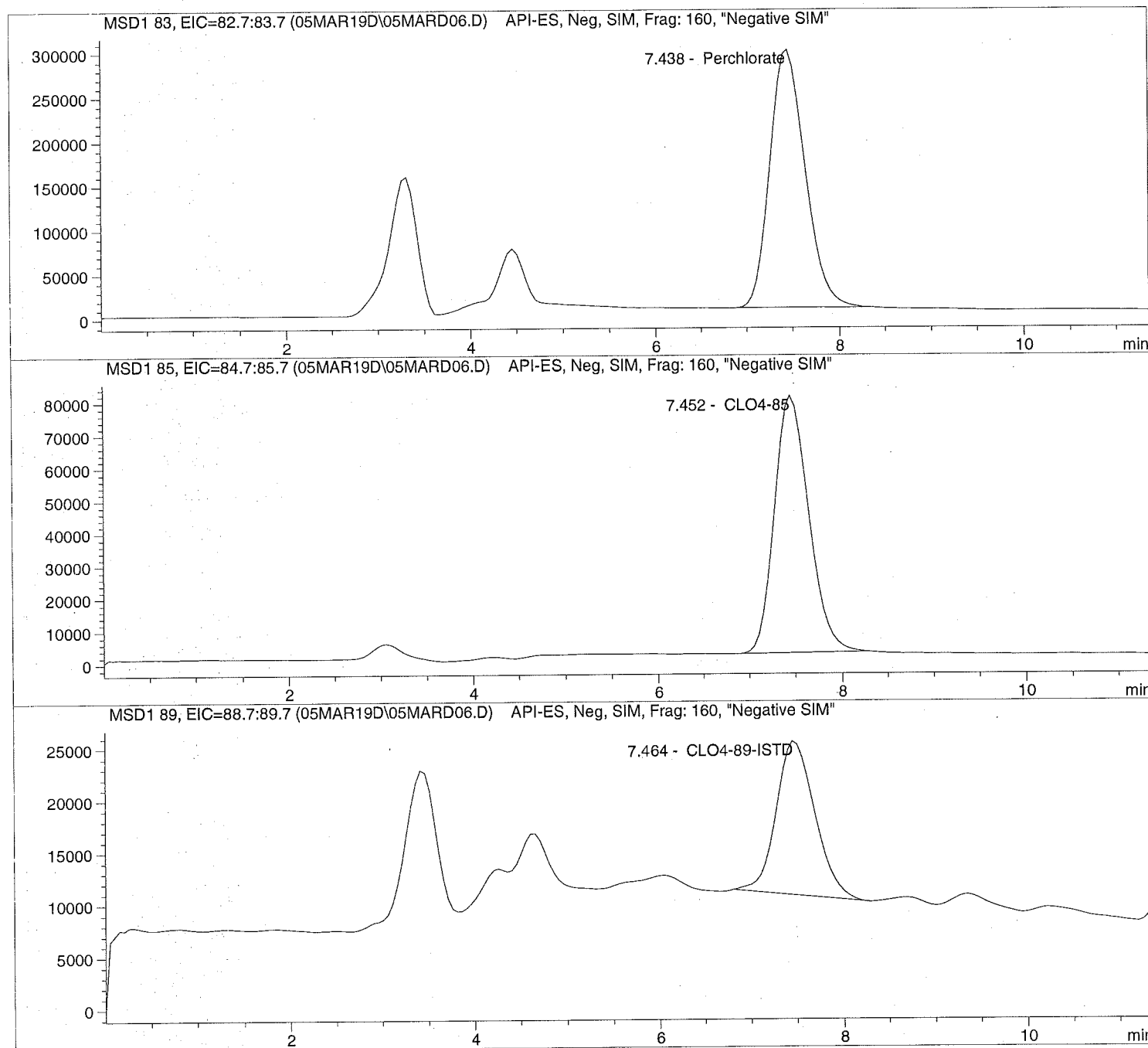
=====
*** End of Report ***

Injection Date: 3/05/2019 09:54:04
Sample Name: 1906112001
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 09:54:04      Seg Line: 6  
Sample Name: 1906112001                 Location: Vial 76  
Acq Operator: TNB                       Inj. No.: 1  
                                           Inj. Vol.: 20 µl  
=====
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.438	PBA	7711270.5	51.2515	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.452	PBA	2027855.1	51.1033	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PB	426473.5	5.0000	CLO4-89-ISTD

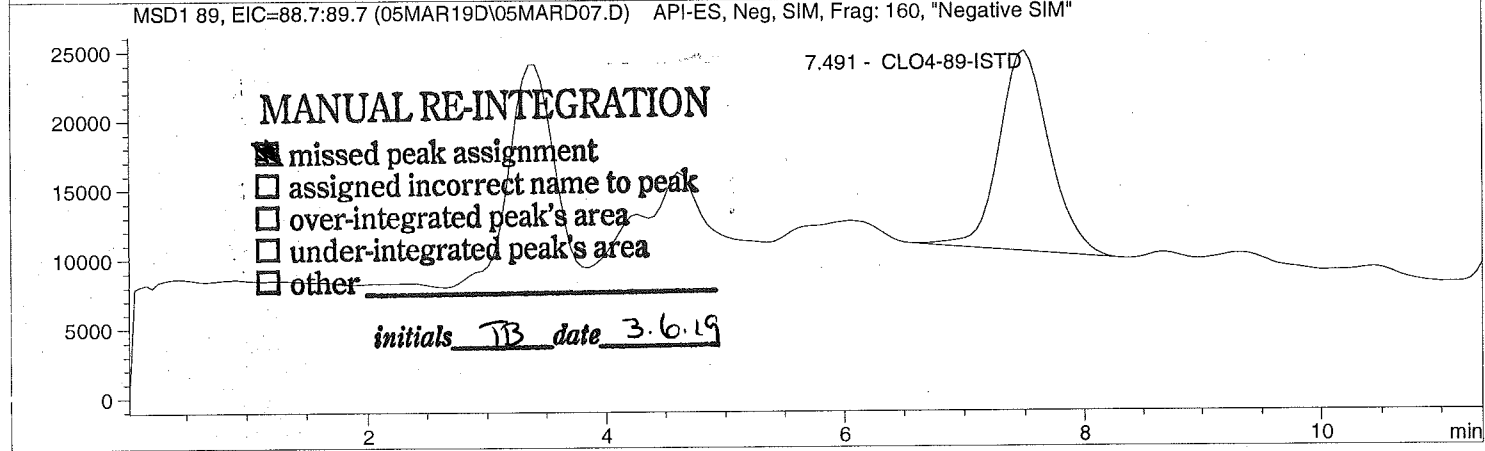
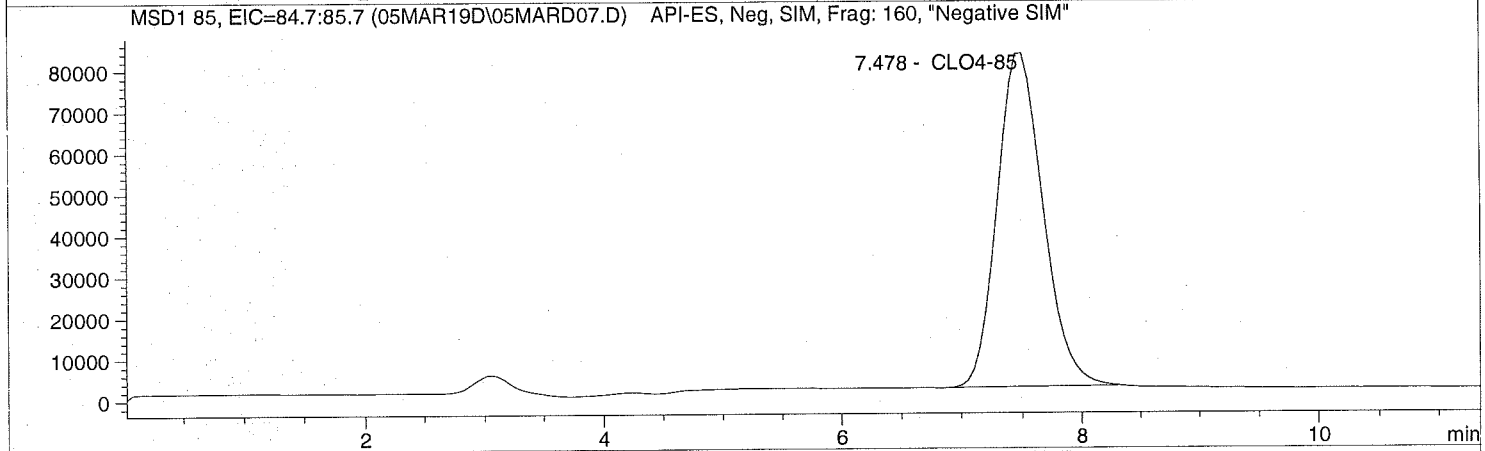
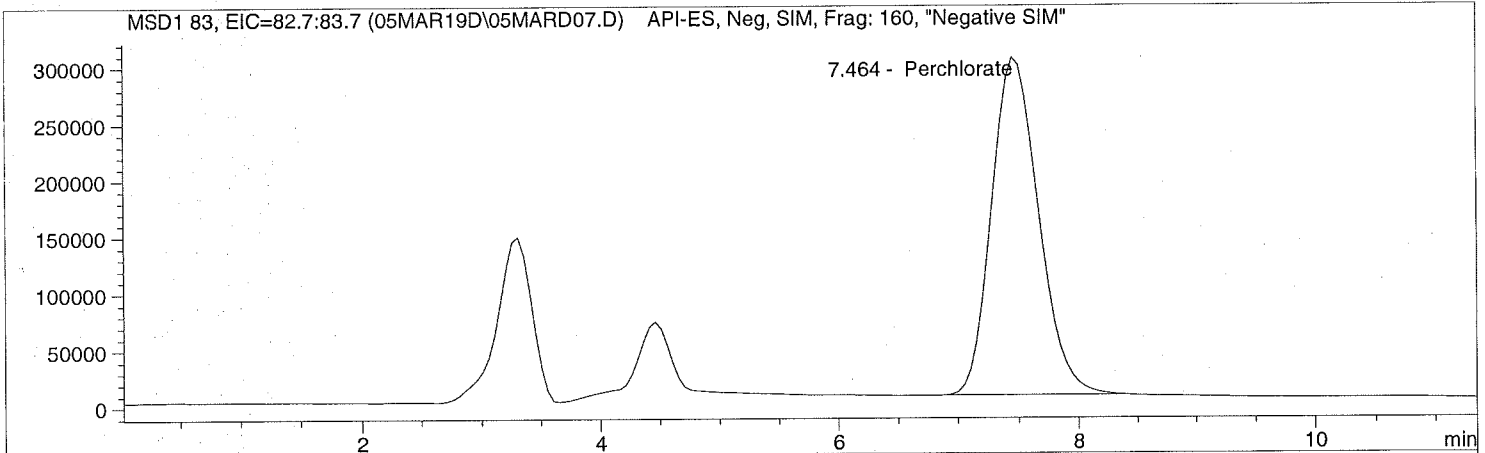
=====
*** End of Report ***
=====

Injection Date: 3/05/2019 10:07:11
Sample Name: 1906112002 MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 10:07:11      Seq Line: 7
Sample Name: 1906112002 MS              Location: Vial 77
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	53.8797	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	54.4680	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.491	MM	419549.0	5.0000	CLO4-89-ISTD

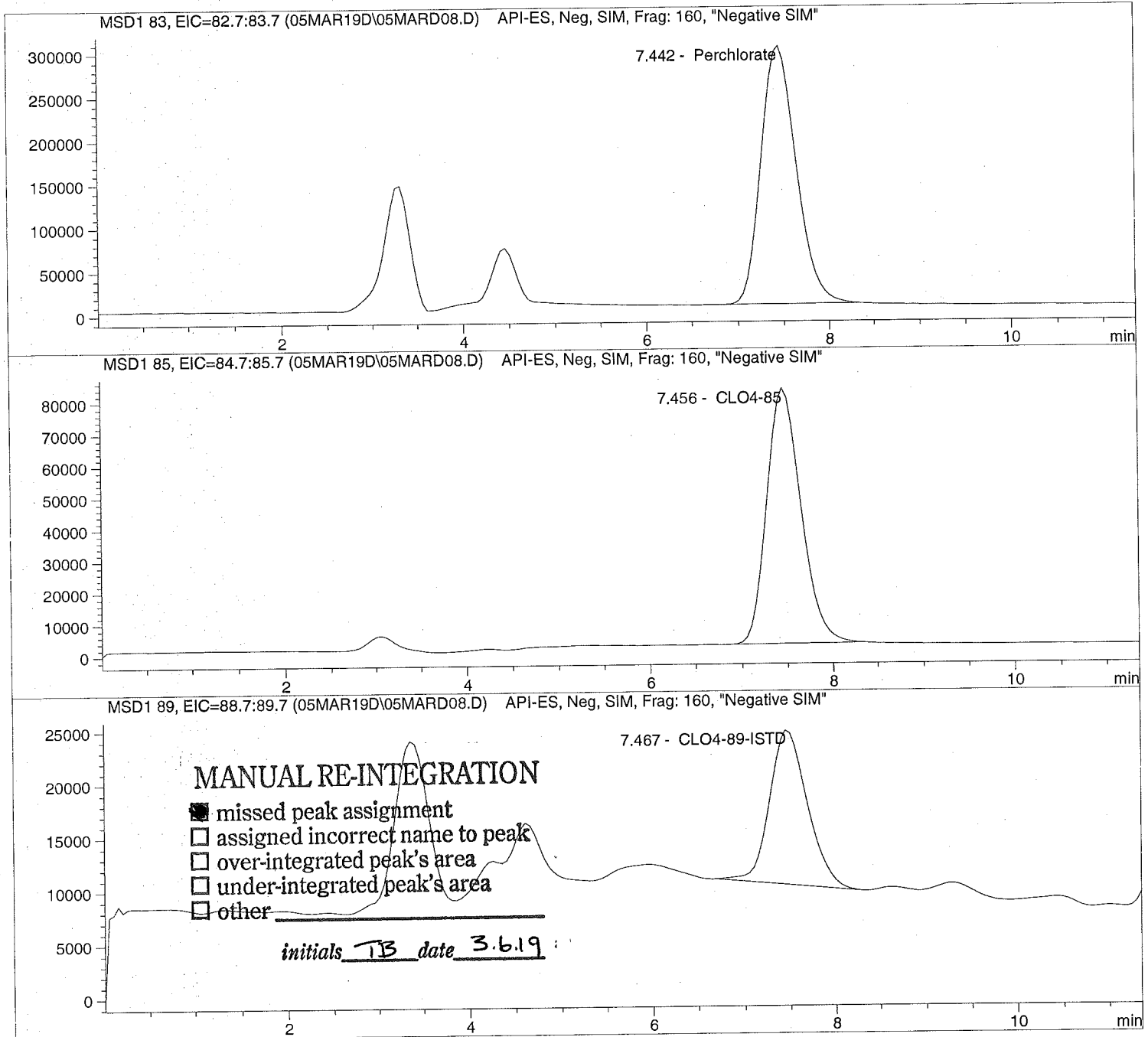
=====
*** End of Report ***

Injection Date: 3/05/2019 10:20:17
Sample Name: 1906112003 MSD
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	53.6994	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	53.9810	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.467	MM	416709.1	5.0000	CLO4-89-ISTD

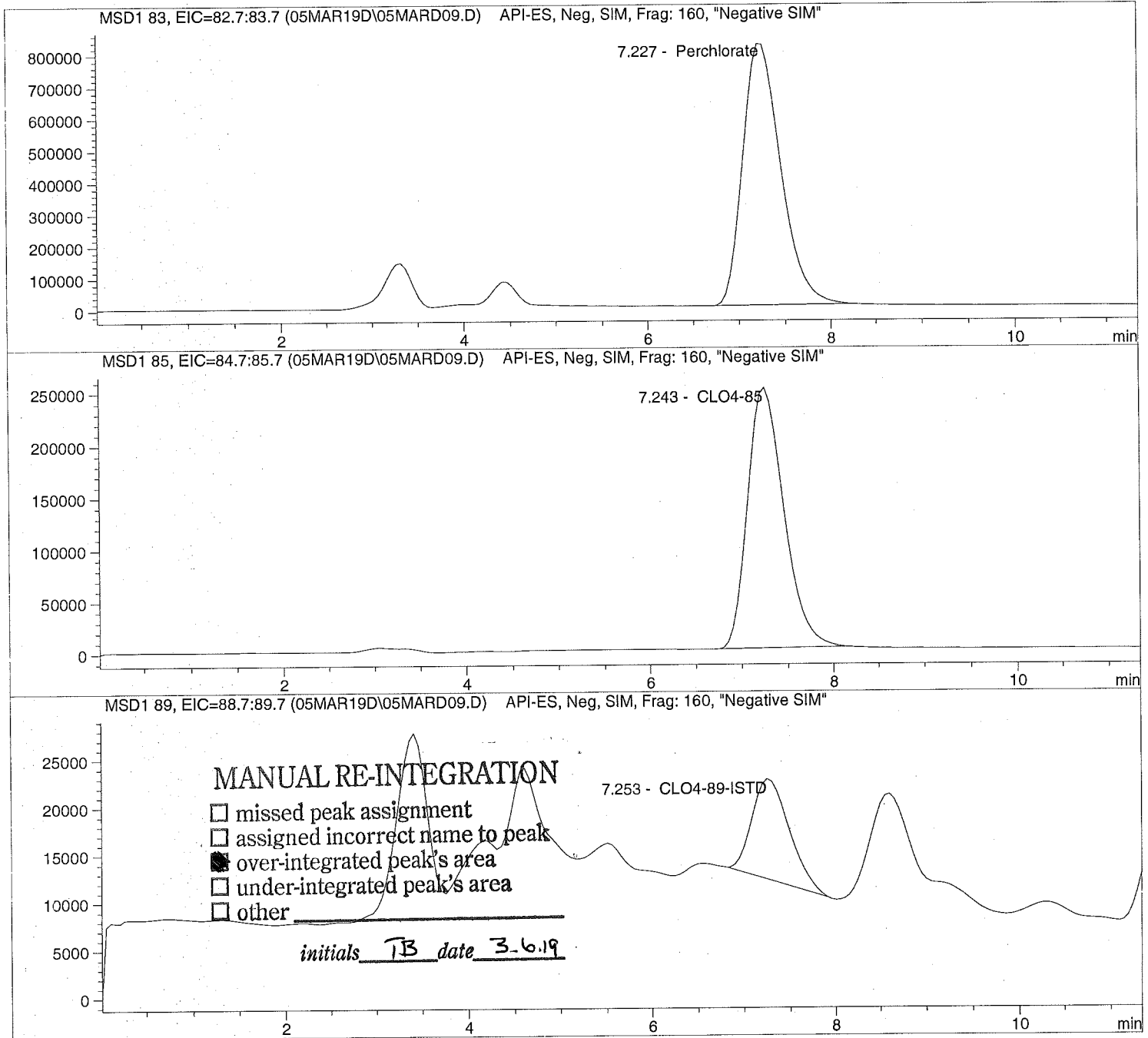
*** End of Report ***

Injection Date: 3/05/2019 10:33:21
Sample Name: 1906112004
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 10:33:21      Seq Line:          9
Sample Name:    1906112004              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	172.3652	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	180.4596	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.253	MM	295705.2	5.0000	CLO4-89-ISTD

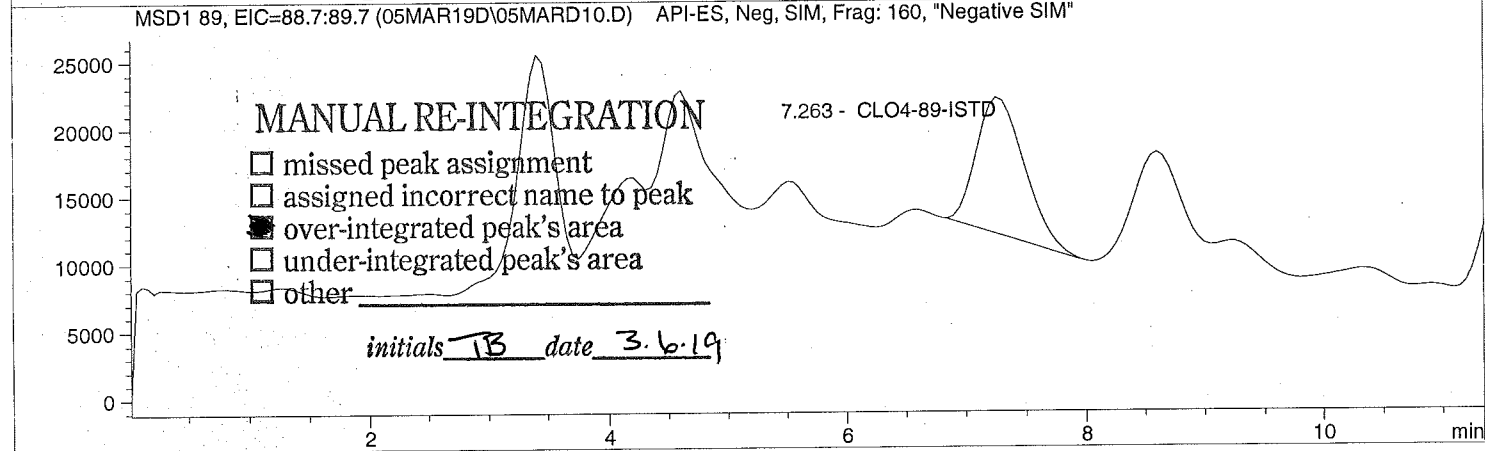
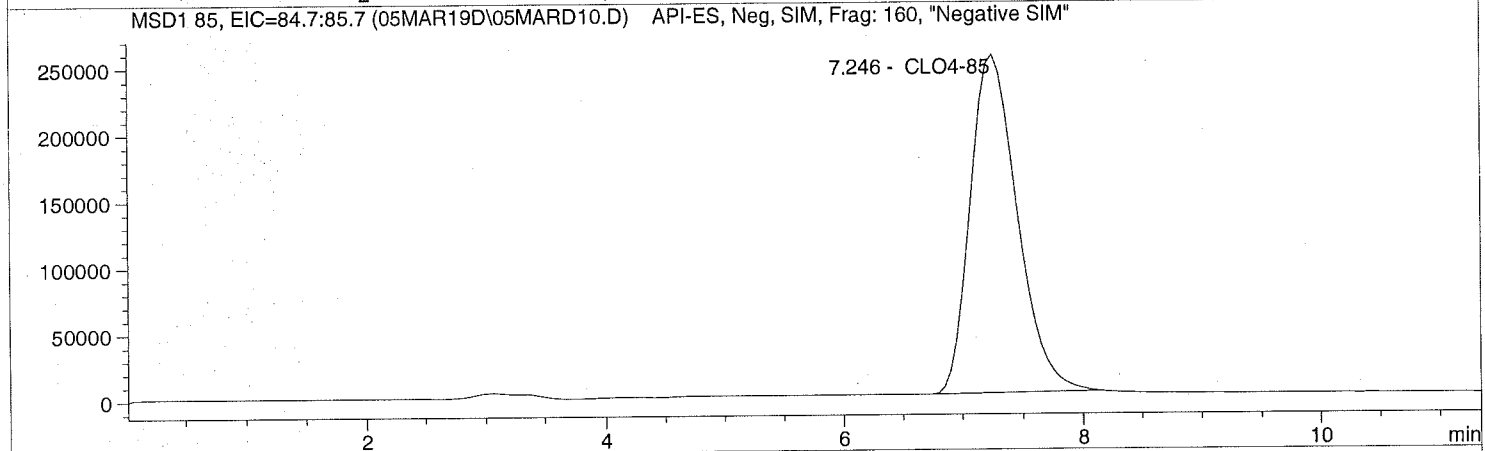
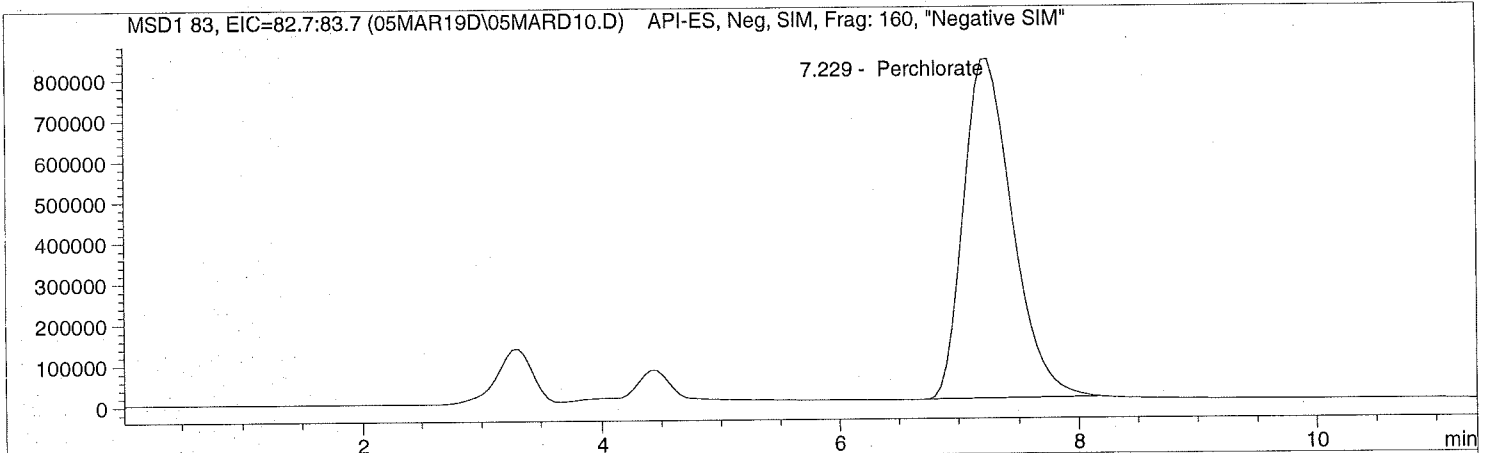
=====
*** End of Report ***

Injection Date: 3/05/2019 10:46:26
Sample Name: 1906112005
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```

=====
Injection Date: 3/05/2019 10:46:26      Seq Line:      10
Sample Name:    1906112005              Location:      Vial 80
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	180.3857	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	190.3438	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	MM	275946.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

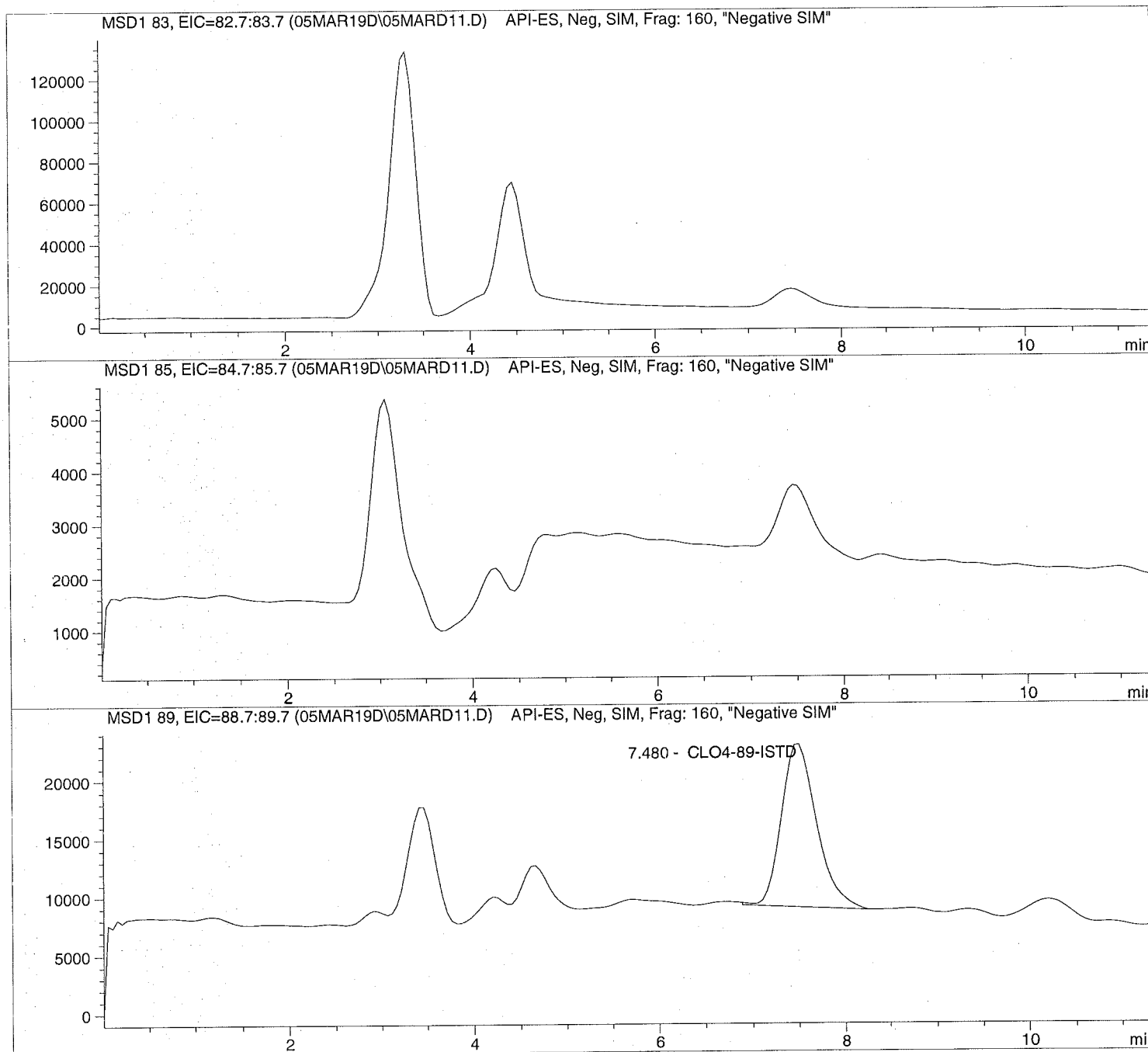
```

Injection Date: 3/05/2019 10:59:36
Sample Name: 1906112006
Acq Operator: TNB

Seq Line: 11
Location: Vial 81
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:59:36 Seq Line: 11
Sample Name: 1906112006 Location: Vial 81
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.480	BBA	373575.4	5.0000	CLO4-89-ISTD

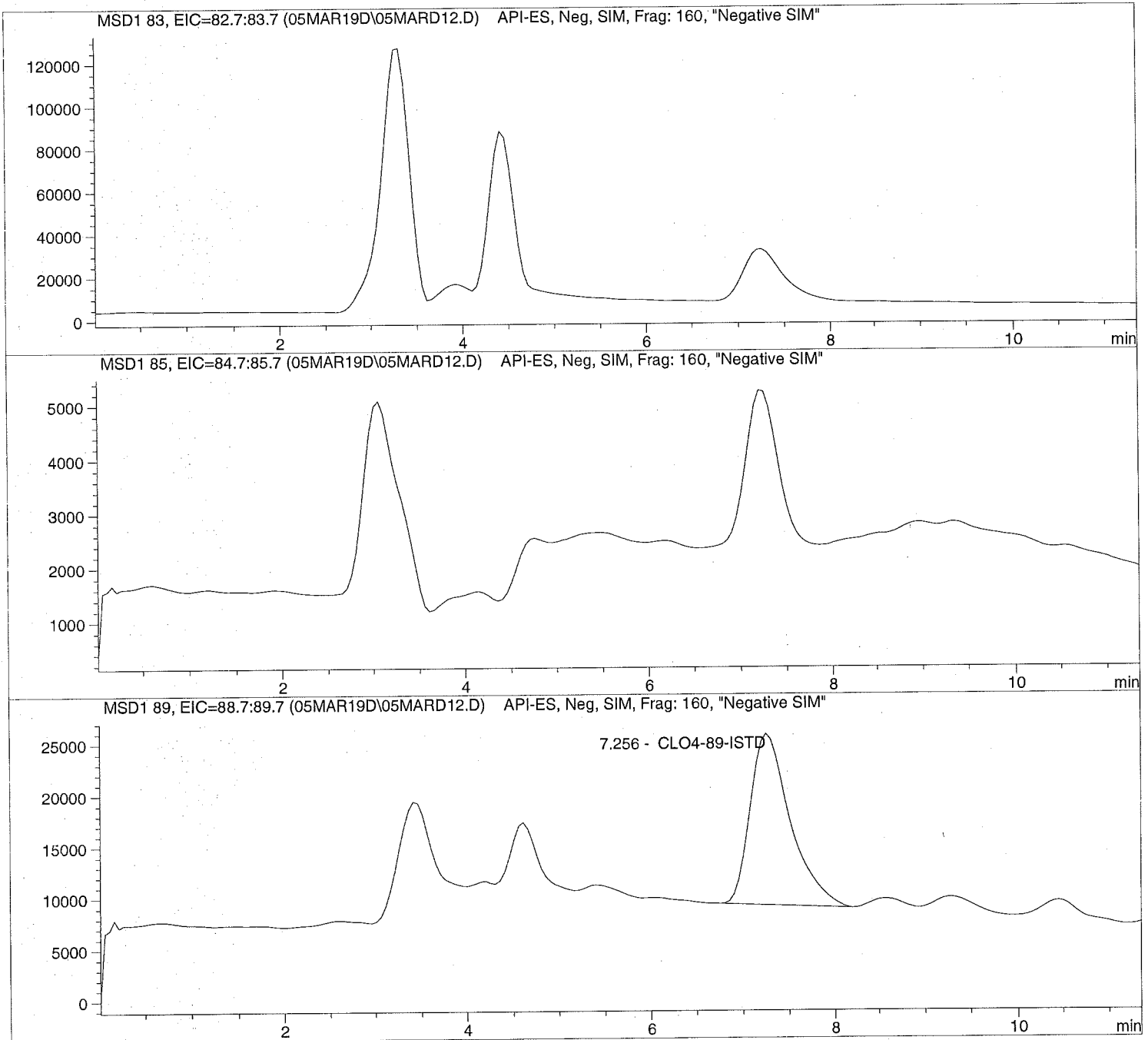
*** End of Report ***

Injection Date: 3/05/2019 11:12:28
Sample Name: 1906112007
Acq Operator: TNB

Seq Line: 12
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 11:12:28      Seq Line: 12  
Sample Name: 1906112007                Location: Vial 82  
Acq Operator: TNB                      Inj. No.: 1  
                                         Inj. Vol.: 20 µl
```

```
Acq. Method: CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By: Signal  
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am  
Multiplier: 1.000000  
Dilution: 1.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.256	PB	500532.9	5.0000	CLO4-89-ISTD

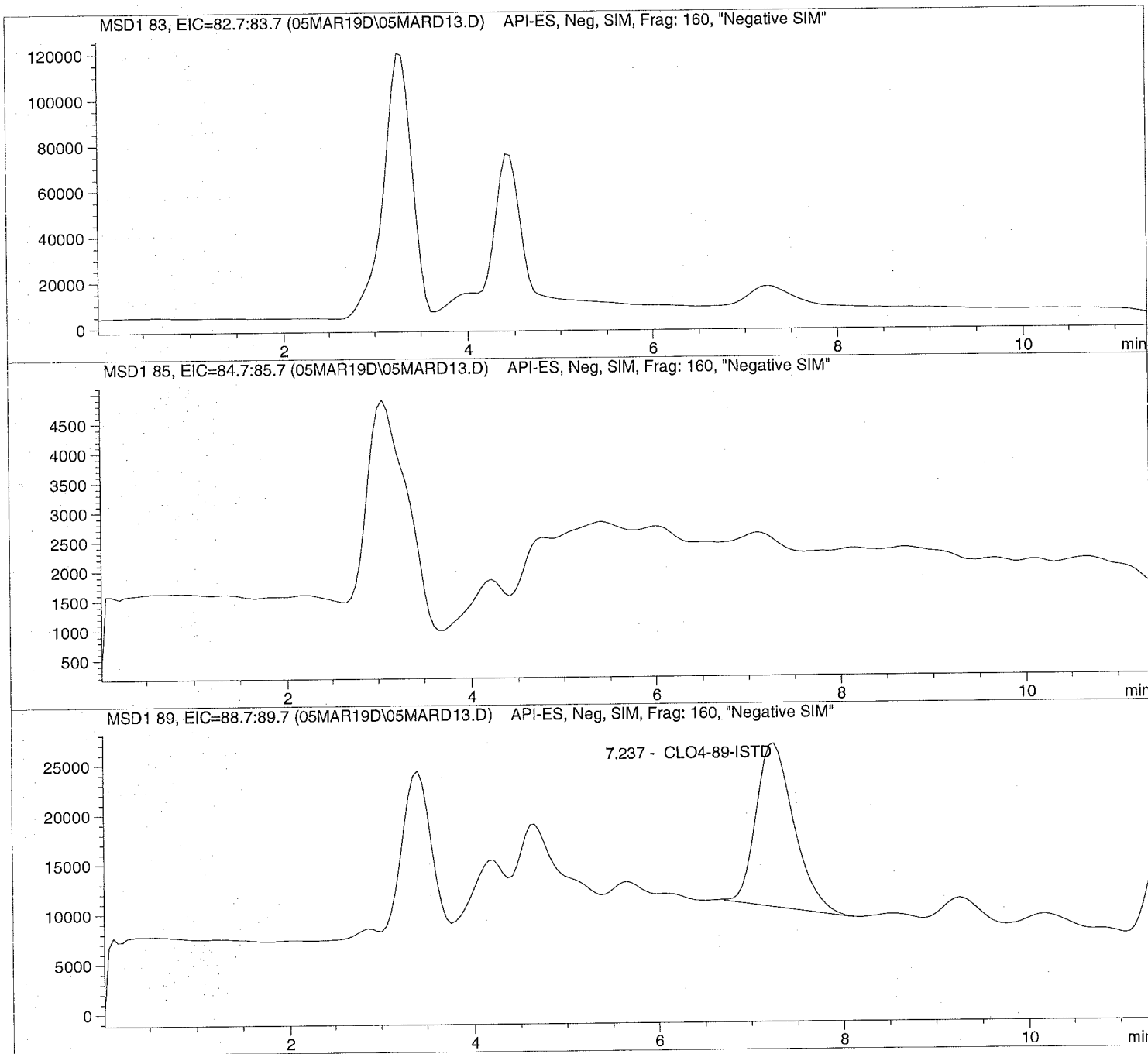
=====
*** End of Report ***

Injection Date: 3/05/2019 11:26:23
Sample Name: 1906112008
Acq Operator: TNB

Seq Line: 13
Location: Vial 83
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```

=====
Injection Date:  3/05/2019  11:26:23      Seq Line:      13
Sample Name:    1906112008                Location:      Vial 83
Acq Operator:   TNB                      Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.237	BBA	465121.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

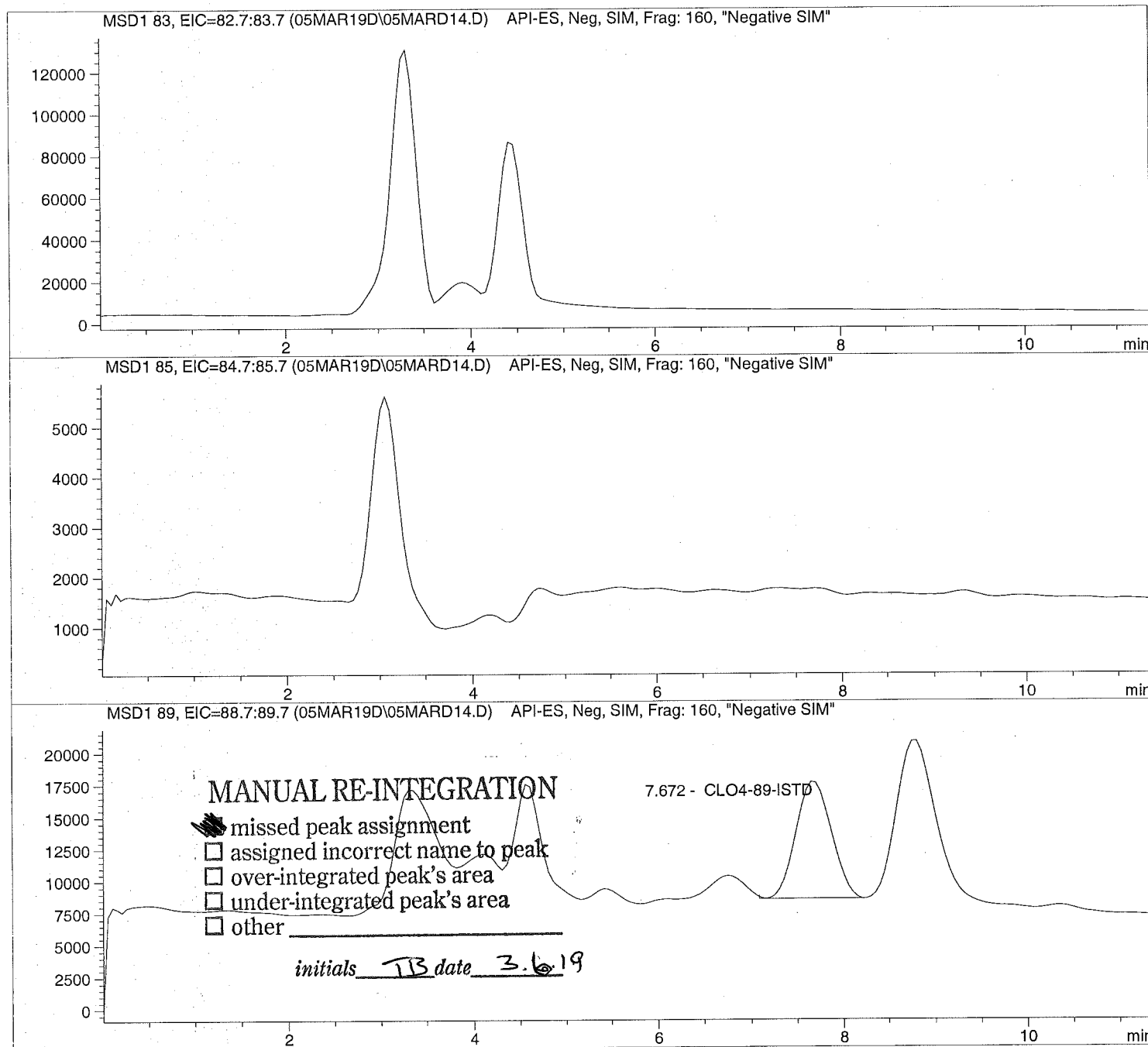
```

Injection Date: 3/05/2019 11:39:24
Sample Name: 1906112009
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```

=====
Injection Date:  3/05/2019  11:39:24      Seq Line:      14
Sample Name:    1906112009                Location:      Vial 84
Acq Operator:   TNB                       Inj. No.:     1
                                           Inj. Vol.:    20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

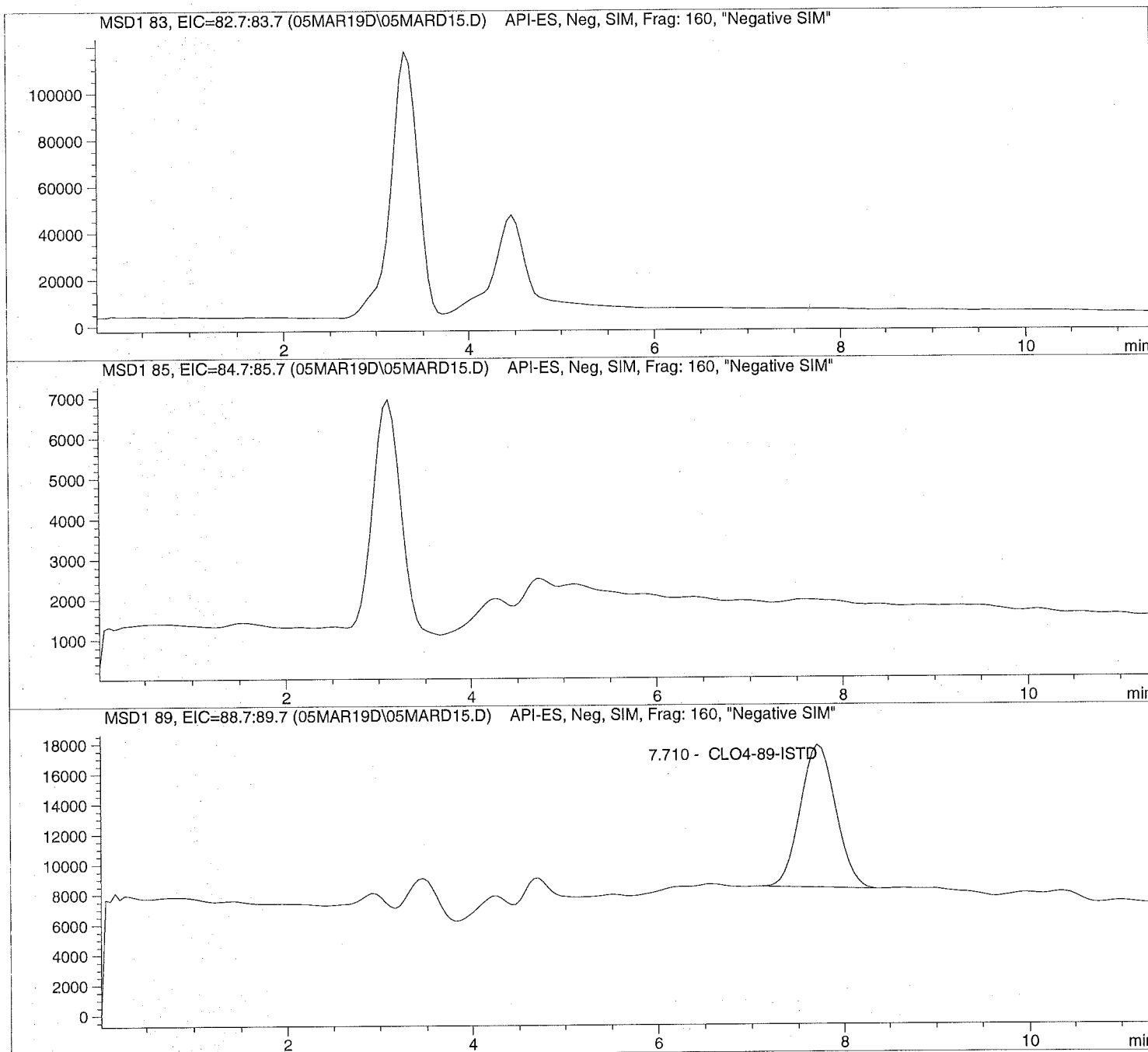
```

Injection Date: 3/05/2019 11:52:27
Sample Name: 1906112010
Acq Operator: TNB

Seq Line: 15
Location: Vial 85
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 11:52:27      Seq Line: 15
Sample Name: 1906112010                 Location: Vial 85
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.710	BBA	251865.4	5.0000	CLO4-89-ISTD

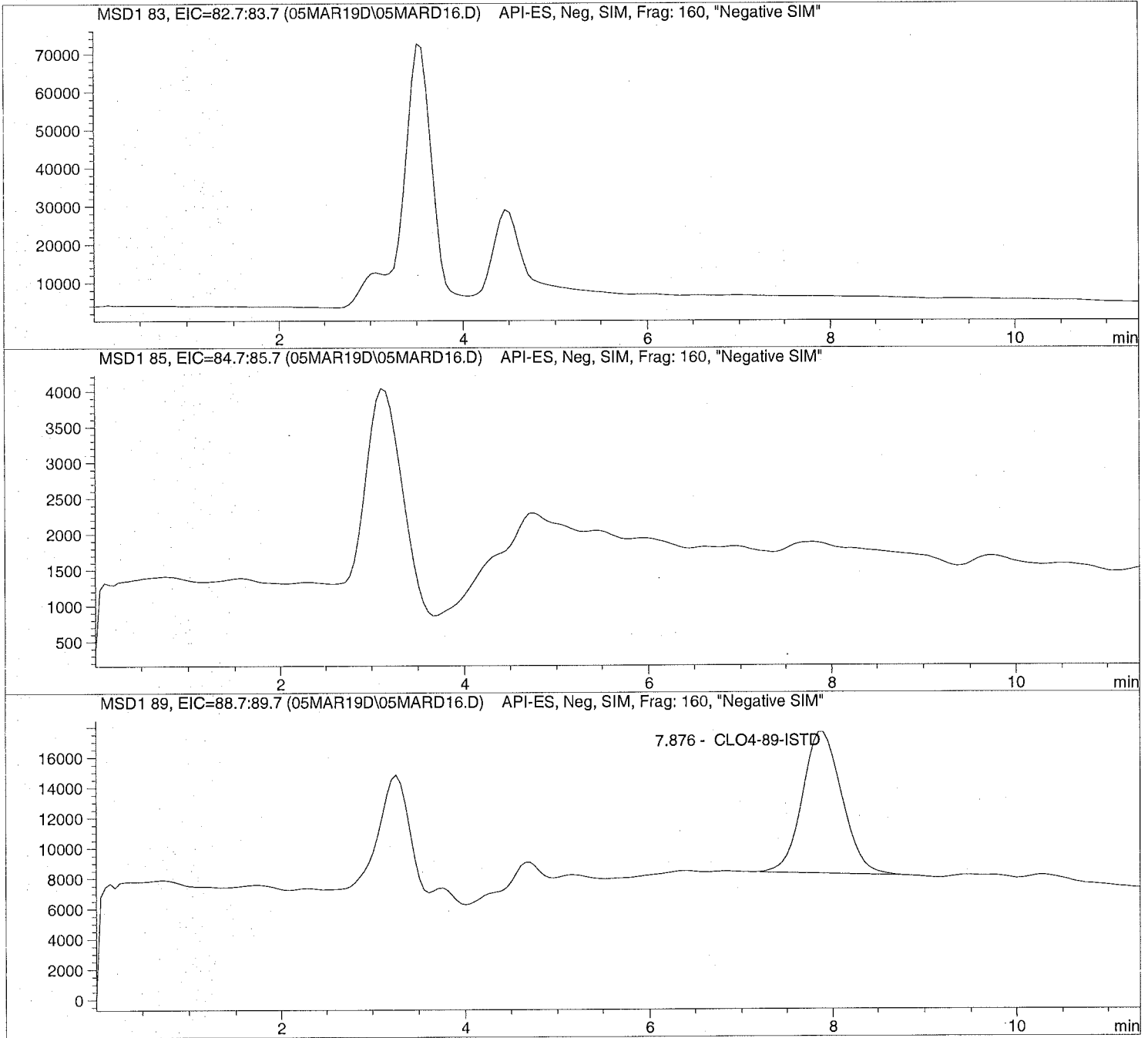
=====
*** End of Report ***

Injection Date: 3/05/2019 12:05:39
Sample Name: 1906112011
Acq Operator: TNB

Seq Line: 16
Location: Vial 86
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date:  3/05/2019  12:05:39          Seq Line:           16
Sample Name:    1906112011                    Location:           Vial 86
Acq Operator:   TNB                           Inj. No.:          1
                                           Inj. Vol.:         20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:      1.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

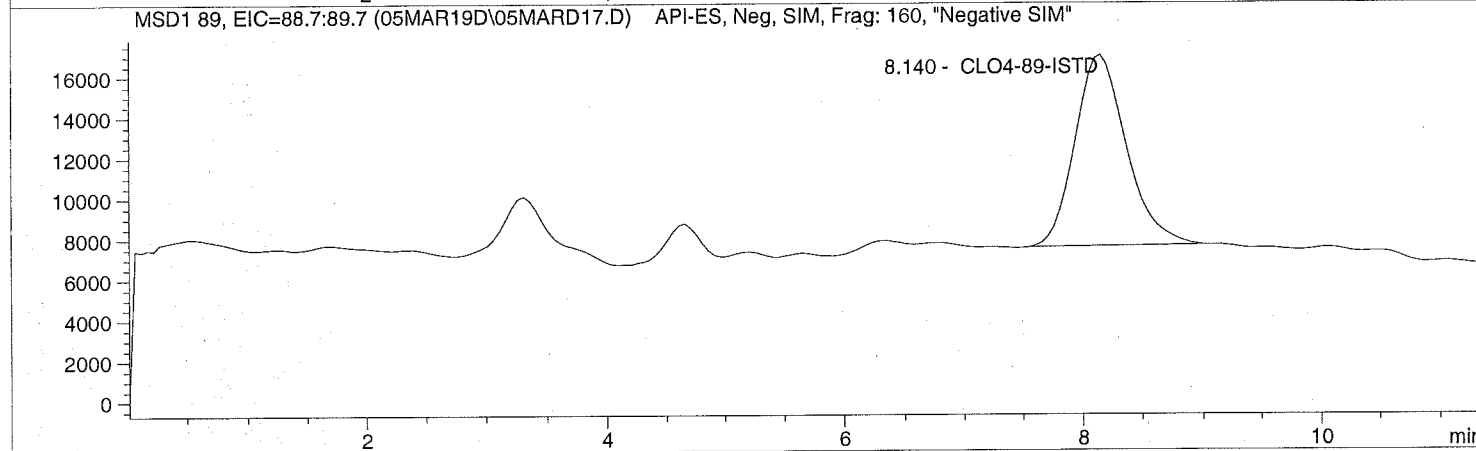
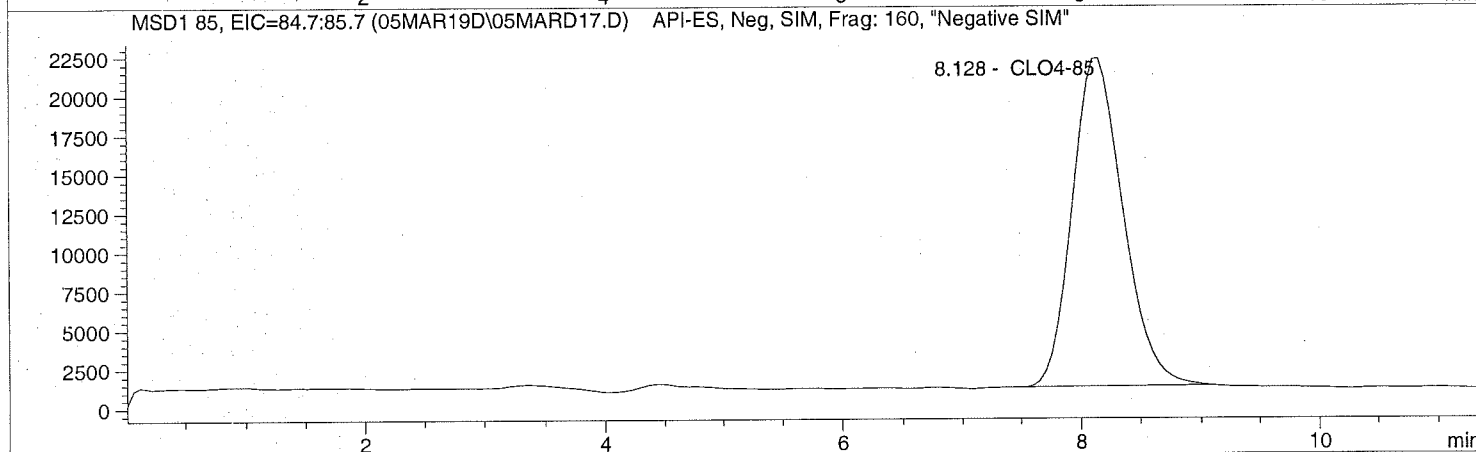
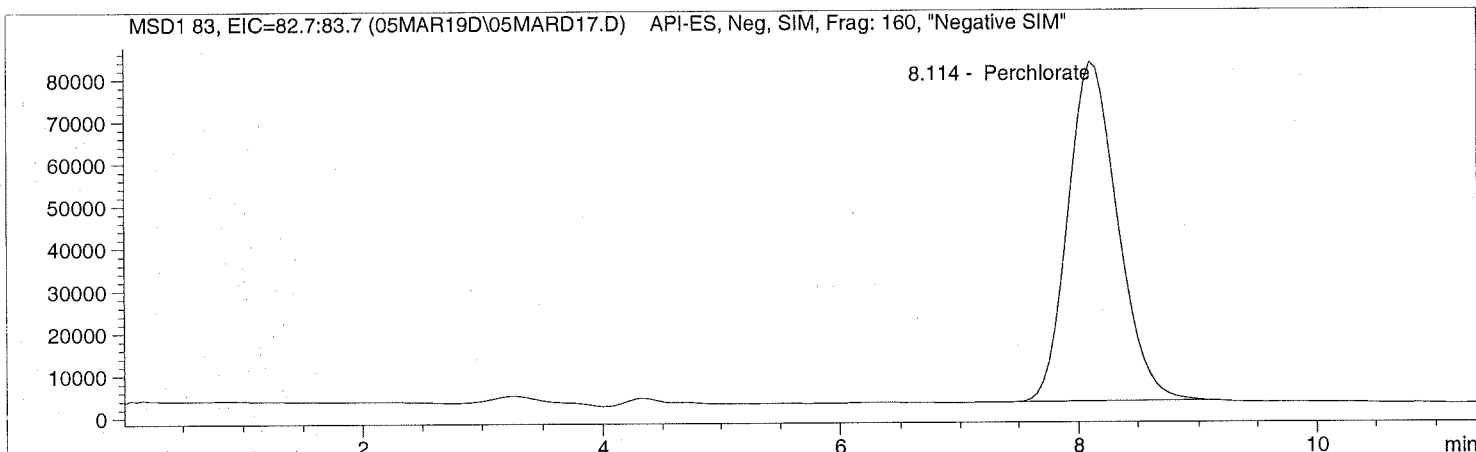
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.876	BBA	280792.2	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

Injection Date: 3/05/2019 12:18:41 Seq Line: 17
Sample Name: 642101 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 12:18:41 Seq Line: 17
Sample Name: 642101 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	2336112.7	25.0199	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.128	PBA	628961.2	25.6341	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.140	PBA	283460.4	5.0000	CLO4-89-ISTD

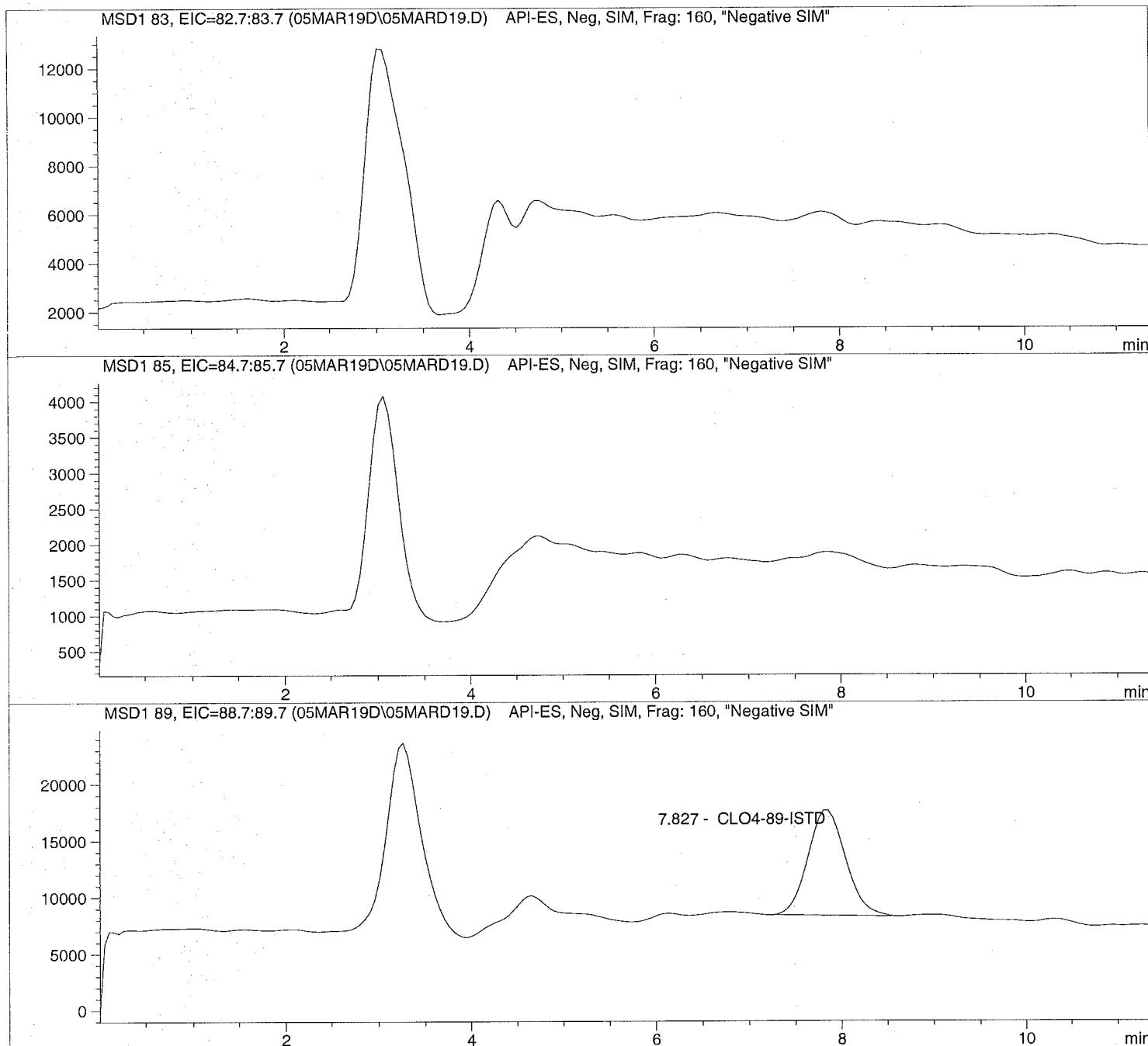
*** End of Report ***

Injection Date: 3/05/2019 12:49:07
Sample Name: 1906332001
Acq Operator: TNB

Seq Line: 19
Location: Vial 88
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 12:49:07      Seq Line: 19
Sample Name: 1906332001                 Location: Vial 88
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.827	PBA	264674.3	5.0000	CLO4-89-ISTD

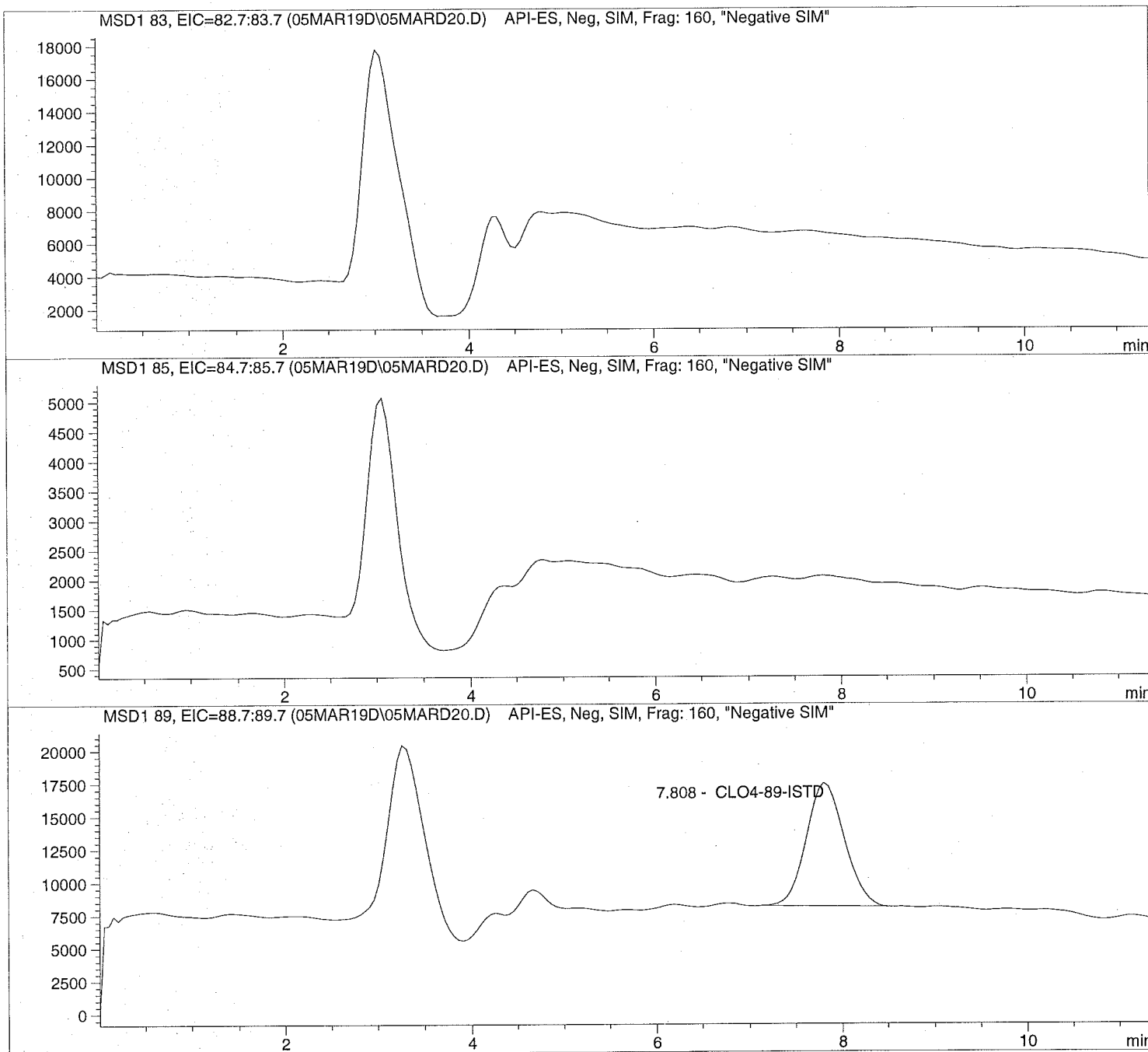
=====
*** End of Report ***

Injection Date: 3/05/2019 13:02:15
Sample Name: 1906334001
Acq Operator: TNB

Seq Line: 20
Location: Vial 89
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 13:02:15      Seq Line:          20
Sample Name:    1906334001              Location:          Vial 89
Acq Operator:   TNB                      Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

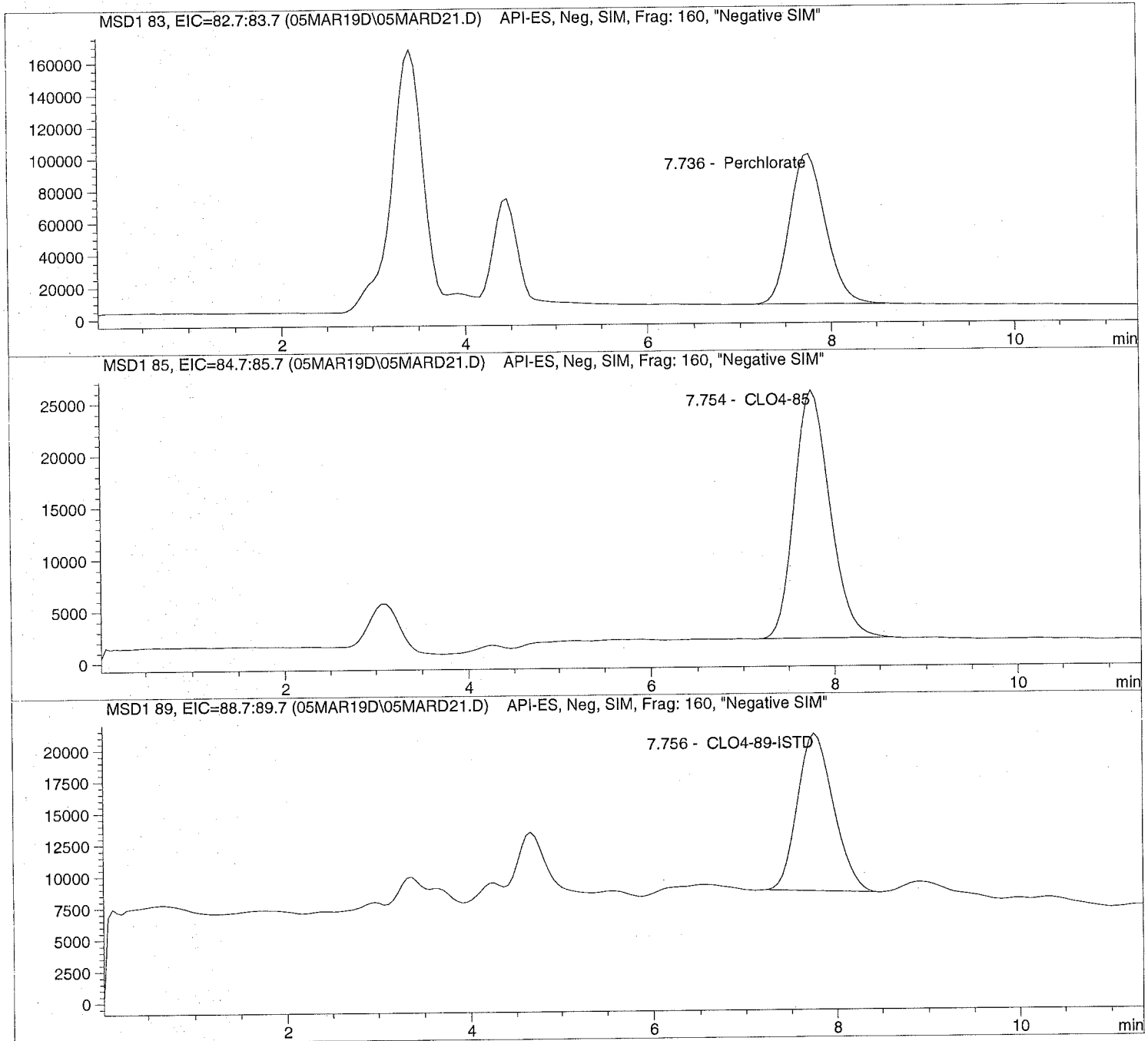
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.808	PBA	265662.0	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

Injection Date: 3/05/2019 13:15:18 Seq Line: 21
Sample Name: 1906112004 10X Location: Vial 90
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 13:15:18      Seq Line: 21
Sample Name: 1906112004 10X             Location: Vial 90
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 10.000000
Sample Amount: 0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.736	PBA	2531259.3	226.4815	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.754	PBA	646870.4	220.9191	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	PB	341607.9	50.0000	CLO4-89-ISTD

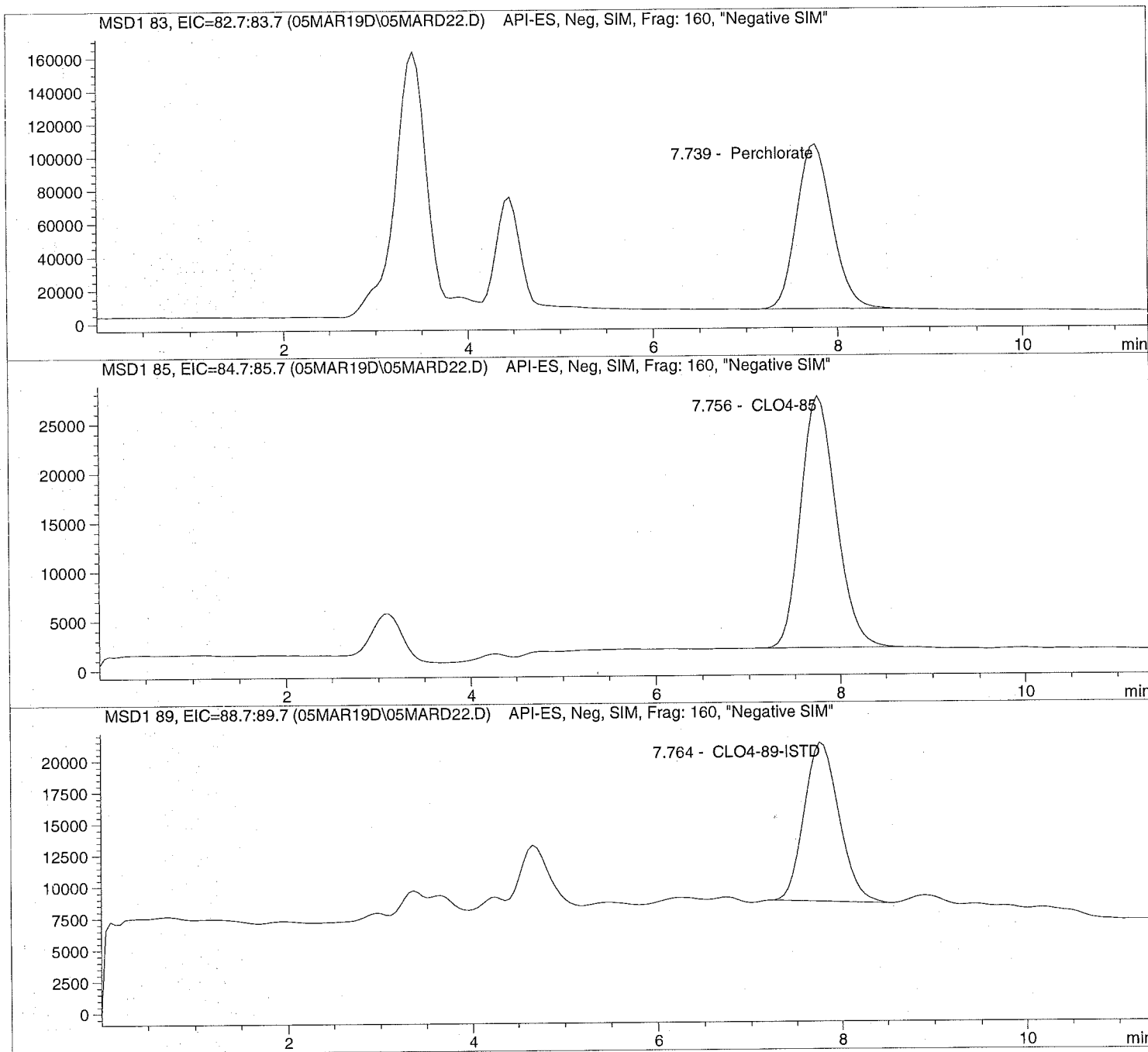
=====
*** End of Report ***

Injection Date: 3/05/2019 13:28:32
Sample Name: 1906112005 10X
Acq Operator: TNB

Seq Line: 22
Location: Vial 91
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 13:28:32      Seq Line:      22  
Sample Name:    1906112005 10X          Location:      Vial 91  
Acq Operator:  TNB                      Inj. No.:     1  
                                           Inj. Vol.:    20 µl
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      10.000000  
Sample Amount: 0.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.739	PBA	2682370.5	241.0371	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	BBA	695643.1	238.4467	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.764	PB	338724.1	50.0000	CLO4-89-ISTD

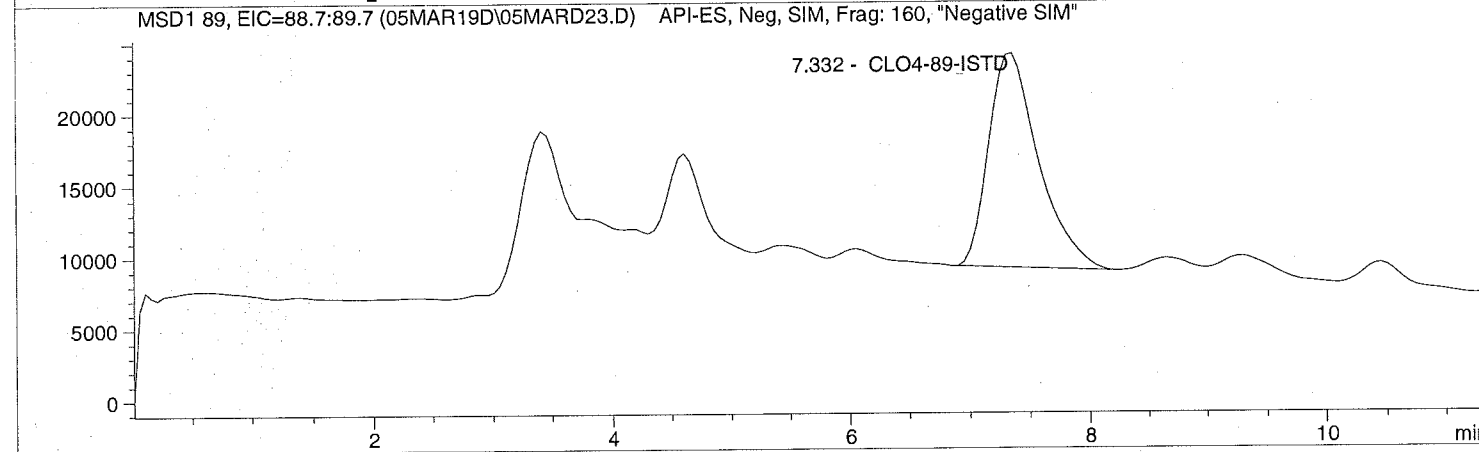
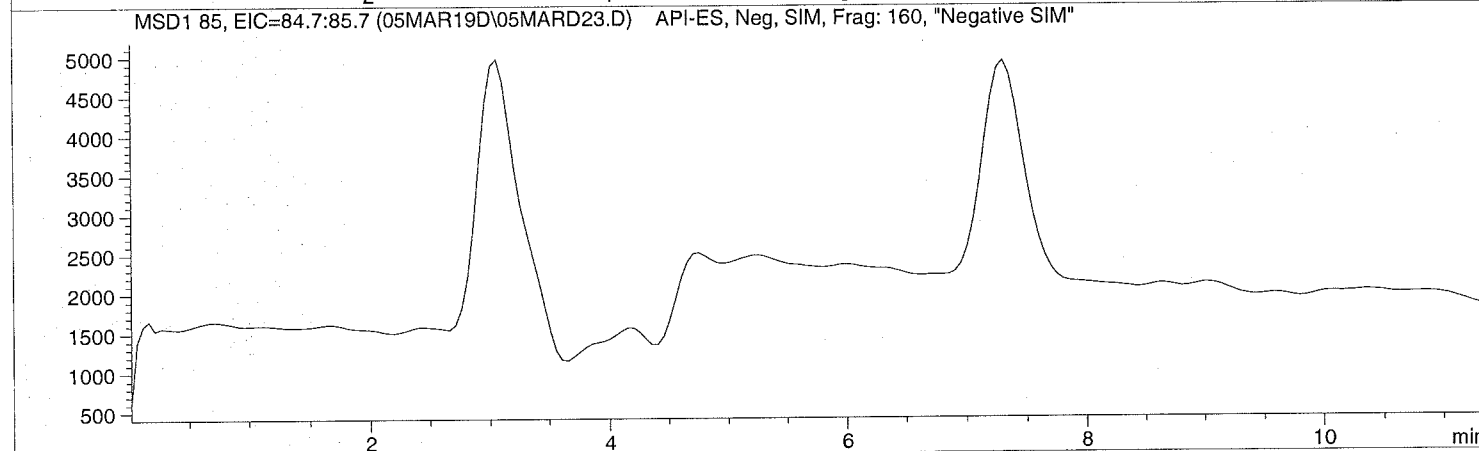
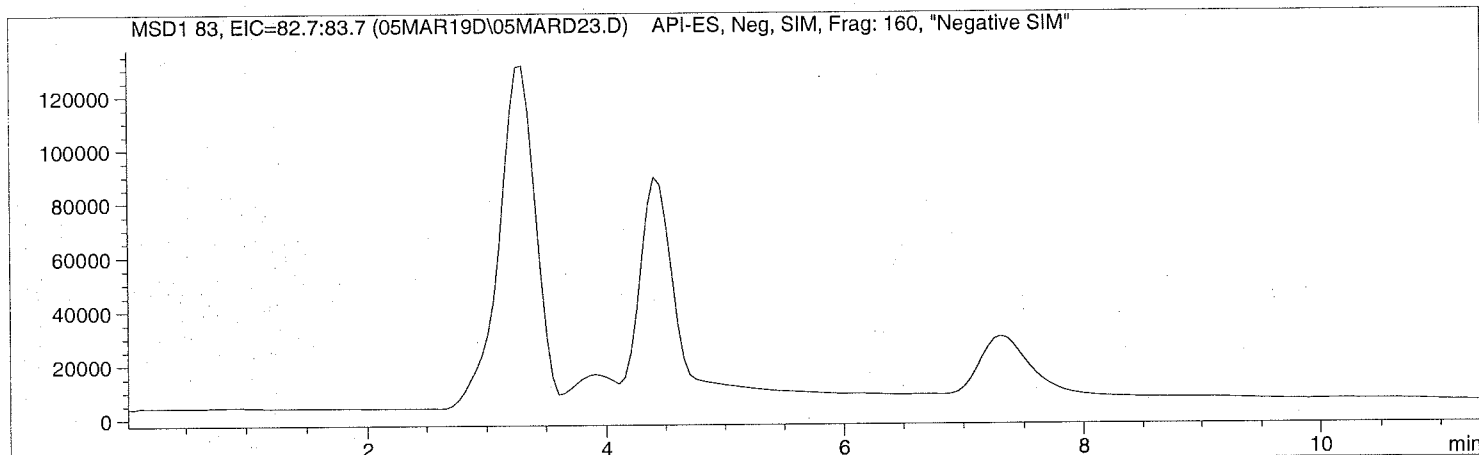
=====
*** End of Report ***

Injection Date: 3/05/2019 13:41:34
Sample Name: 1906112007 RE
Acq Operator: TNB

Seq Line: 23
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 13:41:34 Seq Line: 23
Sample Name: 1906112007 RE Location: Vial 82
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.332	PB	431134.6	5.0000	CLO4-89-ISTD

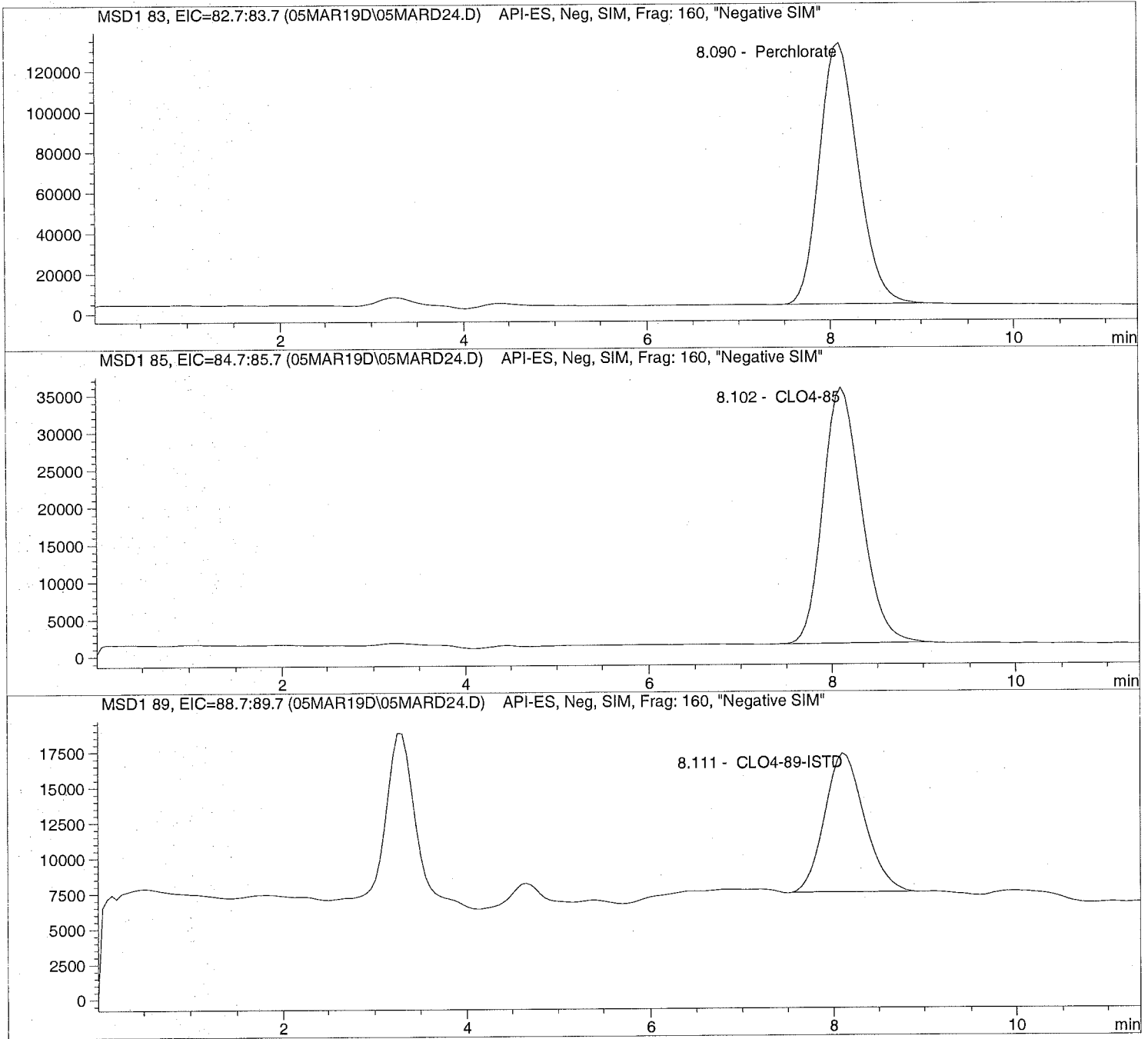
*** End of Report ***

Injection Date: 3/05/2019 13:54:34
Sample Name: 1906330001 100
Acq Operator: TNB

Seq Line: 24
Location: Vial 92
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====
Injection Date: 3/05/2019 13:54:34      Seq Line:      24
Sample Name:    1906330001 100          Location:      Vial 92
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       100.000000
Sample Amount:  0.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.090	PBA	3773628.7	3708.6567	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	PBA	1010205.2	3771.6748	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.111	PBA	298984.6	500.0000	CLO4-89-ISTD

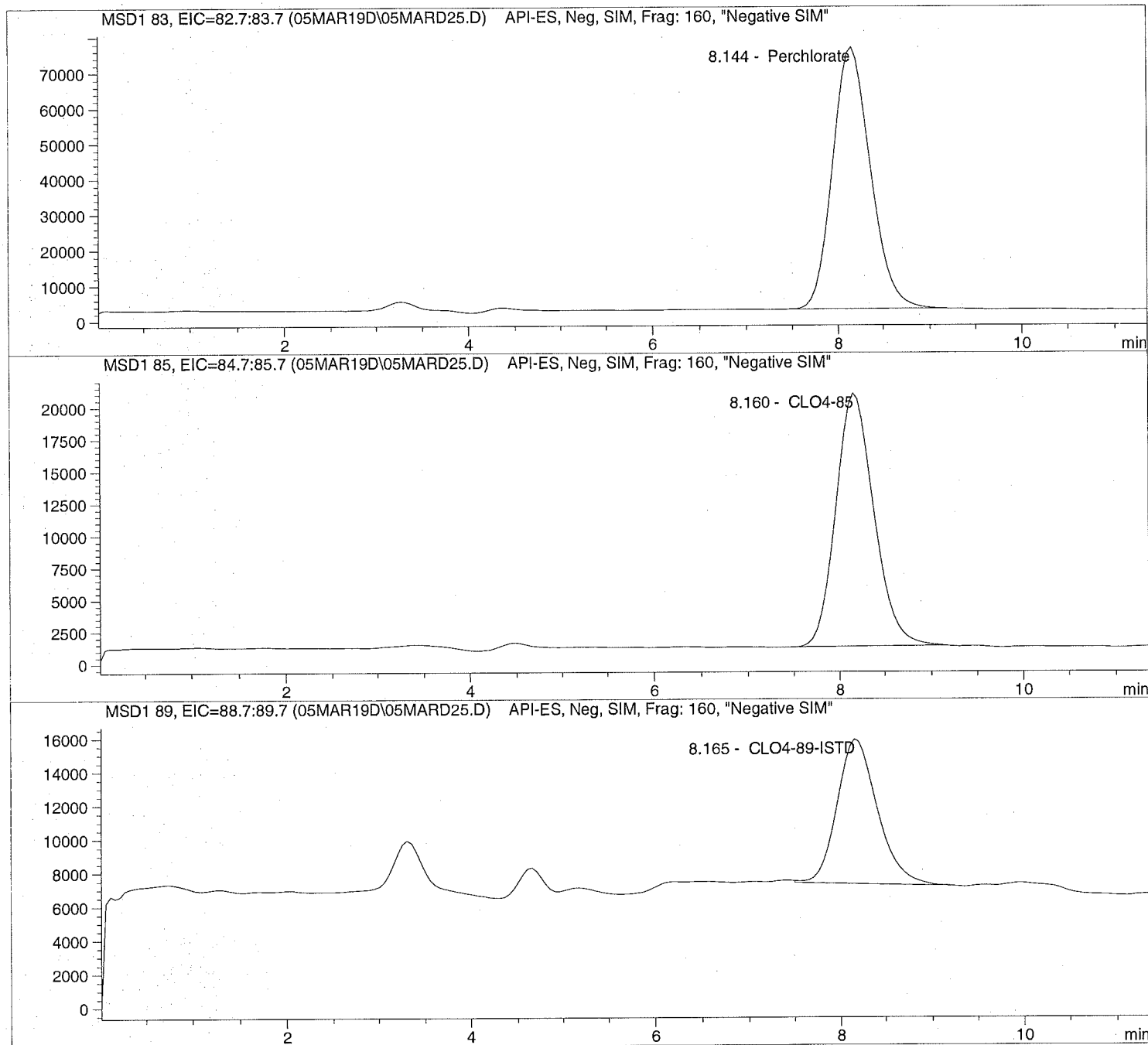
=====
*** End of Report ***

Injection Date: 3/05/2019 14:07:36
Sample Name: 642102 CCV@25
Acq Operator: TNB

Seq Line: 25
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```
=====  
Injection Date: 3/05/2019 14:07:36      Seq Line:          25  
Sample Name:    642102  CCV@25          Location:          Vial 71  
Acq Operator:  TNB                      Inj. No.:         1  
                                           Inj. Vol.:        20 µl
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 25.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.144	PBA	2157871.5	24.4553	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.160	PBA	583711.5	25.1665	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.165	BBA	268305.4	5.0000	CLO4-89-ISTD

=====
*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

**Initial
Calibration**

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	8.94006e4	7.889	9.89924e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.97443e5	8.114	2.26028
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	4.79370e5	7.828	4.65688
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	9.30136e5	7.904	9.14998
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.81067e6	7.793	25.52636
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	5.66830e6	7.976	51.07439
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	8.69624e6	7.886	74.30603
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	1.01141e6	7.988	9.46019

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.26121e4	7.914	9.98836e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	5.53134e4	8.127	2.11360
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.39247e5	7.842	4.91261
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.54396e5	7.923	9.39034
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	7.35969e5	7.811	25.48268
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.47152e6	7.993	50.35774
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.32809e6	7.900	74.72233
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.81230e5	8.007	9.87858

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.41443e5	7.900	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.99651e5	8.132	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.38646e5	7.853	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	3.25154e5	7.925	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	3.33799e5	7.819	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	3.14712e5	7.999	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	3.13909e5	7.908	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	3.41503e5	8.005	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

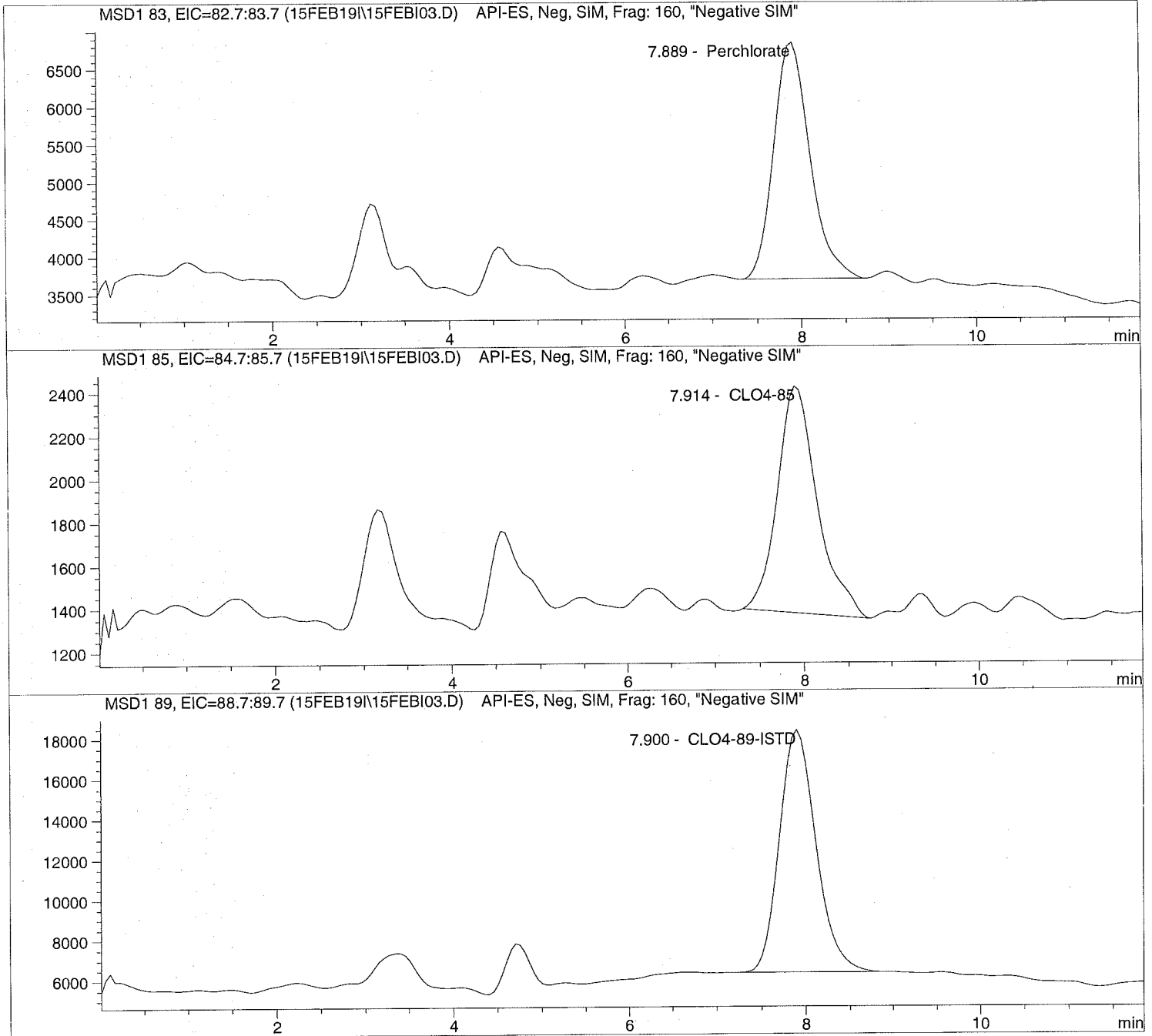
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ .20ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Injection Date: 2/15/2019 09:51:42
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====
Injection Date:  2/15/2019  09:51:42      Seq Line:           3
Sample Name:     CLO4@ 1.0ug/L           Location:           Vial 73
Acq Operator:    TNB                      Inj. No.:          1
                                           Inj. Vol.:         25 µl
=====
```

```
Acq. Method:     CLO4-AQN.M
Analysis Method:  C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:    2/19/2019  09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:       Signal
Calib. Data Modified:  Tue, 19. Feb. 2019,09:07:33 am
Multiplier:      1.000000
Dilution:        1.000000
Sample Amount:   1.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.889	PBA	89400.6	0.9899	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.914	BBA	32612.1	0.9988	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	BBA	341443.2	5.0000	CLO4-89-ISTD

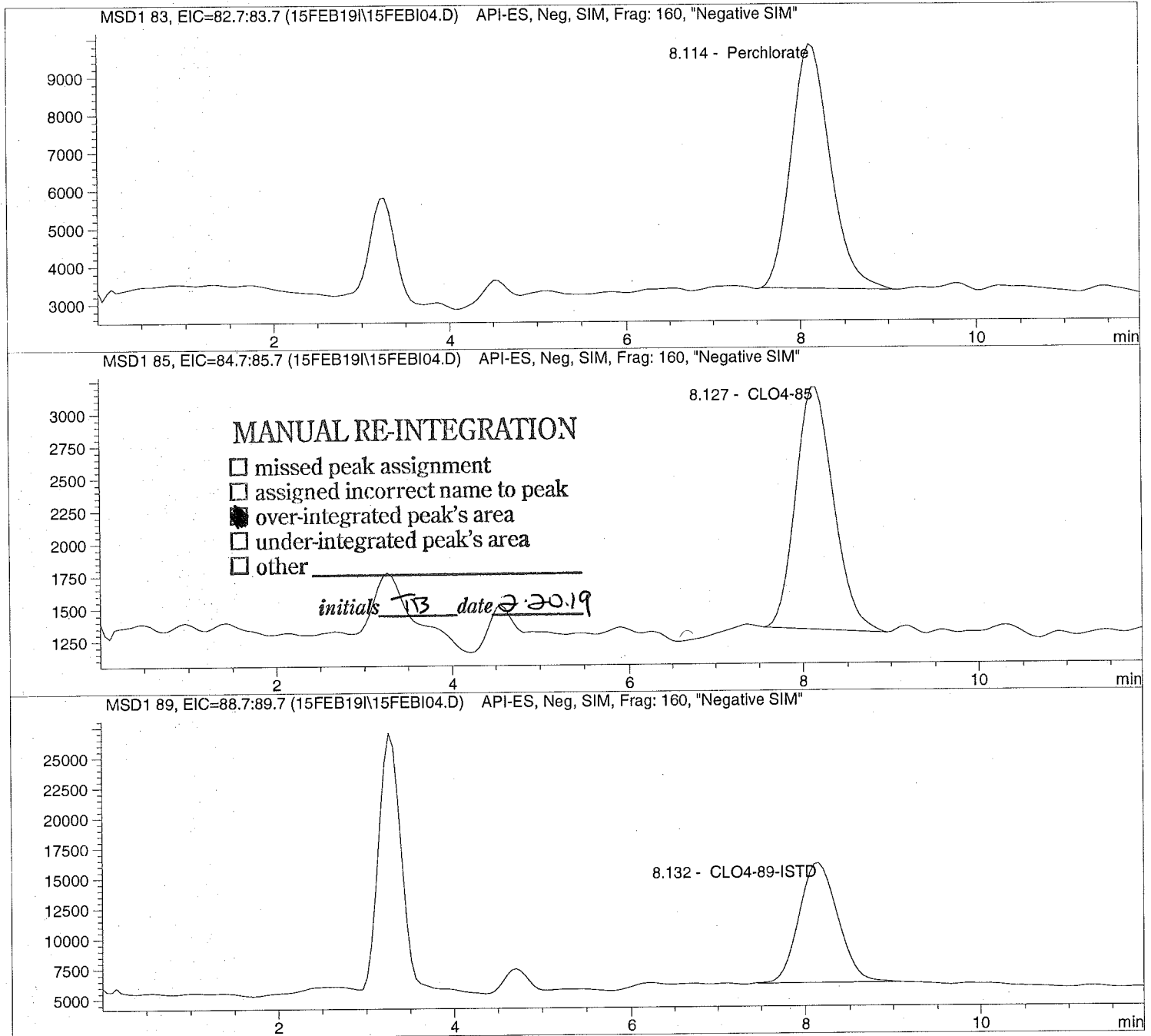
=====
*** End of Report ***

Injection Date: 2/15/2019 10:05:24
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Injection Date: 2/15/2019 10:05:24 Seq Line: 4
Sample Name: CLO4@ 2.0ug/L Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 2.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	MM	55313.4	2.1136	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

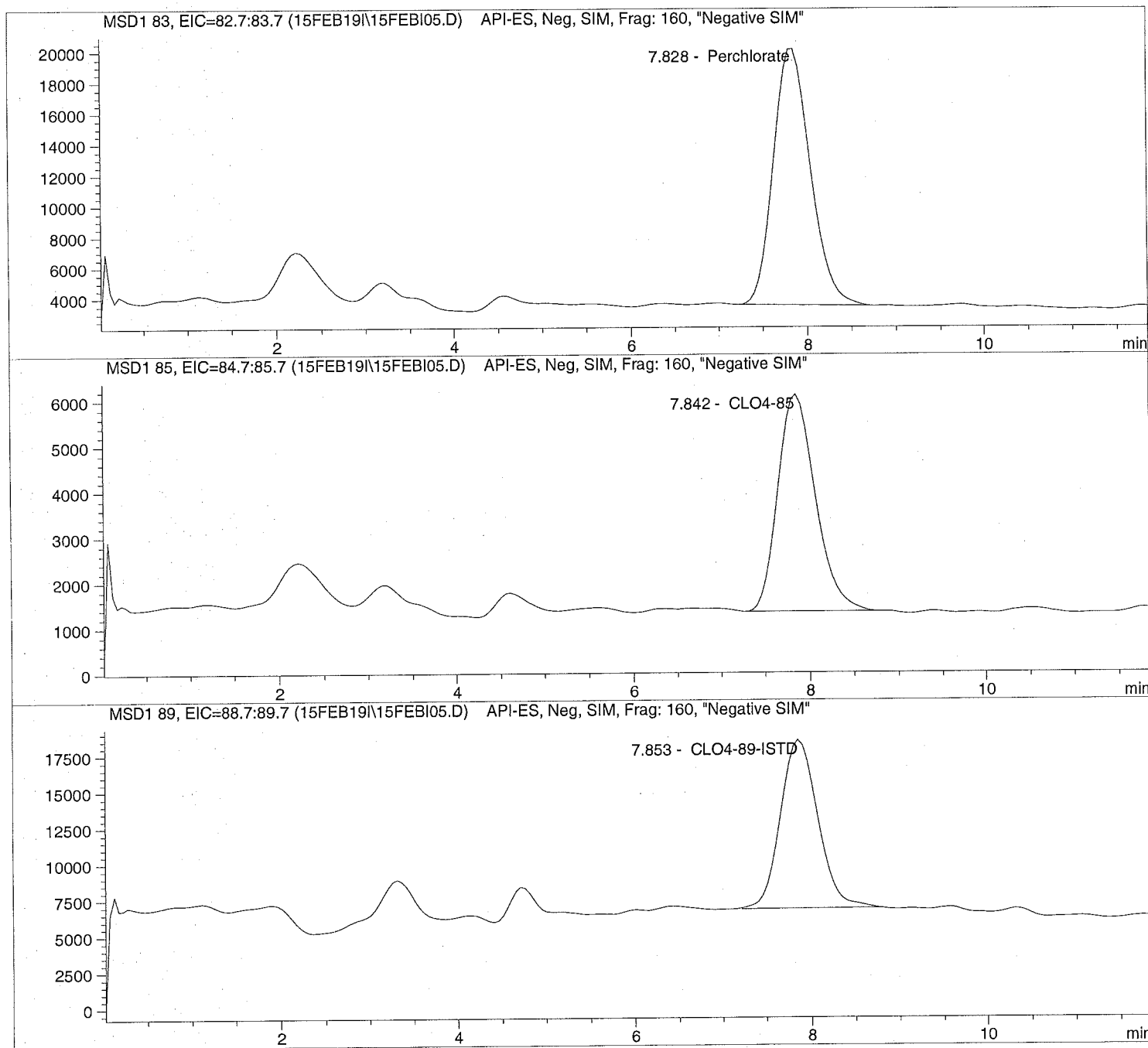
*** End of Report ***

Injection Date: 2/15/2019 11:42:56
Sample Name: CLO4@ 5.0ug/L
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis




```
=====
Injection Date: 2/15/2019 11:42:56      Seq Line: 5
Sample Name:    CLO4@ 5.0ug/L           Location:  Vial 75
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

===== Sample Information =====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====
```

===== LCMS Results =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.828	PBA	479370.4	4.6569	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.842	PBA	139246.9	4.9126	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.853	PBA	338646.3	5.0000	CLO4-89-ISTD

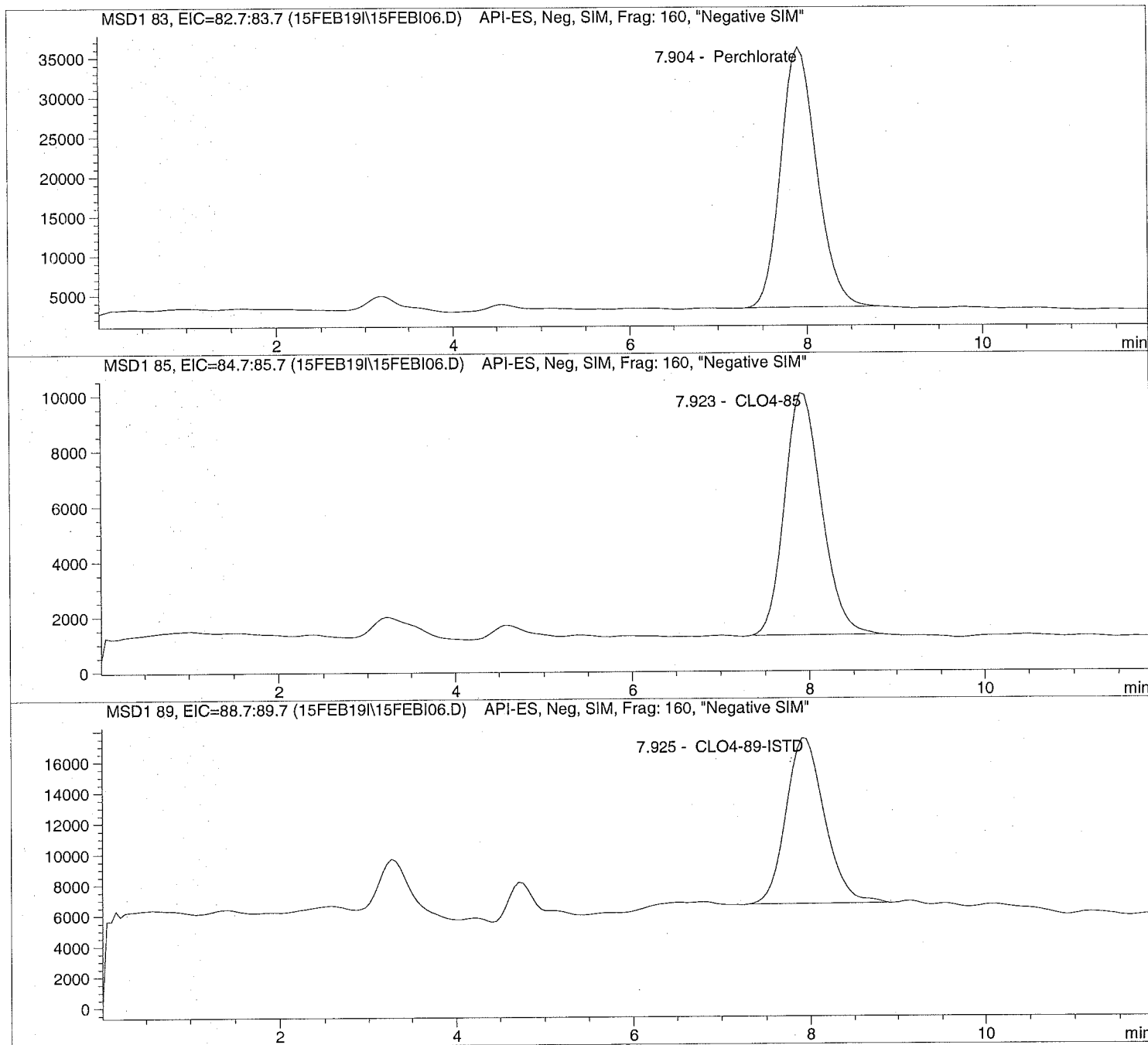
=====
*** End of Report ***

Injection Date: 2/15/2019 11:56:38
Sample Name: CLO4@ 10.ug/L
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====
Injection Date: 2/15/2019 11:56:38      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis

Sample Information

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.904	PBA	930135.8	9.1500	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.923	BBA	254395.6	9.3903	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.925	PBA	325154.4	5.0000	CLO4-89-ISTD

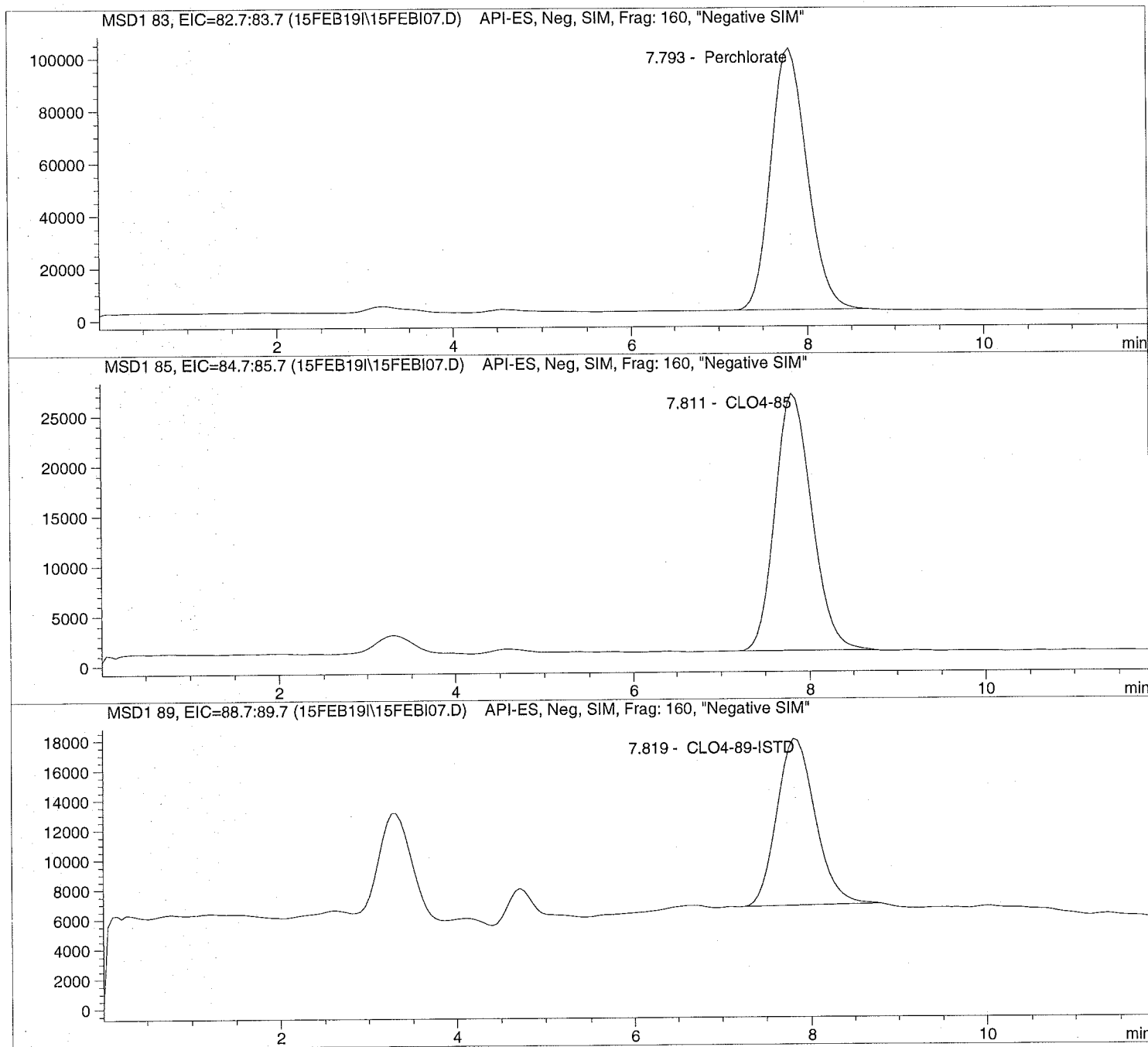
*** End of Report ***

Injection Date: 2/15/2019 12:10:22
Sample Name: CLO4@ 25.ug/L
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 12:10:22      Seq Line: 7  
Sample Name:    CLO4@ 25.ug/L           Location:  Vial 77  
Acq Operator:   TNB                     Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:09:20
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 25.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.793	PBA	2810669.2	25.5264	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.811	BBA	735968.9	25.4827	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

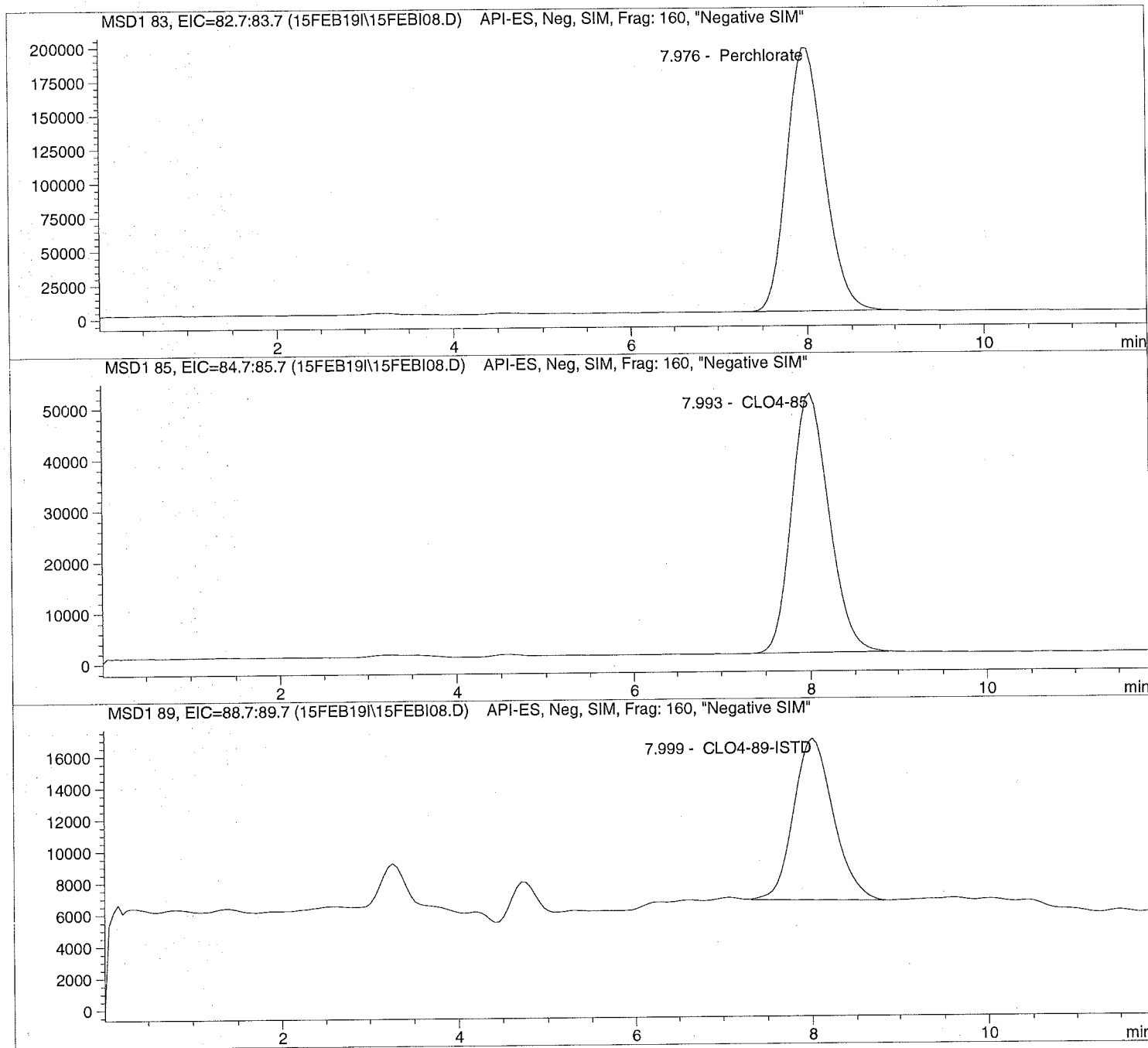
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.819	PBA	333799.0	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

=====
Injection Date: 2/15/2019 12:24:06 Seq Line: 8
Sample Name: CLO4@ 50.ug/L Location: Vial 78
Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis
=====

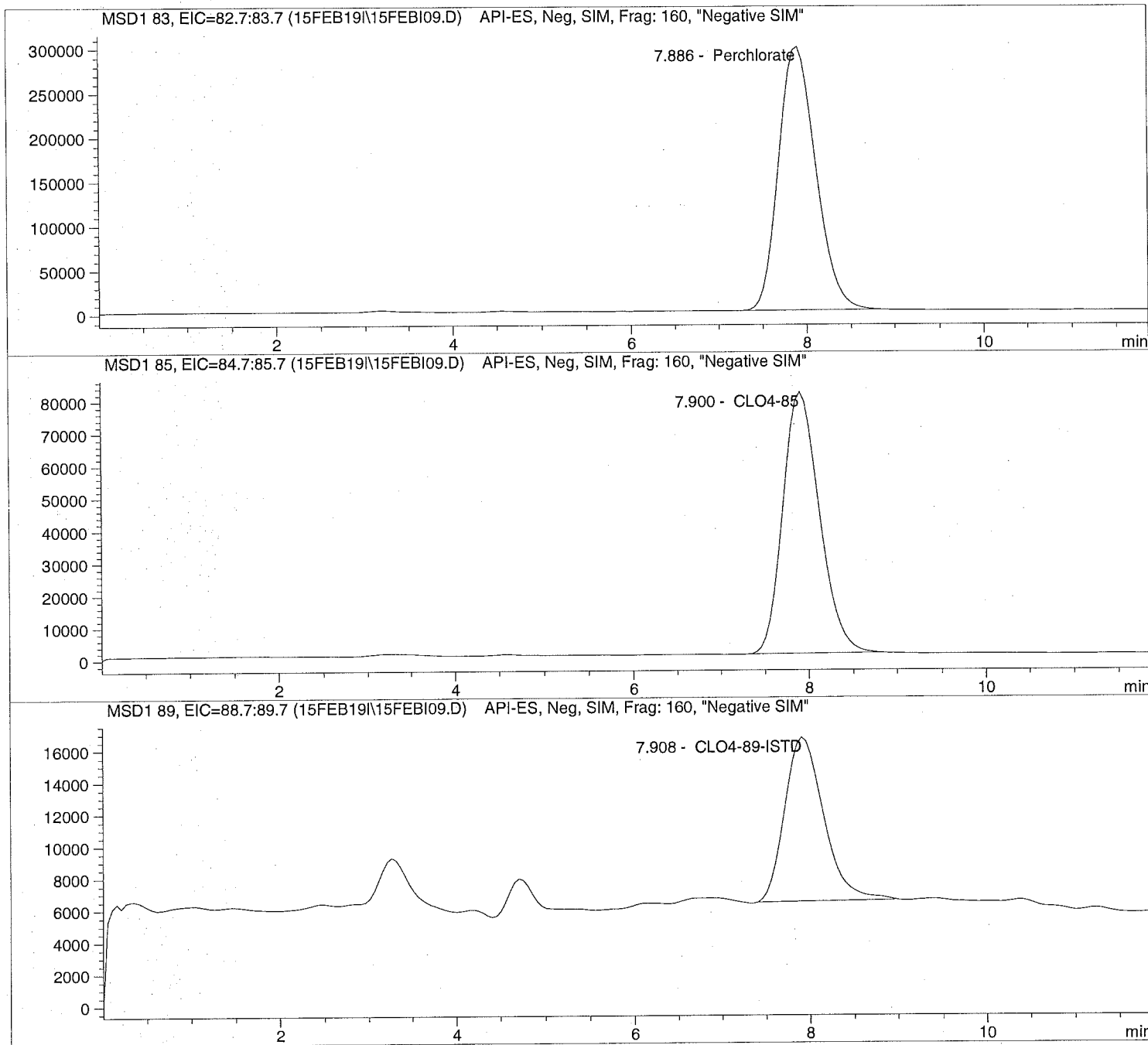


Injection Date: 2/15/2019 12:37:48
Sample Name: CLO4@ 75.ug/L
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis




```
=====  
Injection Date: 2/15/2019 12:37:48      Seq Line: 9  
Sample Name:    CLO4@ 75.ug/L           Location:  Vial 79  
Acq Operator:   TNB                     Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:09:20  
=====
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 75.000  
=====
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.886	PBA	8696239.0	74.3060	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	PBA	2328089.5	74.7223	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.908	PBA	313908.9	5.0000	CLO4-89-ISTD

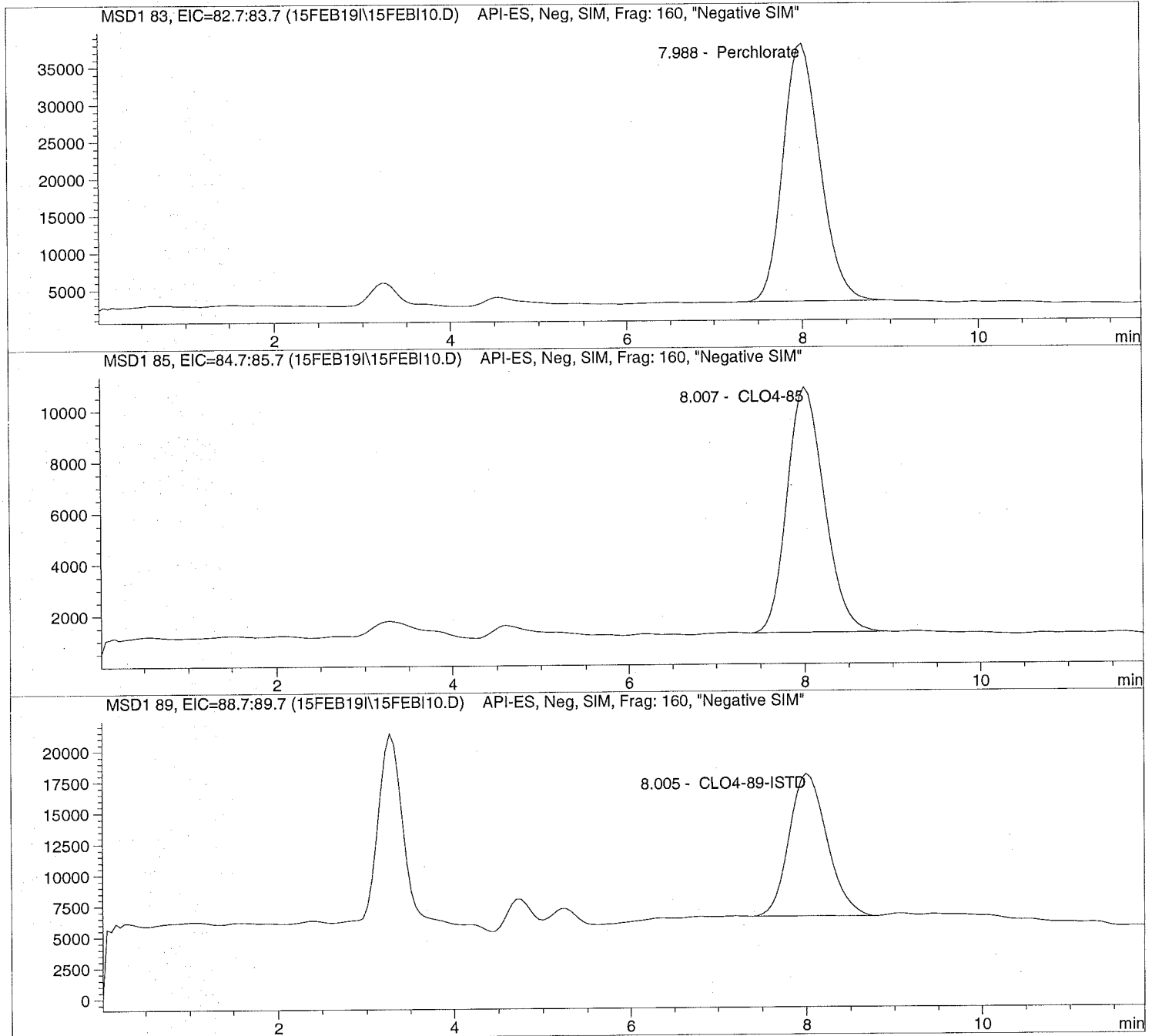
=====
*** End of Report ***
=====

Injection Date: 2/15/2019 12:51:29
Sample Name: ICAL Verf@10ug/L
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 12:51:29      Seq Line: 10  
Sample Name:    ICAL Verf@10ug/L        Location:  Vial 80  
Acq Operator:  TNB                      Inj. No.: 1  
                                           Inj. Vol.: 25 µl  
=====
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:  2/19/2019 09:09:20
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 10.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.988	BBA	1011409.8	9.4602	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.007	BBA	281229.9	9.8786	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.005	BBA	341503.2	5.0000	CLO4-89-ISTD

=====
*** End of Report ***
=====



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

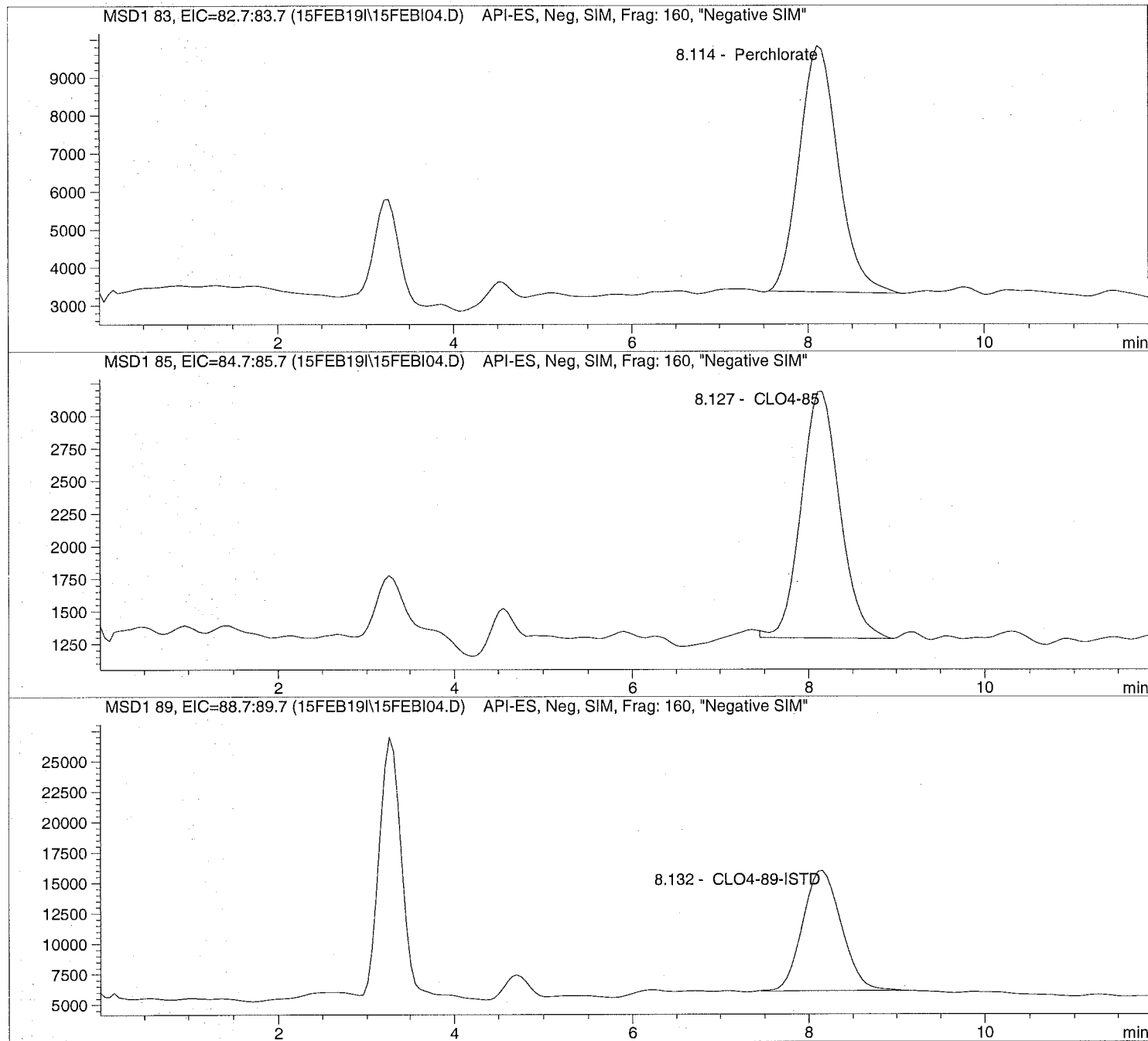
Unmodified

Injection Date: 2/15/2019 10:05:24
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:12:36

Perchlorate analysis



```
=====  
Injection Date: 2/15/2019 10:05:24      Seq Line:      4  
Sample Name:    CLO4@ 2.0ug/L           Location:      Vial 74  
Acq Operator:   TNB                     Inj. No.:     1  
                                           Inj. Vol.:    25 µl
```

```
Acq. Method:    CLO4-AQN.M  
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M  
Last Changed:   2/19/2019 09:12:36
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:      Signal  
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am  
Multiplier:    1.000000  
Dilution:      1.000000  
Sample Amount: 2.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	BBA	57206.1	2.1923	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

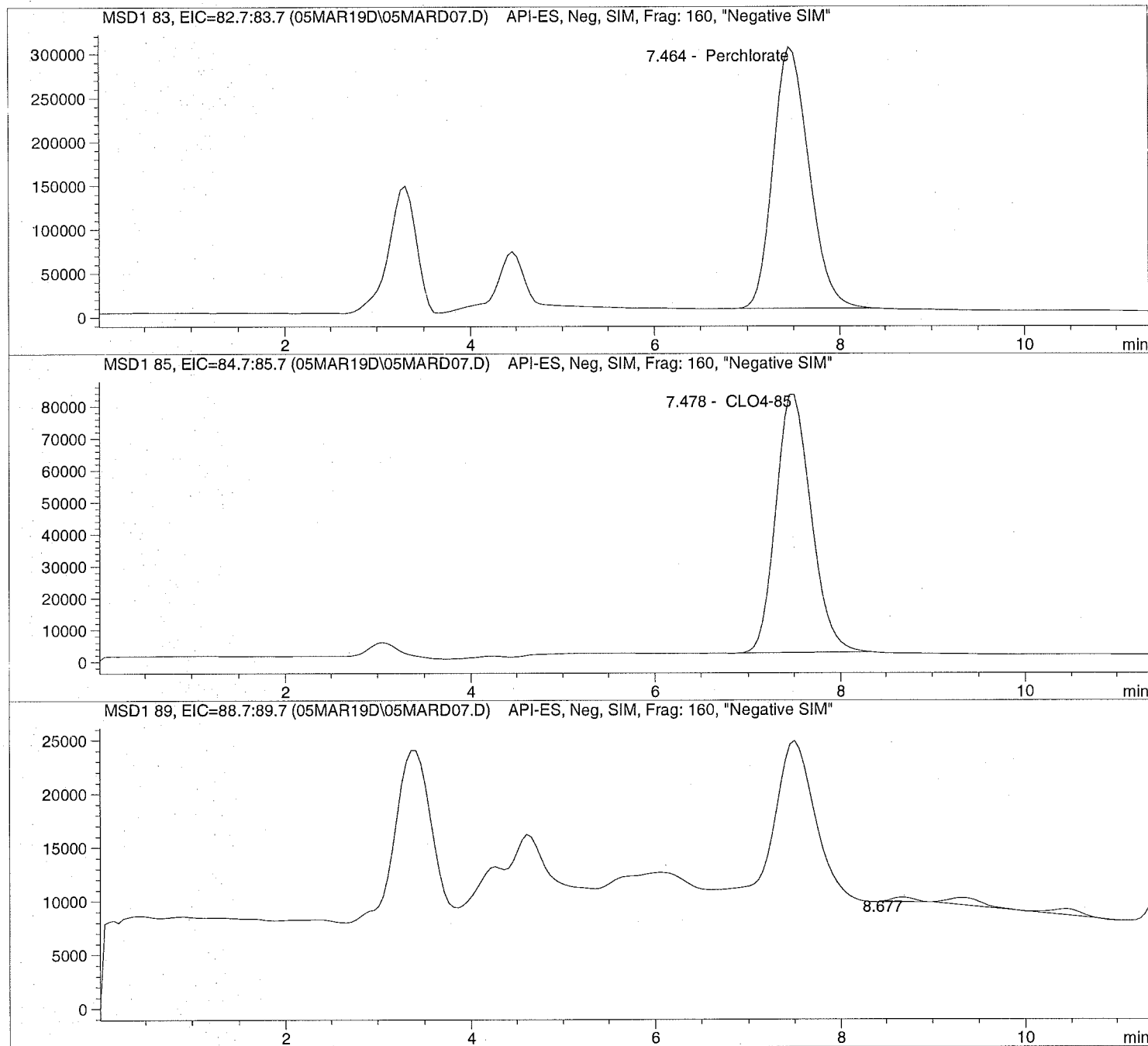
=====
*** End of Report ***

Injection Date: 3/05/2019 10:07:11
Sample Name: 1906112002 MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:07:11 Seq Line: 7
Sample Name: 1906112002 MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	419.8794	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	402.6179	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

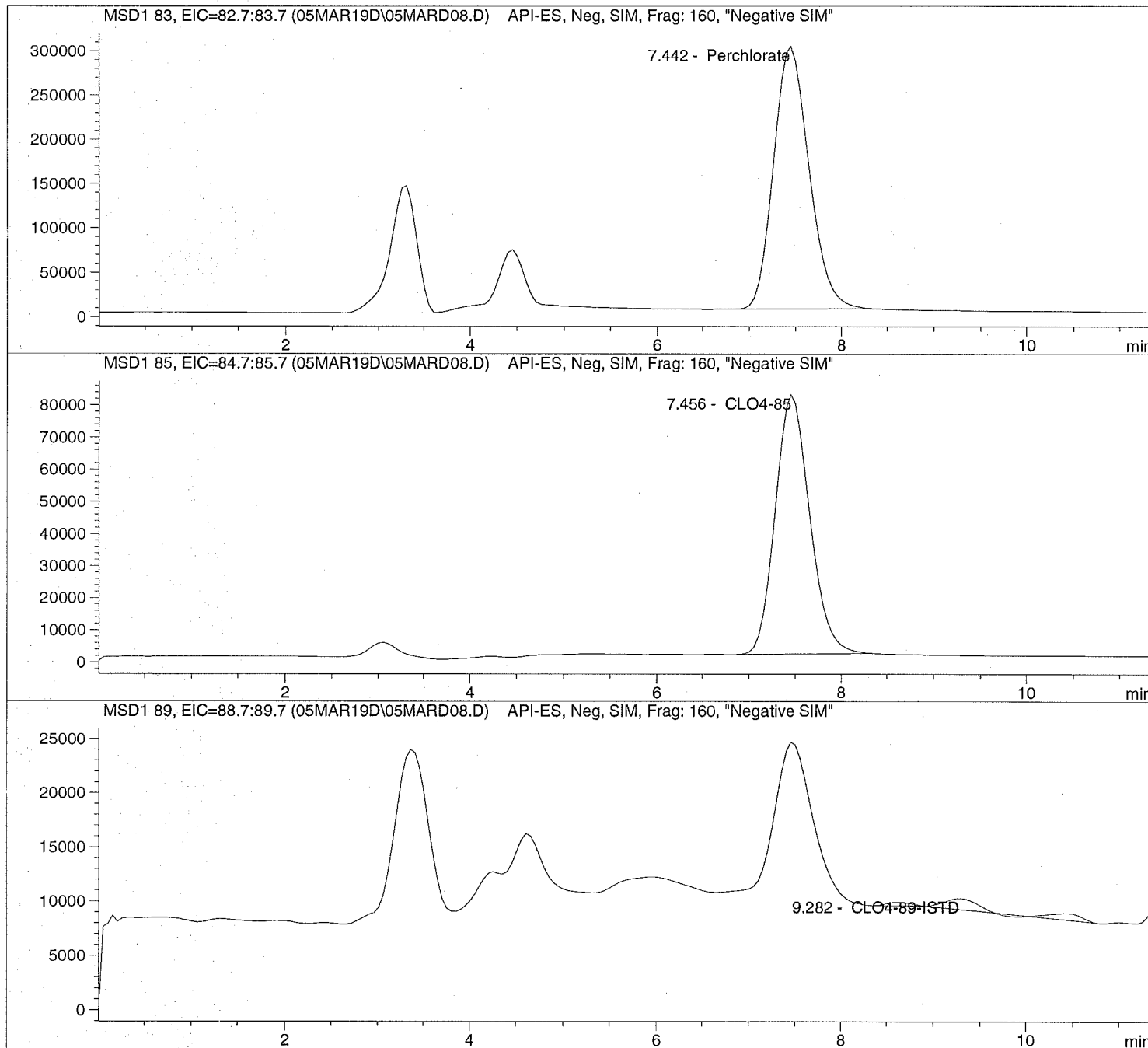
RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.677	BB	7208.6	0.0000	
9.316	VBA	28561.1	5.0000	CLO4-89-ISTD

*** End of Report ***

Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	359.0996	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	344.2233	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.621	VB	5769.9	0.0000	
9.282	VBA	35831.6	5.0000	CLO4-89-ISTD

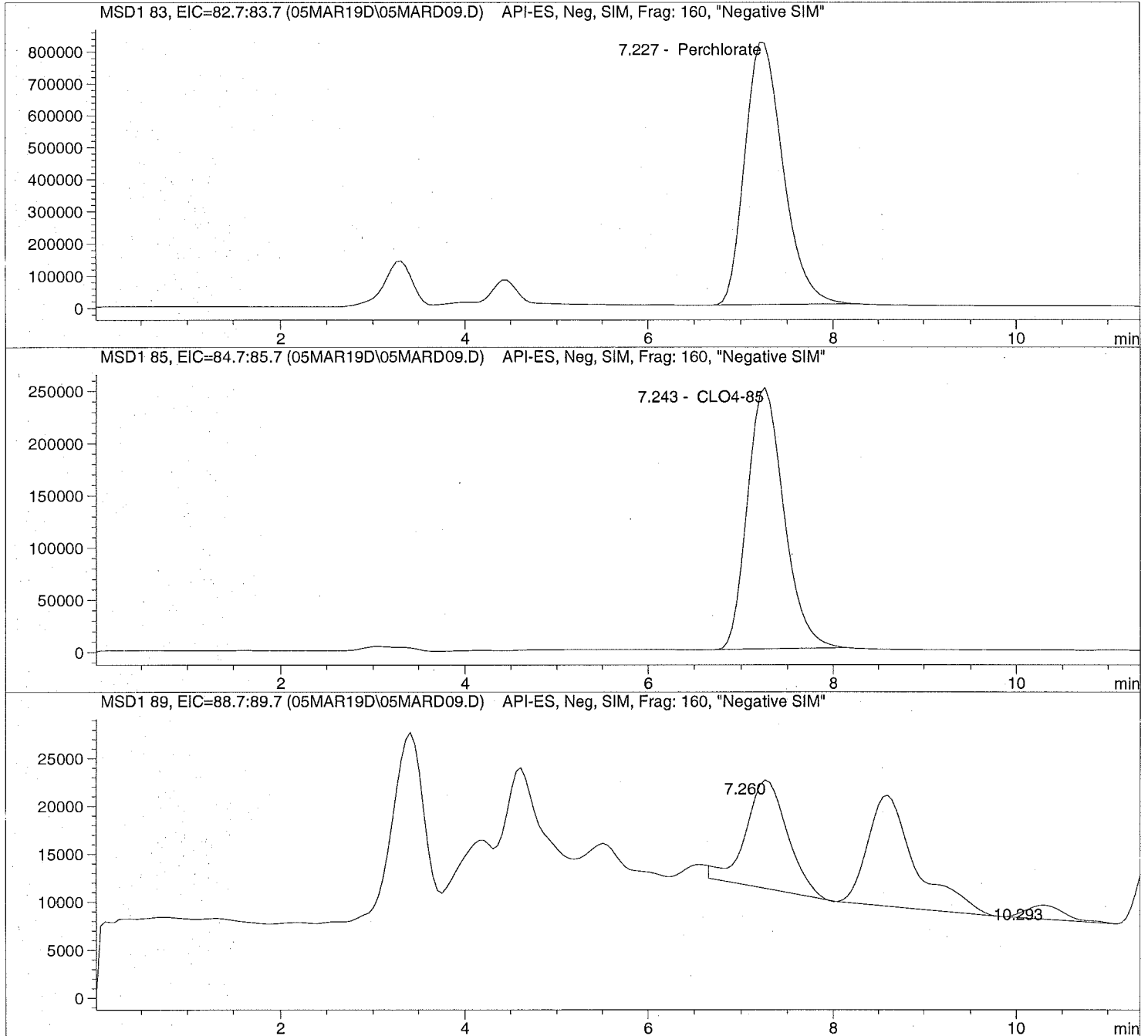
*** End of Report ***

Injection Date: 3/05/2019 10:33:21
Sample Name: 1906112004
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 10:33:21 Seq Line: 9
Sample Name: 1906112004 Location: Vial 79
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	131.1742	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	138.8050	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.260	BB	352149.7	0.0000	
8.589	VBA	421141.9	5.0000	CLO4-89-ISTD
10.293	BBA	41603.7	0.0000	

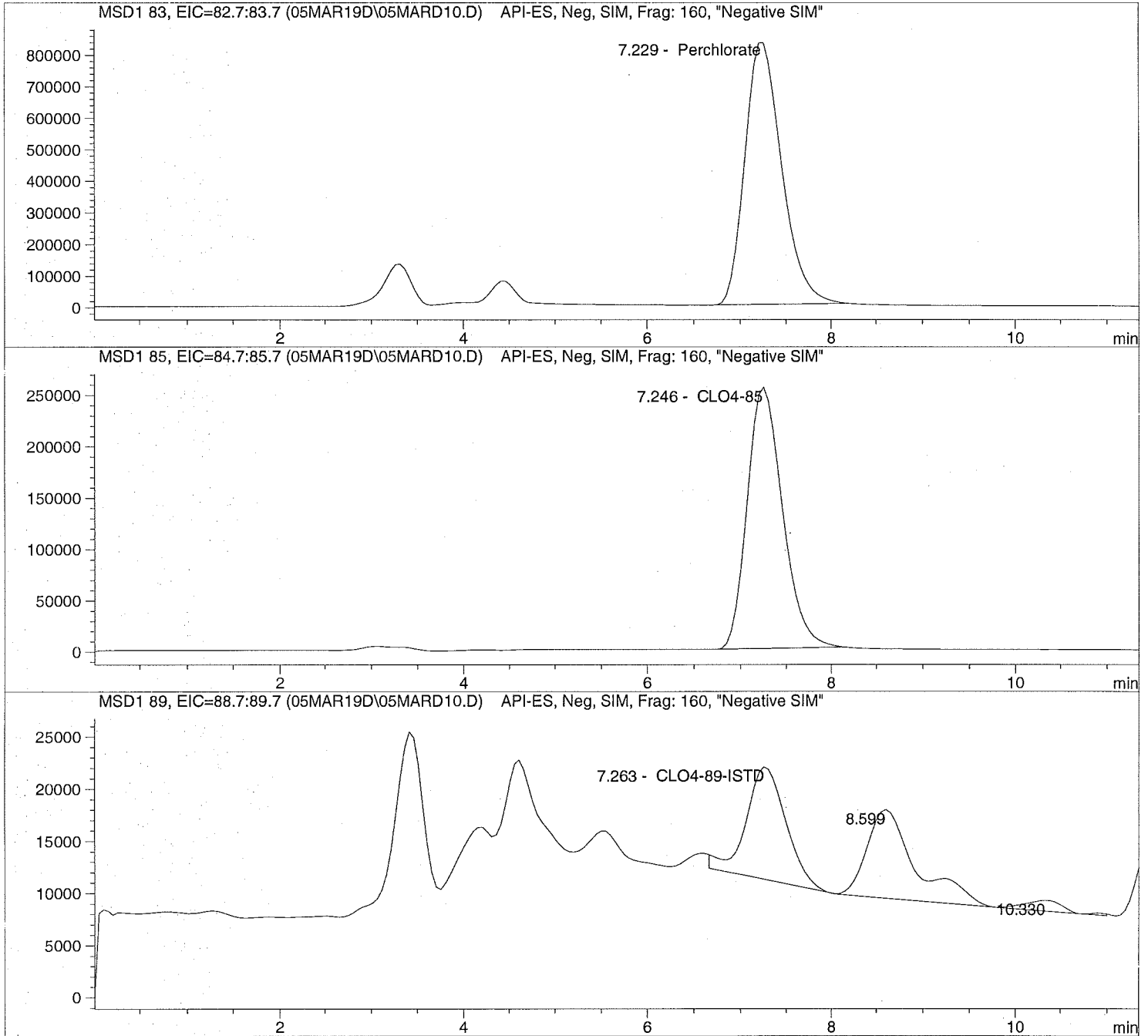
*** End of Report ***

Injection Date: 3/05/2019 10:46:26
Sample Name: 1906112005
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



```

=====
Injection Date: 3/05/2019 10:46:26      Seq Line:           10
Sample Name:   1906112005                Location:           Vial 80
Acq Operator:  TNB                        Inj. No.:          1
                                           Inj. Vol.:         20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	159.0418	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	168.6882	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	BB	325841.0	5.0000	CLO4-89-ISTD
8.599	VB	308921.9	0.0000	
10.330	VBA	30210.3	0.0000	

```

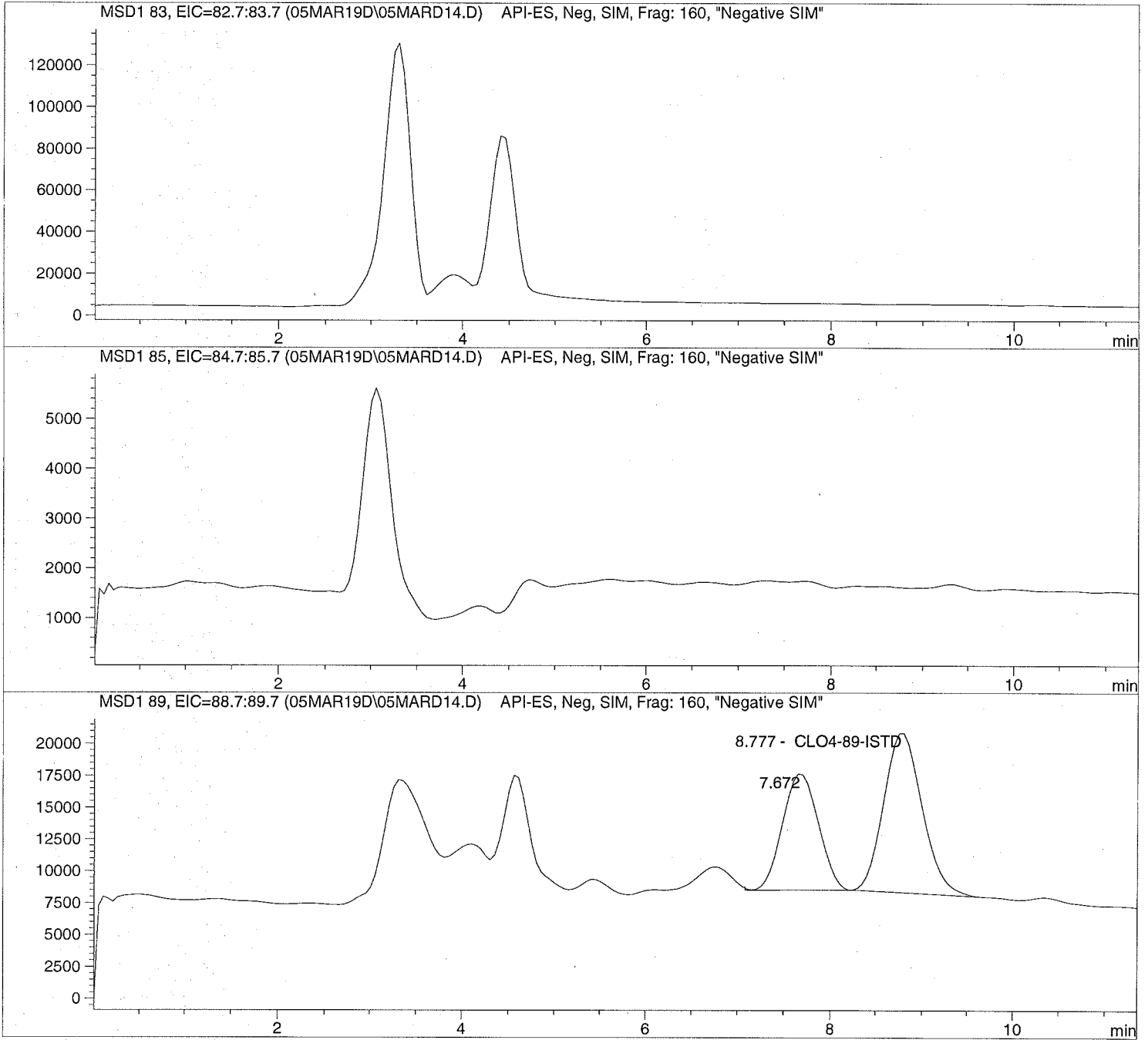
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*** End of Report ***
=====

```

Injection Date: 3/05/2019 11:39:24 Seq Line: 14
Sample Name: 1906112009 Location: Vial 84
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Injection Date: 3/05/2019 11:39:24 Seq Line: 14
Sample Name: 1906112009 Location: Vial 84
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	0.0000	
8.777	VBA	362717.3	5.0000	CLO4-89-ISTD

*** End of Report ***



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 18, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030014**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 01, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

Generated By: DAYNA.FISHER
RJ Modashia
Project Manager

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030014

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030014-01	LH18/24-SP650_022819	Water		28-Feb-2019 14:00	01-Mar-2019 08:50	<input type="checkbox"/>
HS19030014-02	LH18/24-SP650_022819_BIX	Water		28-Feb-2019 14:00	01-Mar-2019 08:50	<input type="checkbox"/>

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
-

Work Order Comments

- The analysis for TOC was subcontracted to ALS Environmental in Kelso, WA. Final Report attached.
-

WetChemistry by Method E350.3**Batch ID: R333997**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

WetChemistry by Method E365.3**Batch ID: R333840**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_022819
 Collection Date: 28-Feb-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030014
 Lab ID:HS19030014-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
AMMONIA AS N BY E350.3(ISE)								Analyst: MZD
Nitrogen, Ammonia (As N)	9.5		0.20	0.20	0.20	mg/L	1	05-Mar-2019 13:40
ORTHO PHOSPHATE (PO4) AS P BY E365.3								Analyst: MZD
Phosphorus, Total Orthophosphate (As P)	1.69		0.100	0.250	0.250	mg/L	10	01-Mar-2019 12:00
SUBCONTRACT ANALYSIS - TOC ANALYSIS								Analyst: SUBK
Subcontract Analysis	See Attached		0	0		NA	1	18-Mar-2019 10:26

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_022819_BIX
 Collection Date: 28-Feb-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19030014
 Lab ID:HS19030014-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA		Analyst: SUB				
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030014

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R333840	Test Name : ORTHO PHOSPHATE (PO4) AS P BY E365.3		Matrix: Water			
HS19030014-01	LH18/24-SP650_022819	28 Feb 2019 14:00			01 Mar 2019 12:00	10
Batch ID R333997	Test Name : AMMONIA AS N BY E350.3(ISE)		Matrix: Water			
HS19030014-01	LH18/24-SP650_022819	28 Feb 2019 14:00			05 Mar 2019 13:40	1
Batch ID R334176	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19030014-02	LH18/24-SP650_022819_BIX	28 Feb 2019 14:00			07 Mar 2019 17:47	1
Batch ID R334688	Test Name : SUBCONTRACT ANALYSIS - TOC ANALYSIS		Matrix: Water			
HS19030014-01	LH18/24-SP650_022819	28 Feb 2019 14:00			18 Mar 2019 10:26	1

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030014

QC BATCH REPORT NEW

Batch ID:	R333840 (0)	Instrument:	UV-2450	Method:	ORTHO PHOSPHATE (PO4) AS P BY E365.3					
MBLK	Sample ID: MBLK-333840	Units: mg/L		Analysis Date: 01-Mar-2019 12:00						
Client ID:		Run ID: UV-2450_333840		SeqNo: 4971681	PrepDate:	DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.0250	0.0250							U	
LCS	Sample ID: LCS-333840	Units: mg/L		Analysis Date: 01-Mar-2019 12:00						
Client ID:		Run ID: UV-2450_333840		SeqNo: 4971682	PrepDate:	DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.227	0.0250	0.25	0	90.8	85 - 115				
MS	Sample ID: HS19030014-01MS	Units: mg/L		Analysis Date: 01-Mar-2019 12:00						
Client ID: LH18/24-SP650_022819		Run ID: UV-2450_333840		SeqNo: 4971684	PrepDate:	DF: 10				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	4.17	0.250	2.5	1.69	99.2	80 - 120				
MSD	Sample ID: HS19030014-01MSD	Units: mg/L		Analysis Date: 01-Mar-2019 12:00						
Client ID: LH18/24-SP650_022819		Run ID: UV-2450_333840		SeqNo: 4971685	PrepDate:	DF: 10				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	4.21	0.250	2.5	1.69	101	80 - 120	4.17	0.955	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030014

QC BATCH REPORT NEW

Batch ID: R333997 (0)		Instrument: WetChem_HS		Method: AMMONIA AS N BY E350.3(ISE)					
MBLK	Sample ID: MBLK-333997	Units: mg/L		Analysis Date: 05-Mar-2019 13:40					
Client ID:	Run ID: WetChem_HS_333997	SeqNo: 4975026		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Nitrogen, Ammonia (As N)	0.20	0.20							U
LCS	Sample ID: LCS-333997	Units: mg/L		Analysis Date: 05-Mar-2019 13:40					
Client ID:	Run ID: WetChem_HS_333997	SeqNo: 4975027		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Nitrogen, Ammonia (As N)	9.903	0.20	10	0	99.0	80 - 120			
MS	Sample ID: HS19021330-01MS	Units: mg/L		Analysis Date: 05-Mar-2019 13:40					
Client ID:	Run ID: WetChem_HS_333997	SeqNo: 4975029		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Nitrogen, Ammonia (As N)	10.68	0.20	10	0.2419	104	80 - 120			
MSD	Sample ID: HS19021330-01MSD	Units: mg/L		Analysis Date: 05-Mar-2019 13:40					
Client ID:	Run ID: WetChem_HS_333997	SeqNo: 4975030		PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual
Nitrogen, Ammonia (As N)	10.59	0.20	10	0.2419	103	80 - 120	10.68	0.846	20

The following samples were analyzed in this batch: HS19030014-01

ALS Houston, US

Date: 18-Mar-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19030014	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

ALS Houston, US

Date: 18-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030014

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19030014-01	LH18/24-SP650_022819	Login	3/1/2019 10:40:31 AM	JRM	WET371
HS19030014-01	LH18/24-SP650_022819	Login	3/1/2019 10:40:31 AM	JRM	WET371
HS19030014-01	LH18/24-SP650_022819	Login	3/1/2019 10:40:31 AM	JRM	Sub
HS19030014-02	LH18/24-SP650_022819_BIX	Login	3/1/2019 10:40:31 AM	JRM	Sub

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030014

Date/Time Received: **01-Mar-2019 08:50**
 Received by: **JRM**

Checklist completed by: Jared R. Makan 1-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 1-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.8c/1.8c UC/C IR25
 Cooler(s)/Kit(s): 43795
 Date/Time sample(s) sent to storage: 03/01/2019 10:45

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd. Suite 210 Houston, TX. 77099 (281) 530-5656 ATTN: R.J Modshia

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001			Analyses												Remarks (Preservatives, etc.)	Lab I.D.#
Job: GROUNDWATER TREATMENT PLANT WEEKLY SAMPLES																			
Prepared By: Scott Beesinger						P.O. Number													
Field Sample I.D.	Sample Matrix	Date / Time	MS / MSD	No. OF CONTAINERS	AMMONIA-N	TOTAL ORGANIC CARBON	ORTHO-PHOSPHATE	PERCHLORATE											
LH18/24-SP650_022819	Water	02/28/19 / 14:00		2	X	X												H2SO4	
LH18/24-SP650_022819	Water	02/28/19 / 14:00		1			X											NONE	
LH18/24-SP650_022819_BIX	Water	02/28/19 / 14:00		1				X										NONE	
Additional Remarks: Standard TAT on all parameters																			
Relinquished By: <i>Scott Beesinger</i>	Date 02/28/19	Time 14:30	Received By: J. WALKER	Date 3/1/19	Time 08:50	Relinquished By:	Date	Time	Received By:	Date	Time								

Received At Lab By:										For Lab Use Only									
			Date	Time	Airbill No.	Opened By:			Date	Time	Temp of Container	Seal No.	Condition						
Remarks: Cooler 43795 Temp 1.8 MLRS CFO.O																			

HS19030014
Bhate Environmental Associates, Inc.
118/24 Longhorn GW Treatment Plant Weekly Sample



TRK# 4380 9530 9423
0221


FRI - 01 MAR 10:30A
PRIORITY OVERNIGHT

AB SGRA

77099
TX-US
IAH



3646008 01Mar 00:38 AFWM 547C2/0F20/A1/LC

 ALS Environmental 10450 Stancliff Rd., Suite 210 Houston, Texas 77009 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CLIST Date: 2/29/19 Name: SCOTT B Company: E.H.A.
	(Empty space for additional information)

BODY SEAL	Seal Broken By:
Time: 1430	JM
E. SINGAR	Date:
ET	3/01/19



ALS Environmental
ALS Group USA, Corp
1317 South 13th Avenue
Kelso, WA 98626
T : +1 360 577 7222
F : +1 360 636 1068
www.alsglobal.com

March 18, 2019

Analytical Report for Service Request No: K1901895

RJ Modashia
ALS Laboratory Group
10450 Stancliff Road
Suite 210
Houston, TX 77099-4338

RE: HS19030014 / HS19030014

Dear RJ,

Enclosed are the results of the sample(s) submitted to our laboratory March 05, 2019
For your reference, these analyses have been assigned our service request number **K1901895**.

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. The test results meet requirements of the current NELAP standards, where applicable, and except as noted in the laboratory case narrative provided. For a specific list of NELAP-accredited analytes, refer to the certifications section at www.alsglobal.com. All results are intended to be considered in their entirety, and ALS Group USA Corp. dba ALS Environmental (ALS) is not responsible for use of less than the complete report. Results apply only to the items submitted to the laboratory for analysis and individual items (samples) analyzed, as listed in the report.

Please contact me if you have any questions. My extension is 3350. You may also contact me via email at Kelley.Lovejoy@alsglobal.com.

Respectfully submitted,

ALS Group USA, Corp. dba ALS Environmental

Kelley Lovejoy

Kelley Lovejoy
Project Manager



ALS Environmental
ALS Group USA, Corp
1317 South 13th Avenue
Kelso, WA 98626
T : +1 360 577 7222
F : +1 360 636 1068
www.alsglobal.com

Table of Contents

Acronyms

Qualifiers

State Certifications, Accreditations, And Licenses

Case Narrative

Chain of Custody

General Chemistry

Raw Data

 General Chemistry

Acronyms

ASTM	American Society for Testing and Materials
A2LA	American Association for Laboratory Accreditation
CARB	California Air Resources Board
CAS Number	Chemical Abstract Service registry Number
CFC	Chlorofluorocarbon
CFU	Colony-Forming Unit
DEC	Department of Environmental Conservation
DEQ	Department of Environmental Quality
DHS	Department of Health Services
DOE	Department of Ecology
DOH	Department of Health
EPA	U. S. Environmental Protection Agency
ELAP	Environmental Laboratory Accreditation Program
GC	Gas Chromatography
GC/MS	Gas Chromatography/Mass Spectrometry
LOD	Limit of Detection
LOQ	Limit of Quantitation
LUFT	Leaking Underground Fuel Tank
M	Modified
MCL	Maximum Contaminant Level is the highest permissible concentration of a substance allowed in drinking water as established by the USEPA.
MDL	Method Detection Limit
MPN	Most Probable Number
MRL	Method Reporting Limit
NA	Not Applicable
NC	Not Calculated
NCASI	National Council of the Paper Industry for Air and Stream Improvement
ND	Not Detected
NIOSH	National Institute for Occupational Safety and Health
PQL	Practical Quantitation Limit
RCRA	Resource Conservation and Recovery Act
SIM	Selected Ion Monitoring
TPH	Total Petroleum Hydrocarbons
tr	Trace level is the concentration of an analyte that is less than the PQL but greater than or equal to the MDL.

Inorganic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- E The result is an estimate amount because the value exceeded the instrument calibration range.
- J The result is an estimated value.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.
- H The holding time for this test is immediately following sample collection. The samples were analyzed as soon as possible after receipt by the laboratory.

Metals Data Qualifiers

- # The control limit criteria is not applicable. See case narrative.
- J The result is an estimated value.
- E The percent difference for the serial dilution was greater than 10%, indicating a possible matrix interference in the sample.
- M The duplicate injection precision was not met.
- N The Matrix Spike sample recovery is not within control limits. See case narrative.
- S The reported value was determined by the Method of Standard Additions (MSA).
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- W The post-digestion spike for furnace AA analysis is out of control limits, while sample absorbance is less than 50% of spike absorbance.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- + The correlation coefficient for the MSA is less than 0.995.
- Q See case narrative. One or more quality control criteria was outside the limits.

Organic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- A A tentatively identified compound, a suspected aldol-condensation product.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- C The analyte was qualitatively confirmed using GC/MS techniques, pattern recognition, or by comparing to historical data.
- D The reported result is from a dilution.
- E The result is an estimated value.
- J The result is an estimated value.
- N The result is presumptive. The analyte was tentatively identified, but a confirmation analysis was not performed.
- P The GC or HPLC confirmation criteria was exceeded. The relative percent difference is greater than 40% between the two analytical results.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a chromatographic interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.

Additional Petroleum Hydrocarbon Specific Qualifiers

- F The chromatographic fingerprint of the sample matches the elution pattern of the calibration standard.
- L The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of lighter molecular weight constituents than the calibration standard.
- H The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of heavier molecular weight constituents than the calibration standard.
- O The chromatographic fingerprint of the sample resembles an oil, but does not match the calibration standard.
- Y The chromatographic fingerprint of the sample resembles a petroleum product eluting in approximately the correct carbon range, but the elution pattern does not match the calibration standard.
- Z The chromatographic fingerprint does not resemble a petroleum product.

**ALS Group USA Corp. dba ALS Environmental (ALS) - Kelso
State Certifications, Accreditations, and Licenses**

Agency	Web Site	Number
Alaska DEH	http://dec.alaska.gov/eh/lab/cs/csapproval.htm	UST-040
Arizona DHS	http://www.azdhs.gov/lab/license/env.htm	AZ0339
Arkansas - DEQ	http://www.adeq.state.ar.us/techsvs/labcert.htm	88-0637
California DHS (ELAP)	http://www.cdph.ca.gov/certlic/labs/Pages/ELAP.aspx	2795
DOD ELAP	http://www.denix.osd.mil/edqw/Accreditation/AccreditedLabs.cfm	L16-58-R4
Florida DOH	http://www.doh.state.fl.us/lab/EnvLabCert/WaterCert.htm	E87412
Hawaii DOH	http://health.hawaii.gov/	-
ISO 17025	http://www.pjlabs.com/	L16-57
Louisiana DEQ	http://www.deq.louisiana.gov/page/la-lab-accreditation	03016
Maine DHS	http://www.maine.gov/dhhs/	WA01276
Minnesota DOH	http://www.health.state.mn.us/accreditation	053-999-457
Nevada DEP	http://ndep.nv.gov/bsdw/labservice.htm	WA01276
New Jersey DEP	http://www.nj.gov/dep/enforcement/oqa.html	WA005
New York - DOH	https://www.wadsworth.org/regulatory/elap	12060
North Carolina DEQ	https://deq.nc.gov/about/divisions/water-resources/water-resources-data/water-sciences-home-page/laboratory-certification-branch/non-field-lab-certification	605
Oklahoma DEQ	http://www.deq.state.ok.us/CSDnew/labcert.htm	9801
Oregon – DEQ (NELAP)	http://public.health.oregon.gov/LaboratoryServices/EnvironmentalLaboratoryAccreditation/Pages/index.aspx	WA100010
South Carolina DHEC	http://www.scdhec.gov/environment/EnvironmentalLabCertification/	61002
Texas CEQ	http://www.tceq.texas.gov/field/qa/env_lab_accreditation.html	T104704427
Washington DOE	http://www.ecy.wa.gov/programs/eap/labs/lab-accreditation.html	C544
Wyoming (EPA Region 8)	https://www.epa.gov/region8-waterops/epa-region-8-certified-drinking-water	-
Kelso Laboratory Website	www.alsglobal.com	NA

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. A complete listing of specific NELAP-certified analytes, can be found in the certification section at www.ALSGlobal.com or at the accreditation bodies web site.

Please refer to the certification and/or accreditation body's web site if samples are submitted for compliance purposes. The states highlighted above, require the analysis be listed on the state certification if used for compliance purposes and if the method/analyte is offered by that state.



Case Narrative

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Client: ALS Environmental - US
Project: HS19030014
Sample Matrix: Water

Service Request: K1901895
Date Received: 03/05/2019

CASE NARRATIVE

All analyses were performed consistent with the quality assurance program of ALS Environmental. This report contains analytical results for samples designated for Tier IV validation deliverables including summary forms and all of the associated raw data for each of the analyses. When appropriate to the method, method blank results have been reported with each analytical test.

Sample Receipt:

One water sample was received for analysis at ALS Environmental on 03/05/2019. The sample was received in good condition and consistent with the accompanying chain of custody form. The sample was stored in a refrigerator at 4°C upon receipt at the laboratory.

General Chemistry:

No significant anomalies were noted with this analysis.

Approved by *Kelley Lovejoy*

Date 03/18/2019



Chain of Custody

ALS Environmental—Kelso Laboratory
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K190189500934810

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10855

SUBCONTRACT TO:

ALS Environmental Kelso
1317 S. 13th Avenue
Kelso, WA 98626

Phone: +1 360 501 3312

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030014
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030014-01	LH18/24-SP650_022819	Water	28 Feb 2019 14:00
TOC Analysis for DOD Level IV			15 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: [Signature]
Received By: [Signature]
Cooler ID(s): _____

Date/Time: 3-4-19 1800
Date/Time: 3/5/19 1145
Temperature(s): _____

ALS IS YOUR RIGHT PARTNER



Cooler Receipt and Preservation Form

Client ALS - Houston Service Request K19 01895
 Received: 3/5/19 Opened: 3/5/19 By: BR Unloaded: 3/5/19 By: BR

1. Samples were received via? USPS Fed Ex UPS DHL PDX Courier Hand Delivered
 2. Samples were received in: (circle) Cooler Box Envelope Other NA
 3. Were custody seals on coolers? NA (Y) N If yes, how many and where? 1 front
 If present, were custody seals intact? (Y) N If present, were they signed and dated? (Y) N

Raw Cooler Temp	Corrected Cooler Temp	Raw Temp Blank	Corrected Temp Blank	Corr. Factor	Thermometer ID	Cooler/COC ID	Tracking Number	NA	Filed
-0.2	-0.4	4.6	4.4	-0.2	300	NA	480978312804		

4. Packing material: Inserts Baggies Bubble Wrap Gel Packs Wet Ice Dry Ice Sleeves
 5. Were custody papers properly filled out (ink, signed, etc.)? NA (Y) N
 6. Were samples received in good condition (temperature, unbroken)? Indicate in the table below. NA (Y) N
 If applicable, tissue samples were received: Frozen Partially Thawed Thawed
 7. Were all sample labels complete (i.e analysis, preservation, etc.)? NA (Y) N
 8. Did all sample labels and tags agree with custody papers? Indicate major discrepancies in the table on page 2. NA (Y) N
 9. Were appropriate bottles/containers and volumes received for the tests indicated? NA (Y) N
 10. Were the pH-preserved bottles (*see SMO GEN SOP*) received at the appropriate pH? Indicate in the table below NA Y N
 11. Were VOA vials received without headspace? Indicate in the table below. NA Y N
 12. Was C12/Res negative? NA Y N

Sample ID on Bottle	Sample ID on COC	Identified by:

Sample ID	Bottle Count	Bottle Type	Out of Temp	Head-space	Broke	pH	Reagent	Volume added	Reagent Lot Number	Initials	Time

Notes, Discrepancies, & Resolutions:

temp discrepancy due to sample & ice being separate from blank.



General Chemistry

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Analytical Report

Client: ALS Environmental - US
Project: HS19030014/HS19030014
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1901895
Date Collected: 02/28/19
Date Received: 03/5/19
Units: mg/L
Basis: NA

Carbon, Total Organic

Sample Name	Lab Code	Result	LOQ	LOD	MDL	Dil.	Date Analyzed	Q
LH18/24-SP650_022819	K1901895-001	5.10	0.50	0.20	0.07	1	03/13/19 12:59	
Method Blank	K1901895-MB	0.32 J	0.50	0.20	0.07	1	03/13/19 12:26	

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030014/HS19030014
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1901895
Date Collected: 02/28/19
Date Received: 03/05/19

Units: mg/L
Basis: NA

Replicate Sample Summary
Carbon, Total Organic

Sample Name:	Lab Code:	LOQ	LOD	MDL	Sample Result	Duplicate Result	Average	RPD	RPD Limit	Date Analyzed
LH18/24-SP650_022819	K1901895-001DUP	0.50	0.20	0.07	5.10	4.90	5.00	4	10	03/13/19

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030014/HS19030014
Sample Matrix: Water

Service Request: K1901895
Date Collected: N/A
Date Received: N/A
Date Analyzed: 03/13/19
Date Extracted: NA

Matrix Spike Summary
Carbon, Total Organic

Sample Name: Batch QC
Lab Code: K1901955-001
Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA

Matrix Spike
K1901955-001MS

<u>Analyte Name</u>	<u>Sample Result</u>	<u>Result</u>	<u>Spike Amount</u>	<u>% Rec</u>	<u>% Rec Limits</u>
Carbon, Total Organic	3.02	29.7	25.0	107	83-117

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030014/HS19030014
Sample Matrix: Water

Service Request: K1901895
Date Analyzed: 03/13/19
Date Extracted: NA

Lab Control Sample Summary
Carbon, Total Organic

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA
Analysis Lot: 628234

Sample Name	Lab Code	Result	Spike Amount	% Rec	% Rec Limits
Lab Control Sample	K1901895-LCS	25.8	25.0	103	83-117

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030014/HS19030014

Service Request: K1901895**Continuing Calibration Verification (CCV) Summary****Carbon, Total Organic****Analysis Method:** SM 5310 C**Units:** mg/L

	Analysis		Date	True	Measured	Percent	Acceptance Limits
	Lot	Lab Code	Analyzed	Value	Value	Recovery	
CCV1	628234	KQ1903351-05	03/13/19 06:29	25.0	24.7	99	90-110
CCV2	628234	KQ1903351-06	03/13/19 11:53	25.0	24.4	98	90-110
CCV3	628234	KQ1903351-07	03/13/19 17:00	25.0	24.6	98	90-110
CCV4	628234	KQ1903351-08	03/14/19 00:30	25.0	24.3	97	90-110

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030014/HS19030014

Service Request: K1901895

Continuing Calibration Blank (CCB) Summary
Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis Lot	Lab Code	Date Analyzed	LOQ	LOD	MDL	Result	Q
CCB1	628234	KQ1903351-01	03/13/19 06:46	0.50	0.20	0.07	ND	U
CCB2	628234	KQ1903351-02	03/13/19 12:09	0.50	0.20	0.07	ND	U
CCB3	628234	KQ1903351-03	03/13/19 17:17	0.50	0.20	0.07	ND	U
CCB4	628234	KQ1903351-04	03/14/19 00:47	0.50	0.20	0.07	ND	U



Raw Data

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General Chemistry

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Work Request # ^{Original} () K1902025, 2074, 1653, 1698, 1701, 1719, 1940, 1830, 1875, 1895, 1926, 1955, 1956, 2084, 1725, 1786, 1941
 Tier: II II II II II II I II II IV I IV IV IV II IV II
 Date Analyzed: 3/12/2019
 Analyst: BCD
 Analysis: TOC/DOC Run # TOC: 628233
628234
DOC: 628232
628235

**DATA QUALITY REPORT
INORGANICS**

Explain any "no" responses to questions below, and any corrective actions in the comments section below.

1. Is the method name and number correct and appropriate? yes/no/NA
2. Holding times met for all analyses and for all samples? yes/no/NA
3. Are calculations correct? yes/no/NA
4. Is the reporting basis correct? (Dry Weight) yes/no/NA
5. All quality control criteria met? yes/no/NA
6. Is the calibration curve correlation coefficient ≥ 0.995 ? yes/no
7. MBs, CCVs, CCBs, LCSs, Dups, and Spikes, analyzed at proper frequency? yes/no/NA
8. Are ICVs, CCVs, and CCBs all within acceptance limits? yes/no/NA
9. Are results for methods blanks all ND? yes/no/NA
10. Are all QC samples within acceptance criteria? (LCS % rec, MS/DMS % rec, DUP or MS/DMS RPDs, etc.) yes/no/NA
11. Are all exceptions explained? yes/no/NA
12. Have all applicable service requests been reviewed? yes/no/NA
13. Are all samples labeled correctly? yes/no/NA
14. Have all instructions on the service request been followed? (e.g. Special MRLs, QC on a specific sample, Form V) yes/no/NA
15. Are detection limits and units reported correctly? yes/no/NA
16. Is the unused space on the benchsheet crossed out? yes/no/NA
17. Was analysis turned in by the due date? (n-2) (If not record SR#) yes/no/NA

Q12D 3/15/19

COMMENTS: *K1902074-1/2, 1653-1/2/3/4/5/6, 1719-1/2/3/4/5/6/7, 1698-1/2/3, 1701-1, and 1940-1 sent for RA due to high CV.
 K1902084-1/1/1, 1926-3/1/1 report a high % RSD due to non-homogeneous dirty sample.
 K1901975-2/2/4, 1926-1/1/1, 1941-1/1/1/1 report a high % RSD, but those samples are less than six the MRL
 K1901926-2/1/1, 1941-1 require a minimum dilution due to dirty samples.
 K1901941-7/1/1's report a high % RSD and low recovery due to dirty sample interference.*

Final Approved by: *H. [Signature]* Date: 03/15/19 DQREPORT

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628232

Method/Testcode: SM 5310 C/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC? T
K1902025-002	Carbon, Dissolved Organic (DOC)	N/A		Water	3.24 mg/L	10 ml	3.24 mg/L	1	0.07	0.50			3/12/19 17:50	N II
K1902025-004	Carbon, Dissolved Organic (DOC)	N/A		Water	2.46 mg/L	10 ml	2.46 mg/L	1	0.07	0.50			3/12/19 18:55	N II
K1902074-001	Carbon, Dissolved Organic (DOC)	N/A		Effluent	1.39 mg/L	10 ml	1.39 mg/L	1	0.07	0.50			3/12/19 20:01	N II
K1902074-002	Carbon, Dissolved Organic (DOC)	N/A		Water	1.49 mg/L	10 ml	1.49 mg/L	1	0.07	0.50			3/12/19 20:33	N II
KQ1903346-01	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/12/19 14:04	N II
KQ1903346-02	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.09 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/12/19 19:44	N II
KQ1903346-03	Carbon, Dissolved Organic (DOC)	CCB		Water	1.40 mg/L	10 ml	1.40 mg/L	1	0.07	0.50			3/13/19 01:22	N II
KQ1903346-04	Carbon, Dissolved Organic (DOC)	CCV		Water	24.82 mg/L	10 ml	24.8 mg/L	1			99		3/12/19 13:47	N II
KQ1903346-05	Carbon, Dissolved Organic (DOC)	CCV		Water	25.19 mg/L	10 ml	25.2 mg/L	1			101		3/12/19 19:27	N II
KQ1903346-06	Carbon, Dissolved Organic (DOC)	CCV		Water	24.85 mg/L	10 ml	24.8 mg/L	1			99		3/13/19 01:06	N II
KQ1903346-07	Carbon, Dissolved Organic (DOC)	MS	K1902025-002	Water	30.63 mg/L	10 ml	30.6 mg/L	1	0.07	0.50	110		3/12/19 18:22	N II
KQ1903346-08	Carbon, Dissolved Organic (DOC)	DUP	K1902025-002	Water	3.13 mg/L	10 ml	3.13 mg/L	1	0.07	0.50		3	3/12/19 17:50	N II
KQ1903346-09	Carbon, Dissolved Organic (DOC)	DUP	K1902025-004	Water	2.68 mg/L	10 ml	2.68 mg/L	1	0.07	0.50		9	3/12/19 18:55	N II
KQ1903346-10	Carbon, Dissolved Organic (DOC)	DUP	K1902074-001	Effluent	1.77 mg/L	10 ml	1.77 mg/L	1	0.07	0.50		24*	3/12/19 20:01	N II
KQ1903346-11	Carbon, Dissolved Organic (DOC)	DUP	K1902074-002	Water	1.47 mg/L	10 ml	1.47 mg/L	1	0.07	0.50		1	3/12/19 20:33	N II
KQ1903346-12	Carbon, Dissolved Organic (DOC)	MB		Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/12/19 14:21	N II
KQ1903346-13	Carbon, Dissolved Organic (DOC)	LCS		Water	25.83 mg/L	10 ml	25.8 mg/L	1	0.07	0.50	103		3/12/19 14:37	N II

03/15/19
[Handwritten Signature]

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628233 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1901653-001	Carbon, Total Organic	N/A		Brackish Water	0.91 mg/L	10 ml	1.8 mg/L	2	0.2	1.0			3/12/19 21:05	N	II
K1901653-002	Carbon, Total Organic	N/A		Brackish Water	0.45 mg/L	10 ml	0.9 mg/L J	2	0.2	1.0			3/12/19 21:37	N	II
K1901653-003	Carbon, Total Organic	N/A		Brackish Water	0.76 mg/L	10 ml	1.5 mg/L	2	0.2	1.0			3/12/19 22:09	N	II
K1901653-004	Carbon, Total Organic	N/A		Brackish Water	0.82 mg/L	10 ml	1.6 mg/L	2	0.2	1.0			3/12/19 22:41	N	II
K1901653-005	Carbon, Total Organic	N/A		Brackish Water	0.43 mg/L	10 ml	0.9 mg/L J	2	0.2	1.0			3/12/19 23:45	Y	II
K1901653-006	Carbon, Total Organic	N/A		Brackish Water	0.57 mg/L	10 ml	1.1 mg/L	2	0.2	1.0			3/12/19 23:13	N	II
K1901698-001	Carbon, Total Organic	N/A		Water	0.48 mg/L	10 ml	5.0 mg/L U	10	0.7	5.0			3/13/19 07:03	N	II
K1901698-002	Carbon, Total Organic	N/A		Water	2.16 mg/L	10 ml	21.6 mg/L	10	0.7	5.0			3/13/19 08:23	N	II
K1901698-003	Carbon, Total Organic	N/A		Water	-0.18 mg/L	10 ml	5.0 mg/L U	10	0.7	5.0			3/13/19 08:55	N	II
K1901701-001	Carbon, Total Organic	N/A		Water	7.35 mg/L	10 ml	73.5 mg/L	10	0.7	5.0			3/13/19 09:28	N	II
K1901719-001	Carbon, Total Organic	N/A		Brackish Water	0.60 mg/L	10 ml	1.2 mg/L	2	0.2	1.0			3/13/19 02:12	N	II
K1901719-002	Carbon, Total Organic	N/A		Brackish Water	0.87 mg/L	10 ml	1.7 mg/L	2	0.2	1.0			3/13/19 02:44	N	II
K1901719-003	Carbon, Total Organic	N/A		Brackish Water	18.64 mg/L	10 ml	37.3 mg/L	2	0.2	1.0			3/13/19 03:17	N	II
K1901719-004	Carbon, Total Organic	N/A		Brackish Water	1.15 mg/L	10 ml	4.6 mg/L	4	0.3	2.0			3/13/19 03:49	N	II
K1901719-005	Carbon, Total Organic	N/A		Brackish Water	1.95 mg/L	10 ml	3.9 mg/L	2	0.2	1.0			3/13/19 04:21	N	II
K1901719-006	Carbon, Total Organic	N/A		Brackish Water	0.76 mg/L	10 ml	1.5 mg/L	2	0.2	1.0			3/13/19 04:53	N	II
K1901719-007	Carbon, Total Organic	N/A		Brackish Water	6.40 mg/L	10 ml	25.6 mg/L	4	0.3	2.0			3/13/19 05:25	N	II
K1901940-001	Carbon, Total Organic	N/A		Drinking Water	0.27 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/13/19 10:32	N	I
KQ1903349-02	Carbon, Total Organic	CCB		Brackish Water	-0.09 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/12/19 19:44	N	II
KQ1903349-03	Carbon, Total Organic	CCB		Brackish Water	1.40 mg/L	10 ml	1.40 mg/L	1	0.07	0.50			3/13/19 01:22	N	II
KQ1903349-04	Carbon, Total Organic	CCB		Brackish Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/13/19 06:46	N	II
KQ1903349-05	Carbon, Total Organic	CCB		Brackish Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/13/19 12:09	N	II
KQ1903349-07	Carbon, Total Organic	CCV		Brackish Water	25.19 mg/L	10 ml	25.2 mg/L	1			101		3/12/19 19:27	N	II
KQ1903349-08	Carbon, Total Organic	CCV		Brackish Water	24.85 mg/L	10 ml	24.8 mg/L	1			99		3/13/19 01:06	N	II
KQ1903349-09	Carbon, Total Organic	CCV		Brackish Water	24.74 mg/L	10 ml	24.7 mg/L	1			99		3/13/19 06:29	N	II
KQ1903349-10	Carbon, Total Organic	CCV		Brackish Water	24.42 mg/L	10 ml	24.4 mg/L	1			98		3/13/19 11:53	N	II
KQ1903349-11	Carbon, Total Organic	LCS		Brackish Water	26.52 mg/L	10 ml	26.5 mg/L	1	0.07	0.50	106		3/13/19 01:56	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Printed 3/15/19 10:11

Results Summary

03/15/19
[Signature]

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628233 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1903349-12	Carbon, Total Organic	MB		Brackish Water	0.12 mg/L	10 ml	0.12 mg/L J	1	0.07	0.50			3/13/19 01:39	N	II
KQ1903349-13	Carbon, Total Organic	MS	K1901653-005	Brackish Water	20.19 mg/L	10 ml	40.4 mg/L	2	0.2	1.0	79*		3/13/19 00:17	N	II
KQ1903349-14	Carbon, Total Organic	MS	K1901698-001	Water	27.78 mg/L	10 ml	278 mg/L	10	0.7	5.0	111		3/13/19 07:35	N	II
KQ1903349-15	Carbon, Total Organic	DUP	K1901653-001	Brackish Water	0.82 mg/L	10 ml	1.6 mg/L	2	0.2	1.0		11*	3/12/19 21:05	N	II
KQ1903349-16	Carbon, Total Organic	DUP	K1901653-002	Brackish Water	0.67 mg/L	10 ml	1.3 mg/L	2	0.2	1.0		39*	3/12/19 21:37	N	II
KQ1903349-17	Carbon, Total Organic	DUP	K1901653-003	Brackish Water	0.67 mg/L	10 ml	1.3 mg/L	2	0.2	1.0		12*	3/12/19 22:09	N	II
KQ1903349-18	Carbon, Total Organic	DUP	K1901653-004	Brackish Water	0.83 mg/L	10 ml	1.7 mg/L	2	0.2	1.0		<1	3/12/19 22:41	N	II
KQ1903349-19	Carbon, Total Organic	DUP	K1901653-006	Brackish Water	0.44 mg/L	10 ml	0.9 mg/L J	2	0.2	1.0		26*	3/12/19 23:13	N	II
KQ1903349-20	Carbon, Total Organic	DUP	K1901653-005	Brackish Water	0.34 mg/L	10 ml	0.7 mg/L J	2	0.2	1.0		23*	3/12/19 23:45	N	II
KQ1903349-21	Carbon, Total Organic	DUP	K1901719-001	Brackish Water	0.44 mg/L	10 ml	0.9 mg/L J	2	0.2	1.0		29*	3/13/19 02:12	N	II
KQ1903349-22	Carbon, Total Organic	DUP	K1901719-002	Brackish Water	0.82 mg/L	10 ml	1.6 mg/L	2	0.2	1.0		6	3/13/19 02:44	N	II
KQ1903349-23	Carbon, Total Organic	DUP	K1901719-003	Brackish Water	18.78 mg/L	10 ml	37.6 mg/L	2	0.2	1.0		<1	3/13/19 03:17	N	II
KQ1903349-24	Carbon, Total Organic	DUP	K1901719-004	Brackish Water	1.06 mg/L	10 ml	4.3 mg/L	4	0.3	2.0		8	3/13/19 03:49	N	II
KQ1903349-25	Carbon, Total Organic	DUP	K1901719-005	Brackish Water	1.80 mg/L	10 ml	3.6 mg/L	2	0.2	1.0		8	3/13/19 04:21	N	II
KQ1903349-26	Carbon, Total Organic	DUP	K1901719-006	Brackish Water	0.68 mg/L	10 ml	1.4 mg/L	2	0.2	1.0		12*	3/13/19 04:53	N	II
KQ1903349-27	Carbon, Total Organic	DUP	K1901719-007	Brackish Water	6.26 mg/L	10 ml	25.1 mg/L	4	0.3	2.0		2	3/13/19 05:25	N	II
KQ1903349-28	Carbon, Total Organic	DUP	K1901698-001	Water	0.52 mg/L	10 ml	5.2 mg/L	10	0.7	5.0		NC	3/13/19 07:03	N	II
KQ1903349-29	Carbon, Total Organic	DUP	K1901698-002	Water	2.10 mg/L	10 ml	21.0 mg/L	10	0.7	5.0		3	3/13/19 08:23	N	II
KQ1903349-30	Carbon, Total Organic	DUP	K1901698-003	Water	-0.18 mg/L	10 ml	5.0 mg/L U	10	0.7	5.0		NC	3/13/19 08:55	N	II
KQ1903349-31	Carbon, Total Organic	DUP	K1901701-001	Water	7.35 mg/L	10 ml	73.5 mg/L	10	0.7	5.0		<1	3/13/19 09:28	N	II
KQ1903349-32	Carbon, Total Organic	DUP	K1901940-001	Drinking Water	0.37 mg/L	10 ml	0.37 mg/L J	1	0.07	0.50		NC	3/13/19 10:32	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628234 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1901830-001	Carbon, Total Organic	N/A		Drinking Water	0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/13/19 16:12	N	II
K1901875-001	Carbon, Total Organic	N/A		Water	5.56 mg/L	10 ml	556 mg/L	100	7	50			3/13/19 17:34	N	II
K1901875-002	Carbon, Total Organic	N/A		Water	0.61 mg/L	10 ml	61 mg/L	100	7	50			3/13/19 18:06	N	II
K1901875-003	Carbon, Total Organic	N/A		Water	1.34 mg/L	10 ml	134 mg/L	100	7	50			3/13/19 18:38	N	II
K1901895-001	Carbon, Total Organic	N/A		Water	5.10 mg/L	10 ml	5.10 mg/L	1	0.07	0.50			3/13/19 12:59	N	IV
K1901926-001	Carbon, Total Organic	N/A		Ground Water	0.60 mg/L	10 ml	60 mg/L	100	7	50			3/13/19 19:43	N	I
K1901926-002	Carbon, Total Organic	N/A		Ground Water	0.07 mg/L	10 ml	50 mg/L U	100	7	50			3/13/19 20:15	N	I
K1901926-003	Carbon, Total Organic	N/A		Ground Water	2.38 mg/L	10 ml	2.38 mg/L	1	0.07	0.50			3/13/19 20:47	N	I
K1901955-001	Carbon, Total Organic	N/A		Water	3.02 mg/L	10 ml	3.02 mg/L	1	0.07	0.50			3/13/19 11:04	Y	IV
K1901956-001	Carbon, Total Organic	N/A		Water	30.12 mg/L	10 ml	30.1 mg/L	1	0.07	0.50			3/13/19 13:31	N	IV
K1901956-002	Carbon, Total Organic	N/A		Water	31.32 mg/L	10 ml	31.3 mg/L	1	0.07	0.50			3/13/19 14:04	N	IV
K1902084-001	Carbon, Total Organic	N/A		Water	4.41 mg/L	10 ml	4.41 mg/L	1	0.07	0.50			3/13/19 14:36	N	IV
K1902084-002	Carbon, Total Organic	N/A		Water	3.20 mg/L	10 ml	3.20 mg/L	1	0.07	0.50			3/13/19 15:08	N	IV
K1902084-003	Carbon, Total Organic	N/A		Water	3.31 mg/L	10 ml	3.31 mg/L	1	0.07	0.50			3/13/19 15:40	N	IV
KQ1903351-01	Carbon, Total Organic	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/13/19 06:46	N	IV
KQ1903351-02	Carbon, Total Organic	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/13/19 12:09	N	IV
KQ1903351-03	Carbon, Total Organic	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/13/19 17:17	N	IV
KQ1903351-04	Carbon, Total Organic	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/14/19 00:47	N	IV
KQ1903351-05	Carbon, Total Organic	CCV		Water	24.74 mg/L	10 ml	24.7 mg/L	1					3/13/19 06:29	N	IV
KQ1903351-06	Carbon, Total Organic	CCV		Water	24.42 mg/L	10 ml	24.4 mg/L	1					3/13/19 11:53	N	IV
KQ1903351-07	Carbon, Total Organic	CCV		Water	24.59 mg/L	10 ml	24.6 mg/L	1					3/13/19 17:00	N	IV
KQ1903351-08	Carbon, Total Organic	CCV		Water	24.30 mg/L	10 ml	24.3 mg/L	1					3/14/19 00:30	N	IV
KQ1903351-09	Carbon, Total Organic	MB		Water	0.32 mg/L	10 ml	0.32 mg/L J	1	0.07	0.50			3/13/19 12:26	N	IV
KQ1903351-10	Carbon, Total Organic	LCS		Water	25.77 mg/L	10 ml	25.8 mg/L	1	0.07	0.50	103		3/13/19 12:43	N	IV
KQ1903351-11	Carbon, Total Organic	MS	K1901955-001	Water	29.69 mg/L	10 ml	29.7 mg/L	1	0.07	0.50	107		3/13/19 11:36	N	IV
KQ1903351-12	Carbon, Total Organic	DUP	K1901955-001	Water	3.04 mg/L	10 ml	3.04 mg/L	1	0.07	0.50		<1	3/13/19 11:04	N	IV
KQ1903351-13	Carbon, Total Organic	DUP	K1901895-001	Water	4.90 mg/L	10 ml	4.90 mg/L	1	0.07	0.50		4	3/13/19 12:59	N	IV
KQ1903351-14	Carbon, Total Organic	DUP	K1901956-001	Water	30.97 mg/L	10 ml	31.0 mg/L	1	0.07	0.50		3	3/13/19 13:31	N	IV
KQ1903351-15	Carbon, Total Organic	DUP	K1901956-002	Water	31.78 mg/L	10 ml	31.8 mg/L	1	0.07	0.50		1	3/13/19 14:04	N	IV
KQ1903351-16	Carbon, Total Organic	DUP	K1902084-001	Water	3.61 mg/L	10 ml	3.61 mg/L	1	0.07	0.50		20*	3/13/19 14:36	N	IV
KQ1903351-17	Carbon, Total Organic	DUP	K1902084-002	Water	2.91 mg/L	10 ml	2.91 mg/L	1	0.07	0.50		10	3/13/19 15:08	N	IV
KQ1903351-18	Carbon, Total Organic	DUP	K1902084-003	Water	3.23 mg/L	10 ml	3.23 mg/L	1	0.07	0.50		2	3/13/19 15:40	N	IV
KQ1903351-19	Carbon, Total Organic	DUP	K1901830-001	Drinking Water	0.15 mg/L	10 ml	0.15 mg/L J	1	0.07	0.50		NC	3/13/19 16:12	N	II
KQ1903351-20	Carbon, Total Organic	DUP	K1901875-001	Water	5.53 mg/L	10 ml	553 mg/L	100	7	50		<1	3/13/19 17:34	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628234 Method/Testcode: SM 5310 C/TOC T

<u>Lab Code</u>	<u>Target Analytes</u>	<u>QC</u>	<u>Parent Sample</u>	<u>Matrix</u>	<u>Raw Result</u>	<u>Sample Amt.</u>	<u>Final Result</u>	<u>Dil</u>	<u>MDL</u>	<u>POL</u>	<u>% Rec</u>	<u>% RSD</u>	<u>Date Analyzed</u>	<u>QC?</u>	<u>Tier</u>
KQ1903351-21	Carbon, Total Organic	DUP	K1901875-002	Water	0.55 mg/L	10 ml	55 mg/L	100	7	50		11*	3/13/19 18:06	N	II
KQ1903351-22	Carbon, Total Organic	DUP	K1901875-003	Water	1.23 mg/L	10 ml	123 mg/L	100	7	50		8	3/13/19 18:38	N	II
KQ1903351-23	Carbon, Total Organic	DUP	K1901926-001	Ground Water	0.45 mg/L	10 ml	45 mg/L J	100	7	50		27*	3/13/19 19:43	N	I
KQ1903351-24	Carbon, Total Organic	DUP	K1901926-002	Ground Water	3.00 mg/L	10 ml	300 mg/L	100	7	50		NC	3/13/19 20:15	N	I
KQ1903351-25	Carbon, Total Organic	DUP	K1901926-003	Ground Water	1.92 mg/L	10 ml	1.92 mg/L	1	0.07	0.50		21*	3/13/19 20:47	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

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Results Summary

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Analytical Results Summary

00934827

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628235 Method/Testcode: 9060/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1901725-001	Carbon, Dissolved Organic (DOC)	N/A		Water	2.68 mg/L	10 ml	10.7 mg/L	4	0.3	2.0			3/13/19 21:19	N	II
K1901725-002	Carbon, Dissolved Organic (DOC)	N/A		Water	1.59 mg/L	10 ml	6.4 mg/L	4	0.3	2.0			3/13/19 22:22	N	II
K1901725-003	Carbon, Dissolved Organic (DOC)	N/A		Water	2.36 mg/L	10 ml	9.4 mg/L	4	0.3	2.0			3/13/19 23:26	Y	II
K1901786-001	Carbon, Dissolved Organic (DOC)	N/A		Water	0.28 mg/L	10 ml	0.28 mg/L	J 1	0.07	0.50			3/14/19 03:13	N	IV
K1901786-002	Carbon, Dissolved Organic (DOC)	N/A		Water	1.50 mg/L	10 ml	750 mg/L	500	40	250			3/14/19 04:16	Y	IV
K1901941-001	Carbon, Dissolved Organic (DOC)	N/A		Water	0.18 mg/L	10 ml	1.8 mg/L	J 10	0.7	5.0			3/14/19 06:56	N	II
K1901941-002	Carbon, Dissolved Organic (DOC)	N/A		Water	0.56 mg/L	10 ml	5.6 mg/L	10	0.7	5.0			3/14/19 08:00	N	II
K1901941-003	Carbon, Dissolved Organic (DOC)	N/A		Water	0.58 mg/L	10 ml	5.8 mg/L	10	0.7	5.0			3/14/19 09:37	Y	II
K1901941-004	Carbon, Dissolved Organic (DOC)	N/A		Water	1.13 mg/L	10 ml	1.13 mg/L	1	0.07	0.50			3/14/19 12:16	N	II
K1901941-005	Carbon, Dissolved Organic (DOC)	N/A		Water	-0.04 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 13:20	N	II
KQ1903347-03	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/13/19 17:17	N	II
KQ1903347-04	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 00:47	N	II
KQ1903347-05	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 09:20	N	II
KQ1903347-06	Carbon, Dissolved Organic (DOC)	CCV		Water	24.59 mg/L	10 ml	24.6 mg/L	1					3/13/19 17:00	N	II
KQ1903347-07	Carbon, Dissolved Organic (DOC)	CCV		Water	24.30 mg/L	10 ml	24.3 mg/L	1					3/14/19 00:30	N	II
KQ1903347-08	Carbon, Dissolved Organic (DOC)	CCV		Water	24.38 mg/L	10 ml	24.4 mg/L	1					3/14/19 09:03	N	II
KQ1903347-09	Carbon, Dissolved Organic (DOC)	MB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 01:03	N	II
KQ1903347-10	Carbon, Dissolved Organic (DOC)	LCS		Water	25.49 mg/L	10 ml	25.5 mg/L	1	0.07	0.50	102		3/14/19 01:20	N	II
KQ1903347-24	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 16:48	N	II
KQ1903347-25	Carbon, Dissolved Organic (DOC)	CCV		Water	24.23 mg/L	10 ml	24.2 mg/L	1					3/14/19 16:32	N	II
KQ1903347-26	Carbon, Dissolved Organic (DOC)	MB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 14:56	N	II
KQ1903347-27	Carbon, Dissolved Organic (DOC)	MB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 14:56	N	II
KQ1903347-28	Carbon, Dissolved Organic (DOC)	MB		Water	-0.18 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/14/19 14:56	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

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Results Summary

03/15/19
[Handwritten Signature]

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628235 Method/Testcode: 9060/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1903347-29	Carbon, Dissolved Organic LCS (DOC)			Water	25.26 mg/L	10 ml	25.3 mg/L	1	0.07	0.50	101		3/14/19 15:44	N	II
KQ1903347-30	Carbon, Dissolved Organic LCS (DOC)			Water	25.55 mg/L	10 ml	25.6 mg/L	1	0.07	0.50	102		3/14/19 15:44	N	II
KQ1903347-31	Carbon, Dissolved Organic LCS (DOC)			Water	25.77 mg/L	10 ml	25.8 mg/L	1	0.07	0.50	103		3/14/19 15:44	N	II
KQ1903347-32	Carbon, Dissolved Organic MS (DOC)		K1901725-003	Water	28.75 mg/L	10 ml	115 mg/L	4	0.3	2.0	106		3/14/19 01:37	N	II
KQ1903347-33	Carbon, Dissolved Organic MS (DOC)		K1901725-003	Water	28.99 mg/L	10 ml	116 mg/L	4	0.3	2.0	107		3/14/19 01:37	N	II
KQ1903347-34	Carbon, Dissolved Organic MS (DOC)		K1901725-003	Water	28.87 mg/L	10 ml	115 mg/L	4	0.3	2.0	106		3/14/19 01:37	N	II
KQ1903347-35	Carbon, Dissolved Organic MS (DOC)		K1901725-003	Water	28.98 mg/L	10 ml	116 mg/L	4	0.3	2.0	106		3/14/19 01:37	N	II
KQ1903347-36	Carbon, Dissolved Organic MS (DOC)		K1901786-002	Water	27.55 mg/L	10 ml	13800 mg/L	500	40	250	104		3/14/19 05:20	N	IV
KQ1903347-37	Carbon, Dissolved Organic MS (DOC)		K1901786-002	Water	27.78 mg/L	10 ml	13900 mg/L	500	40	250	105		3/14/19 05:20	N	IV
KQ1903347-38	Carbon, Dissolved Organic MS (DOC)		K1901786-002	Water	27.59 mg/L	10 ml	13800 mg/L	500	40	250	104		3/14/19 05:20	N	IV
KQ1903347-39	Carbon, Dissolved Organic MS (DOC)		K1901786-002	Water	27.80 mg/L	10 ml	13900 mg/L	500	40	250	105		3/14/19 05:20	N	IV
KQ1903347-40	Carbon, Dissolved Organic MS (DOC)		K1901941-003	Water	27.52 mg/L	10 ml	275 mg/L	10	0.7	5.0	108		3/14/19 10:41	N	II
KQ1903347-41	Carbon, Dissolved Organic MS (DOC)		K1901941-003	Water	27.04 mg/L	10 ml	270 mg/L	10	0.7	5.0	106		3/14/19 10:41	N	II
KQ1903347-42	Carbon, Dissolved Organic MS (DOC)		K1901941-003	Water	0.62 mg/L	10 ml	6.2 mg/L	10	0.7	5.0	0*		3/14/19 10:41	N	II
KQ1903347-43	Carbon, Dissolved Organic MS (DOC)		K1901941-003	Water	-0.06 mg/L	10 ml	5.0 mg/L U	10	0.7	5.0	-2*		3/14/19 10:41	N	II
KQ1903347-44	Carbon, Dissolved Organic DUP (DOC)		K1901725-001	Water	2.35 mg/L	10 ml	9.4 mg/L	4	0.3	2.0		13	3/13/19 21:19	N	II
KQ1903347-45	Carbon, Dissolved Organic TRP (DOC)		K1901725-001	Water	2.23 mg/L	10 ml	8.9 mg/L	4	0.3	2.0		10	3/13/19 21:19	N	II
KQ1903347-46	Carbon, Dissolved Organic QUAD (DOC)		K1901725-001	Water	2.23 mg/L	10 ml	8.9 mg/L	4	0.3	2.0		9	3/13/19 21:19	N	II
KQ1903347-47	Carbon, Dissolved Organic DUP (DOC)		K1901725-002	Water	1.53 mg/L	10 ml	6.1 mg/L	4	0.3	2.0		4	3/13/19 22:22	N	II
KQ1903347-48	Carbon, Dissolved Organic TRP (DOC)		K1901725-002	Water	1.42 mg/L	10 ml	5.7 mg/L	4	0.3	2.0		6	3/13/19 22:22	N	II
KQ1903347-49	Carbon, Dissolved Organic QUAD (DOC)		K1901725-002	Water	1.52 mg/L	10 ml	6.1 mg/L	4	0.3	2.0		5	3/13/19 22:22	N	II
KQ1903347-50	Carbon, Dissolved Organic DUP (DOC)		K1901725-003	Water	2.34 mg/L	10 ml	9.4 mg/L	4	0.3	2.0		<1	3/13/19 23:26	N	II
KQ1903347-51	Carbon, Dissolved Organic TRP (DOC)		K1901725-003	Water	2.42 mg/L	10 ml	9.7 mg/L	4	0.3	2.0		2	3/13/19 23:26	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Printed 3/15/19 11:12

Results Summary

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Page 2 of 3

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 628235 Method/Testcode: 9060/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1903347-52	Carbon, Dissolved Organic (DOC)	QUAD	K1901725-003	Water	2.40 mg/L	10 ml	9.6 mg/L	4	0.3	2.0		1	3/13/19 23:26	N	II
KQ1903347-53	Carbon, Dissolved Organic (DOC)	DUP	K1901786-001	Water	0.28 mg/L	10 ml	0.28 mg/L	J 1	0.07	0.50		<1	3/14/19 03:13	N	IV
KQ1903347-54	Carbon, Dissolved Organic (DOC)	TRP	K1901786-001	Water	0.28 mg/L	10 ml	0.28 mg/L	J 1	0.07	0.50		<1	3/14/19 03:13	N	IV
KQ1903347-55	Carbon, Dissolved Organic (DOC)	QUAD	K1901786-001	Water	0.23 mg/L	10 ml	0.23 mg/L	J 1	0.07	0.50		9	3/14/19 03:13	N	IV
KQ1903347-56	Carbon, Dissolved Organic (DOC)	DUP	K1901786-002	Water	1.56 mg/L	10 ml	780 mg/L	500	40	250		4	3/14/19 04:16	N	IV
KQ1903347-57	Carbon, Dissolved Organic (DOC)	TRP	K1901786-002	Water	1.55 mg/L	10 ml	770 mg/L	500	40	250		2	3/14/19 04:16	N	IV
KQ1903347-58	Carbon, Dissolved Organic (DOC)	QUAD	K1901786-002	Water	1.55 mg/L	10 ml	780 mg/L	500	40	250		2	3/14/19 04:16	N	IV
KQ1903347-59	Carbon, Dissolved Organic (DOC)	DUP	K1901941-001	Water	0.11 mg/L	10 ml	1.1 mg/L	J 10	0.7	5.0		48*	3/14/19 06:56	N	II
KQ1903347-60	Carbon, Dissolved Organic (DOC)	TRP	K1901941-001	Water	0.16 mg/L	10 ml	1.6 mg/L	J 10	0.7	5.0		24*	3/14/19 06:56	N	II
KQ1903347-61	Carbon, Dissolved Organic (DOC)	QUAD	K1901941-001	Water	0.16 mg/L	10 ml	1.6 mg/L	J 10	0.7	5.0		20	3/14/19 06:56	N	II
KQ1903347-62	Carbon, Dissolved Organic (DOC)	DUP	K1901941-002	Water	0.58 mg/L	10 ml	5.8 mg/L	10	0.7	5.0		4	3/14/19 08:00	N	II
KQ1903347-63	Carbon, Dissolved Organic (DOC)	TRP	K1901941-002	Water	0.51 mg/L	10 ml	5.1 mg/L	10	0.7	5.0		6	3/14/19 08:00	N	II
KQ1903347-64	Carbon, Dissolved Organic (DOC)	QUAD	K1901941-002	Water	0.51 mg/L	10 ml	5.1 mg/L	10	0.7	5.0		7	3/14/19 08:00	N	II
KQ1903347-65	Carbon, Dissolved Organic (DOC)	DUP	K1901941-003	Water	0.51 mg/L	10 ml	5.1 mg/L	10	0.7	5.0		13	3/14/19 09:37	N	II
KQ1903347-66	Carbon, Dissolved Organic (DOC)	TRP	K1901941-003	Water	0.54 mg/L	10 ml	5.4 mg/L	10	0.7	5.0		6	3/14/19 09:37	N	II
KQ1903347-67	Carbon, Dissolved Organic (DOC)	QUAD	K1901941-003	Water	0.53 mg/L	10 ml	5.3 mg/L	10	0.7	5.0		5	3/14/19 09:37	N	II
KQ1903347-68	Carbon, Dissolved Organic (DOC)	DUP	K1901941-004	Water	1.09 mg/L	10 ml	1.09 mg/L	1	0.07	0.50		4	3/14/19 12:16	N	II
KQ1903347-69	Carbon, Dissolved Organic (DOC)	TRP	K1901941-004	Water	1.12 mg/L	10 ml	1.12 mg/L	1	0.07	0.50		2	3/14/19 12:16	N	II
KQ1903347-70	Carbon, Dissolved Organic (DOC)	QUAD	K1901941-004	Water	1.07 mg/L	10 ml	1.07 mg/L	1	0.07	0.50		3	3/14/19 12:16	N	II
KQ1903347-71	Carbon, Dissolved Organic (DOC)	DUP	K1901941-005	Water	0.02 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50		NC	3/14/19 13:20	N	II
KQ1903347-72	Carbon, Dissolved Organic (DOC)	TRP	K1901941-005	Water	0.00 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50		NC	3/14/19 13:20	N	II
KQ1903347-73	Carbon, Dissolved Organic (DOC)	QUAD	K1901941-005	Water	0.01 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50		NC	3/14/19 13:20	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

TOC: 628233,
628234
DOC: 628232,
628235

Schedule: 03122019

Version: 8

Instrument: Fusion1

Last Saved by: Fusion1 (Fusion1)

Last Saved on: 2019/03/14 09:36 - Thursday

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps
(Clean)	Clean	Clean		1
(Clean)	Clean	Clean		1
(Clean)	Clean	Clean		1
(Blank)	Blank	Reagent/Acid Blank		1
D	Sample	RB	Extended Reaction 021711 (Extended Reaction 021711)	1
B	Check Standard	[TOC] CCV 021711 [25 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
D	Check Standard	[TOC] CCB 021711 [0.0 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
1	Sample	MB1	Extended Reaction 021711 (Extended Reaction 021711)	1
C	Check Standard	[TOC] LCS ER [25.0 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
2	Sample	ICS	Extended Reaction 021711 (Extended Reaction 021711)	1
3	Sample	RB	Extended Reaction 021711 (Extended Reaction 021711)	2
4	Sample	LOD	Extended Reaction 021711 (Extended Reaction 021711)	4
5	Sample	LOQ	Extended Reaction 021711 (Extended Reaction 021711)	4
6	Sample	K1902025-002.01 doc	Extended Reaction 021711 (Extended Reaction 021711)	2
7	Sample	K1902025-002.01 ms doc	Extended Reaction 021711 (Extended Reaction 021711)	1
8	Sample	RB	Extended Reaction 021711 (Extended Reaction 021711)	1
9	Sample	K1902025-004.01 doc	Extended Reaction 021711 (Extended Reaction 021711)	2
B	Check Standard	[TOC] CCV 021711 [25 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
D	Check Standard	[TOC] CCB 021711 [0.0 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
10	Sample	K1902074-001.03 doc	Extended Reaction 021711 (Extended Reaction 021711)	2
11	Sample	K1902074-002.03 doc	Extended Reaction 021711 (Extended Reaction 021711)	2
12	Sample	K1901653-001.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
13	Sample	K1901653-002.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
14	Sample	K1901653-003.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
15	Sample	K1901653-004.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
16	Sample	K1901653-006.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
17	Sample	K1901653-005.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
18	Sample	K1901653-005.08 ms 2x	Extended Reaction 021711 (Extended Reaction 021711)	1
19	Sample	RB	Extended Reaction 021711 (Extended Reaction 021711)	2
B	Check Standard	[TOC] CCV 021711 [25 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
D	Check Standard	[TOC] CCB 021711 [0.0 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
20	Sample	MB2	Extended Reaction 021711 (Extended Reaction 021711)	1
C	Check Standard	[TOC] LCS ER [25.0 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
21	Sample	K1901719-001.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
22	Sample	K1901719-002.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
23	Sample	K1901719-003.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
24	Sample	K1901719-004.08 4x	Extended Reaction 021711 (Extended Reaction 021711)	2
25	Sample	K1901719-005.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
26	Sample	K1901719-006.08 2x	Extended Reaction 021711 (Extended Reaction 021711)	2
27	Sample	K1901719-007.08 4x	Extended Reaction 021711 (Extended Reaction 021711)	2
28	Sample	RB	Extended Reaction 021711 (Extended Reaction 021711)	2
B	Check Standard	[TOC] CCV 021711 [25 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
D	Check Standard	[TOC] CCB 021711 [0.0 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1
29	Sample	K1901698-001.10 10x	Extended Reaction 021711 (Extended Reaction 021711)	2
30	Sample	K1901698-001.10 ms 10x	Extended Reaction 021711 (Extended Reaction 021711)	1
31	Sample	RB	Extended Reaction 021711 (Extended Reaction 021711)	2
32	Sample	K1901698-002.10 10x	Extended Reaction 021711 (Extended Reaction 021711)	2
33	Sample	K1901698-003.10 10x	Extended Reaction 021711 (Extended Reaction 021711)	2
34	Sample	K1901701-001.05 10x	Extended Reaction 021711 (Extended Reaction 021711)	2
35	Sample	RB	Extended Reaction 021711 (Extended Reaction 021711)	2
36	Sample	K1901940-001.14	Extended Reaction 021711 (Extended Reaction 021711)	2
37	Sample	K1901955-001.04	Extended Reaction 021711 (Extended Reaction 021711)	2
38	Sample	K1901955-001.04 ms	Extended Reaction 021711 (Extended Reaction 021711)	1
B	Check Standard	[TOC] CCV 021711 [25 ppm]	Extended Reaction 021711 (Extended Reaction 021711)	1

Printed on: March 15, 2019 09:13:09

Page 1

Blank substitution
0.18324

03/15/19
Fusion1

Schedule: 03122019

Sample No.	Sample Type	Sample	Method (1) (Description)	Range
38	Check Standard	TCC/COB 021711 (1.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1
39	Check Standard	TCC/LCS ER (25.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1
41	Sample	K150155-021711	Extended Reaction 021711 (Extended Reaction 021711)	2
43	Sample	K150156-021711	Extended Reaction 021711 (Extended Reaction 021711)	2
45	Sample	K150154-021711	Extended Reaction 021711 (Extended Reaction 021711)	2
47	Sample	FB	Extended Reaction 021711 (Extended Reaction 021711)	1
49	Check Standard	TCC/CALB 021711 (1.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1
51	Sample	K150157-021711 100x	Extended Reaction 021711 (Extended Reaction 021711)	2
53	Sample	FB	Extended Reaction 021711 (Extended Reaction 021711)	2
55	Sample	K150158-021711 100x	Extended Reaction 021711 (Extended Reaction 021711)	2
56	Sample	K150152-021711 10 disc 4x	Extended Reaction 021711 (Extended Reaction 021711)	4
57	Sample	K150153-021711 10 disc 4x	Extended Reaction 021711 (Extended Reaction 021711)	4
59	Check Standard	TCC/LCS 021711 (1.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1
61	Check Standard	TCC/LCS ER (25.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1
63	Sample	FB	Extended Reaction 021711 (Extended Reaction 021711)	2
65	Sample	K150159-021711 24 disc 500x	Extended Reaction 021711 (Extended Reaction 021711)	4
67	Sample	FB	Extended Reaction 021711 (Extended Reaction 021711)	4
69	Sample	FB	Extended Reaction 021711 (Extended Reaction 021711)	2
71	Sample	K150161-021711 10 disc 100x	Extended Reaction 021711 (Extended Reaction 021711)	4
73	Check Standard	TCC/COB 021711 (1.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1
75	Sample	K150162-021711 10 ms disc 10x	Extended Reaction 021711 (Extended Reaction 021711)	4
77	Sample	K150164-021711 21 disc	Extended Reaction 021711 (Extended Reaction 021711)	4
79	Sample	FB	Extended Reaction 021711 (Extended Reaction 021711)	2
81	Check Standard	TCC/LCS ER (25.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1
83	Check Standard	TCC/COB 021711 (1.0 ppm)	Extended Reaction 021711 (Extended Reaction 021711)	1

Fusion Report - 03122019

Tuesday, March 12, 2019 11:18 AM

(View - Reps, Unused Reps, Meta-Data, Signature, History)
 Printed on 2019/03/15 09:13 - Friday

Report Summary Information

Company Location: Gen Chem Lab
 Schedule Name: 03122019
 Instrument Name: Fusion1
 Report Version: 1 of 1
 Report Creation by Operators (schedule version): Fusion1 (Fusion1) (v2)
 Fusion1 (Fusion1) (v3)
 Fusion1 (Fusion1) (v4)
 Fusion1 (Fusion1) (v5)
 Fusion1 (Fusion1) (v6)
 Fusion1 (Fusion1) (v7)
 Fusion1 (Fusion1) (v8)

Engine 1.1.5.1
 Version:
 Firmware 1.2.0696
 Version:
 Connection: RS232 COM1

Comment:

Report Results

03/15/19
[Signature]

Sample Type: Clean From Schedule Version 2

Pos	Analysis Type	Sample ID	Start Time
◆ (clean)		Clean	2019/03/12 11:18

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	13.77	16.68	2.90	49.56	05:25
2	TC Clean	6.82	9.74	2.91	50.07	04:01
3	TC Clean	2.12	5.09	2.97	50.10	03:46
4	TC Clean	1.33	4.45	3.12	50.06	03:48

Sample Type: Clean From Schedule Version 3

Pos	Analysis Type	Sample ID	Start Time
◆ (clean)		Clean	2019/03/12 11:49

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	12.97	15.64	2.67	49.74	05:22

2	TC Clean	3.98	6.84	2.85	50.09	04:04
3	TC Clean	1.46	4.35	2.90	49.99	03:48
4	TC Clean	1.50	4.33	2.83	50.07	03:47

Sample Type: Clean From Schedule Version 4

Pos	Analysis Type	Sample ID			Start Time	
◊ (clean)		Clean			2019/03/12 12:14	

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	12.71	15.54	2.84	49.74	05:11
2	TC Clean	4.13	6.91	2.78	50.05	04:04
3	TC Clean	1.52	4.47	2.96	50.08	03:46
4	TC Clean	1.51	4.39	2.89	50.03	03:47

Sample Type: Blank (Creating v1235) From Schedule Version 5

Pos	Analysis Type	Sample ID			Start Time	
◊ (blank)		Reagent/Acid Blank			2019/03/12 12:36	

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	12.63	15.40	2.77	49.75	05:12
2	TC Clean	3.84	6.88	3.04	50.04	04:04
3	TC Clean	1.64	4.62	2.98	50.11	03:50
4	TC Clean	1.35	4.41	3.06	50.06	03:47
5	Reagent Blank	3.56	6.53	2.96	50.11	05:06
6	Acid Blank	0.88	3.61	2.74	49.77	05:31

Sample Type: Sample From Schedule Version 6

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ D	TOC	RB	0.6073 ppm	0.0000 ppm	0.0000%	2019/03/12 13:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6073	6.0733	14.64	17.43	2.79	50.18	12:31

Dilution **Blank Contribution** **Method** **Calibration**

1:10 (TC) 10.2853 (IC) Extended Reaction Extended Reaction
 (v1235) 021711 (v4) 021711 (v27)

Sample Type: Check Standard --> CCV 021711

From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	25.0023 ppm (PASS)	0.0000 ppm	0%	2019/03/12 13:47

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.0023	250.0233	190.44	193.17	2.74	50.20	12:33

Completion State Success - Criteria met.
Success Action Do Nothing
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB 021711

From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0072 ppm (PASS)	0.0000 ppm	0%	2019/03/12 14:04

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0072	0.0724	11.22	14.21	2.99	50.23	12:33

Completion State Success - Criteria met.
Success Action Do Nothing
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)
STD Conc - Pos D 0 ppmC

Sample Type: Sample

From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 1	TOC	MB1	0.0032 ppm	0.0000 ppm	0.0000%	2019/03/12 14:21

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0032	0.0317	10.31	13.27	2.96	50.22	12:32

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Sample Type: Check Standard --> LCS ER

From Schedule Version 7

Pos	BAT	Concentration	Dil	Sample ID	Min / Max	Result	Std. Dev.	RSD	Start Time
-----	-----	---------------	-----	-----------	-----------	--------	-----------	-----	------------

			(ppm)			(% dev)				
♦	C	TOC	25.0000	1:1	[TOC] LCS ER [25.0 ppm]	0 / infinity (NA / NA)	26.0129 ppm (PASS)	0.0000 ppm	0%	2019/03/12 14:37

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.0129	260.1289	197.68	200.43	2.74	50.24	12:31

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos C</u>
Success - Criteria met.	Do Nothing	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)	25 ppmC

Sample Type: Sample From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	2	TOC	ICS	0.3560 ppm	0.0000 ppm	0.0000%	2019/03/12 14:54

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3560	3.5601	12.84	15.77	2.93	50.25	12:31

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	3	TOC	RB	0.0149 ppm	0.0085 ppm	56.9800%	2019/03/12 15:11

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0089	0.0889	10.35	13.32	2.97	50.28	12:27
2	TOC	0.0209	0.2088	10.43	13.34	2.91	50.30	12:25

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	4	TOC	LOD	0.2108 ppm	0.0188 ppm	8.9000%	2019/03/12 15:43

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2132	2.1320	11.81	14.68	2.87	50.26	12:30
2	TOC	0.2044	2.0442	11.75	14.60	2.85	50.29	12:27
3	TOC	0.1903	1.9033	11.65	14.61	2.96	50.33	12:27
4	TOC	0.2351	2.3510	11.97	14.81	2.84	50.34	12:26

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time

5	TOC	LOQ	0.5628 ppm	0.0291 ppm	5.1700%	2019/03/12 16:47
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Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5804	5.8041	14.45	17.40	2.95	50.33	12:28
2	TOC	0.5214	5.2142	14.02	16.98	2.95	50.30	12:26
3	TOC	0.5856	5.8557	14.48	17.21	2.72	50.24	12:29
4	TOC	0.5638	5.6382	14.33	17.18	2.85	50.22	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
6	TOC	K1902025-002.01 doc	3.3727 ppm	0.0780 ppm	2.3100%	2019/03/12 17:50

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.4279	34.2786	34.86	37.51	2.65	50.19	12:26
2	TOC	3.3175	33.1754	34.07	36.87	2.80	50.19	12:26

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
7	TOC	K1902025-002.01 ms doc	30.8157 ppm	0.0000 ppm	0.0000%	2019/03/12 18:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	30.8157	308.1574	231.24	234.27	3.03	50.21	12:31

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
8	TOC	RB	0.2175 ppm	0.0000 ppm	0.0000%	2019/03/12 18:39

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2175	2.1753	11.84	14.55	2.70	50.25	12:32

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
9	TOC	K1902025-004.01 doc	2.7506 ppm	0.1556 ppm	5.6600%	2019/03/12 18:55

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.6406	26.4058	29.22	31.96	2.74	50.25	12:28
2	TOC	2.8607	28.6065	30.80	33.64	2.85	50.30	12:29

Dilution 1:10	Blank Contribution (TC) 10.2853 (IC) (v1235)	Method Extended Reaction 021711 (v4)	Calibration Extended Reaction 021711 (v27)
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Sample Type: Check Standard --> CCV 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
* B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	25.3758 ppm (PASS)	0.0000 ppm	0%	2019/03/12 19:27

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.3758	253.7582	193.12	195.99	2.88	50.24	12:31

Completion State Success - Criteria met.	Success Action Do Nothing	Method Extended Reaction 021711 (v4)	Calibration Extended Reaction 021711 (v27)	STD Conc - Pos B 50 ppmC
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Sample Type: Check Standard --> CCB 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
* D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0965 ppm (PASS)	0.0000 ppm	0%	2019/03/12 19:44

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0965	0.9650	11.86	14.69	2.83	50.27	12:32

Completion State Success - Criteria met.	Success Action Do Nothing	Method Extended Reaction 021711 (v4)	Calibration Extended Reaction 021711 (v27)	STD Conc - Pos D 0 ppmC
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Sample Type: Sample From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
* 10	TOC	K1902074-001.03 doc	1.7628 ppm	0.2659 ppm	15.0800%	2019/03/12 20:01

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.5748	15.7479	21.58	24.45	2.87	50.26	12:27
2	TOC	1.9508	19.5079	24.27	27.17	2.90	50.26	12:25

Dilution 1:10	Blank Contribution (TC) 10.2853 (IC) (v1235)	Method Extended Reaction 021711 (v4)	Calibration Extended Reaction 021711 (v27)
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Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time

11	TOC	K1902074-002.03 doc	1.6615 ppm	0.0107 ppm	0.6400%	2019/03/12 20:33
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Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.6691	16.6907	22.25	25.19	2.93	50.26	12:28
2	TOC	1.6540	16.5401	22.14	25.05	2.91	50.25	12:25

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
12	TOC	K1901653-001.08 2x	1.0492 ppm	0.0680 ppm	6.4900%	2019/03/12 21:05

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.0973	10.9727	18.15	21.05	2.89	50.27	12:28
2	TOC	1.0010	10.0104	17.46	20.31	2.85	50.24	12:25

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
13	TOC	K1901653-002.08 2x	0.7410 ppm	0.1522 ppm	20.5300%	2019/03/12 21:37

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6334	6.3341	14.83	17.75	2.92	50.24	12:30
2	TOC	0.8486	8.4860	16.37	19.26	2.89	50.25	12:25

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
14	TOC	K1901653-003.08 2x	0.9012 ppm	0.0625 ppm	6.9400%	2019/03/12 22:09

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.9454	9.4539	17.06	19.88	2.81	50.17	12:26
2	TOC	0.8570	8.5697	16.43	19.22	2.79	50.15	12:28

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
15	TOC	K1901653-004.08 2x	1.0064 ppm	0.0052 ppm	0.5200%	2019/03/12 22:41

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.0027	10.0271	17.48	20.33	2.86	50.11	12:26
2	TOC	1.0101	10.1010	17.53	20.22	2.70	50.11	12:26

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
16	TOC	K1901653-006.08 2x	0.6871 ppm	0.0919 ppm	13.3800%	2019/03/12 23:13

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7521	7.5209	15.68	18.47	2.80	50.04	12:30
2	TOC	0.6221	6.2211	14.75	17.59	2.84	50.06	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
17	TOC	K1901653-005.08 2x	0.5681 ppm	0.0627 ppm	11.0400%	2019/03/12 23:45

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6125	6.1249	14.68	17.60	2.92	50.07	12:26
2	TOC	0.5238	5.2379	14.04	16.99	2.94	50.07	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
18	TOC	K1901653-005.08 ms 2x	20.3764 ppm	0.0000 ppm	0.0000%	2019/03/13 00:17

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	20.3764	203.7641	156.39	159.30	2.91	50.07	12:31

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
19	TOC	RB	0.1439 ppm	0.1221 ppm	84.8500%	2019/03/13 00:34

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2302	2.3022	11.94	14.85	2.91	50.05	12:28
2	TOC	0.0576	0.5756	10.70	13.48	2.79	50.05	12:25

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Sample Type: Check Standard --> CCV 021711

From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	25.0316 ppm (PASS)	0.0000 ppm	0%	2019/03/13 01:06

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.0316	250.3162	190.65	193.50	2.86	50.00	12:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)	50 ppmC

Sample Type: Check Standard --> CCB 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	1.5818 ppm (PASS)	0.0000 ppm	0%	2019/03/13 01:22

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	1.5818	15.8179	22.51	25.41	2.90	49.97	12:28

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)	0 ppmC

Sample Type: Sample From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 20	TOC	MB2	0.3071 ppm	0.0000 ppm	0.0000%	2019/03/13 01:39

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3071	3.0706	12.49	15.33	2.85	50.01	12:32

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Sample Type: Check Standard --> LCS ER From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ C	TOC	25.0000	1:1	[TOC] LCS ER [25.0 ppm]	0 / infinity (NA / NA)	26.7013 ppm (PASS)	0.0000 ppm	0%	2019/03/13 01:56

Pos	Base Analysis	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
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Type	Result	Std. Dev.	RSD	Start Time
C TOC	25.0 ppm	1	26.7013	267.0128
			202.62	205.45
			2.83	49.99
				12:34

Completion State Success - Criteria met.
Success Action Do Nothing
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)
STD Conc - Pos C 25 ppmC

Sample Type: Sample

From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
21	TOC	K1901719-001.08 2x	0.7029 ppm	0.1074 ppm	15.2800%	2019/03/13 02:12

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7789	7.7887	15.87	18.82	2.95	49.97	12:29
2	TOC	0.6270	6.2699	14.78	17.74	2.96	49.97	12:25

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
22	TOC	K1901719-002.08 2x	1.0319 ppm	0.0336 ppm	3.2600%	2019/03/13 02:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.0557	10.5571	17.86	20.72	2.87	49.99	12:31
2	TOC	1.0081	10.0815	17.51	20.36	2.84	49.99	12:29

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
23	TOC	K1901719-003.08 2x	18.8966 ppm	0.0982 ppm	0.5200%	2019/03/13 03:17

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	18.8271	188.2710	145.28	148.28	3.00	49.99	12:28
2	TOC	18.9660	189.6600	146.28	149.06	2.78	49.98	12:26

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
24	TOC	K1901719-004.08 4x	1.2885 ppm	0.0590 ppm	4.5800%	2019/03/13 03:49

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.3302	13.3017	19.82	22.63	2.80	49.97	12:29
2	TOC	1.2468	12.4677	19.22	22.11	2.88	49.97	12:26

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
25	TOC	K1901719-005.08 2x	2.0573 ppm	0.1055 ppm	5.1300%	2019/03/13 04:21

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.1320	21.3195	25.57	28.37	2.79	49.98	12:25
2	TOC	1.9827	19.8273	24.50	27.32	2.82	49.97	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
26	TOC	K1901719-006.08 2x	0.9032 ppm	0.0597 ppm	6.6100%	2019/03/13 04:53

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.9454	9.4539	17.06	19.90	2.84	49.94	12:26
2	TOC	0.8610	8.6101	16.46	19.27	2.81	49.94	12:28

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
27	TOC	K1901719-007.08 4x	6.5144 ppm	0.0958 ppm	1.4700%	2019/03/13 05:25

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.5821	65.8212	57.48	60.26	2.78	49.93	12:29
2	TOC	6.4467	64.4670	56.51	59.26	2.75	49.95	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
28	TOC	RB	0.1038 ppm	0.0677 ppm	65.2700%	2019/03/13 05:57

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1517	1.5170	11.37	14.10	2.72	49.96	12:28
2	TOC	0.0559	0.5589	10.69	13.47	2.79	49.98	12:28

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Sample Type: Check Standard --> CCV 021711

From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	24.9213 ppm (PASS)	0.0000 ppm	0%	2019/03/13 06:29

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.9213	249.2130	189.86	192.85	3.00	49.95	12:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)	50 ppmC

Sample Type: Check Standard --> CCB 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/13 06:46

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0000	0.0000	10.27	13.13	2.86	49.93	12:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)	0 ppmC

Sample Type: Sample From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 29	TOC	K1901698-001.10 10x	0.6847 ppm	0.0235 ppm	3.4300%	2019/03/13 07:03

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6681	6.6813	15.08	17.98	2.90	49.94	12:25
2	TOC	0.7013	7.0133	15.31	18.04	2.72	49.91	12:29

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 30	TOC	K1901698-001.10 ms 10x	27.9616 ppm	0.0000 ppm	0.0000%	2019/03/13 07:35

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	27.9616	279.6160	210.78	213.50	2.72	49.89	12:34

Dilution	Blank Contribution	Method	Calibration
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1:10 (TC) 10.2853 (IC) Extended Reaction Extended Reaction
(v1235) 021711 (v4) 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
31	TOC	RB	0.0186 ppm	0.0263 ppm	141.4200%	2019/03/13 07:51

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0372	0.3720	10.55	13.36	2.81	49.87	12:26
2	TOC	0.0000	0.0000	9.65	12.43	2.78	49.87	12:30

Dilution 1:10 Blank Contribution (TC) 10.2853 (IC) Method Extended Reaction Calibration Extended Reaction
(v1235) 021711 (v4) 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
32	TOC	K1901698-002.10 10x	2.3102 ppm	0.0416 ppm	1.8000%	2019/03/13 08:23

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.3396	23.3962	27.06	29.62	2.56	49.84	12:29
2	TOC	2.2808	22.8076	26.64	29.43	2.79	49.83	12:27

Dilution 1:10 Blank Contribution (TC) 10.2853 (IC) Method Extended Reaction Calibration Extended Reaction
(v1235) 021711 (v4) 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
33	TOC	K1901698-003.10 10x	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/13 08:55

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	10.22	13.01	2.79	49.79	12:27
2	TOC	0.0000	0.0000	9.50	12.34	2.84	49.82	12:30

Dilution 1:10 Blank Contribution (TC) 10.2853 (IC) Method Extended Reaction Calibration Extended Reaction
(v1235) 021711 (v4) 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
34	TOC	K1901701-001.05 10x	7.5363 ppm	0.0025 ppm	0.0300%	2019/03/13 09:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.5380	75.3801	64.34	67.11	2.78	49.83	12:33
2	TOC	7.5345	75.3453	64.31	66.97	2.66	49.82	12:25

Dilution 1:10 Blank Contribution (TC) 10.2853 (IC) Method Extended Reaction Calibration Extended Reaction
(v1235) 021711 (v4) 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
35	TOC	RB	0.0090 ppm	0.0128 ppm	141.4200%	2019/03/13 10:00

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0181	0.1809	10.42	13.25	2.84	49.79	12:29
2	TOC	0.0000	0.0000	9.92	12.56	2.64	49.86	12:28

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
36	TOC	K1901940-001.14	0.5054 ppm	0.0677 ppm	13.3900%	2019/03/13 10:32

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4575	4.5754	13.57	16.34	2.77	49.95	12:24
2	TOC	0.5532	5.5322	14.25	17.03	2.77	50.00	12:28

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
37	TOC	K1901955-001.04	3.2123 ppm	0.0090 ppm	0.2800%	2019/03/13 11:04

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.2060	32.0597	33.27	36.15	2.88	50.04	12:29
2	TOC	3.2187	32.1866	33.36	36.05	2.69	50.02	12:25

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
38	TOC	K1901955-001.04 ms	29.8771 ppm	0.0000 ppm	0.0000%	2019/03/13 11:36

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	29.8771	298.7714	224.51	227.19	2.68	50.02	12:32

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Sample Type: Check Standard --> CCV 021711

From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	24.6039 ppm (PASS)	0.0000 ppm	0%	2019/03/13 11:53

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time

B	TOC	25 ppm	1	24.6039	246.0388	187.58	190.34	2.76	50.03	12:31
Completion State		Success Action		Method		Calibration		STD Conc - Pos B		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		50 ppmC		

Sample Type: Check Standard --> CCB 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/13 12:09

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0000	0.0000	9.36	12.06	2.71	50.03	12:31

Completion State		Success Action		Method		Calibration		STD Conc - Pos D		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		0 ppmC		

Sample Type: Sample From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 39	TOC	MB3	0.5022 ppm	0.0000 ppm	0.0000%	2019/03/13 12:26

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5022	5.0217	13.89	16.89	3.00	50.05	12:27

Dilution		Blank Contribution		Method		Calibration	
1:10		(TC) 10.2853 (IC) (v1235)		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)	

Sample Type: Check Standard --> LCS ER From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ C	TOC	25.0000	1:1	[TOC] LCS ER [25.0 ppm]	0 / infinity (NA / NA)	25.9545 ppm (PASS)	0.0000 ppm	0%	2019/03/13 12:43

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	25.9545	259.5445	197.26	199.89	2.62	50.14	12:31

Completion State		Success Action		Method		Calibration		STD Conc - Pos C		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		25 ppmC		

Sample Type: Sample

From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
40	TOC	K1901895-001.02	5.1831 ppm	0.1352 ppm	2.6100%	2019/03/13 12:59

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.2787	52.7869	48.14	50.91	2.77	50.09	12:30
2	TOC	5.0875	50.8748	46.76	49.60	2.84	50.09	12:31

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
41	TOC	K1901956-001.01	30.7278 ppm	0.6021 ppm	1.9600%	2019/03/13 13:31

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	30.3021	303.0209	227.56	230.45	2.89	50.11	12:30
2	TOC	31.1535	311.5352	233.66	236.68	3.01	50.11	12:26

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
42	TOC	K1901956-002.01	31.7343 ppm	0.3230 ppm	1.0200%	2019/03/13 14:04

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	31.5059	315.0595	236.19	238.98	2.79	50.09	12:26
2	TOC	31.9627	319.6269	239.47	242.42	2.95	50.10	12:25

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
43	TOC	K1902084-001.01	4.1932 ppm	0.5599 ppm	13.3500%	2019/03/13 14:36

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	4.5892	45.8918	43.19	46.06	2.87	50.12	12:28
2	TOC	3.7973	37.9730	37.51	40.30	2.79	50.08	12:28

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
44	TOC	K1902084-002.01	3.2410 ppm	0.2059 ppm	6.3500%	2019/03/13 15:08

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time

1	TOC	3.3866	33.8658	34.57	37.52	2.95	50.07	12:27
2	TOC	3.0954	30.9537	32.48	35.34	2.86	50.07	12:26

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
45	TOC	K1902084-003.01	3.4533 ppm	0.0571 ppm	1.6500%	2019/03/13 15:40

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.4937	34.9368	35.34	38.13	2.79	50.08	12:28
2	TOC	3.4129	34.1293	34.76	37.69	2.93	50.09	12:25

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
46	TOC	K1901830-001.01	0.3771 ppm	0.0594 ppm	15.7400%	2019/03/13 16:12

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4191	4.1905	13.29	16.24	2.95	50.10	12:28
2	TOC	0.3351	3.3509	12.69	15.65	2.96	50.08	12:30

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
47	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/13 16:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	9.78	12.81	3.02	50.06	12:32

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Sample Type: Check Standard --> CCV 021711

From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (-NA/-NA)	24.7687 ppm (PASS)	0.0000 ppm	0%	2019/03/13 17:00

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.7687	247.6873	188.76	191.85	3.09	50.04	12:31

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos B</u>
Success - Criteria met.	Do Nothing	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)	50 ppmC

Sample Type: Check Standard --> CCB 021711

From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/13 17:17

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0000	0.0000	10.34	13.11	2.77	50.04	12:30

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos D</u>
Success - Criteria met.	Do Nothing	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)	0 ppmC

Sample Type: Sample

From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 48	TOC	K1901875-001.01 100x	5.7317 ppm	0.0220 ppm	0.3800%	2019/03/13 17:34

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.7473	57.4729	51.50	54.23	2.73	50.05	12:31
2	TOC	5.7162	57.1619	51.27	54.15	2.88	50.06	12:26

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 49	TOC	K1901875-002.01 100x	0.7626 ppm	0.0459 ppm	6.0100%	2019/03/13 18:06

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7950	7.9505	15.99	18.93	2.94	50.07	12:31
2	TOC	0.7302	7.3020	15.52	18.55	3.03	50.08	12:27

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 50	TOC	K1901875-003.01 100x	1.4672 ppm	0.0729 ppm	4.9700%	2019/03/13 18:38

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.5187	15.1873	21.18	23.86	2.69	50.11	12:25

2	TOC	1.4157	14.1566	20.44	23.41	2.98	50.16	12:27
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Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
51	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/13 19:10

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	9.32	12.01	2.69	50.18	12:29
2	TOC	0.0000	0.0000	9.46	12.18	2.72	50.23	12:27

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
52	TOC	K1901926-001.03 100x	0.7080 ppm	0.0997 ppm	14.0800%	2019/03/13 19:43

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7785	7.7845	15.87	18.57	2.70	50.25	12:28
2	TOC	0.6375	6.3745	14.86	17.72	2.87	50.27	12:27

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
53	TOC	K1901926-002.03 100x	1.7196 ppm	2.0677 ppm	120.2400%	2019/03/13 20:15

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2576	2.5755	12.13	14.94	2.81	50.34	12:29
2	TOC	3.1817	31.8170	33.10	35.99	2.89	50.37	12:26

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
54	TOC	K1901926-003.03	2.3354 ppm	0.3256 ppm	13.9400%	2019/03/13 20:47

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.5657	25.6569	28.68	31.57	2.89	50.34	12:28
2	TOC	2.1052	21.0518	25.38	28.28	2.90	50.34	12:25

Dilution 1:10
Blank Contribution (TC) 10.2853 (IC) (v1235)
Method Extended Reaction 021711 (v4)
Calibration Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
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55	TOC	K1901725-001.10 doc 4x	2.5546 ppm	0.2135 ppm	8.3600%	2019/03/13 21:19
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Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.8639	28.6386	30.82	33.59	2.77	50.32	12:28
2	TOC	2.5300	25.2998	28.43	31.27	2.84	50.31	12:32
3	TOC	2.4103	24.1032	27.57	30.39	2.82	50.30	12:27
4	TOC	2.4142	24.1423	27.60	30.31	2.71	50.31	12:28

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
56	TOC	K1901725-002.10 doc 4x	1.6993 ppm	0.0709 ppm	4.1700%	2019/03/13 22:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.7744	17.7437	23.01	25.83	2.82	50.28	12:28
2	TOC	1.7124	17.1244	22.56	25.35	2.79	50.27	12:26
3	TOC	1.6034	16.0338	21.78	24.57	2.79	50.27	12:27
4	TOC	1.7070	17.0701	22.52	25.49	2.96	50.26	12:28

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
57	TOC	K1901725-003.10 doc 4x	2.5634 ppm	0.0352 ppm	1.3700%	2019/03/13 23:26

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.5424	25.4240	28.52	31.42	2.90	50.27	12:30
2	TOC	2.5266	25.2664	28.40	31.27	2.87	50.27	12:30
3	TOC	2.6035	26.0348	28.95	31.75	2.80	50.25	12:30
4	TOC	2.5812	25.8117	28.79	31.61	2.81	50.25	12:29

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 10.2853 (IC) (v1235)	Extended Reaction 021711 (v4)	Extended Reaction 021711 (v27)

Sample Type: Check Standard --> CCV 021711							From Schedule Version 7			
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Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	24.4881 ppm (PASS)	0.0000 ppm	0%	2019/03/14 00:30

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
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B	TOC	25 ppm	1	24.4881	244.8812	186.75	189.60	2.85	50.24	12:30
Completion State		Success Action		Method		Calibration		STD Conc - Pos B		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		50 ppmC		

Sample Type: Check Standard --> CCB 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/14 00:47

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0000	0.0000	9.68	12.36	2.68	50.23	12:34

Completion State		Success Action		Method		Calibration		STD Conc - Pos D		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		0 ppmC		

Sample Type: Sample From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 58	TOC	MB4	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/14 01:03

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	9.20	12.05	2.85	50.24	12:31

Dilution		Blank Contribution		Method		Calibration	
1:10		(TC) 10.2853 (IC) (v1235)		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)	

Sample Type: Check Standard --> LCS ER From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ C	TOC	25.0000	1:1	[TOC] LCS ER [25.0 ppm]	0 / infinity (NA / NA)	25.6773 ppm (PASS)	0.0000 ppm	0%	2019/03/14 01:20

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	25.6773	256.7734	195.28	198.09	2.81	50.27	12:27

Completion State		Success Action		Method		Calibration		STD Conc - Pos C		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		25 ppmC		

Sample Type: Sample

From Schedule Version 7

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
59	TOC	K1901725-003.10 ms doc 4x	29.0821 ppm	0.1112 ppm	0.3800%	2019/03/14 01:37

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	28.9358	289.3576	217.76	220.70	2.93	50.24	12:31
2	TOC	29.1710	291.7103	219.45	222.32	2.87	50.25	12:29
3	TOC	29.0556	290.5556	218.62	221.49	2.87	50.24	12:26
4	TOC	29.1660	291.6601	219.41	222.29	2.88	50.27	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
60	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/14 02:41

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	10.17	13.05	2.89	50.24	12:29
2	TOC	0.0000	0.0000	10.19	13.10	2.91	50.23	12:31

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
61	TOC	K1901786-001.23 doc	0.4524 ppm	0.0253 ppm	5.6000%	2019/03/14 03:13

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4656	4.6563	13.62	16.61	2.98	50.24	12:26
2	TOC	0.4658	4.6577	13.62	16.40	2.77	50.24	12:27
3	TOC	0.4638	4.6382	13.61	16.41	2.80	50.28	12:29
4	TOC	0.4144	4.1445	13.26	16.18	2.92	50.25	12:29

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
62	TOC	K1901786-002.24 doc 500x	1.7221 ppm	0.0263 ppm	1.5300%	2019/03/14 04:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.6829	16.8288	22.35	25.27	2.92	50.25	12:29
2	TOC	1.7392	17.3922	22.76	25.58	2.82	50.25	12:29
3	TOC	1.7328	17.3281	22.71	25.63	2.92	50.26	12:30
4	TOC	1.7336	17.3364	22.72	25.57	2.85	50.25	12:30

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
63	TOC	K1901786-002.24 ms doc 500x	27.8649 ppm	0.1281 ppm	0.4600%	2019/03/14 05:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	27.7375	277.3748	209.17	211.97	2.80	50.25	12:28
2	TOC	27.9627	279.6271	210.79	213.60	2.81	50.25	12:29
3	TOC	27.7726	277.7262	209.42	212.37	2.94	50.25	12:25
4	TOC	27.9870	279.8698	210.96	213.70	2.74	50.27	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
64	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/14 06:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	10.13	13.00	2.87	50.25	12:26
2	TOC	0.0000	0.0000	9.43	12.31	2.88	50.26	12:24

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
65	TOC	K1901941-001.10 doc 10x	0.3380 ppm	0.0302 ppm	8.9400%	2019/03/14 06:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3666	3.6661	12.91	15.84	2.92	50.26	12:25
2	TOC	0.2953	2.9535	12.40	15.35	2.94	50.27	12:29
3	TOC	0.3468	3.4681	12.77	15.51	2.73	50.27	12:28
4	TOC	0.3430	3.4304	12.74	15.61	2.86	50.26	12:30

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
66	TOC	K1901941-002.10 doc 10x	0.7222 ppm	0.0354 ppm	4.9000%	2019/03/14 08:00

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7419	7.4191	15.60	18.51	2.91	50.25	12:28
2	TOC	0.7620	7.6199	15.75	18.59	2.84	50.24	12:27

3	TOC	0.6960	6.9603	15.28	18.22	2.94	50.23	12:27
4	TOC	0.6891	6.8905	15.23	18.27	3.05	50.21	12:27
Dilution		Blank Contribution		Method		Calibration		
1:10		(TC) 10.2853 (IC) (v1235)		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		

Sample Type: Check Standard --> CCV 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	24.5647 ppm (PASS)	0.0000 ppm	0%	2019/03/14 09:03

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.5647	245.6469	187.30	190.13	2.83	50.16	12:34

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27) **STD Conc - Pos B** 50 ppmC

Sample Type: Check Standard --> CCB 021711 From Schedule Version 7

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/14 09:20

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0000	0.0000	9.69	12.55	2.86	50.18	12:31

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27) **STD Conc - Pos D** 0 ppmC

Sample Type: Sample From Schedule Version 8

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 67	TOC	K1901941-003.10 doc 10x	0.7214 ppm	0.0289 ppm	4.0000%	2019/03/14 09:37

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7602	7.6018	15.74	18.49	2.75	50.17	12:29
2	TOC	0.6907	6.9073	15.24	18.03	2.79	50.13	12:25
3	TOC	0.7207	7.2071	15.45	18.21	2.76	50.10	12:29
4	TOC	0.7142	7.1416	15.41	18.13	2.72	50.11	12:26

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
68	TOC	K1901941-003.10 ms doc 10x	13.9637 ppm	15.5930 ppm	111.6700%	2019/03/14 10:41

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	27.7028	277.0275	208.92	211.70	2.77	50.08	12:28
2	TOC	27.2261	272.2606	205.50	208.36	2.85	50.08	12:27
3	TOC	0.8041	8.0411	16.05	18.90	2.85	50.08	12:25
4	TOC	0.1217	1.2171	11.16	13.86	2.71	50.07	12:28

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
69	TOC	RB	0.0592 ppm	0.0203 ppm	34.3000%	2019/03/14 11:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0449	0.4487	10.61	13.31	2.70	50.05	12:30
2	TOC	0.0736	0.7360	10.81	13.51	2.70	50.04	12:25

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
70	TOC	K1901941-004.10 doc	1.2890 ppm	0.0289 ppm	2.2400%	2019/03/14 12:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.3179	13.1790	19.74	22.41	2.68	50.01	12:27
2	TOC	1.2758	12.7578	19.43	22.35	2.92	50.02	12:27
3	TOC	1.3073	13.0730	19.66	22.66	3.00	50.06	12:25
4	TOC	1.2550	12.5500	19.28	22.16	2.87	50.04	12:31

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
71	TOC	K1901941-005.10 doc	0.1789 ppm	0.0256 ppm	14.3200%	2019/03/14 13:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1419	1.4194	11.30	14.05	2.74	50.06	12:27
2	TOC	0.1991	1.9912	11.71	14.56	2.85	50.09	12:29
3	TOC	0.1822	1.8224	11.59	14.40	2.80	50.07	12:31
4	TOC	0.1924	1.9242	11.66	14.41	2.74	50.07	12:30

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
72	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/14 14:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	9.84	12.54	2.70	50.08	12:27
2	TOC	0.0000	0.0000	9.71	12.52	2.81	50.09	12:27

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
58	TOC	MB4	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/14 14:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	9.33	12.22	2.89	50.08	12:28
2	TOC	0.0000	0.0000	9.38	12.14	2.76	50.07	12:24
3	TOC	0.0000	0.0000	9.67	12.33	2.66	50.08	12:29

Dilution 1:10 **Blank Contribution** (TC) 10.2853 (IC) (v1235) **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27)

Sample Type: Check Standard --> LCS ER From Schedule Version 8

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
C	TOC	25.0000	1:1	[TOC] LCS ER [25.0 ppm]	0 / infinity (NA / NA)	25.7114 ppm (PASS)	0.2529 ppm	0.98%	2019/03/14 15:44

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	25.4476	254.4764	193.63	196.48	2.85	50.08	12:25
C	TOC	25.0 ppm	2	25.7347	257.3466	195.69	198.43	2.74	50.09	12:28
C	TOC	25.0 ppm	3	25.9518	259.5180	197.25	200.16	2.91	50.08	12:26

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** Extended Reaction 021711 (v4) **Calibration** Extended Reaction 021711 (v27) **STD Conc - Pos C** 25 ppmC

Sample Type: Check Standard --> CCV 021711 From Schedule Version 8

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 021711 [25 ppm]	0 / infinity (NA / NA)	24.4100 ppm	0.0000 ppm	0%	2019/03/14 16:32

(PASS)										
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.4100	244.1002	186.19	188.81	2.62	50.09	12:31
Completion State		Success Action		Method		Calibration		STD Conc - Pos B		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		50 ppmC		

Sample Type: Check Standard --> CCB 021711 From Schedule Version 8

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:2	[TOC] CCB 021711 [0.0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/14 16:48

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0.0 ppm	1	0.0000	0.0000	10.48	13.41	2.93	50.08	12:29

Completion State		Success Action		Method		Calibration		STD Conc - Pos D		
Success - Criteria met.		Do Nothing		Extended Reaction 021711 (v4)		Extended Reaction 021711 (v27)		0 ppmC		

Meta Data Used in this Report

Blanks

Version	Reagent (Abs)	Acid (Abs)	DI IC (Abs)	DI TC (Abs)	DI TOC (Abs)	Save Time	Operator
v1234	1.5067	0.6760	0.0000	0.0000	0.0000	2019/03/11 18:53	Fusion1 (Fusion1)
v1235	1.1880	0.8760	0.0000	0.0000	0.0000	2019/03/12 13:10	Fusion1 (Fusion1)

Calibrations

Name: Extended Reaction 021711 (TOC)			
Version:	v27	Calibration curve formula:	TOC: $y = 7.170x + 11.164$
Ver Creation:	2019/03/11 21:51	r ² value:	TOC: $r^2 = 0.99991$
Comment:			
Operator:	Fusion1 (Fusion1)		
Basic Analysis Type	TOC		

Basic Analysis Type: TOC

Sample ID	Y Raw Value	X Expected	Message	End Time
0.0 ppm	10.4100	0.0000		2019/03/11 20:12
0.50 ppm	14.7740	0.5000		2019/03/11 20:28
1.00 ppm	18.0020	1.0000		2019/03/11 20:44
5.00 ppm	47.2310	5.0000		2019/03/11 21:01
10.0 ppm	85.1320	10.0000		2019/03/11 21:17
25.0 ppm	188.5200	25.0000		2019/03/11 21:33
50.0 ppm	370.1610	50.0000		2019/03/11 21:49

Methods

Name: Extended Reaction 021711 (TOC)

Version: v4
Ver Creation: 2019/01/31 11:21
Comment:

Operator: Fusion1 (Fusion1)

Parameter	Value	Advanced Parameter	Value
SampleVolume	10.0 mL	NeedleRinseVolume	5.0 ml
Dilution	1:10	VialPrimeVolume	2.0 ml
AcidVolume	0.5 ml	ICSamplePrimeVolume	2.0 ml
ReagentVolume	2.0 ml	ICSpurgeRinseVolume	12.0 ml
UVReactorPrerinse	Off	BaselineStabilizeTime	0.70 min
UVReactorPrerinseVolume	5.0	DetectorPressureFlow	150 ml/min
NumberOfUVReactorPrerinses	1	SyringeSpeedWaste	10
ICSpurgeTime	1.00 mins	SyringeSpeedAcid	7
DetectorSweepFlow	500 ml/min	SyringeSpeedReagent	7
PreSpurgeTime	4.00 mins	SyringeSpeedDIWater	7
SystemFlow	500 ml/min	NDIRPressurization	60 psig
		SyringeSpeedSampleDispense	5
		SyringeSpeedSampleAspirate	4
		SyringeSpeedUVDispense	5
		SyringeSpeedUVAspirate	5
		SyringeSpeedICDispense	5
		SyringeSpeedICAspirate	5
		NDIRPressureStabilize	1.75 min
		SampleMixing	Off
		SampleMixingCycles	1
		SampleMixingVolume	10.0
		LowLevelFilterNDIR	Off

Acceptance / Approval

Electronic Signatures

Report Version	User Name	Acceptance	Reason	Date
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Report History

Report History				
Report Version	User Name	System Reason	User Reason	Date
1	Fusion1 (Fusion1)	Schedule completed	Schedule completed	2019/03/14 17:09

StarLIMS Run: 628233, 628234, 628232, 628235
 Analysis: TOC/DOC
 Method: 9060, 415.1, SM 5310 C, 9060A

CCV: 11-GEN-05-76B 50 ppm LCS: 11-GEN-05-74L 25.0 ppm

ICAL Date: 3/6/19

ICAL ID: 11-GEN-05-76H

ICS ID: 11-GEN-05-74A

ICS TV: 25.0 ppm ICS % R = 2

Spike ID: 11-GEN-05-700 0.05 ml of 5000 ppm stock ---> 10.0 ml = 25.0 ppm x dilution factor

Sodium Persulfate: 11-GEN-05-76F

21 % H3PO4: 11-GEN-05-76G

Equipment ID: K-TOC-03

PIPETTE ID: 124276B, 129001F, N11314F, Marge

FILTER ID: NA

Analyzed By: <i>BCD</i>	Date Analyzed: <i>3/12/18</i>
Reviewed By: <i>[Signature]</i>	Date Reviewed: <i>03/15/19</i>



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1905651; 1906112; 1906330;
1906332; 1906334

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2223 (233911)

General Set Information: There were thirteen field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Method QC data: The method blank (LMB 642099) was less than 1/2 the CRDL. The recovery for the LCS (642100) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0 μ l of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.0 μ g/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in μ g/L. Results were calculated in μ g/L by the equation $(A) \times (B)$,

where: A = Analyte concentration from the standard curve (μ g/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level of 4.0 μ g/L. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported. Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEBI04) along with datafiles 05MARD07-10.

Thomas Bosch March 06, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 07, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1906334**

Project ID: HS19030014

Purchase Order: HS19030014

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18-24-SP650_022819_BIX	1906334001	02/28/19	03/02/19	



ANALYTICAL REPORT

Workorder: 34-1906334

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18-24-SP650_022819_BIX	Sampling Site: NA	Collected: 02/28/2019				
Lab ID: 1906334001	Media: 125 mL Nalgene	Received: 03/02/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 13:02	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 233911)

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/06/2019 09:17	/S/ Stephen Brose 03/07/2019 11:04

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1906334

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00934867

Analysis Information

Workorder: 1906334

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2223 (HBN: 233911)
Analyzed By: Thomas Bosch

Blank

LMB: 642099 Analyzed: 03/05/2019 09:26 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 642100 Analyzed: 03/05/2019 09:00 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.11	4.00	103	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1906112001 Analyzed: 03/05/2019 09:54 Dilution: 1 Units: ug/L		MS: 1906112002 Analyzed: 03/05/2019 10:07 Dilution: 1 Units: ug/L				MSD: 1906112003 Analyzed: 03/05/2019 10:20 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	51.0	53.9	4	▲ 65.7	78.8 123.8	53.7	▲ 61.2	0.337	0.0 20.0

Continuing Calibration Verification

CCV: 642096 Analyzed: 03/05/2019 08:44 Units: ug/L Criteria: ± 15%			CCV: 642101 Analyzed: 03/05/2019 12:18 Units: ug/L Criteria: ± 15%			CCV: 642102 Analyzed: 03/05/2019 14:07 Units: ug/L Criteria: ± 15%			
Analyte	Result	Target	% Rec.	Result	Target	% Rec.	Result	Target	% Rec.
Perchlorate	25.0	25.0	100	25.0	25.0	100	24.5	25.0	97.8

Interference Check Sample

ICSA: 642098 Analyzed: 03/05/2019 09:13 Units: ug/L Criteria: ± 30%			
Analyte	Result	Target	% Rec.
Perchlorate	3.93	4.00	98.2

Comments

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.



Quality Control Sample Batch Report

00934868

Analysis Information

Workorder: 1906334

Limits: Client SOW/Contract Specified

Preparation: NA

Analysis: EPA 6850, DoD QSM

Basis: DoD QSM

Batch: NA

Batch: ELMS/2223 (HBN: 233911)

Prepared By: NA

Analyzed By: Thomas Bosch

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/06/2019 13:45	/S/ Stephen Brose 03/07/2019 11:04

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable

18698/#2



1906334

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10856

1906334

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com


INVOICE INFORMATION:

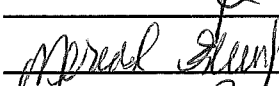
Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030014
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030014-02	LH18/24-SP650_022819_BIX	Water	28 Feb 2019 14:00
SUB_Perch-6850			15 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: 

Received By: 

Cooler ID(s): 9203

Date/Time: 3/1/19 1800.

Date/Time: 3/2/19 / 903

Temperature(s): 3

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: <u>MS Houston</u>		Project/Task/Site: <u>1906334</u>							
Date/Time of Receipt: <u>3/2/19 903</u>		Number of Coolers Received: <u>1</u>							
Condition of Coolers:	Acceptable/Unacceptable	Temperature Control:	Present/Not Included						
Cooler Custody Seals:	Present/Absent/NA	Location Temp Taken:	Control/Between Samples						
Container Custody Seals:	Present/Absent/NA	Are all temperatures within project specific guidelines?	Yes/No/NA						
Ice Present:	Yes/No/NA	VOA Headspace Present?	Yes/No/NA						
	Frozen/Melted/NA								
pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA			
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA			
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA			
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA			
Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	
	1	C19 9203	3 °C	4	C19	°C	7	C19	°C
	2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C	
Taken By: <u>[Signature]</u>		Signature		<u>[Signature]</u>		Printed Name		<u>3/2/19</u>	Date

CLIENT-RELATED INFORMATION

<input type="checkbox"/> Missing Cooler	<input type="checkbox"/> Missing Samples/Bottles	<input type="checkbox"/> Incorrect Preservation	<input type="checkbox"/> Insufficient Sample Volume
<input type="checkbox"/> Cooler Conditions	<input type="checkbox"/> Broken/Leaking Samples	<input type="checkbox"/> pH Criteria Not Met	<input type="checkbox"/> Chain of Custody Problems
<input type="checkbox"/> Missing Paperwork	<input type="checkbox"/> Incorrect Bottle Type	<input type="checkbox"/> Residual Chlorine Present	<input type="checkbox"/> Other:
<input type="checkbox"/> Missing/Incorrect Bottle Labels	<input type="checkbox"/> Cooler Temperatures Out of Range	<input type="checkbox"/> Head Space in Bottles	

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Must Deliver Next Business Day
Time and Tempature Sensitive!

Part #: 159469-434 RIT2 EXP 11/19

ORIGIN ID:SGRA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

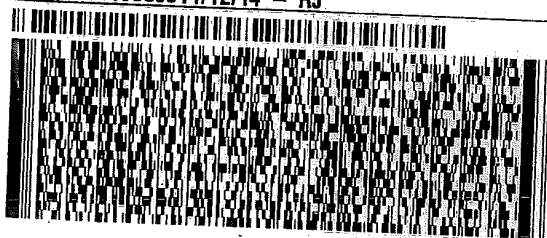
SHIP DATE: 01MAR19
ACTWTG: 8.55 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(801) 266-7700

REF: HS19030011/12/14 - RJ



FedEx
Express



3001/4E30/2C155

FedEx® Saturday Delivery

151966 10/04 MW1



SDR

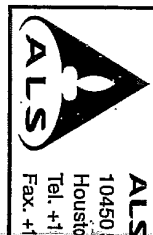
TRK# 4809 7831 2469
0201

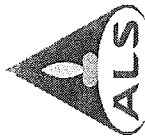
SATURDAY 12:00P
PRIORITY OVERNIGHT

XO BTFA

84123
UT-US SLC

RY **B639** 1
ST **F1** 12:00





Batch Worklist

Batch: ELMS/ 2223 Created: 3/5/2019 08:21 Instrument: HBN: 233911
 Rule: EPA 6850, DoD QSM Water Analyst: T. Bosch Status: WP



- Workorder: 1905651 [ENV_LVL4]
- Workorder: 1906112 [ENV_LVL4]
- Workorder: 1906330 [ENV_LVL4]
- Workorder: 1906332 [ENV_LVL4]
- Workorder: 1906334 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mix	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	642096	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
2	642097	RLVS for HBN 233911 [ELMS/2223]				RLVS	3		E685041C3Q	5311		3/7/2019	
3	642098	ICS for HBN 233911 [ELMS/2223]				ICS	3		E6850.D3Q	5311		3/7/2019	
4	642099	LMB for HBN 233911 [ELMS/2223]				LMB	3		E6850Q413Q	5311		3/7/2019	
5	642100	LCS for HBN 233911 [ELMS/2223]				LCS	3		E6850Q413Q	5311		3/7/2019	
6	1905651001	HS19021158-02/LH18/24-SP650_02				SAMPLE	3	1905651001-A	E6850Q41.3	5480	3/21/2019	3/7/2019	
7	1906112001	EW01_022619				SAMPLE	3	1906112001-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
8	1906112002	EW01_022619MS				MS	3	1906112002-A	E6850Q413Q	5480		3/7/2019	
9	1906112003	EW01_022619MSD				MSD	3	1906112003-A	E6850Q413Q	5480		3/7/2019	
10	1906112004	EW05_022619				SAMPLE	3	1906112004-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
11	1906112005	EW05_022619_FD				FLDDUP	3	1906112005-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
12	1906112006	EW02_022619				SAMPLE	3	1906112006-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
13	1906112007	EW06_022619				SAMPLE	3	1906112007-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
14	1906112008	EW03_022619				SAMPLE	3	1906112008-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
15	1906112009	EW07_022619				SAMPLE	3	1906112009-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
16	642101	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
17	1906112010	EW04_022619				SAMPLE	3	1906112010-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
18	1906112011	EW08_022619				SAMPLE	3	1906112011-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
19	1906330001	LH18/24-SP140_022819				SAMPLE	3	1906330001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
20	1906332001	LH18/24-SP650_022819_BIX				SAMPLE	3	1906332001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
21	1906334001	LH18-24-SP650_022819_BIX				SAMPLE	3	1906334001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
22	642102	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1905651 (001); 1906112 (001-11); 1906330 (001); 1906332 (001); 1906334 (001)
 ELMS Batch/HBN ID: 2223 (233911)
 Prep Date: 03/04/2019 Analysis Date: 03/05/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\05MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by TNB. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 02/15/2019, sequence 15FEB19D.s Offline Quantitation Method: CLO4-DP1.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 3 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 642100; Target = 4.0µg/L. ASTM type II water was used for LMB 642099.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\23911-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEB104) along with datafiles 05MARD07-10.

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
Batch(es)/SDG: <u>E LMS: 2223 HBN: 233911</u>		
Sample Set IDs if Applicable: <u>1905651 / 190912 / 1906330</u> <u>1906332 / 1906334</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR# _____</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850.WKG Std 100.ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK			Description - 6850 Stock AccStd 1,000ug/mL
Standard: 43659		Created By: Thomas Bosch	
MFG: AccuStandard		Create Date: 09/17/2018 09:09AM	
MFG Lot: 218065075		Amount: 100 mL	
Part ID: IC-PER-10X-1		Expires: 07/25/2020	
		Usable: No	
		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT			Description - 6850 Intermdt AccStd 10.ug/mL		
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK			Description: 6850 QC WKG STD 100ug/L		
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdf Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK			Description: -6850 QC Stock STD 1,000ug/mL
Standard: 36748		Created By: Thomas Bosch	
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	
MFG Lot: CP-0860		Amount: 100 mL	
Part ID: ICC-013		Expires: 03/31/2020	
		Usable: Yes	
		Lab Lot: CLO4 QC STOCK	
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	.CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

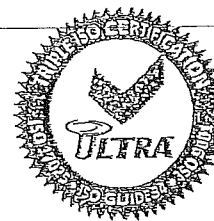
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:
This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:
Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:
The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:
This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:
This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:
Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

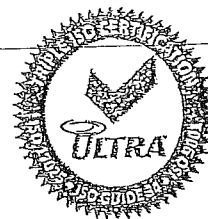
Hazards:
Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:
The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis

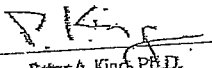


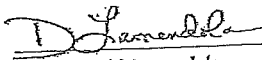
ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:
The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QAVRA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleared Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

For use in routine laboratory analysis.

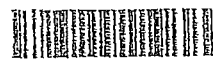


Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:

ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: NaClO₄

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 µg/mL
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 ± 2.8 µg/mL (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount	
*	642096	CCV@25	Vial 71	1	Control	1	2.57589e6	8.017	25.04585
*	642100	QC@4.0	Vial 72	1	Control	2	4.23307e5	8.139	4.10749
*	642098	ICS@4.0	Vial 73	1	Control	3	3.17619e5	7.883	3.92708
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	7.71127e6	7.438	51.25151
*	1906112002	MS	Vial 77	1	Sample	7	8.02696e6	7.464	53.87972
*	1906112003	MSD	Vial 78	1	Sample	8	7.94242e6	7.442	53.69940
*	1906112004		Vial 79	1	Sample	9	2.33017e7	7.227	172.36522
*	1906112005		Vial 80	1	Sample	10	2.30991e7	7.229	180.38568
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.33611e6	8.114	25.01987
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	2.53126e6	7.736	226.48155
*	1906112005	10X	Vial 91	1	Sample	22	2.68237e6	7.739	241.03710
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	3.77363e6	8.090	3708.65666
*	642102	CCV@25	Vial 71	1	Control	25	2.15787e6	8.144	24.45533

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount	
*	642096	CCV@25	Vial 71	1	Control	1	6.79028e5	8.036	25.15984
*	642100	QC@4.0	Vial 72	1	Control	2	1.27412e5	8.156	4.45187
*	642098	ICS@4.0	Vial 73	1	Control	3	9.68721e4	7.906	4.29888
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	2.02786e6	7.452	51.10329
*	1906112002	MS	Vial 77	1	Sample	7	2.14637e6	7.478	54.46797
*	1906112003	MSD	Vial 78	1	Sample	8	2.10991e6	7.456	53.98100
*	1906112004		Vial 79	1	Sample	9	6.79668e6	7.243	180.45962
*	1906112005		Vial 80	1	Sample	10	6.82834e6	7.246	190.34382
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	6.28961e5	8.128	25.63410
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	6.46870e5	7.754	220.91905
*	1906112005	10X	Vial 91	1	Sample	22	6.95643e5	7.756	238.44674
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	1.01021e6	8.102	3771.67475
*	642102	CCV@25	Vial 71	1	Control	25	5.83711e5	8.160	25.16652

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount	
*	642096	CCV@25	Vial 71	1	Control	1	3.12207e5	8.043	5.00000
*	642100	QC@4.0	Vial 72	1	Control	2	3.41038e5	8.157	5.00000
*	642098	ICS@4.0	Vial 73	1	Control	3	2.68237e5	7.902	5.00000
*	642099	LMB	Vial 74	1	Control	4	3.53313e5	8.102	5.00000
*	1905651001		Vial 75	1	Sample	5	3.26356e5	7.780	5.00000
*	1906112001		Vial 76	1	Sample	6	4.26473e5	7.464	5.00000
*	1906112002	MS	Vial 77	1	Sample	7	4.19549e5	7.491	5.00000
*	1906112003	MSD	Vial 78	1	Sample	8	4.16709e5	7.467	5.00000
*	1906112004		Vial 79	1	Sample	9	2.95705e5	7.253	5.00000
*	1906112005		Vial 80	1	Sample	10	2.75946e5	7.263	5.00000
*	1906112006		Vial 81	1	Sample	11	3.73575e5	7.480	5.00000
*	1906112007		Vial 82	1	Sample	12	5.00533e5	7.256	5.00000
*	1906112008		Vial 83	1	Sample	13	4.65121e5	7.237	5.00000
*	1906112009		Vial 84	1	Sample	14	2.43675e5	7.672	5.00000
*	1906112010		Vial 85	1	Sample	15	2.51865e5	7.710	5.00000
*	1906112011		Vial 86	1	Sample	16	2.80792e5	7.876	5.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.83460e5	8.140	5.00000
*	1906332001		Vial 88	1	Sample	19	2.64674e5	7.827	5.00000
*	1906334001		Vial 89	1	Sample	20	2.65662e5	7.808	5.00000
*	1906112004	10X	Vial 90	1	Sample	21	3.41608e5	7.756	50.00000
*	1906112005	10X	Vial 91	1	Sample	22	3.38724e5	7.764	50.00000
*	1906112007	RE	Vial 82	1	Sample	23	4.31135e5	7.332	5.00000
*	1906330001	100	Vial 92	1	Sample	24	2.98985e5	8.111	500.00000
*	642102	CCV@25	Vial 71	1	Control	25	2.68305e5	8.165	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

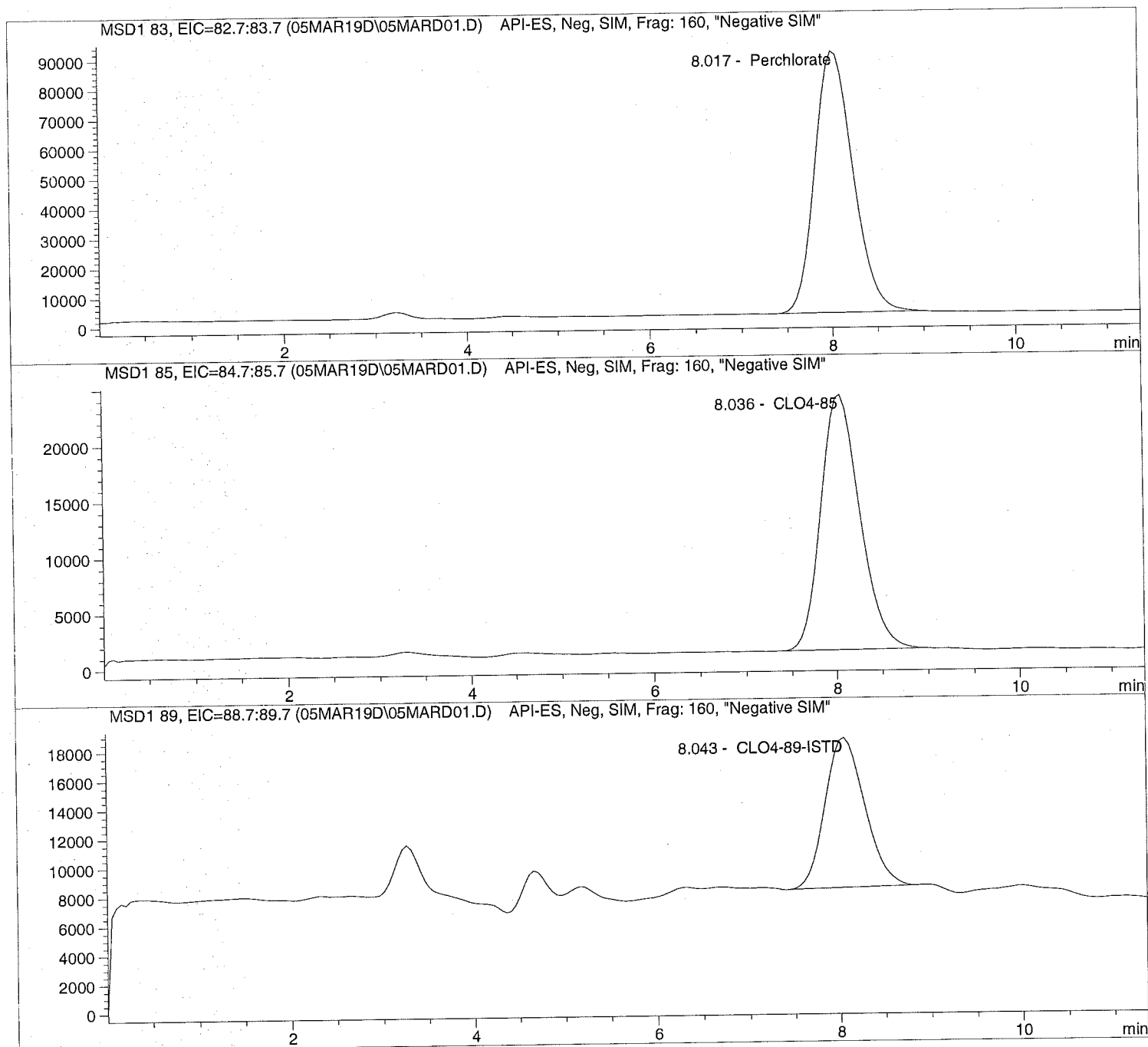
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	642096	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	642100	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	642098	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	642099	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1905651001		CLO4-AQN	1	Sample	
6	Vial 76	1906112001		CLO4-AQN	1	Sample	
7	Vial 77	1906112002	MS	CLO4-AQN	1	Sample	
8	Vial 78	1906112003	MSD	CLO4-AQN	1	Sample	
9	Vial 79	1906112004		CLO4-AQN	1	Sample	
10	Vial 80	1906112005		CLO4-AQN	1	Sample	
11	Vial 81	1906112006		CLO4-AQN	1	Sample	
12	Vial 82	1906112007		CLO4-AQN	1	Sample	
13	Vial 83	1906112008		CLO4-AQN	1	Sample	
14	Vial 84	1906112009		CLO4-AQN	1	Sample	
15	Vial 85	1906112010		CLO4-AQN	1	Sample	
16	Vial 86	1906112011		CLO4-AQN	1	Sample	
17	Vial 71	642101	CCV@25	CLO4-AQN	1	Ctrl Samp	
18	Vial 87	1906330001	1K	CLO4-AQN	1	Sample	
19	Vial 88	1906332001		CLO4-AQN	1	Sample	
20	Vial 89	1906334001		CLO4-AQN	1	Sample	
21	Vial 90	1906112004	10X	CLO4-AQN	1	Sample	
22	Vial 91	1906112005	10X	CLO4-AQN	1	Sample	
23	Vial 82	1906112007	RE	CLO4-AQN	1	Sample	
24	Vial 92	1906330001	100	CLO4-AQN	1	Sample	
25	Vial 71	642102	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD01.D Sample Name: 642096 CCV@25

Injection Date: 3/05/2019 08:44:45 Seq Line: 1
Sample Name: 642096 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD01.D Sample Name: 642096 CCV@25

```

=====
Injection Date: 3/05/2019 08:44:45      Seq Line: 1
Sample Name: 642096    CCV@25      Location: Vial 71
Acq Operator: TNB      Inj. No.: 1
                                     Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.017	PBA	2575886.3	25.0459	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.036	PBA	679028.4	25.1598	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.043	PBA	312206.9	5.0000	CLO4-89-ISTD

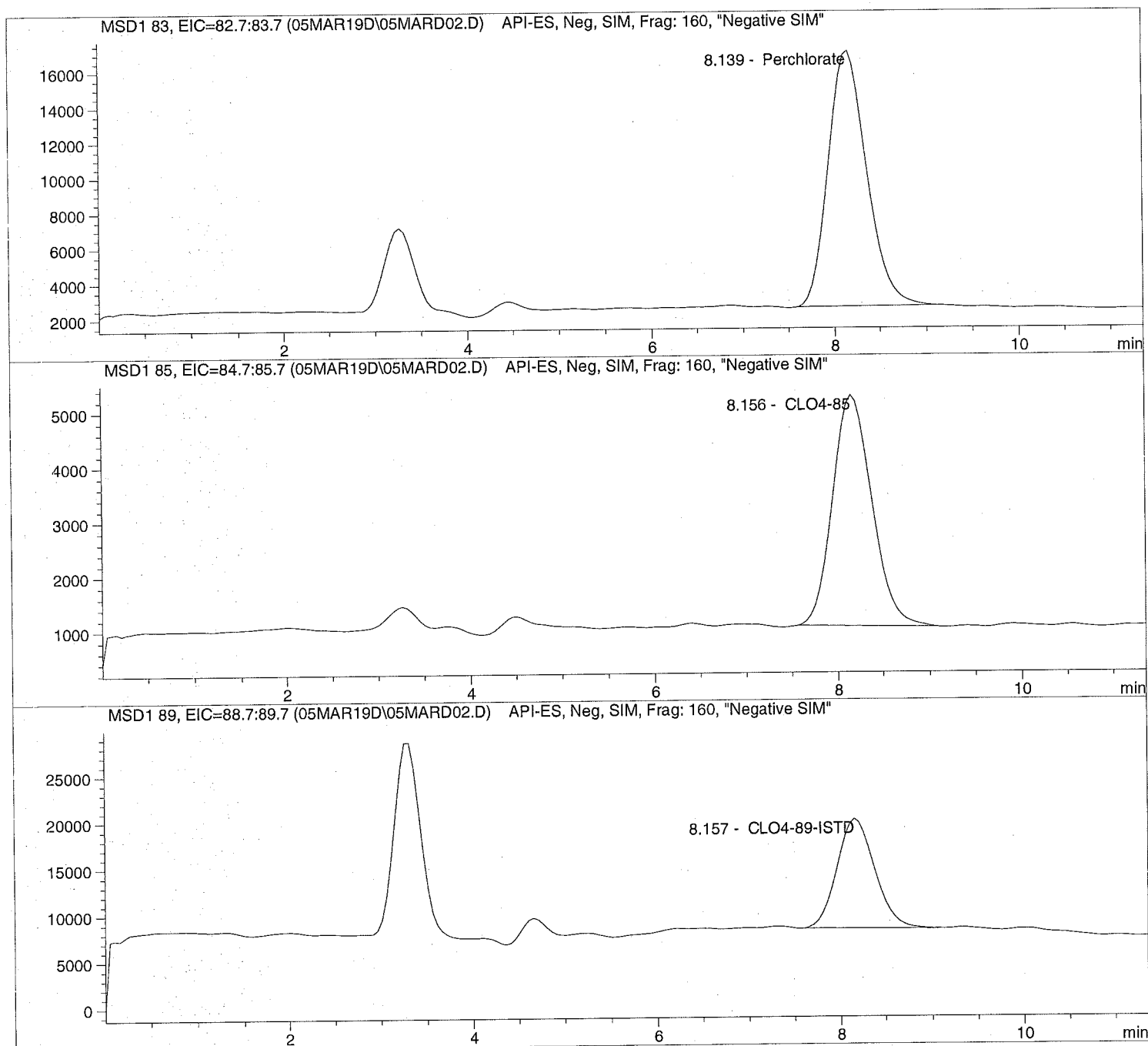
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD02.D Sample Name: 642100 QC@4.0

Injection Date: 3/05/2019 09:00:30 Seq Line: 2
Sample Name: 642100 QC@4.0 Location: Vial 72
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD02.D Sample Name: 642100 QC@4.0

Injection Date: 3/05/2019 09:00:30 Seq Line: 2
 Sample Name: 642100 QC@4.0 Location: Vial 72
 Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
 Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
 Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
 Multiplier: 1.000000
 Dilution: 1.000000
 Sample Amount: 4.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.139	PBA	423307.1	4.1075	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.156	PBA	127412.0	4.4519	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.157	PBA	341038.3	5.0000	CLO4-89-ISTD

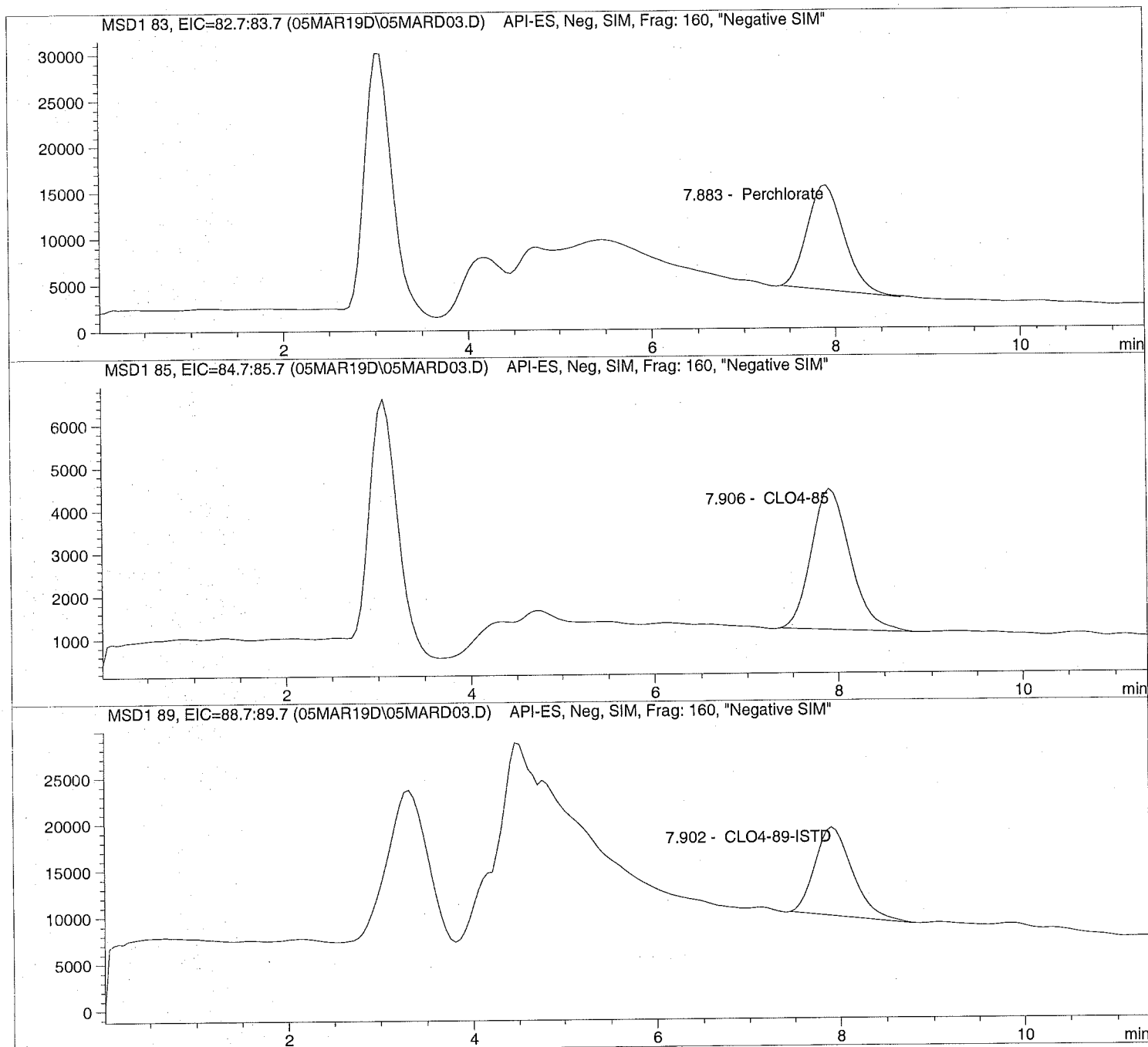
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD03.D Sample Name: 642098 ICS@4.0

Injection Date: 3/05/2019 09:13:34 Seq Line: 3
Sample Name: 642098 ICS@4.0 Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD03.D Sample Name: 642098 ICS@4.0

```

=====
Injection Date: 3/05/2019 09:13:34      Seq Line: 3
Sample Name: 642098 ICS@4.0           Location: Vial 73
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.883	PBA	317618.9	3.9271	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.906	PBA	96872.1	4.2989	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.902	PBA	268236.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

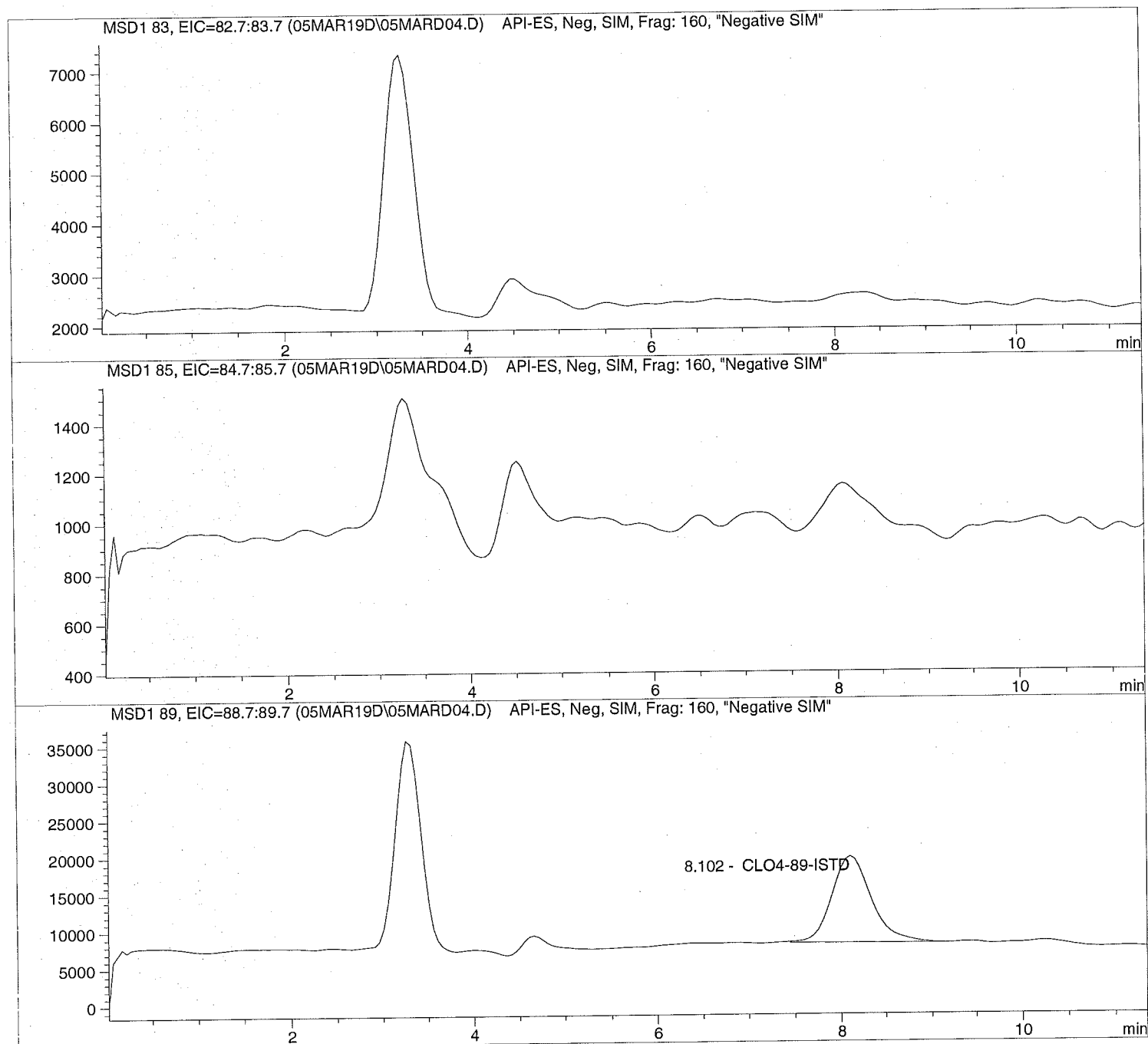
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD04.D Sample Name: 642099 LMB

=====
Injection Date: 3/05/2019 09:26:40 Seq Line: 4
Sample Name: 642099 LMB Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis
=====



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD04.D Sample Name: 642099 LMB

```

=====
Injection Date: 3/05/2019 09:26:40      Seq Line: 4
Sample Name:    642099 LMB              Location:  Vial 74
Acq Operator:   TNB                    Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	BBA	353313.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

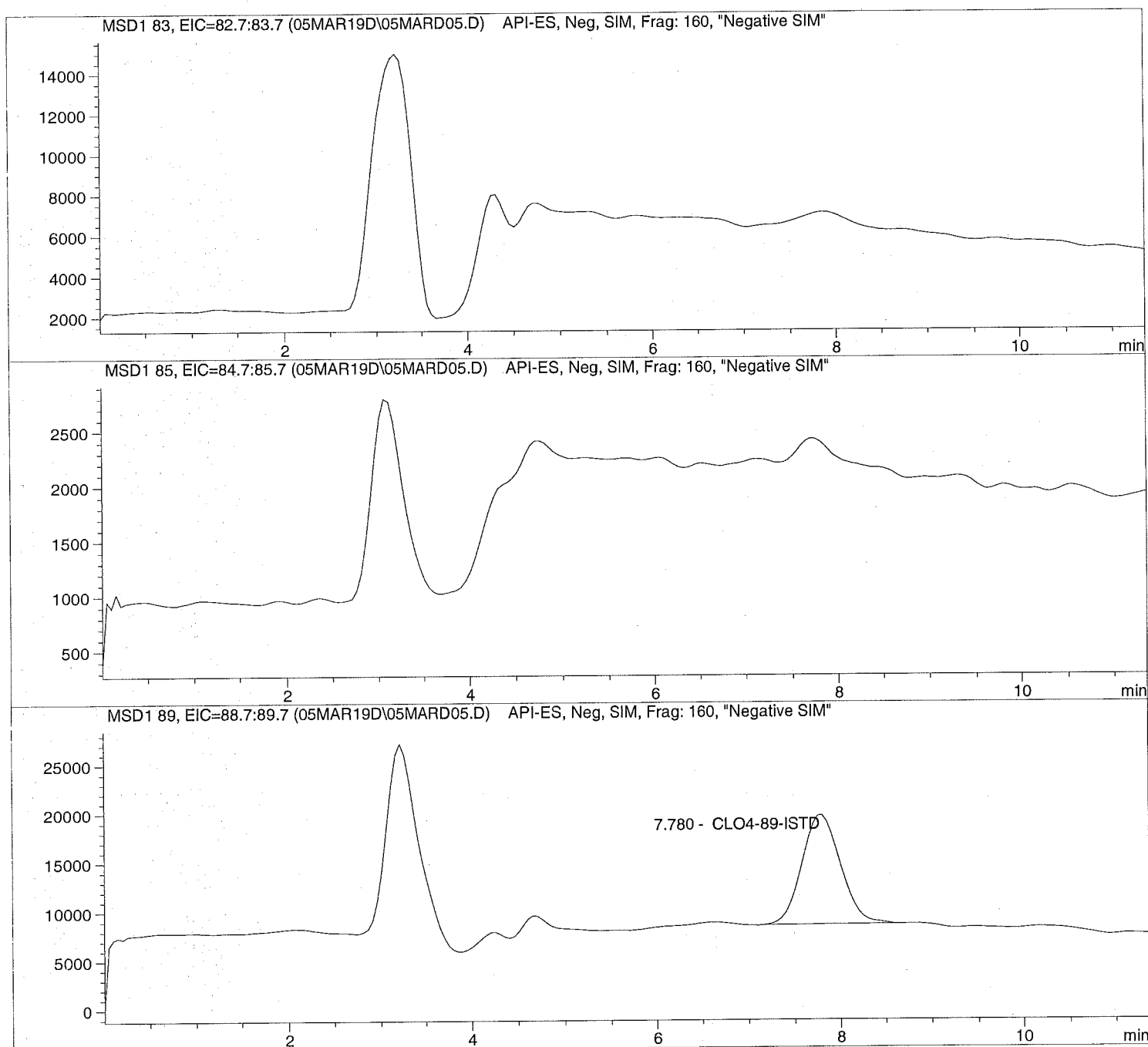
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD05.D Sample Name: 1905651001

```
=====
Injection Date: 3/05/2019 09:40:58      Seq Line:      5
Sample Name:    1905651001              Location:      Vial 75
Acq Operator:   TNB                      Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD05.D Sample Name: 1905651001

```

=====
Injection Date: 3/05/2019 09:40:58      Seq Line:          5
Sample Name:    1905651001              Location:         Vial 75
Acq Operator:   TNB                      Inj. No.:        1
                                           Inj. Vol.:       20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.780	PBA	326356.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

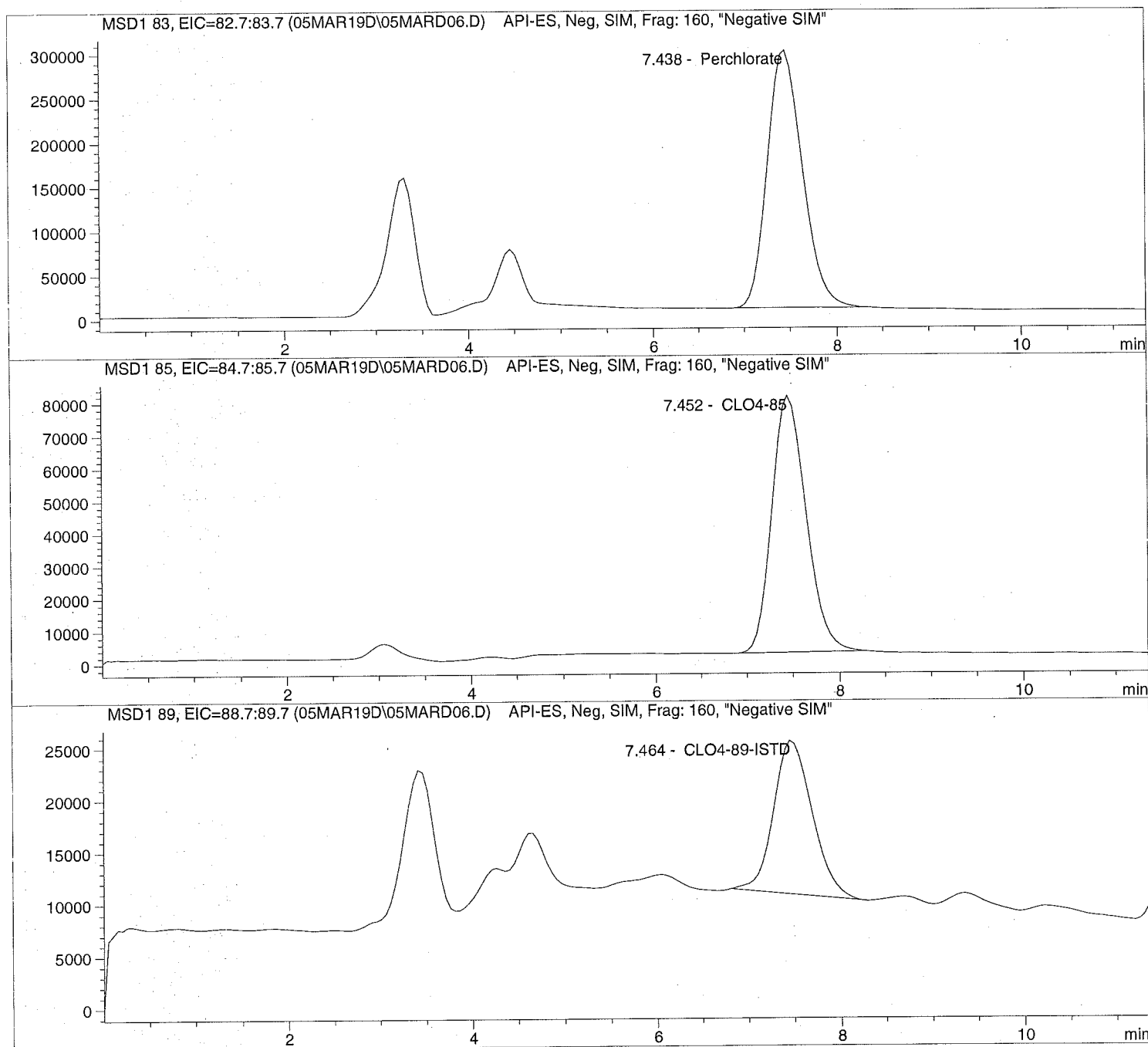
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD06.D Sample Name: 1906112001

```
=====
Injection Date: 3/05/2019 09:54:04      Seq Line: 6
Sample Name: 1906112001                  Location: Vial 76
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD06.D Sample Name: 1906112001

```

=====
Injection Date: 3/05/2019 09:54:04      Seq Line: 6
Sample Name: 1906112001                Location: Vial 76
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.438	PBA	7711270.5	51.2515	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.452	PBA	2027855.1	51.1033	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PB	426473.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

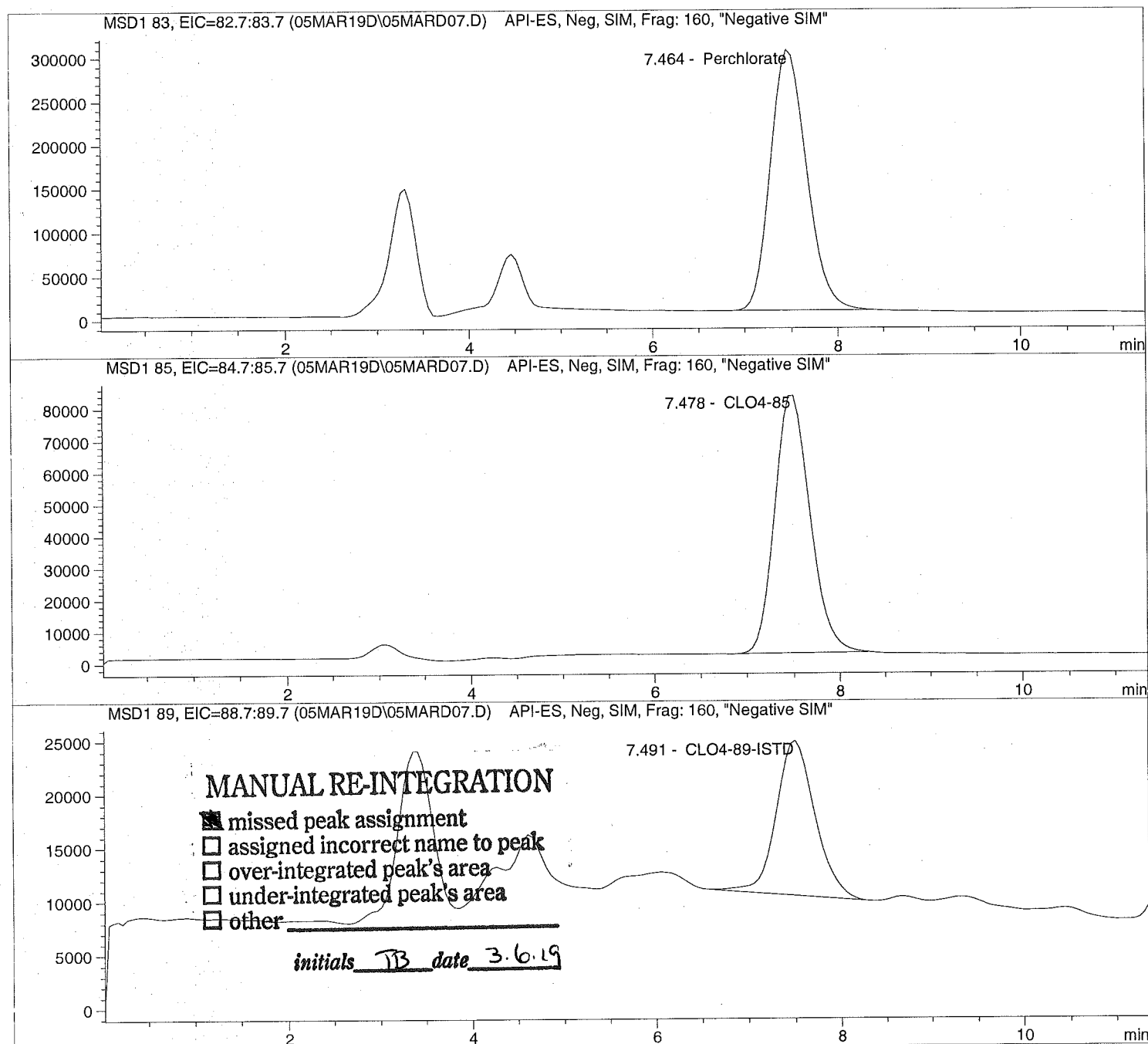
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

=====
 Injection Date: 3/05/2019 10:07:11 Seq Line: 7
 Sample Name: 1906112002 MS Location: Vial 77
 Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
 Last Changed: 2/19/2019 12:13:46

Perchlorate analysis
 =====



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

```

=====
Injection Date: 3/05/2019 10:07:11      Seq Line: 7
Sample Name: 1906112002 MS              Location: Vial 77
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	53.8797	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	54.4680	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.491	MM	419549.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

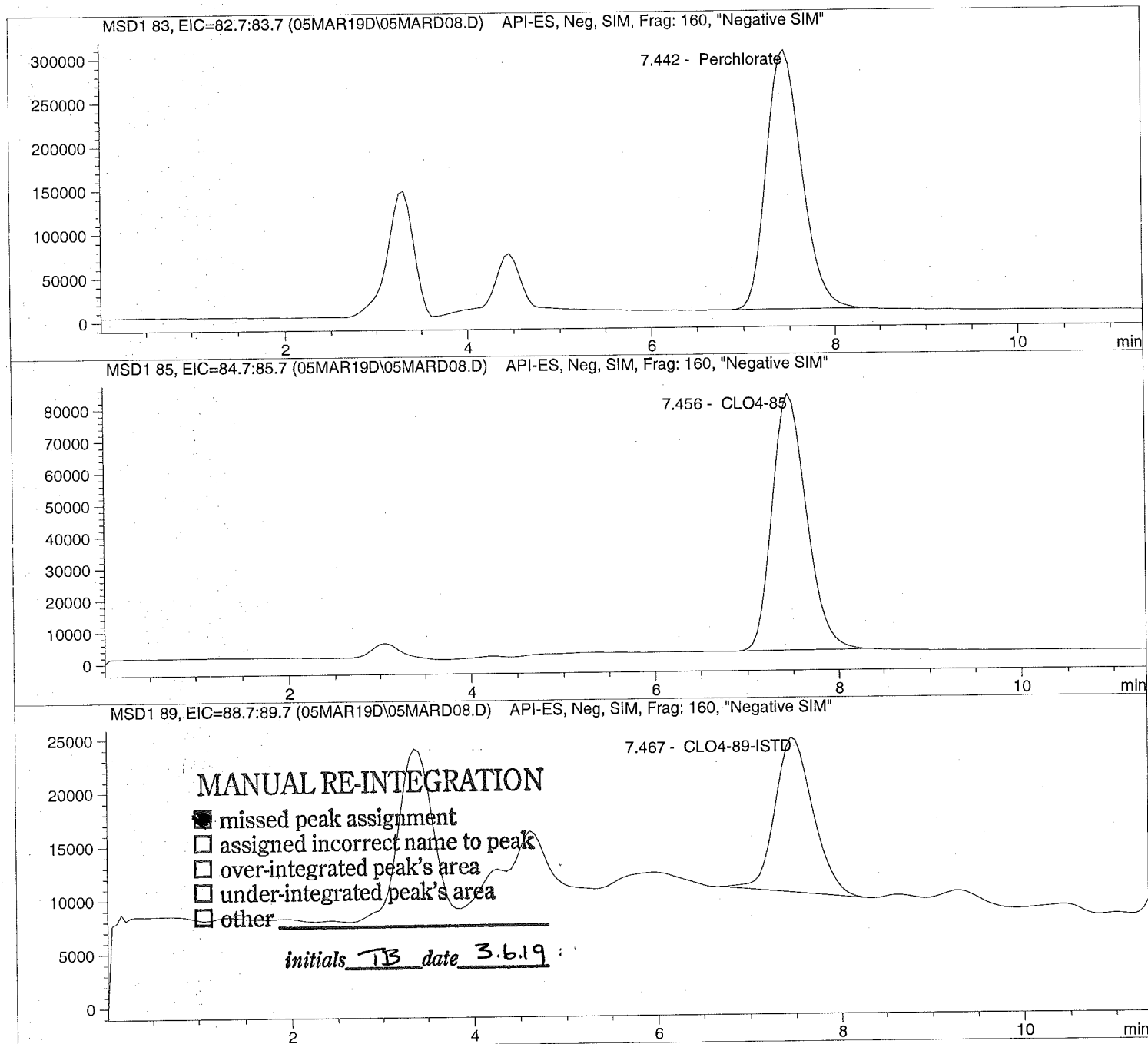
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

```

=====
Injection Date: 3/05/2019 10:20:17      Seq Line:      8
Sample Name:   1906112003  MSD          Location:      Vial 78
Acq Operator:  TNB                    Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	53.6994	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	53.9810	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.467	MM	416709.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D

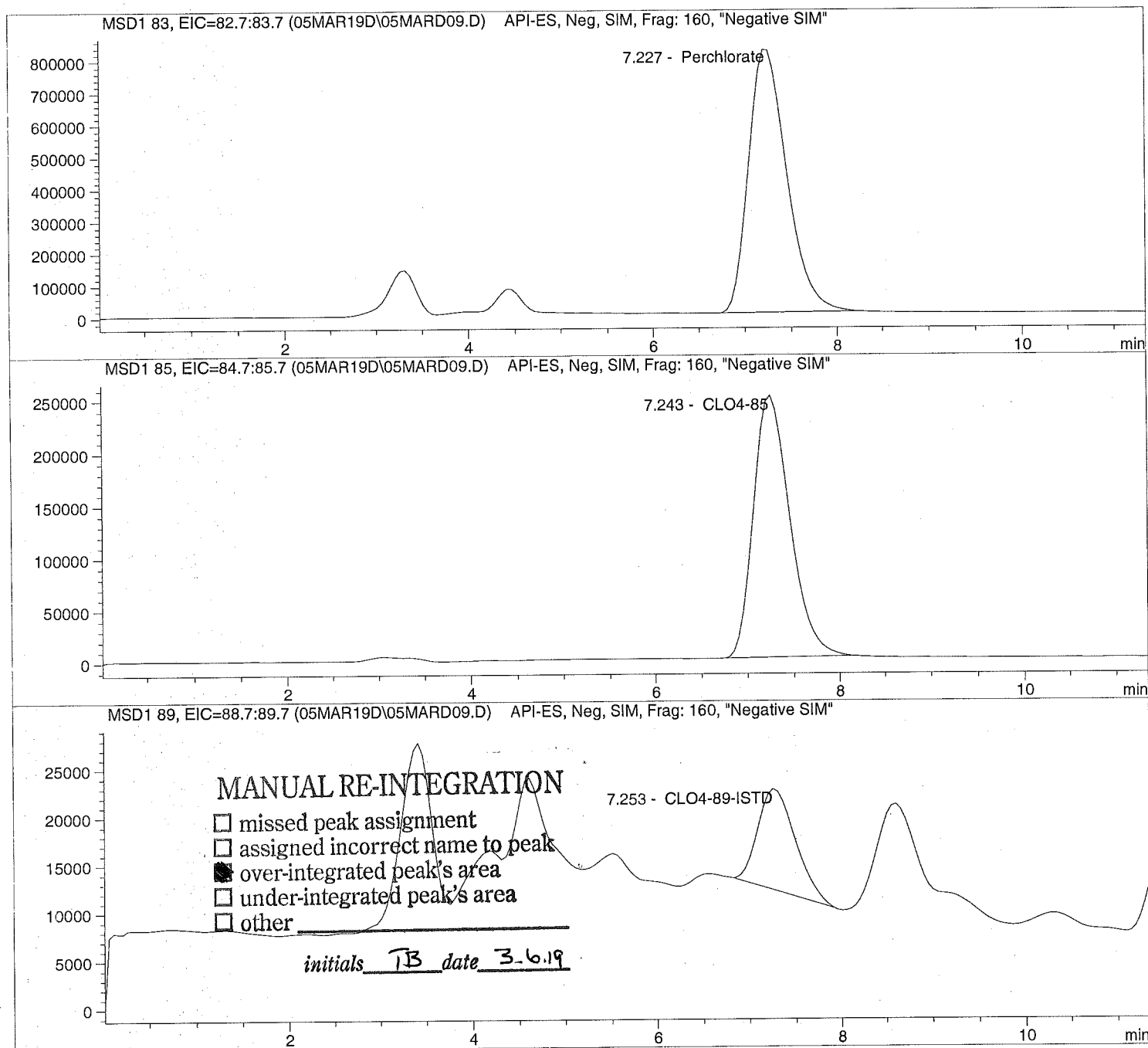
Sample Name: 1906112004

Injection Date: 3/05/2019 10:33:21
Sample Name: 1906112004
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D Sample Name: 1906112004

```

=====
Injection Date: 3/05/2019 10:33:21      Seq Line:          9
Sample Name:    1906112004              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	172.3652	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	180.4596	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.253	MM	295705.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

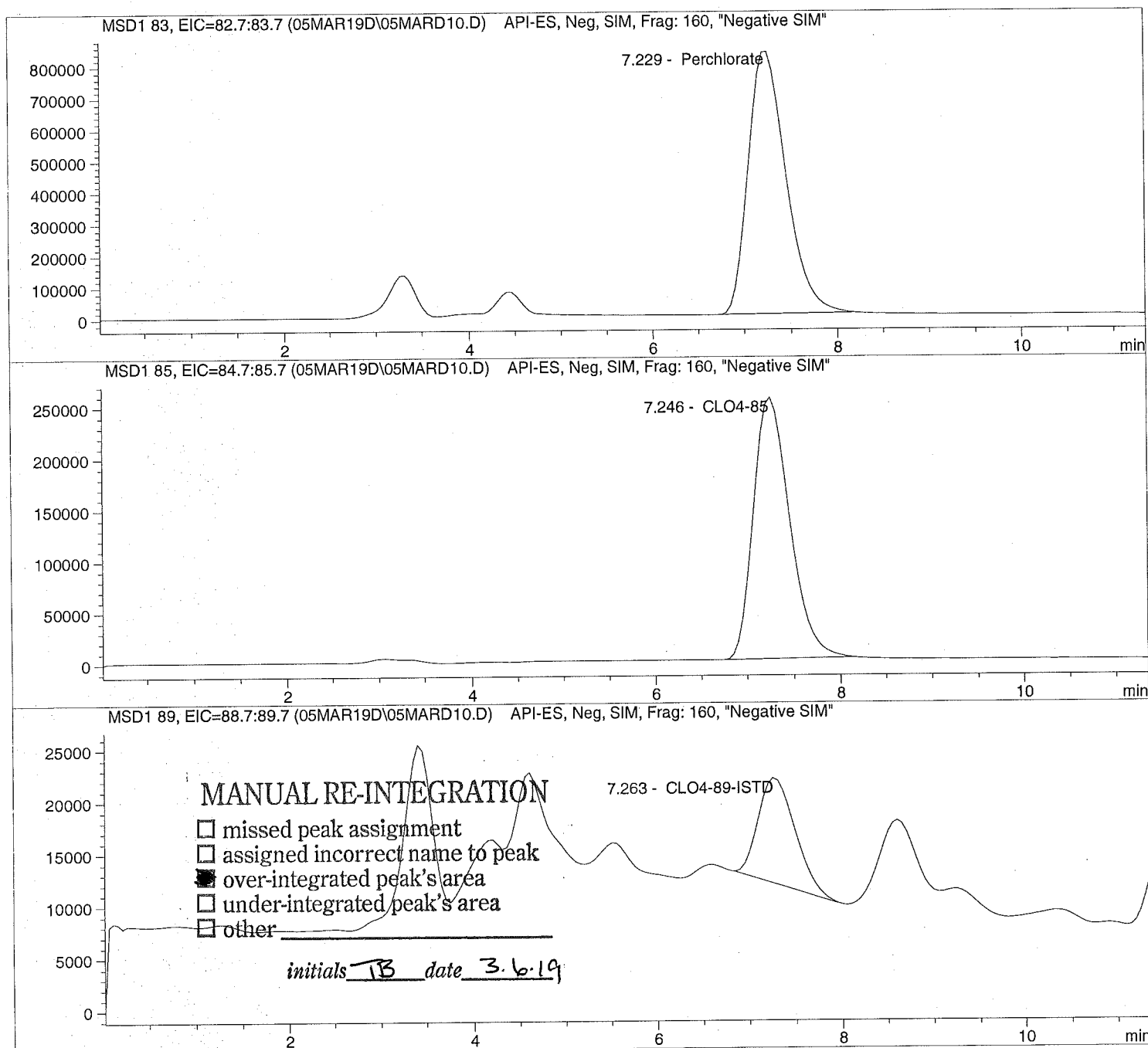
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D Sample Name: 1906112005

=====
Injection Date: 3/05/2019 10:46:26 Seq Line: 10
Sample Name: 1906112005 Location: Vial 80
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D Sample Name: 1906112005

```

=====
Injection Date: 3/05/2019 10:46:26      Seq Line:          10
Sample Name:    1906112005              Location:          Vial 80
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	180.3857	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	190.3438	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	MM	275946.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

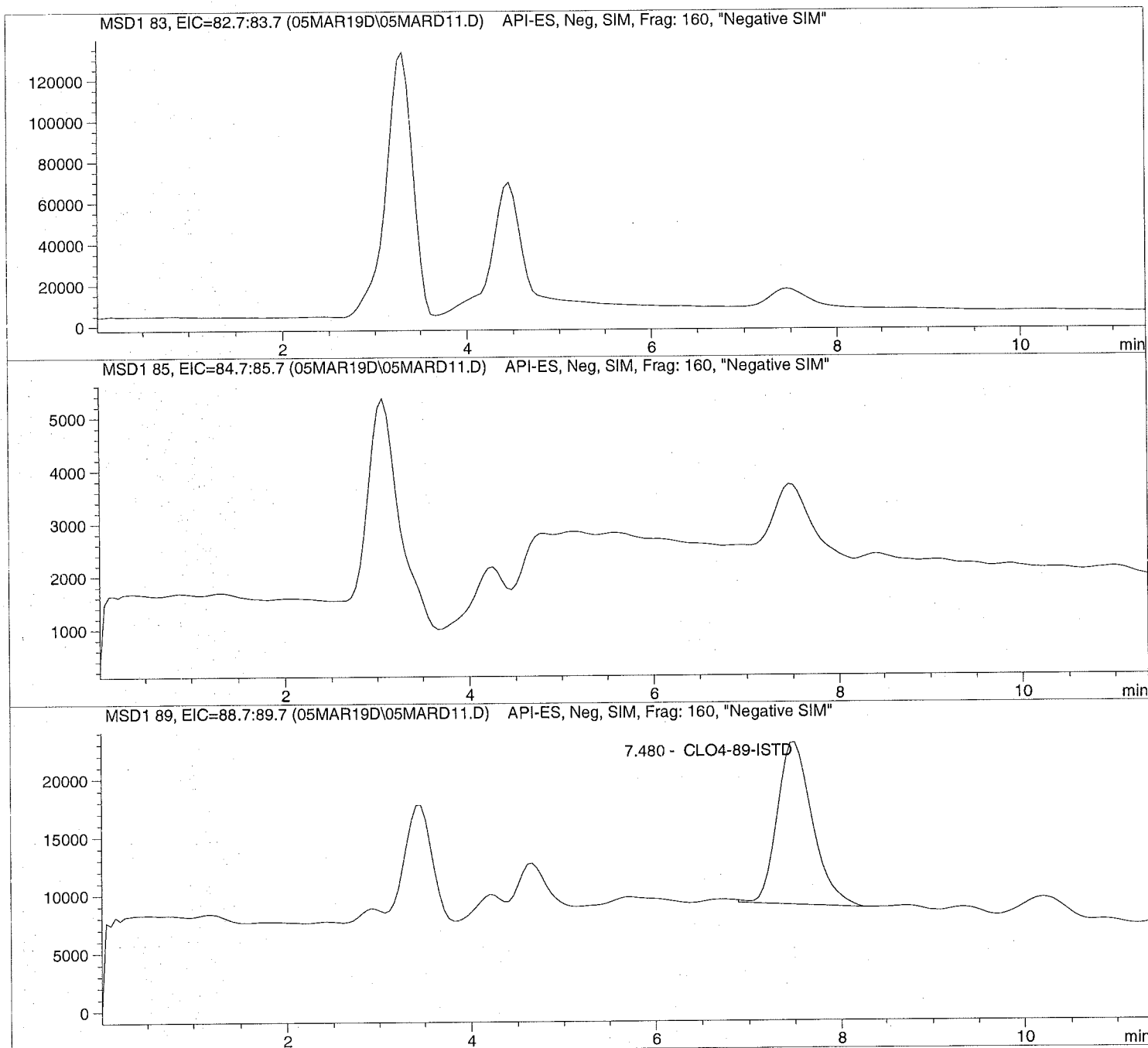
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD11.D Sample Name: 1906112006

```
=====
Injection Date: 3/05/2019 10:59:36      Seq Line:          11
Sample Name:    1906112006              Location:          Vial 81
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD11.D Sample Name: 1906112006

```

=====
Injection Date: 3/05/2019 10:59:36      Seq Line:          11
Sample Name:    1906112006              Location:          Vial 81
Acq Operator:   TNB                      Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.480	BBA	373575.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

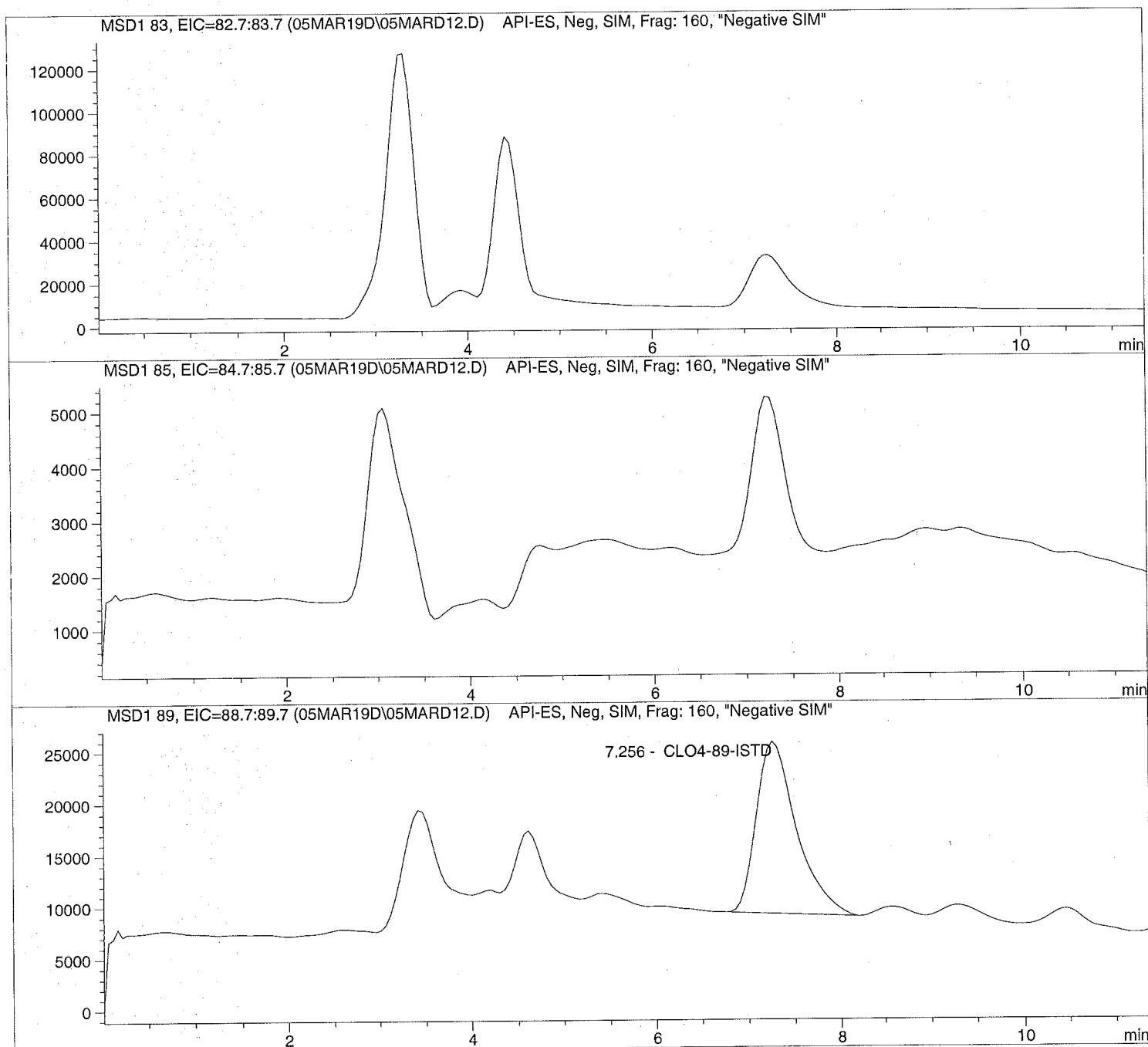
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD12.D Sample Name: 1906112007

```
=====
Injection Date: 3/05/2019 11:12:28 Seq Line: 12
Sample Name: 1906112007 Location: Vial 82
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD12.D Sample Name: 1906112007

```

=====
Injection Date: 3/05/2019 11:12:28      Seq Line: 12
Sample Name: 1906112007                Location: Vial 82
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.256	PB	500532.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

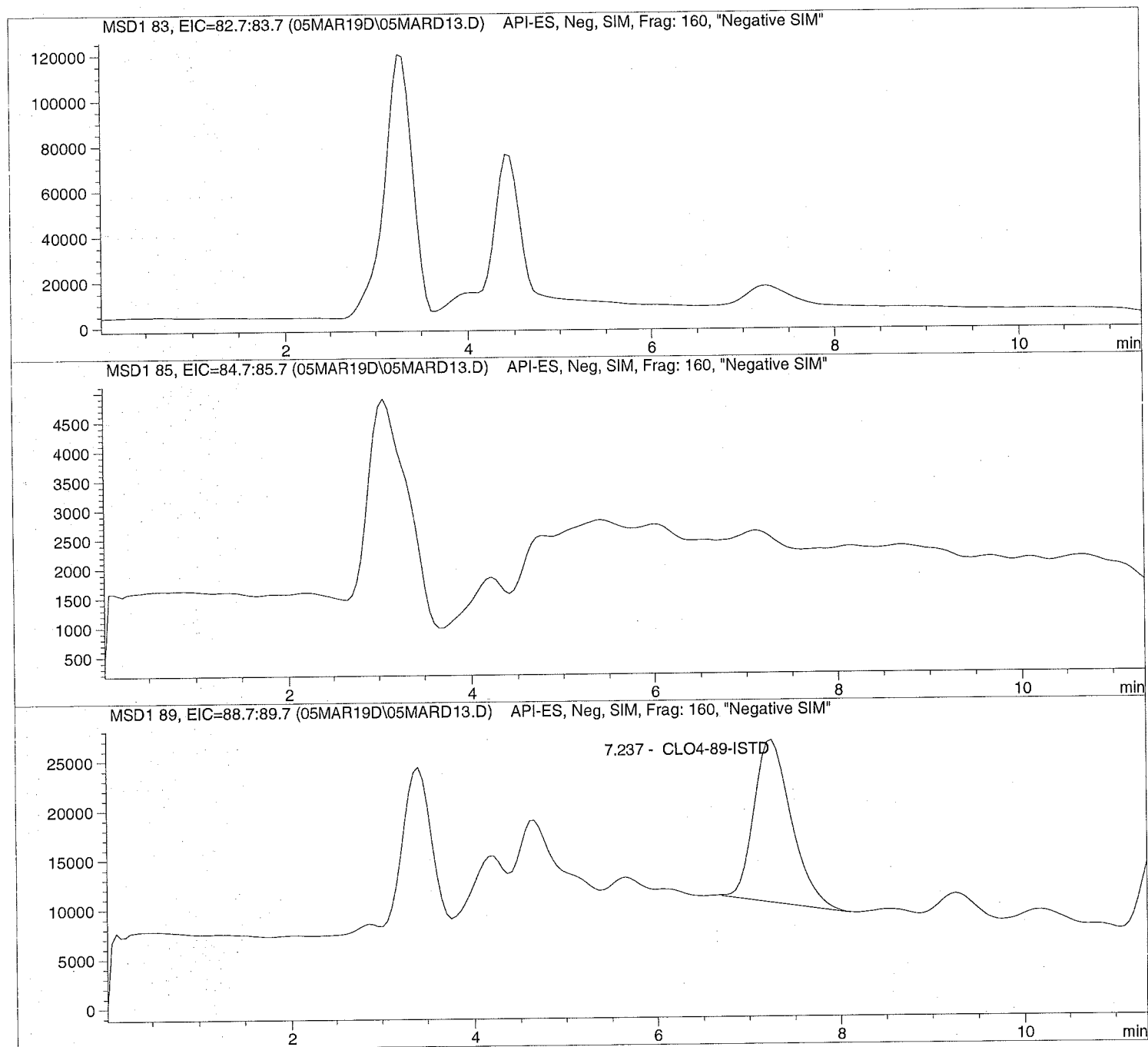
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD13.D Sample Name: 1906112008

```
=====
Injection Date: 3/05/2019 11:26:23      Seq Line:          13
Sample Name:    1906112008              Location:         Vial 83
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD13.D Sample Name: 1906112008

```

=====
Injection Date: 3/05/2019 11:26:23      Seq Line: 13
Sample Name: 1906112008                 Location: Vial 83
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.237	BBA	465121.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D Sample Name: 1906112009

```

=====
Injection Date: 3/05/2019 11:39:24      Seq Line:      14
Sample Name:    1906112009              Location:      Vial 84
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====

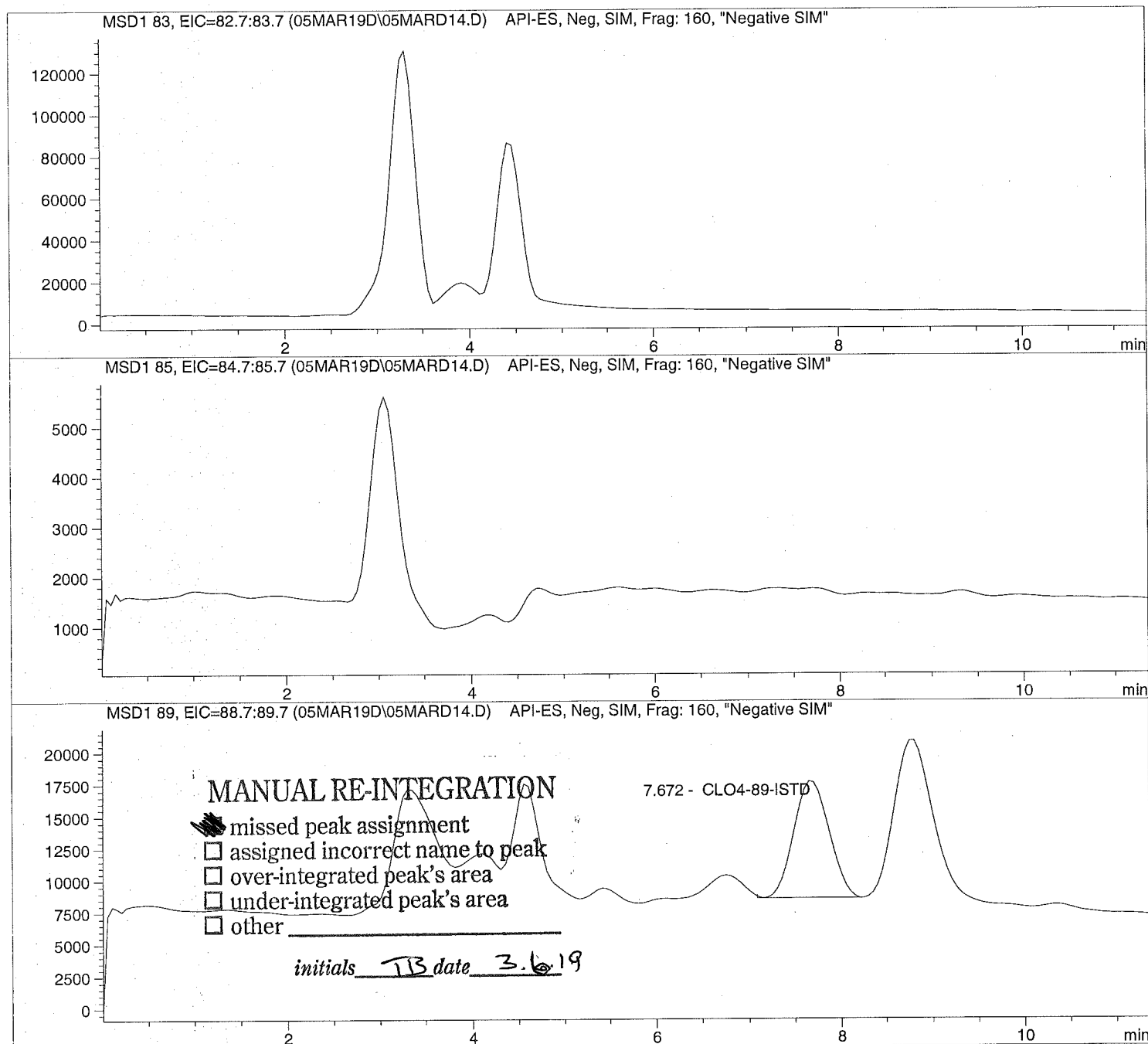
```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D Sample Name: 1906112009

```

=====
Injection Date: 3/05/2019 11:39:24      Seq Line: 14
Sample Name: 1906112009                Location: Vial 84
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

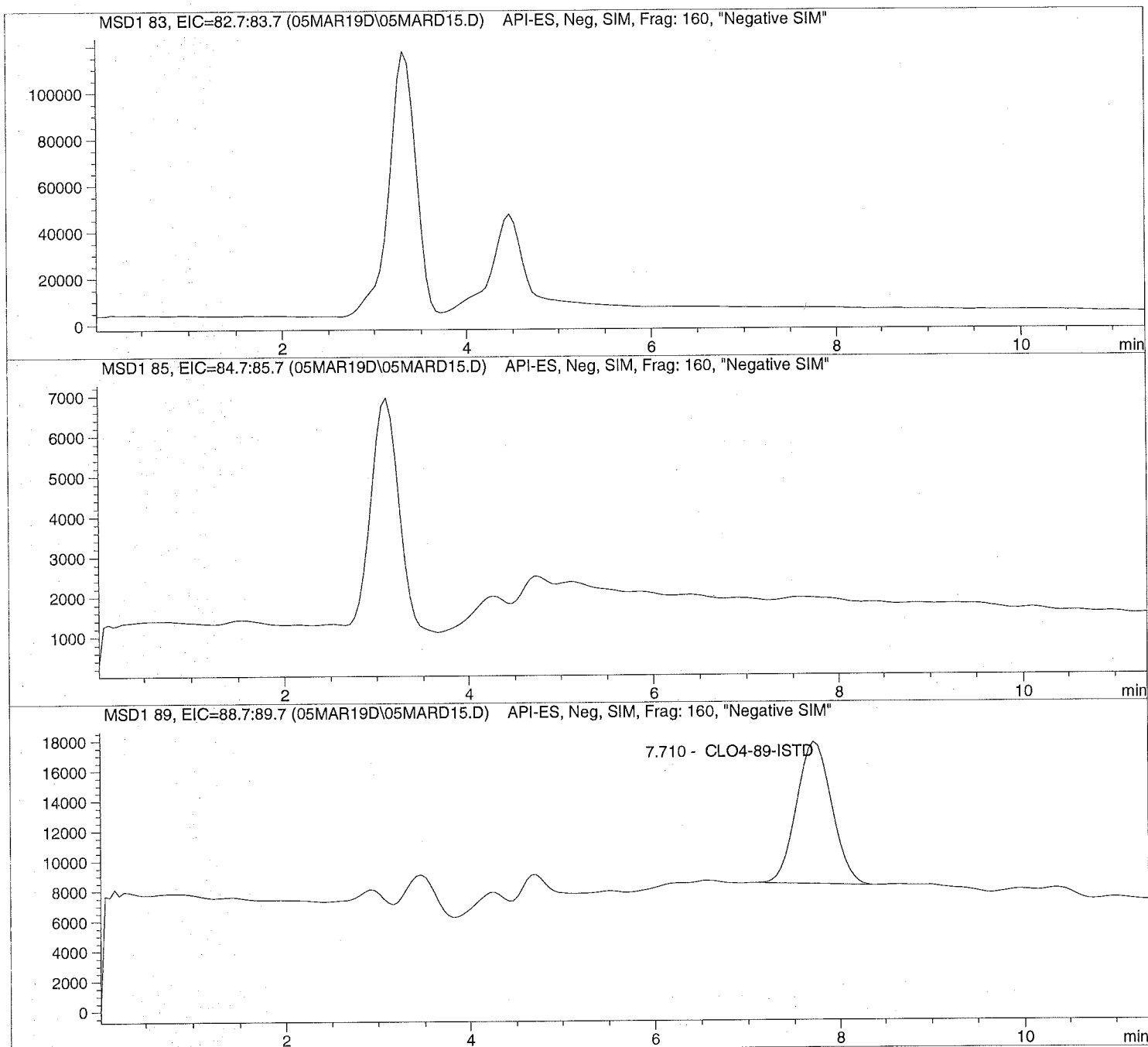
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD15.D Sample Name: 1906112010

```
=====
Injection Date: 3/05/2019 11:52:27      Seq Line:          15
Sample Name:    1906112010              Location:          Vial 85
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD15.D Sample Name: 1906112010

```

=====
Injection Date: 3/05/2019 11:52:27      Seq Line:          15
Sample Name:   1906112010              Location:         Vial 85
Acq Operator:  TNB                      Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.710	BBA	251865.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

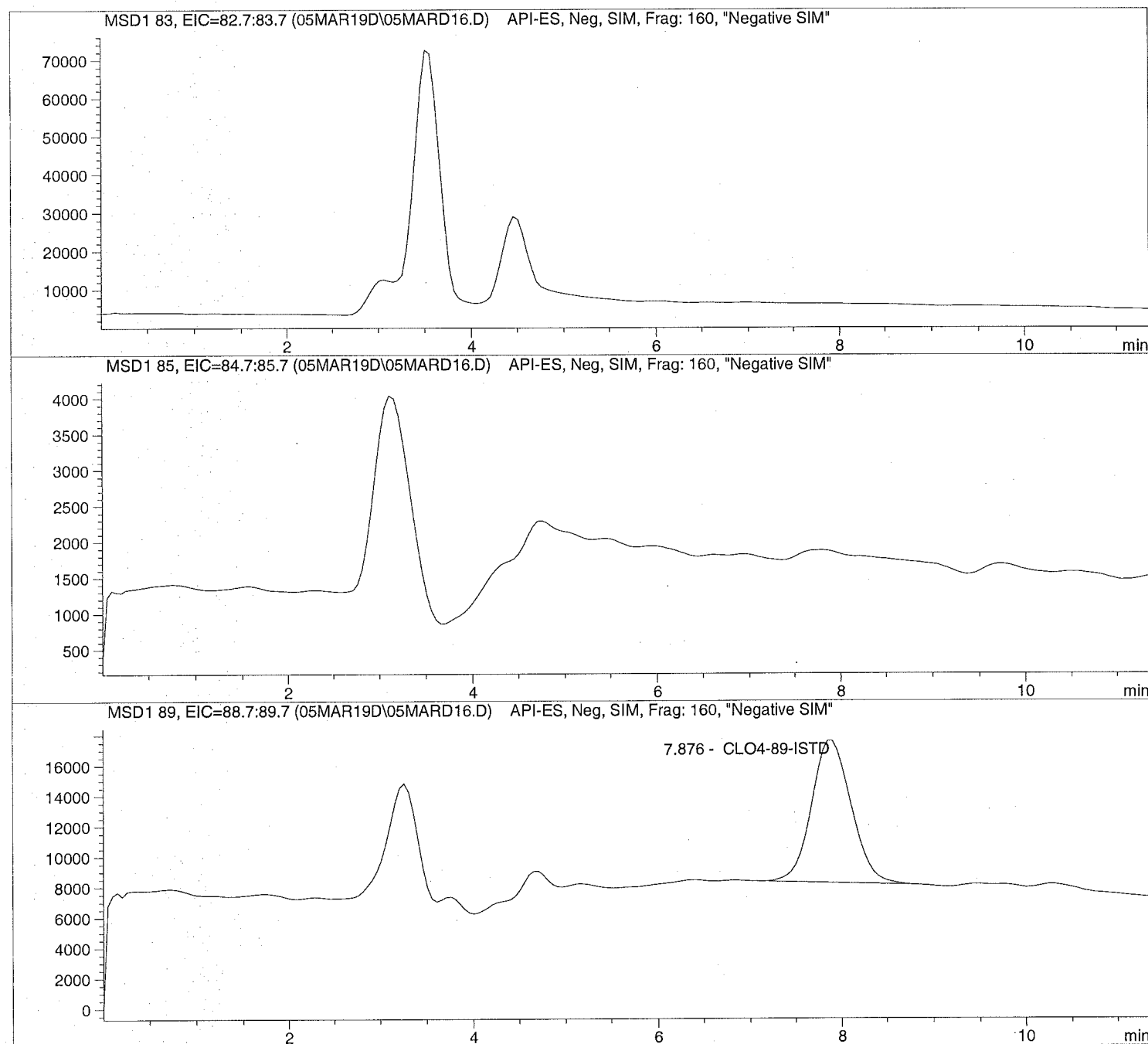
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD16.D Sample Name: 1906112011

```
=====
Injection Date:  3/05/2019  12:05:39      Seq Line:      16
Sample Name:    1906112011                Location:      Vial 86
Acq Operator:   TNB                       Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD16.D Sample Name: 1906112011

```

=====
Injection Date: 3/05/2019 12:05:39      Seq Line:          16
Sample Name:    1906112011              Location:          Vial 86
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:       20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.876	BBA	280792.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

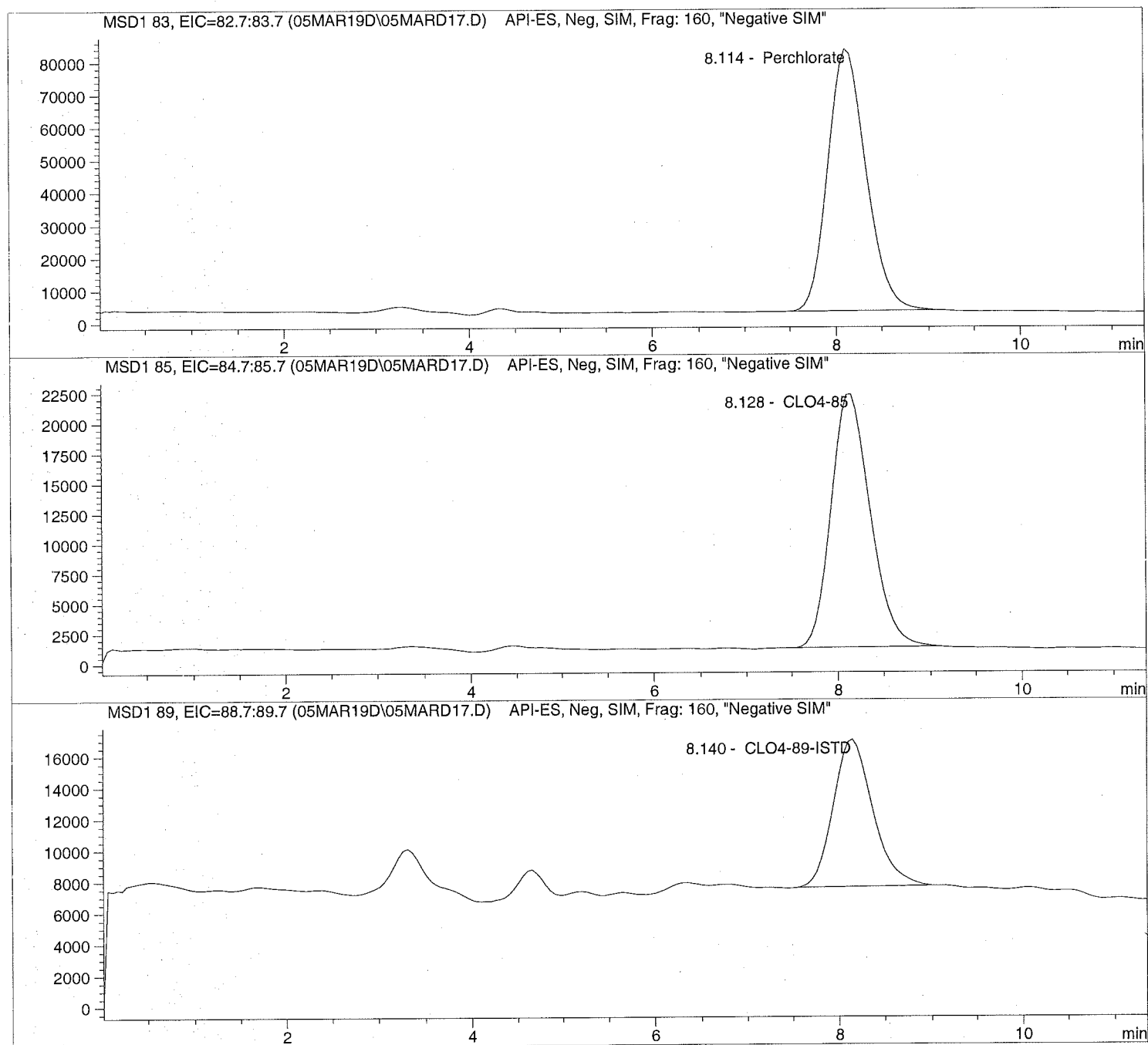
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD17.D Sample Name: 642101 CCV025

```
=====
Injection Date: 3/05/2019 12:18:41 Seq Line: 17
Sample Name: 642101 CCV025 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD17.D Sample Name: 642101 CCV@25

```

=====
Injection Date: 3/05/2019 12:18:41      Seq Line: 17
Sample Name: 642101 CCV@25             Location: Vial 71
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	2336112.7	25.0199	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.128	PBA	628961.2	25.6341	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.140	PBA	283460.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

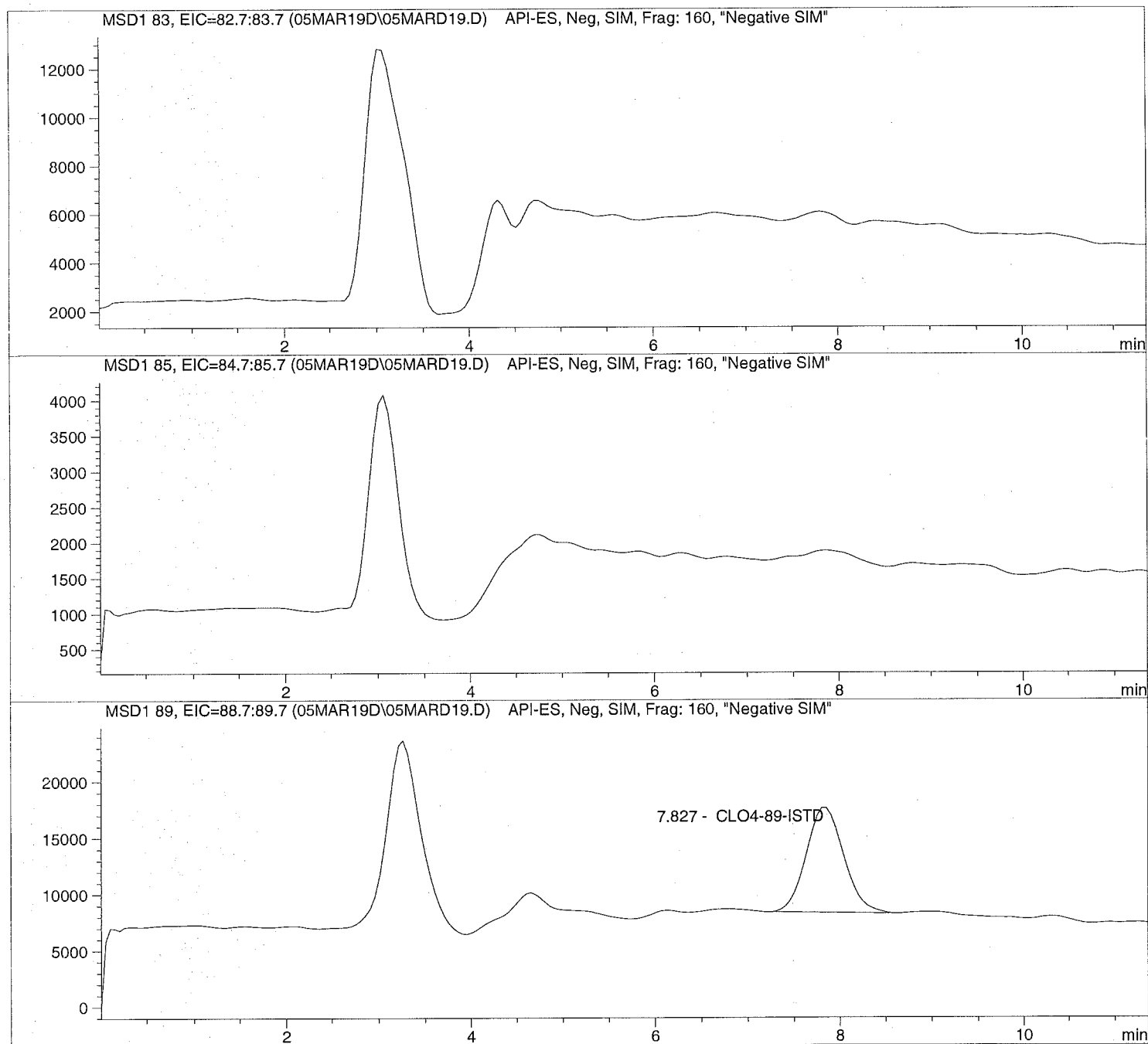
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD19.D Sample Name: 1906332001

```
=====
Injection Date: 3/05/2019 12:49:07      Seq Line:          19
Sample Name:    1906332001              Location:          Vial 88
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD19.D Sample Name: 1906332001

```

=====
Injection Date: 3/05/2019 12:49:07      Seq Line:      19
Sample Name:    1906332001              Location:      Vial 88
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.827	PBA	264674.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD20.D

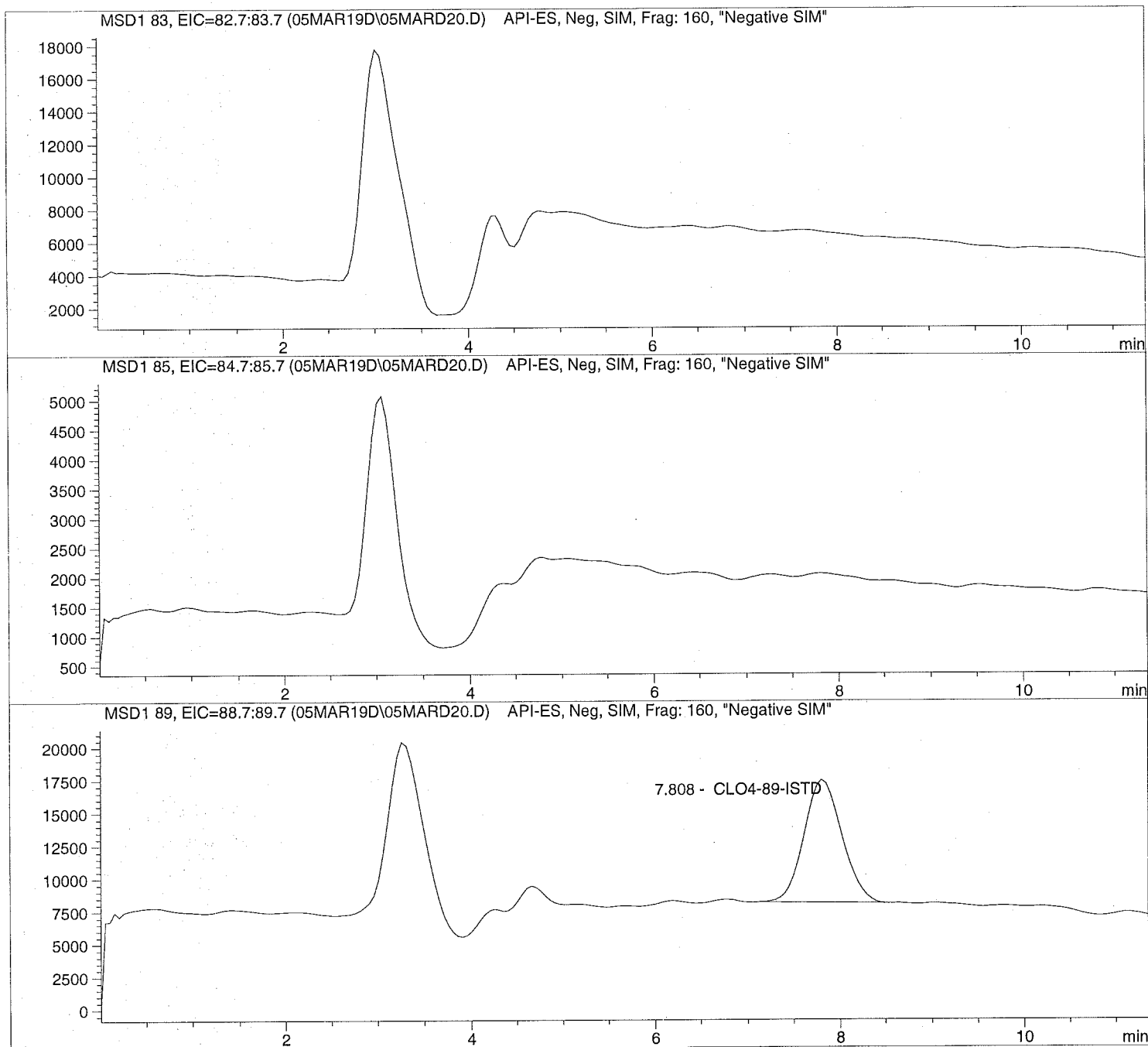
Sample Name: 1906334001

Injection Date: 3/05/2019 13:02:15
Sample Name: 1906334001
Acq Operator: TNB

Seq Line: 20
Location: Vial 89
Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD20.D Sample Name: 1906334001

```

=====
Injection Date: 3/05/2019 13:02:15      Seq Line:          20
Sample Name:   1906334001              Location:          Vial 89
Acq Operator:  TNB                      Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.808	PBA	265662.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

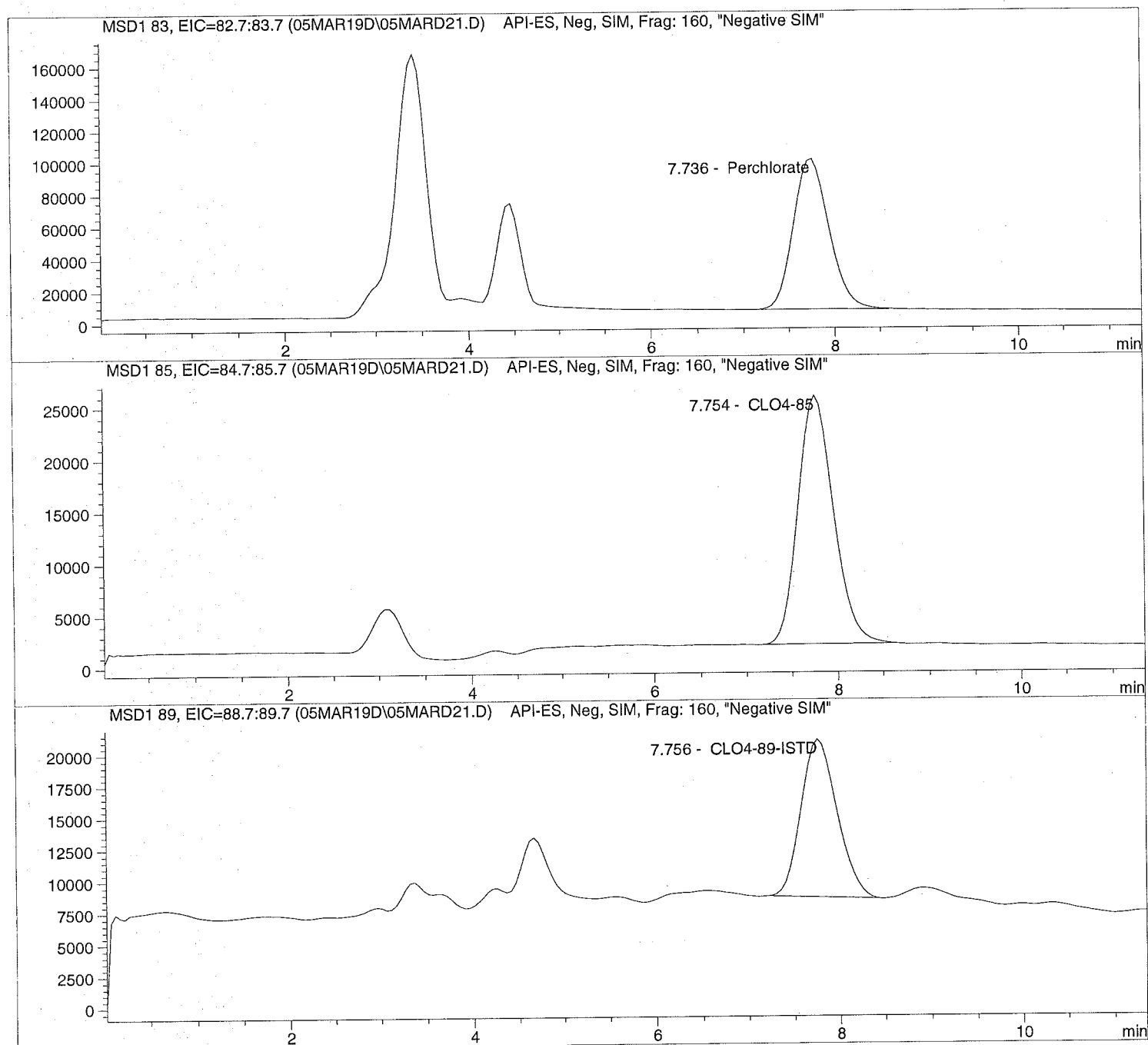
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD21.D Sample Name: 1906112004 10X

=====
Injection Date: 3/05/2019 13:15:18 Seq Line: 21
Sample Name: 1906112004 10X Location: Vial 90
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD21.D Sample Name: 1906112004 10X

```

=====
Injection Date: 3/05/2019 13:15:18      Seq Line:          21
Sample Name:   1906112004 10X          Location:         Vial 90
Acq Operator:  TNB                    Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      10.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.736	PBA	2531259.3	226.4815	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.754	PBA	646870.4	220.9191	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	PB	341607.9	50.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

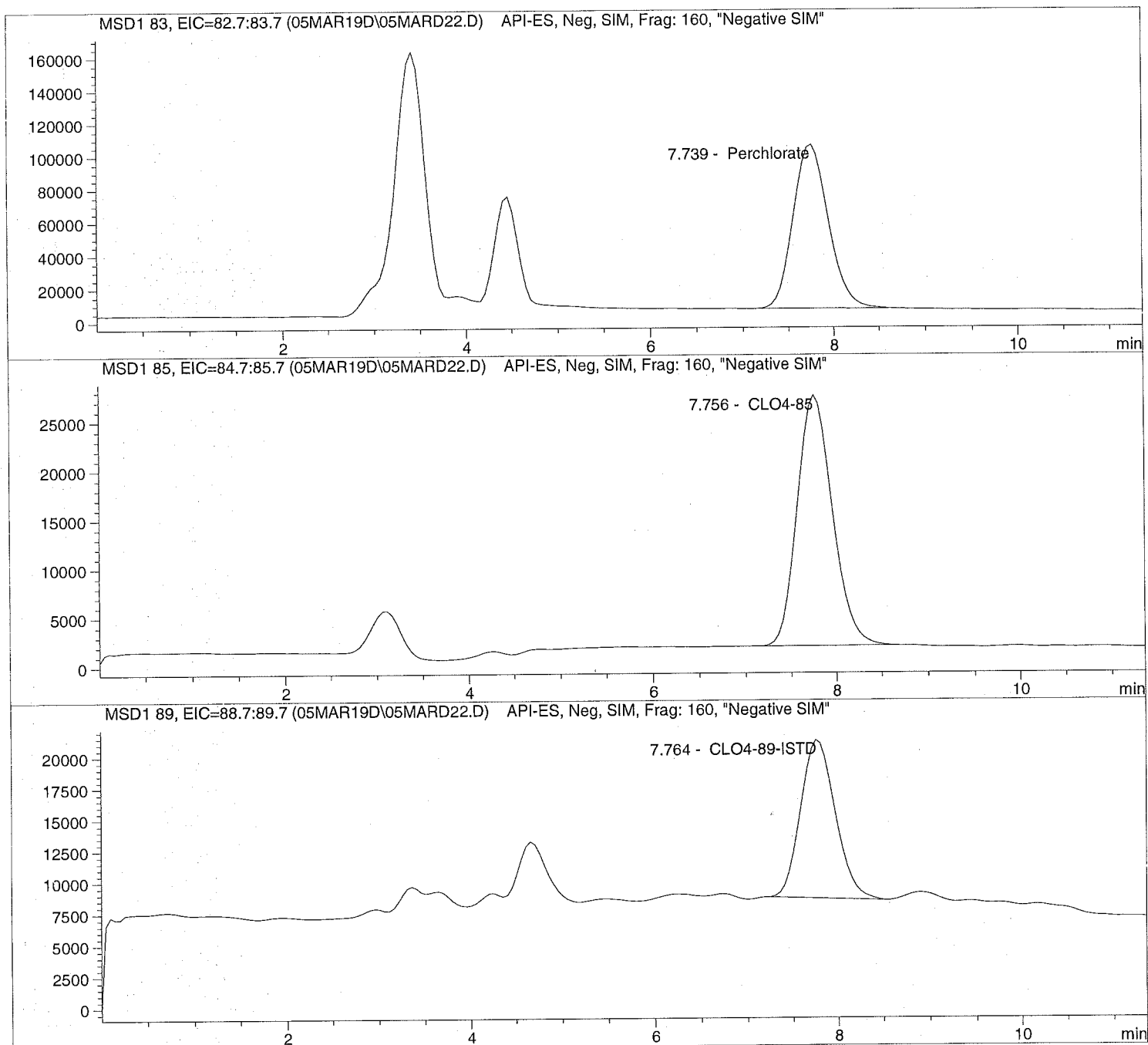
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD22.D Sample Name: 1906112005 10X

=====
Injection Date: 3/05/2019 13:28:32 Seq Line: 22
Sample Name: 1906112005 10X Location: Vial 91
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19\05MARD22.D Sample Name: 1906112005 10X

```

=====
Injection Date: 3/05/2019 13:28:32      Seq Line:      22
Sample Name:   1906112005 10X          Location:     Vial 91
Acq Operator:  TNB                    Inj. No.:    1
                                           Inj. Vol.:   20 µl
  
```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
  
```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:     10.000000
Sample Amount: 0.000
  
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.739	PBA	2682370.5	241.0371	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	BBA	695643.1	238.4467	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.764	PB	338724.1	50.0000	CLO4-89-ISTD

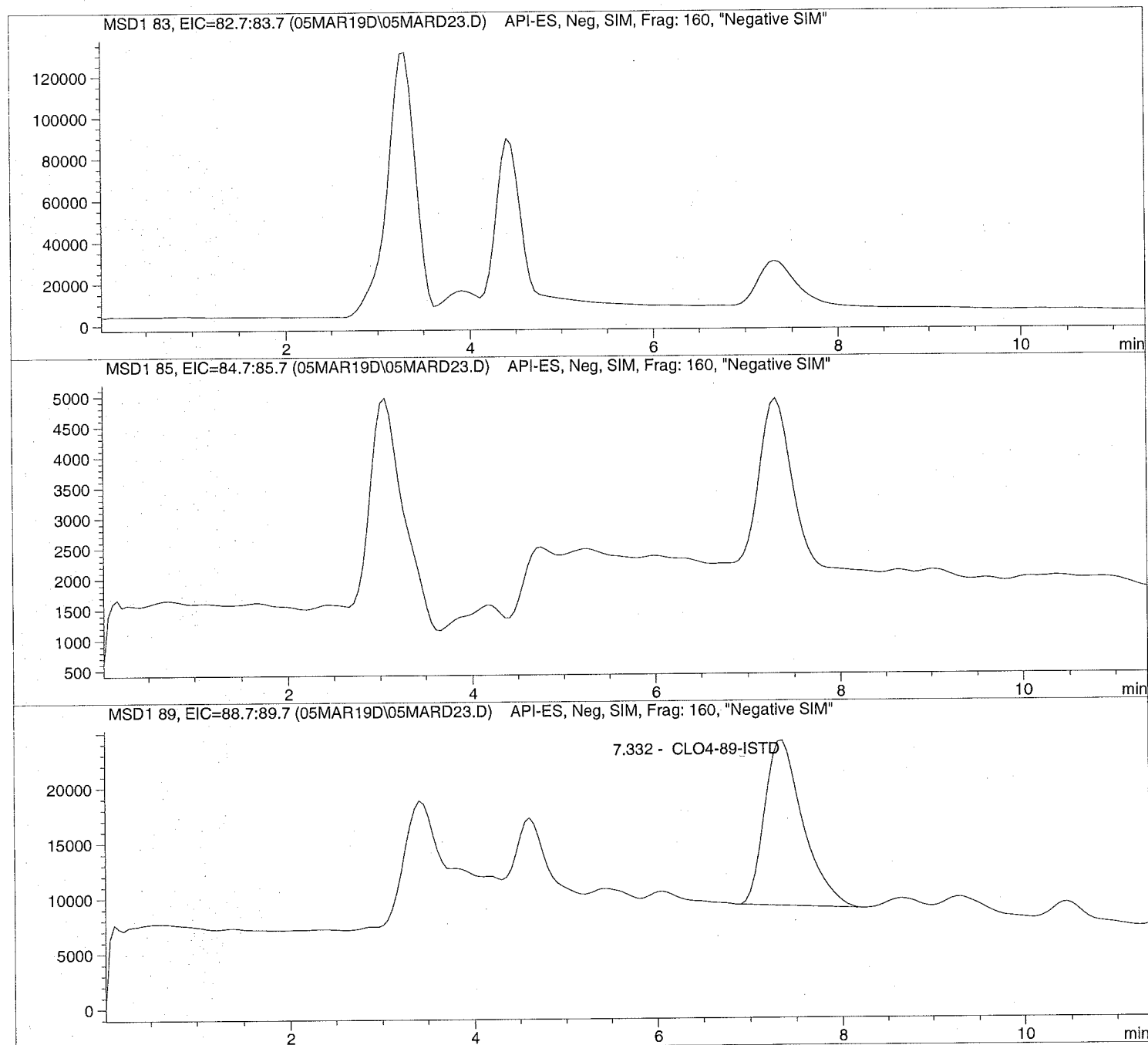
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD23.D Sample Name: 1906112007 RE

```
=====
Injection Date: 3/05/2019 13:41:34      Seq Line:      23
Sample Name:    1906112007 RE           Location:      Vial 82
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD23.D Sample Name: 1906112007 RE

```

=====
Injection Date: 3/05/2019 13:41:34      Seq Line:          23
Sample Name:    1906112007 RE           Location:          Vial 82
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.332	PB	431134.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

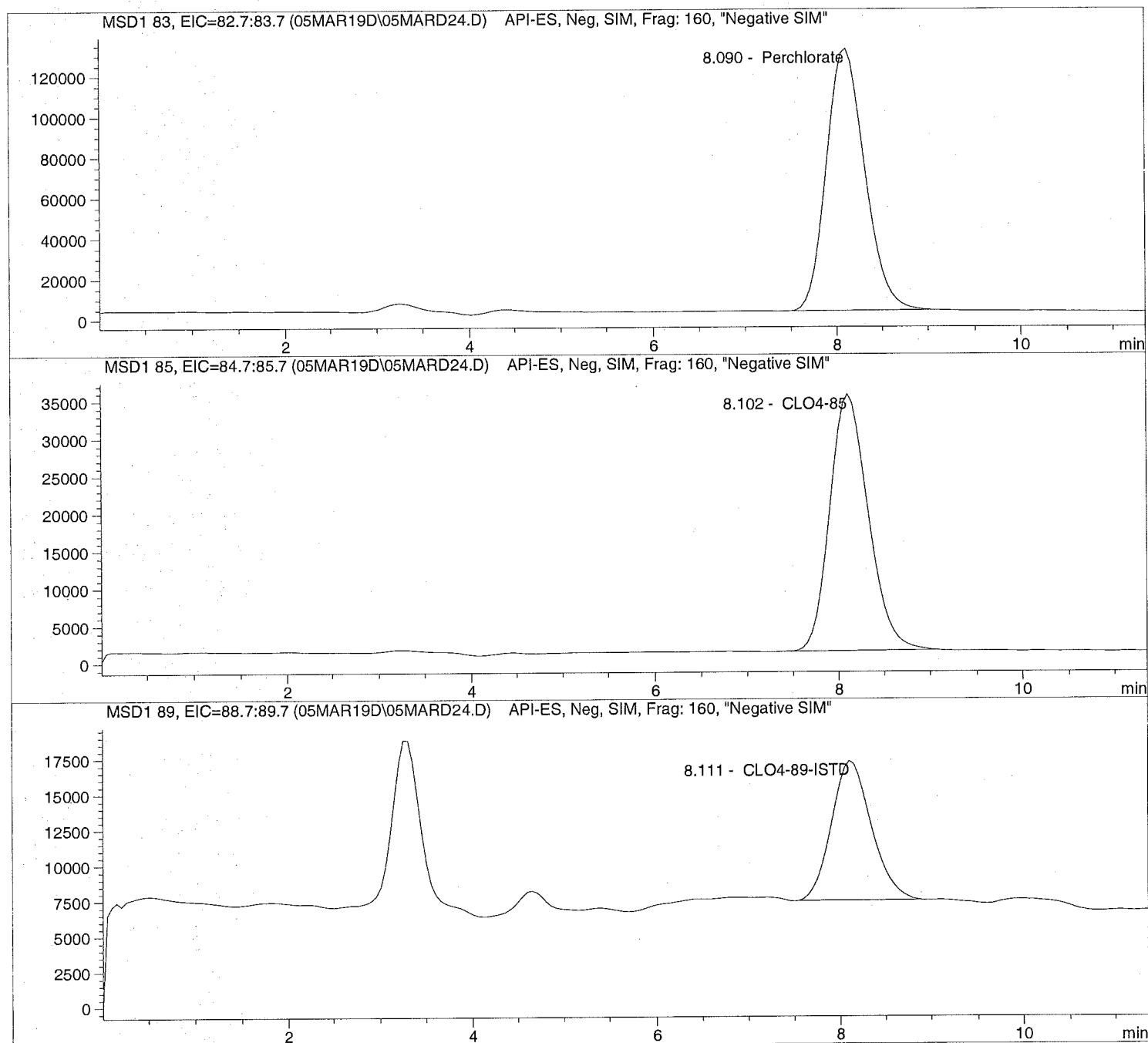
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD24.D Sample Name: 1906330001 100

```
=====
Injection Date: 3/05/2019 13:54:34 Seq Line: 24
Sample Name: 1906330001 100 Location: Vial 92
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD24.D Sample Name: 1906330001 100

```

=====
Injection Date: 3/05/2019 13:54:34      Seq Line:          24
Sample Name:   1906330001 100          Location:         Vial 92
Acq Operator:  TNB                    Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       100.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.090	PBA	3773628.7	3708.6567	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	PBA	1010205.2	3771.6748	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.111	PBA	298984.6	500.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

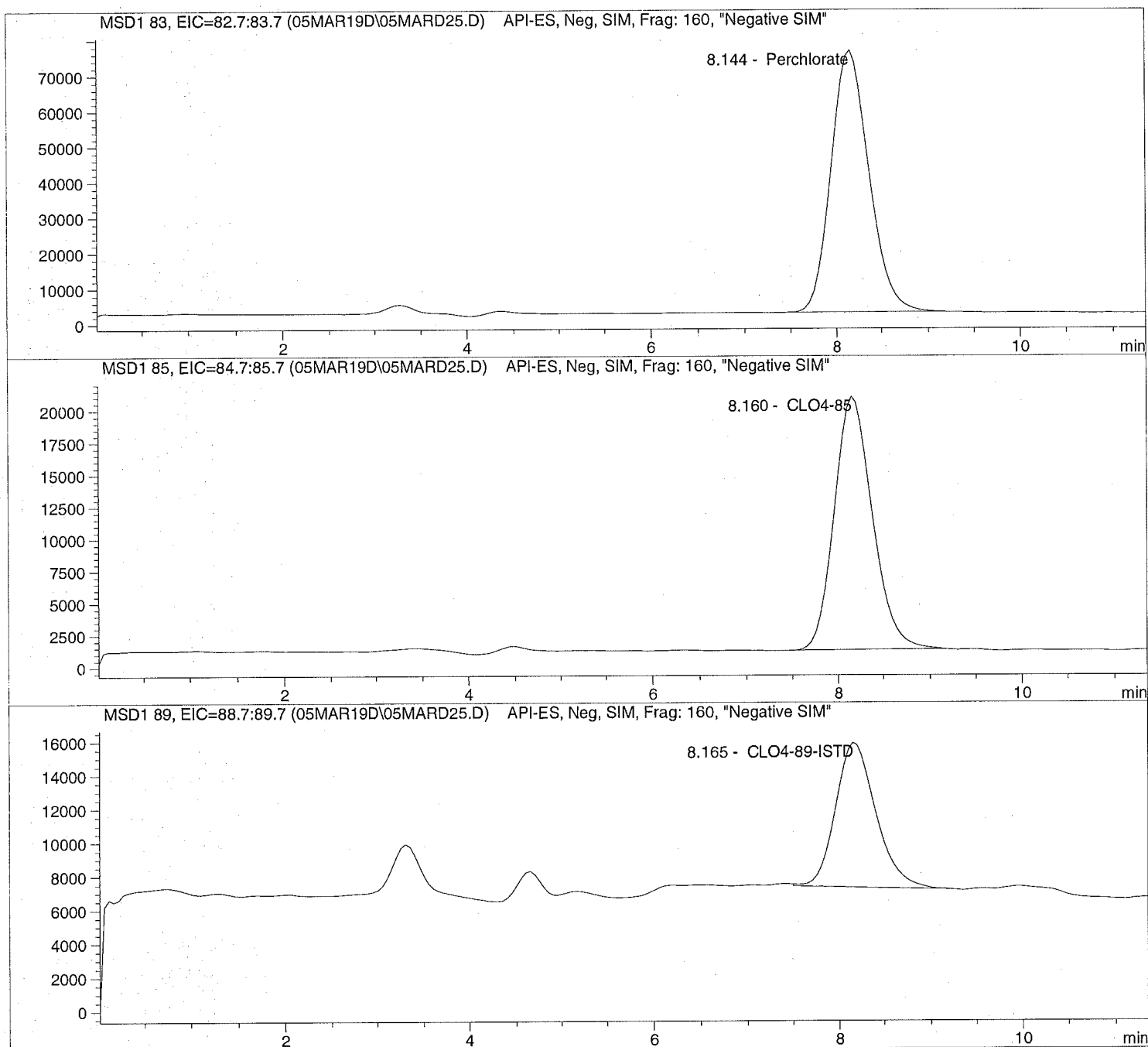
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD25.D Sample Name: 642102 CCV@25

Injection Date: 3/05/2019 14:07:36 Seq Line: 25
Sample Name: 642102 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD25.D Sample Name: 642102 CCV@25

=====
Injection Date: 3/05/2019 14:07:36 Seq Line: 25
Sample Name: 642102 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

=====
Sample Information
=====

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.144	PBA	2157871.5	24.4553	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.160	PBA	583711.5	25.1665	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.165	BBA	268305.4	5.0000	CLO4-89-ISTD

=====
*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

**Initial
Calibration**

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	8.94006e4	7.889	9.89924e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.97443e5	8.114	2.26028
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	4.79370e5	7.828	4.65688
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	9.30136e5	7.904	9.14998
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.81067e6	7.793	25.52636
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	5.66830e6	7.976	51.07439
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	8.69624e6	7.886	74.30603
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	1.01141e6	7.988	9.46019

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.26121e4	7.914	9.98836e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	5.53134e4	8.127	2.11360
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.39247e5	7.842	4.91261
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.54396e5	7.923	9.39034
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	7.35969e5	7.811	25.48268
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.47152e6	7.993	50.35774
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.32809e6	7.900	74.72233
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.81230e5	8.007	9.87858

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.41443e5	7.900	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.99651e5	8.132	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.38646e5	7.853	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	3.25154e5	7.925	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	3.33799e5	7.819	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	3.14712e5	7.999	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	3.13909e5	7.908	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	3.41503e5	8.005	5.00000

*** End of Report ***

Sequence: C:\HPCHEM\1\SEQUENCE\CLO4\2019\FEB\15FEB19I.S

Sequence Table:

Method and Injection Info Part:

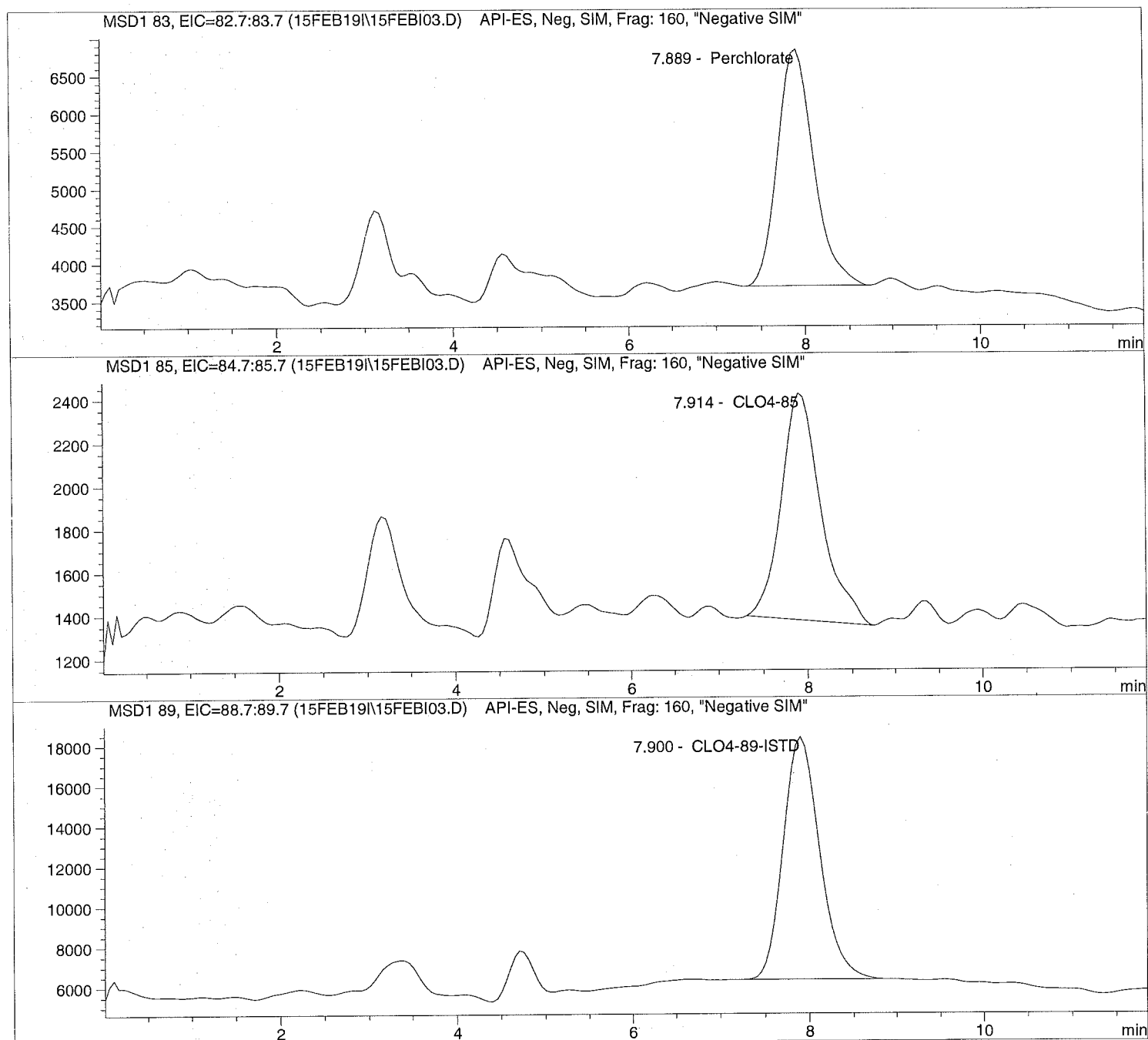
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
1	Vial 71	CLO4@ .20ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI03.D Sample Name: CLO4@ 1.0ug/L

```
=====
Injection Date: 2/15/2019 09:51:42      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L           Location:  Vial 73
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 2/15/2019 09:51:42      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L           Location:  Vial 73
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 1.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.889	PBA	89400.6	0.9899	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.914	BBA	32612.1	0.9988	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	BBA	341443.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

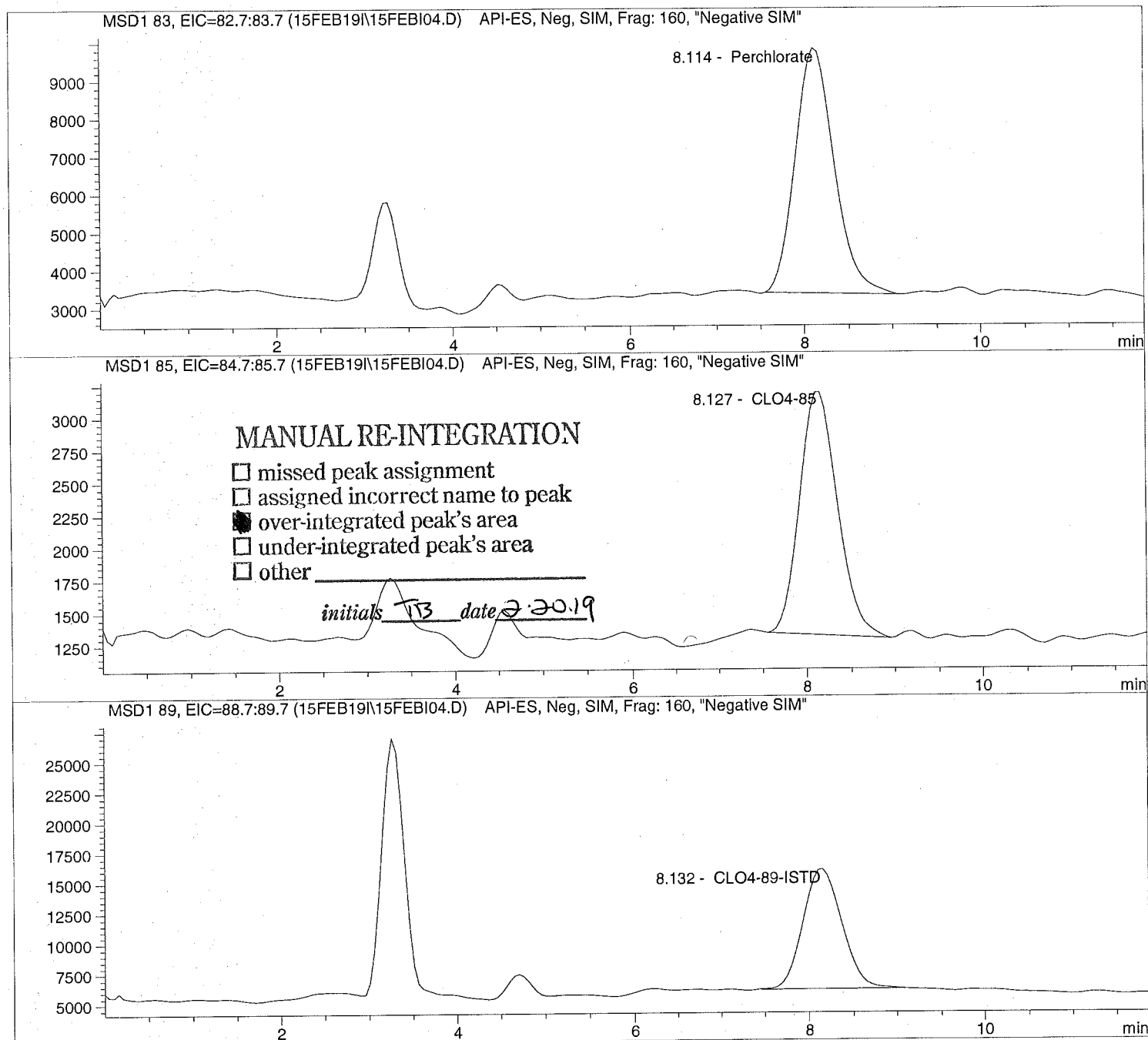
```


Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D Sample Name: CLO4@ 2.0ug/L

=====
Injection Date: 2/15/2019 10:05:24 Seq Line: 4
Sample Name: CLO4@ 2.0ug/L Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 2/15/2019 10:05:24      Seq Line:          4
Sample Name:    CLO4@ 2.0ug/L           Location:          Vial 74
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:       25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	MM	55313.4	2.1136	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

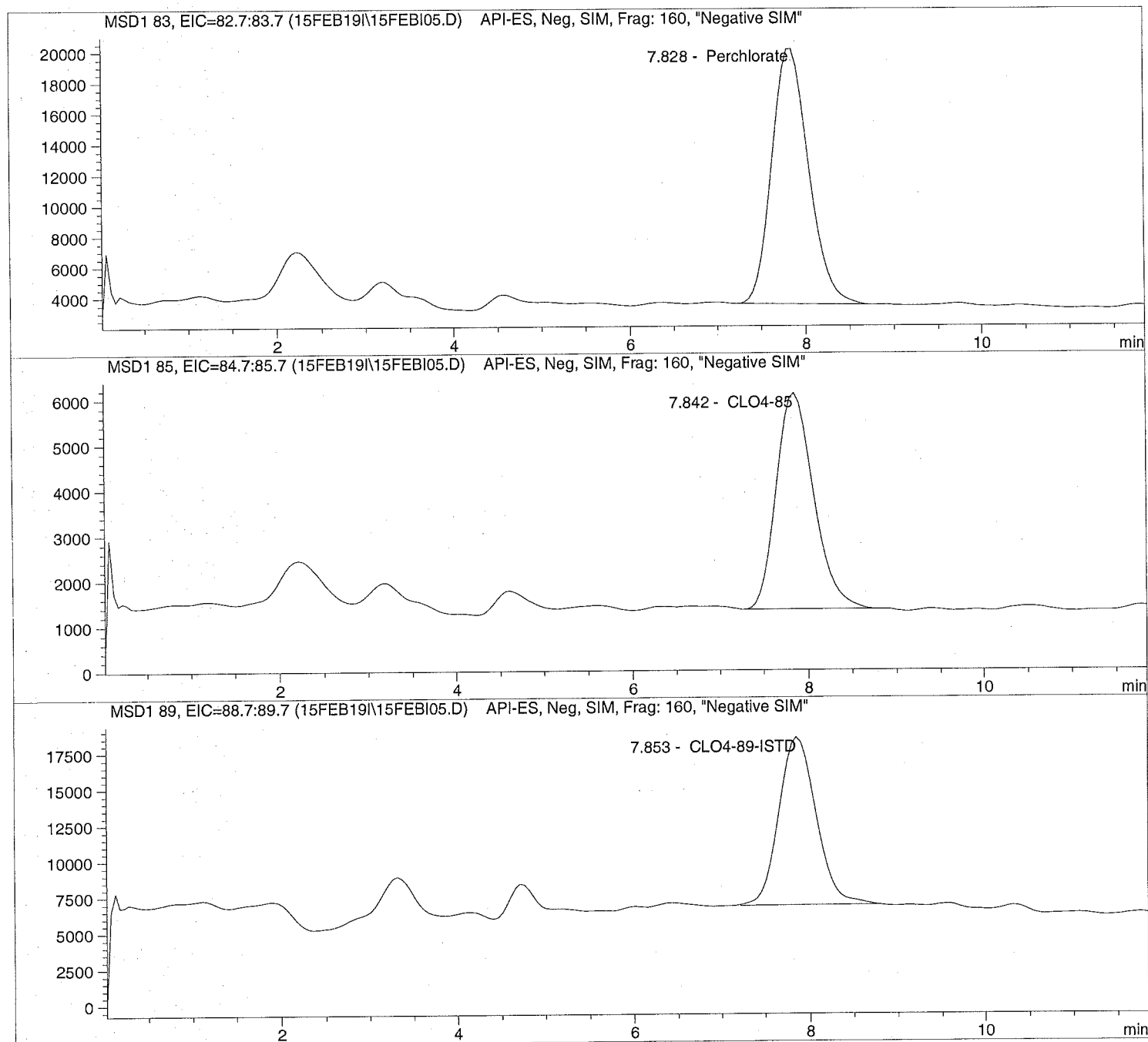
```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI05.D Sample Name: CLO4@ 5.0ug/L

=====
Injection Date: 2/15/2019 11:42:56 Seq Line: 5
Sample Name: CLO4@ 5.0ug/L Location: Vial 75
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 2/15/2019 11:42:56      Seq Line: 5
Sample Name:    CLO4@ 5.0ug/L           Location:  Vial 75
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.828	PBA	479370.4	4.6569	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.842	PBA	139246.9	4.9126	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.853	PBA	338646.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

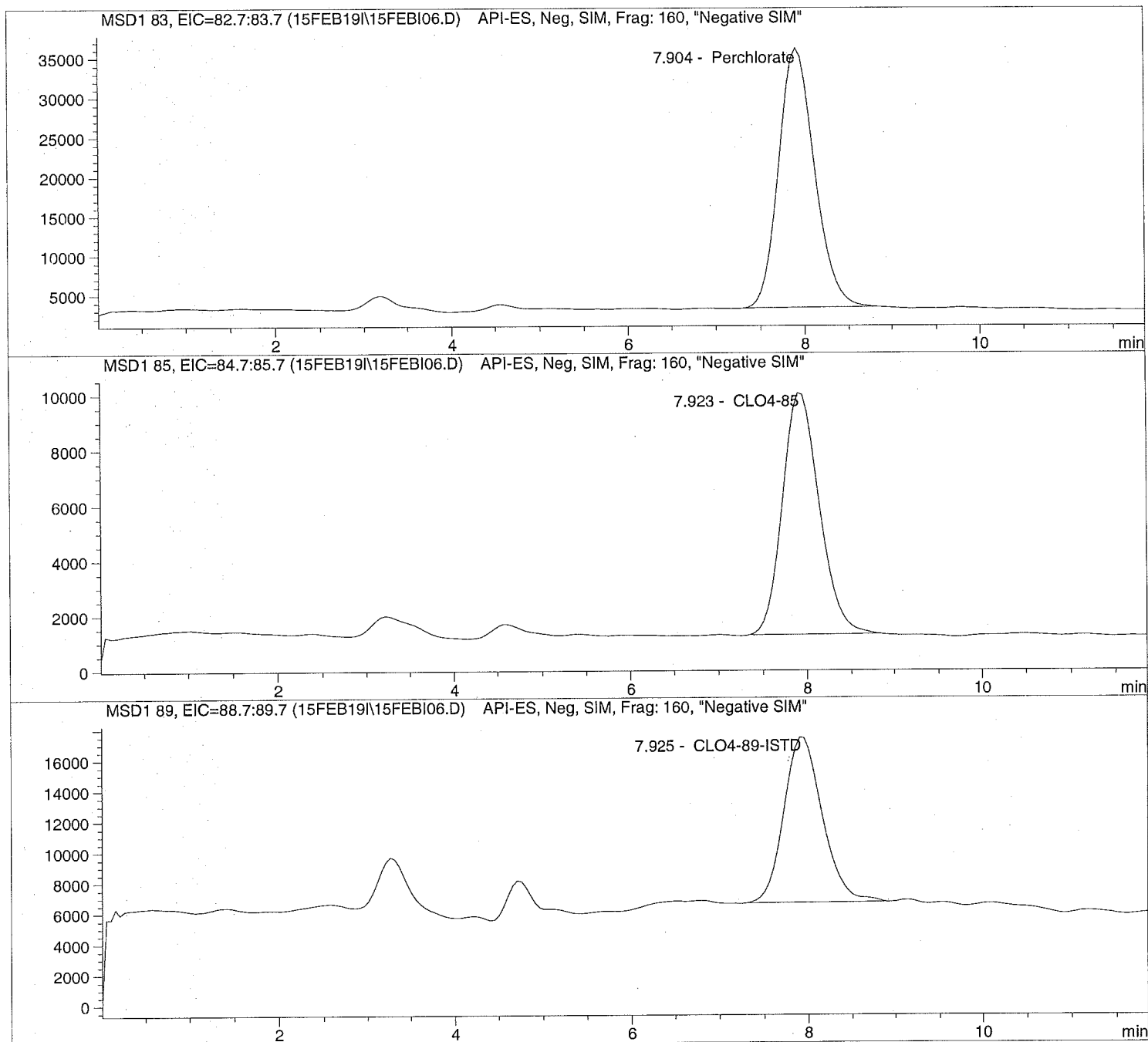
```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI06.D Sample Name: CLO4@ 10.ug/L

```
=====
Injection Date: 2/15/2019 11:56:38      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 2/15/2019 11:56:38      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.904	PBA	930135.8	9.1500	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.923	BBA	254395.6	9.3903	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.925	PBA	325154.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

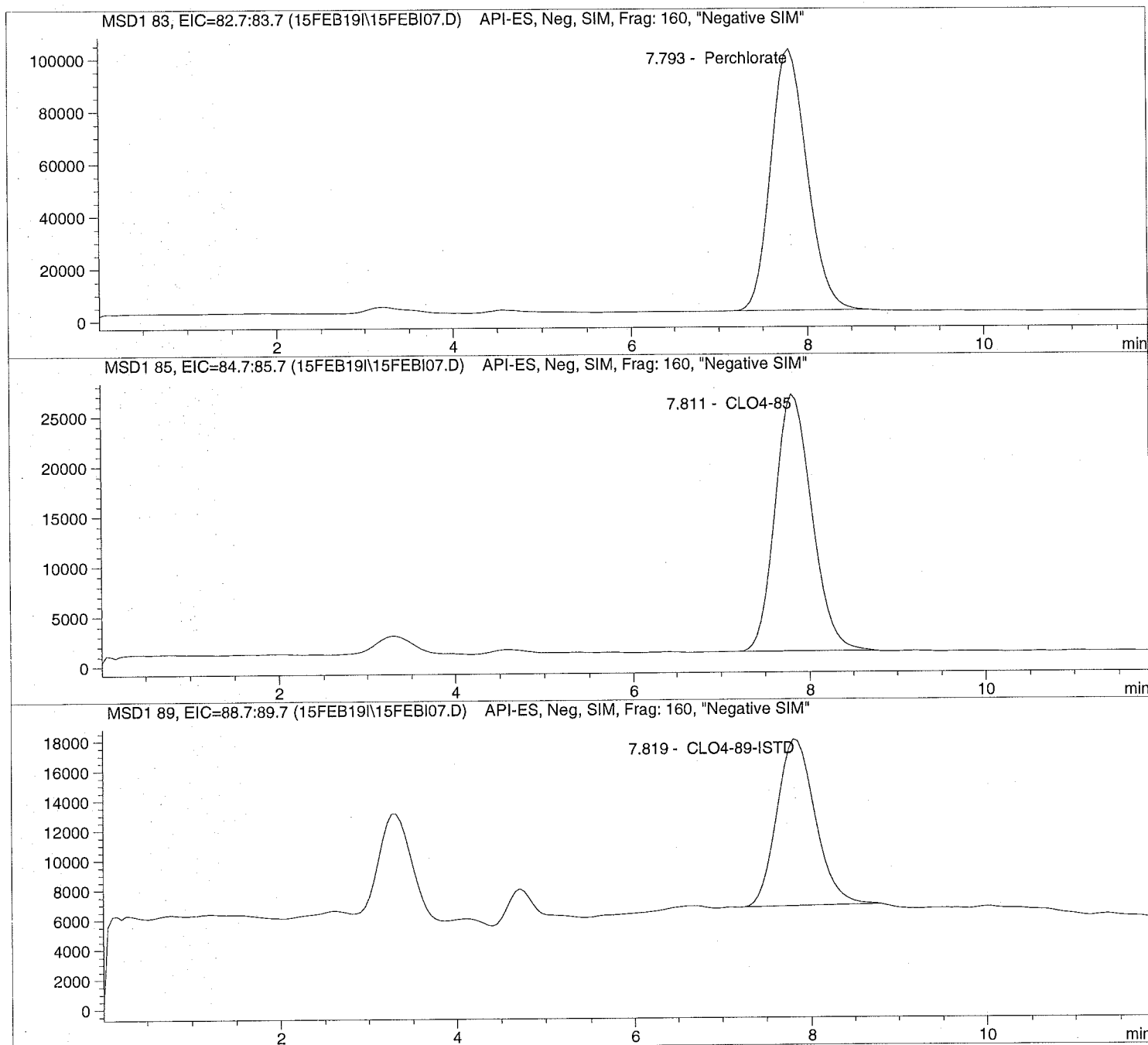
```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI07.D Sample Name: CLO4@ 25.ug/L

=====
Injection Date: 2/15/2019 12:10:22 Seq Line: 7
Sample Name: CLO4@ 25.ug/L Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 2/15/2019 12:10:22      Seq Line: 7
Sample Name:    CLO4@ 25.ug/L           Location:  Vial 77
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 25.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.793	PBA	2810669.2	25.5264	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.811	BBA	735968.9	25.4827	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.819	PBA	333799.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI08.D

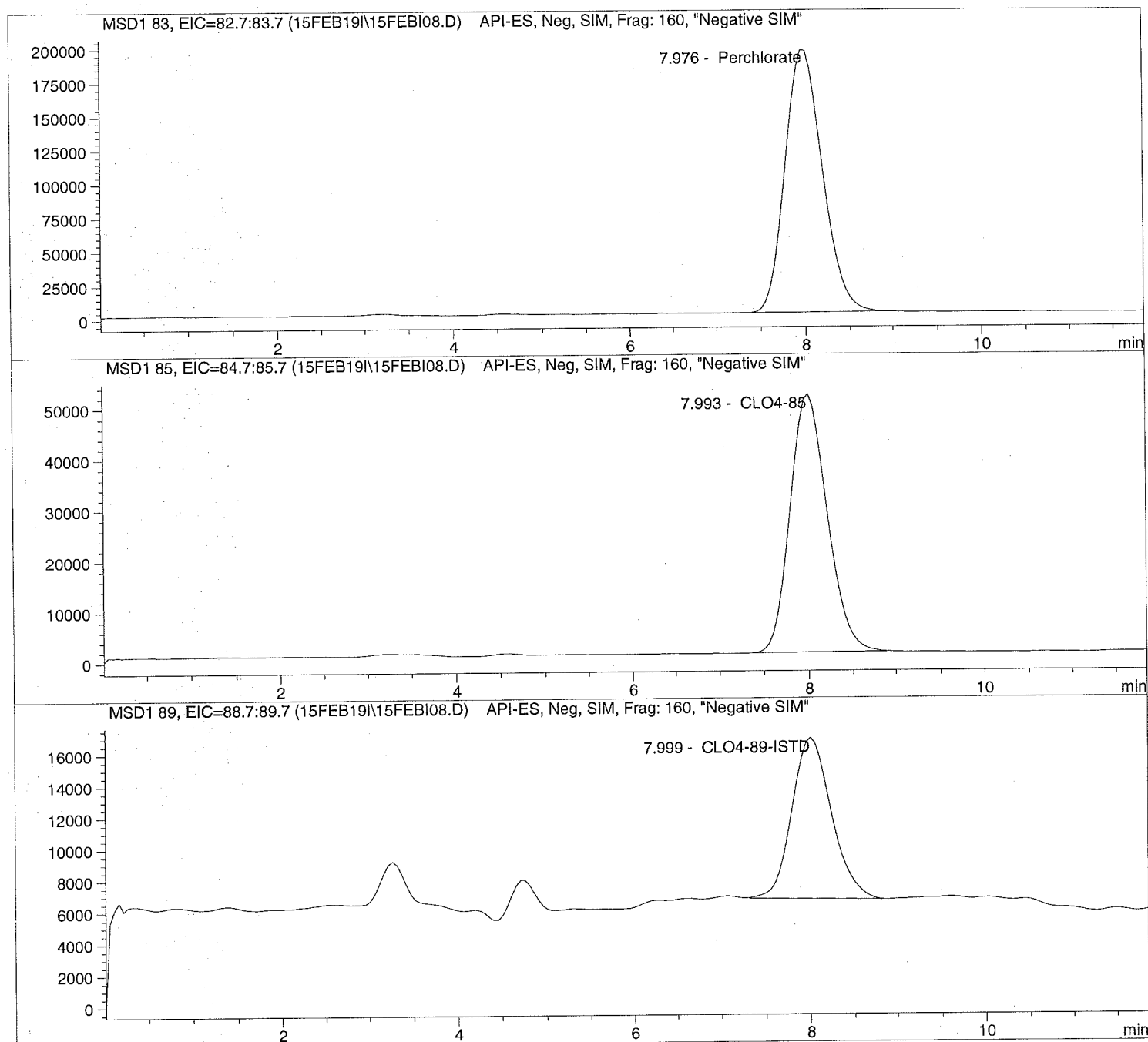
Sample Name: CLO4@ 50.ug/L

Injection Date: 2/15/2019 12:24:06
Sample Name: CLO4@ 50.ug/L
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 2/15/2019 12:24:06      Seq Line:      8
Sample Name:   CLO4@ 50.ug/L           Location:     Vial 78
Acq Operator:  TNB                     Inj. No.:    1
                                           Inj. Vol.:   25 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 50.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.976	PBA	5668301.5	51.0744	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.993	PBA	1471522.9	50.3577	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.999	BBA	314711.8	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 2/15/2019 12:37:48

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

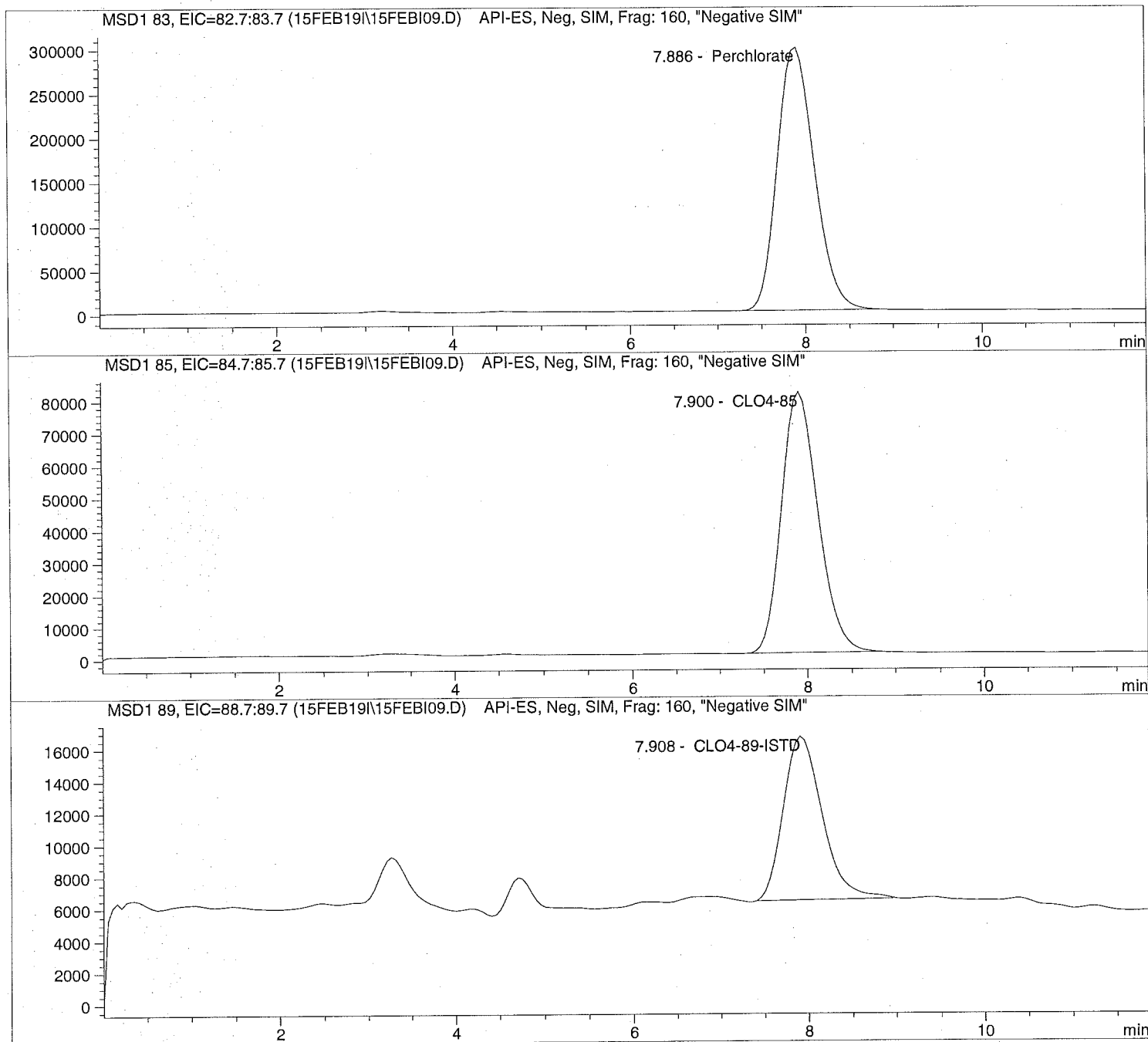
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M

Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date: 2/15/2019 12:37:48      Seq Line:          9
Sample Name:    CLO4@ 75.ug/L           Location:          Vial 79
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  75.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.886	PBA	8696239.0	74.3060	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	PBA	2328089.5	74.7223	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.908	PBA	313908.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

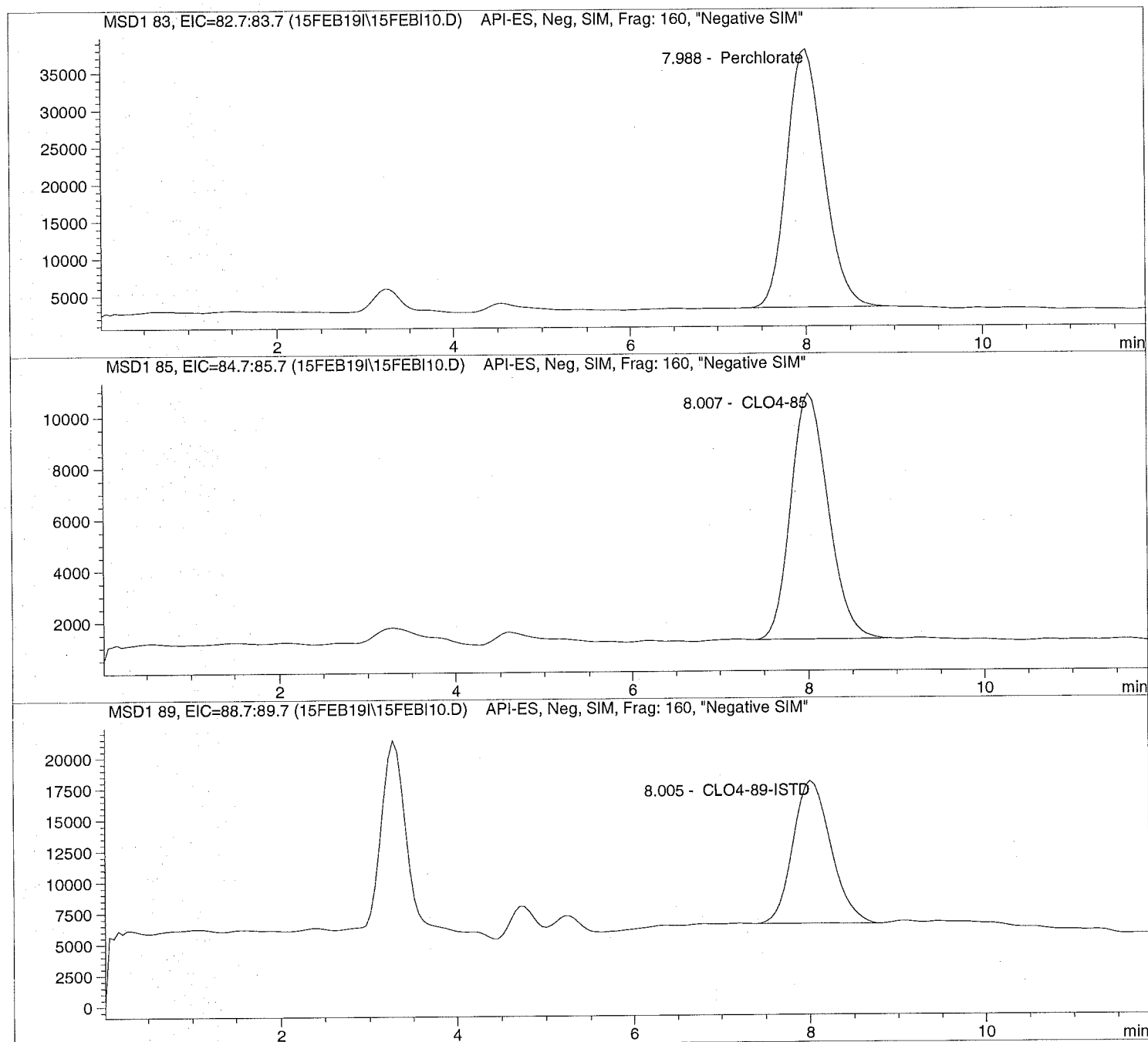
```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI10.D Sample Name: ICAL Verf@10ug/L

=====
Injection Date: 2/15/2019 12:51:29 Seq Line: 10
Sample Name: ICAL Verf@10ug/L Location: Vial 80
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 2/15/2019 12:51:29      Seq Line: 10
Sample Name:    ICAL Verf@10ug/L        Location:  Vial 80
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019, 09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.988	BBA	1011409.8	9.4602	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.007	BBA	281229.9	9.8786	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.005	BBA	341503.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

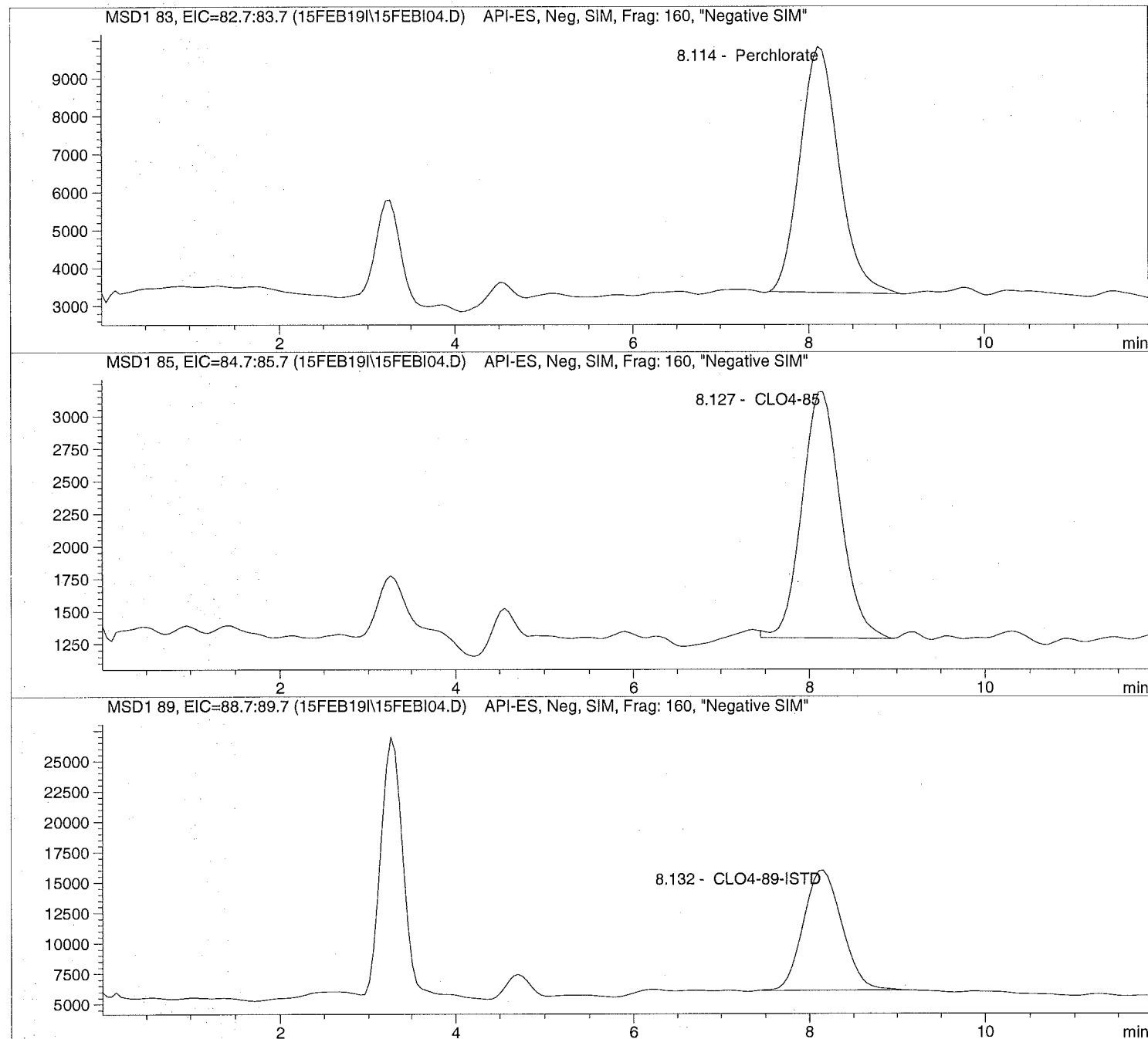
Unmodified

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D Sample Name: CLO4@ 2.0ug/L

=====
Injection Date: 2/15/2019 10:05:24 Seq Line: 4
Sample Name: CLO4@ 2.0ug/L Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:12:36

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D Sample Name: CLO4@ 2.0ug/L

=====
Injection Date: 2/15/2019 10:05:24 Seq Line: 4
Sample Name: CLO4@ 2.0ug/L Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:12:36

Perchlorate analysis

=====
Sample Information
=====

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 2.000

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	BBA	57206.1	2.1923	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

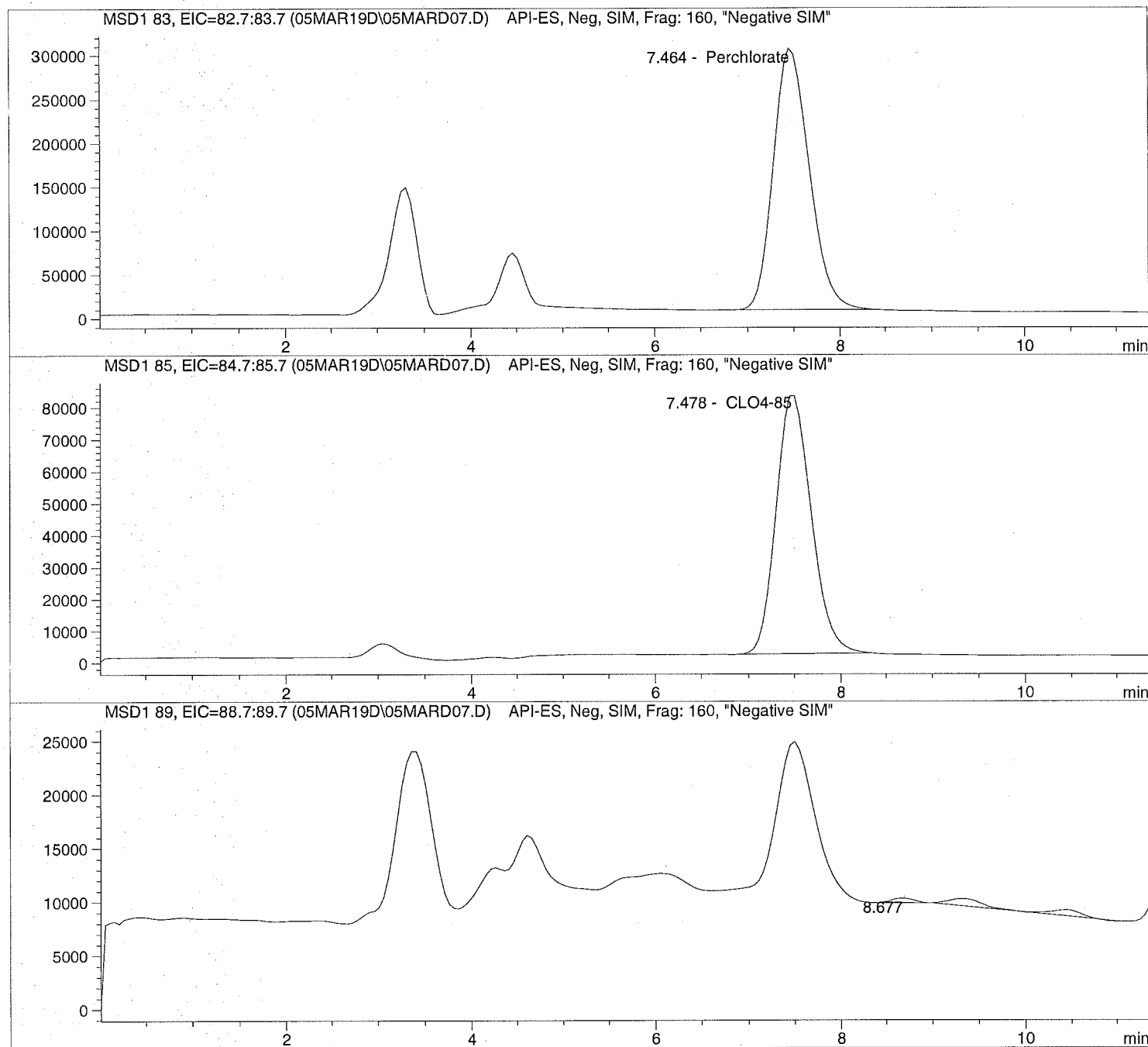
=====
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

=====
Injection Date: 3/05/2019 10:07:11 Seq Line: 7
Sample Name: 1906112002 MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

```

=====
Injection Date: 3/05/2019 10:07:11      Seq Line: 7
Sample Name: 1906112002 MS              Location: Vial 77
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	419.8794	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	402.6179	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.677	BB	7208.6	0.0000	
9.316	VBA	28561.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

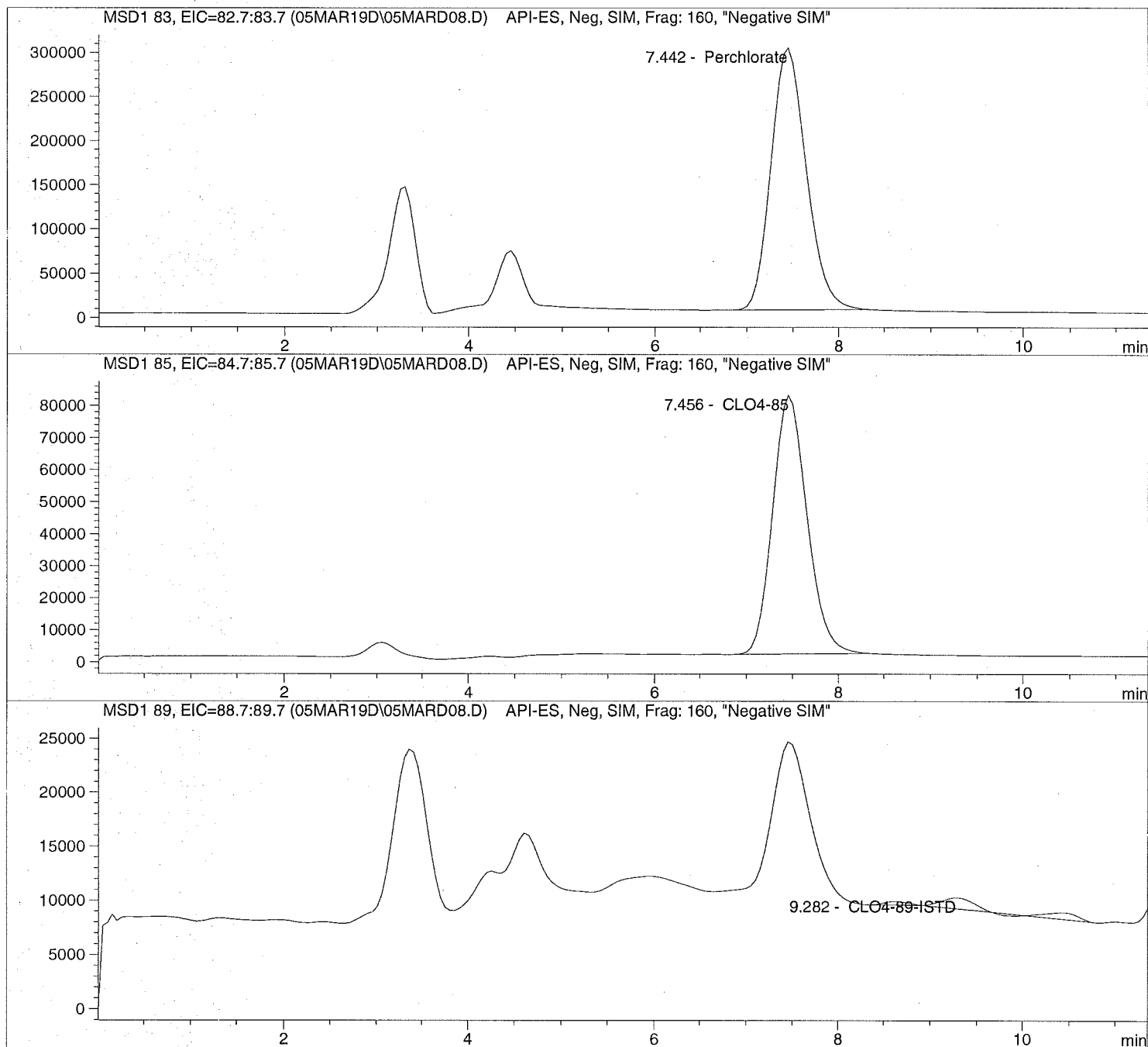
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

=====
Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

```

=====
Injection Date: 3/05/2019 10:20:17      Seq Line:      8
Sample Name:   1906112003  MSD          Location:      Vial 78
Acq Operator:  TNB                    Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	359.0996	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	344.2233	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.621	VB	5769.9	0.0000	
9.282	VBA	35831.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D

Sample Name: 1906112004

Injection Date: 3/05/2019 10:33:21

Seq Line: 9

Sample Name: 1906112004

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

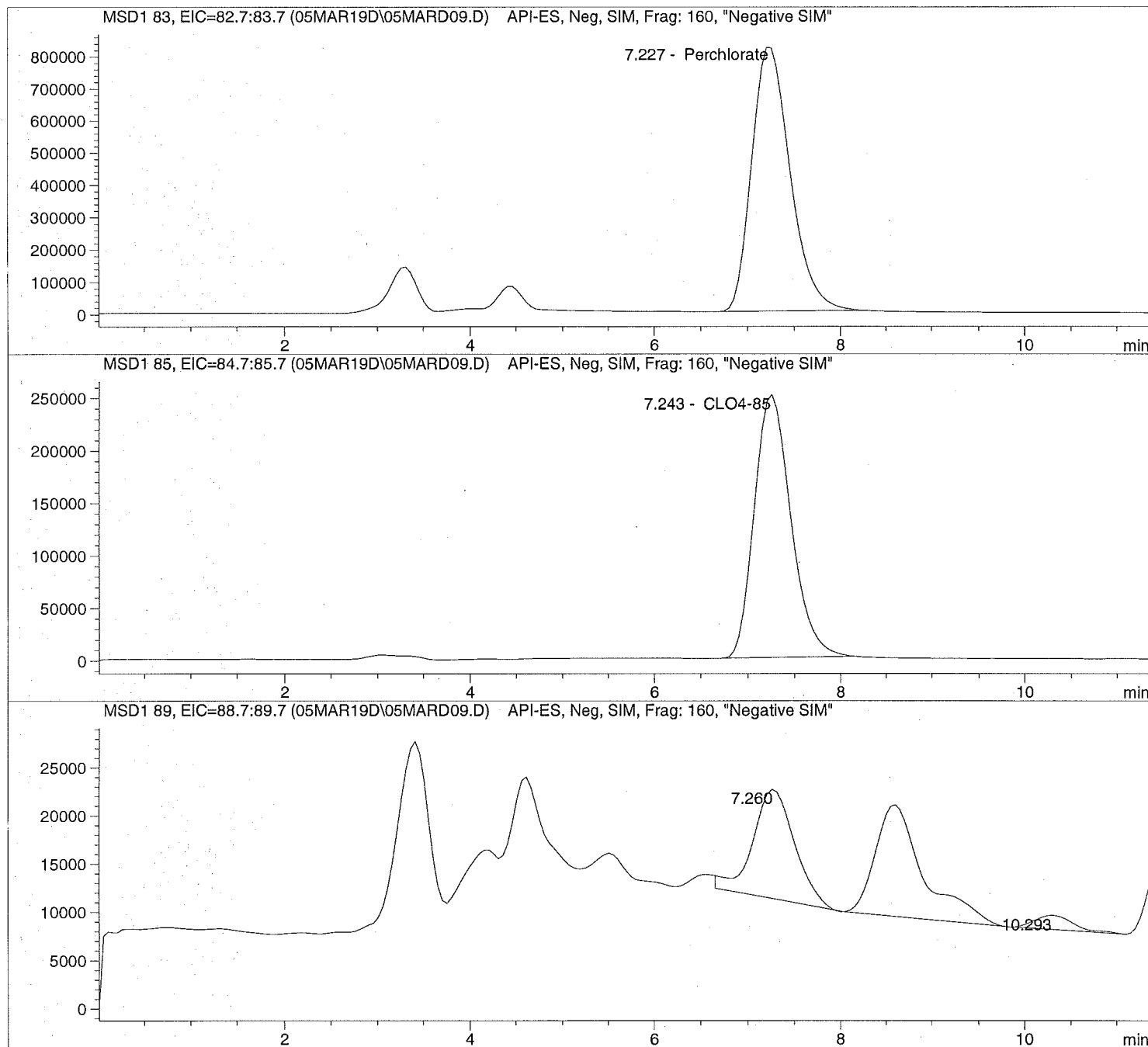
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M

Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D Sample Name: 1906112004

```

=====
Injection Date: 3/05/2019 10:33:21      Seq Line: 9
Sample Name: 1906112004                Location: Vial 79
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	131.1742	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	138.8050	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.260	BB	352149.7	0.0000	
8.589	VBA	421141.9	5.0000	CLO4-89-ISTD
10.293	BBA	41603.7	0.0000	

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D

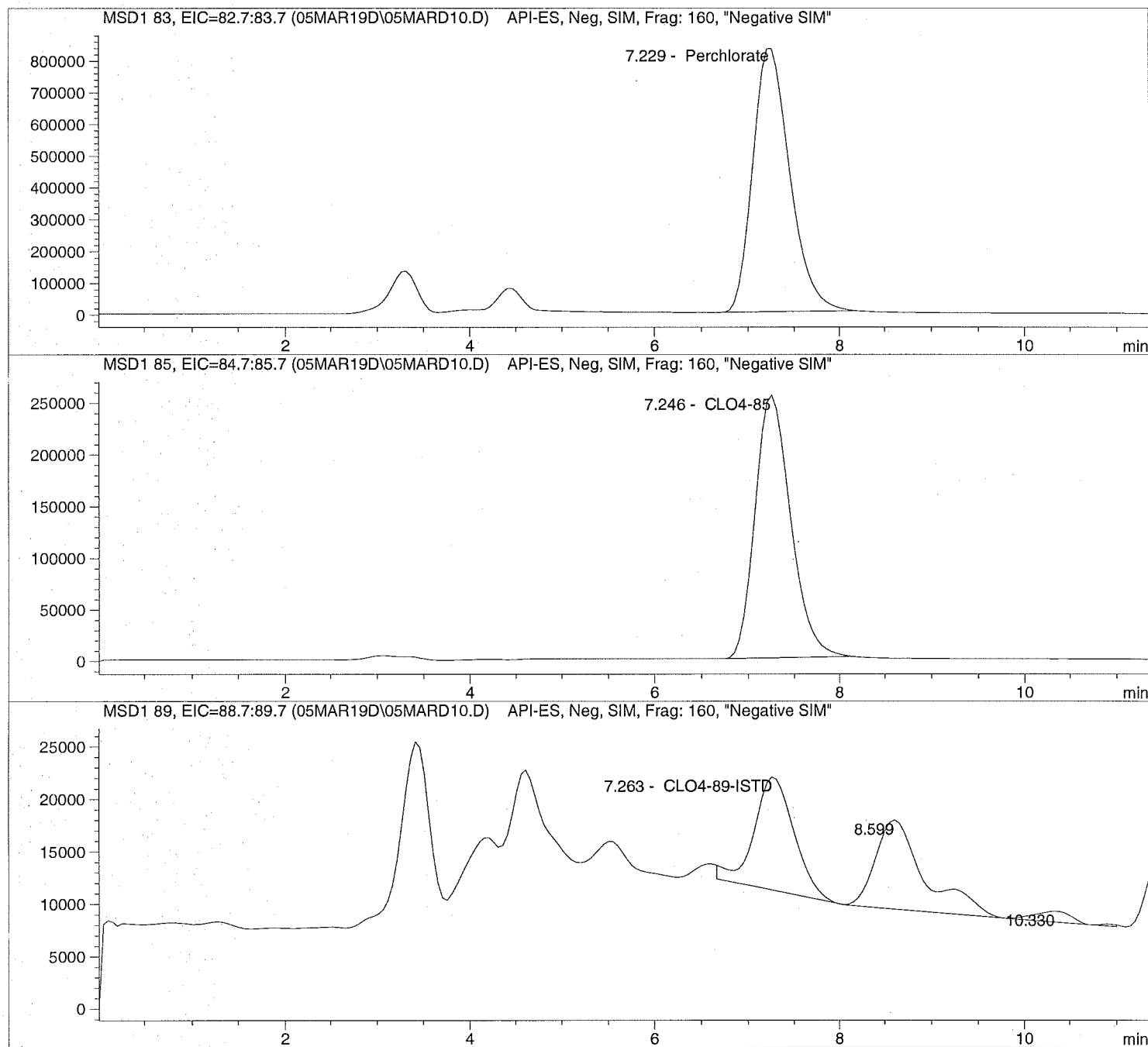
Sample Name: 1906112005

Injection Date: 3/05/2019 10:46:26
Sample Name: 1906112005
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D Sample Name: 1906112005

```

=====
Injection Date: 3/05/2019 10:46:26      Seq Line:          10
Sample Name:    1906112005              Location:          Vial 80
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	159.0418	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	168.6882	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	BB	325841.0	5.0000	CLO4-89-ISTD
8.599	VB	308921.9	0.0000	
10.330	VBA	30210.3	0.0000	

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D

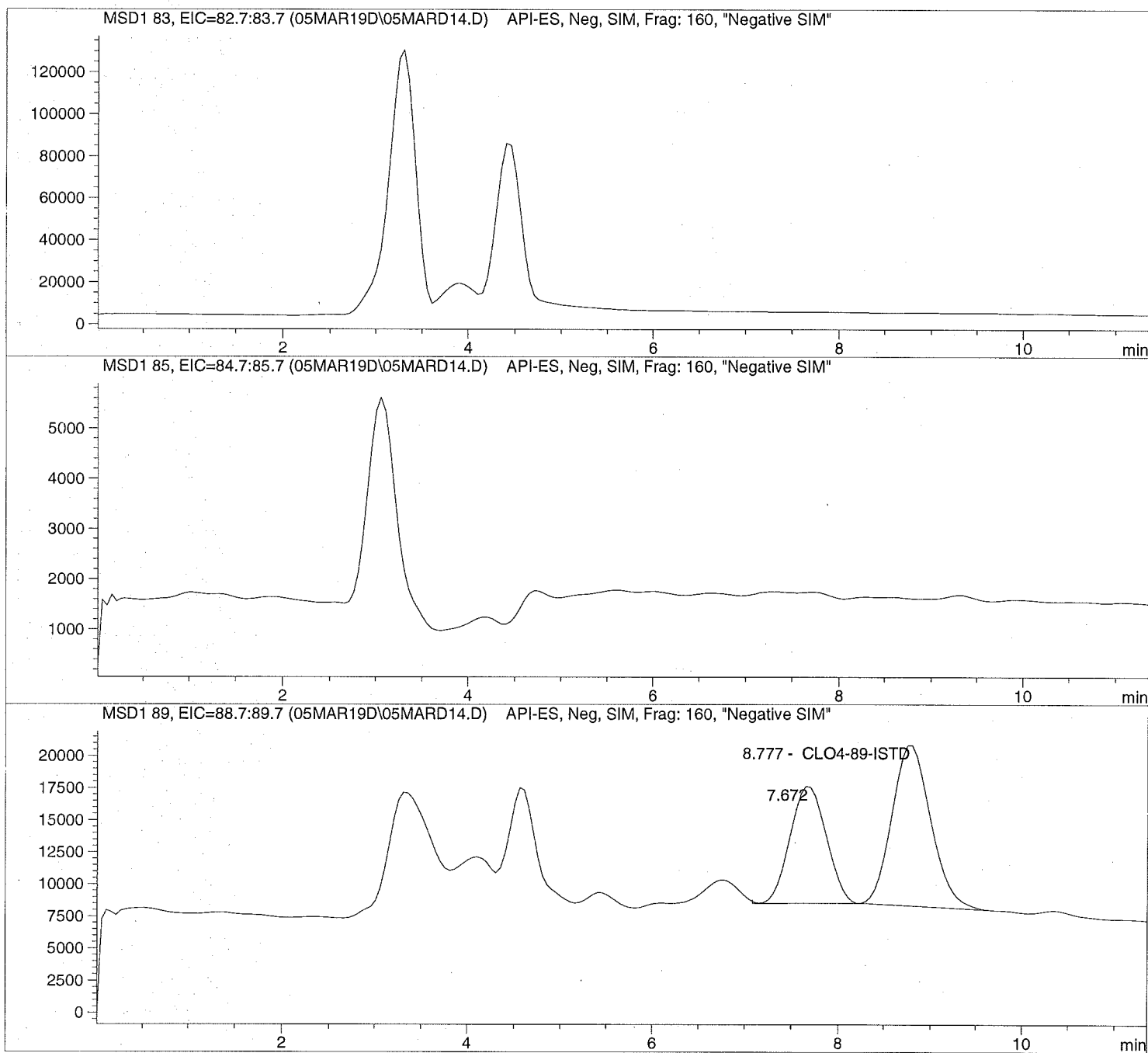
Sample Name: 1906112009

Injection Date: 3/05/2019 11:39:24
Sample Name: 1906112009
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D

Sample Name: 1906112009

```

=====
Injection Date: 3/05/2019 11:39:24      Seq Line:      14
Sample Name:   1906112009              Location:      Vial 84
Acq Operator:  TNB                     Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	0.0000	
8.777	VBA	362717.3	5.0000	CLO4-89-ISTD

```

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*** End of Report ***
=====

```



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 28, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030298**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 07, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030298

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030298-01	LH18/24-SP650_030619	Water		06-Mar-2019 13:00	07-Mar-2019 08:51	<input type="checkbox"/>
HS19030298-02	LH18/24-SP650_030619_BIX	Water		06-Mar-2019 13:00	07-Mar-2019 08:51	<input type="checkbox"/>

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
-

Work Order Comments

- The analysis for TOC was subcontracted to ALS Environmental in Kelso WA. Final report attached.
-

WetChemistry by Method E350.3**Batch ID: R334781**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

WetChemistry by Method E365.3**Batch ID: R334240**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_030619
 Collection Date: 06-Mar-2019 13:00

ANALYTICAL REPORT

WorkOrder:HS19030298
 Lab ID:HS19030298-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
AMMONIA AS N BY E350.3(ISE)								Analyst: KVL
Nitrogen, Ammonia (As N)	5.5		0.20	0.20	0.20	mg/L	1	18-Mar-2019 12:30
ORTHO PHOSPHATE (PO4) AS P BY E365.3								Analyst: MZD
Phosphorus, Total Orthophosphate (As P)	0.480		0.100	0.250	0.250	mg/L	10	08-Mar-2019 11:50
SUBCONTRACT ANALYSIS - TOC ANALYSIS								Analyst: SUBK
Subcontract Analysis	See Attached		0	0		NA	1	28-Mar-2019 08:57

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_030619_BIX
 Collection Date: 06-Mar-2019 13:00

ANALYTICAL REPORT

WorkOrder:HS19030298
 Lab ID:HS19030298-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA		Analyst: SUB				
Subcontract Analysis	See Attached		0	0		NA	1	21-Mar-2019 17:49

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030298

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R334240	Test Name : ORTHO PHOSPHATE (PO4) AS P BY E365.3		Matrix: Water			
HS19030298-01	LH18/24-SP650_030619	06 Mar 2019 13:00			08 Mar 2019 11:50	10
Batch ID R334781	Test Name : AMMONIA AS N BY E350.3(ISE)		Matrix: Water			
HS19030298-01	LH18/24-SP650_030619	06 Mar 2019 13:00			18 Mar 2019 12:30	1
Batch ID R335013	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19030298-02	LH18/24-SP650_030619_BIX	06 Mar 2019 13:00			21 Mar 2019 17:49	1
Batch ID R335428	Test Name : SUBCONTRACT ANALYSIS - TOC ANALYSIS		Matrix: Water			
HS19030298-01	LH18/24-SP650_030619	06 Mar 2019 13:00			28 Mar 2019 08:57	1

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030298

QC BATCH REPORT NEW

Batch ID:	R334240 (0)	Instrument:	UV-2450	Method:	ORTHO PHOSPHATE (PO4) AS P BY E365.3					
MBLK	Sample ID: MBLK-334240	Units: mg/L		Analysis Date: 08-Mar-2019 11:50						
Client ID:		Run ID: UV-2450_334240		SeqNo: 4980068	PrepDate:	DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.0250	0.0250							U	
LCS	Sample ID: LCS-334240	Units: mg/L		Analysis Date: 08-Mar-2019 11:50						
Client ID:		Run ID: UV-2450_334240		SeqNo: 4980069	PrepDate:	DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	0.221	0.0250	0.25	0	88.4	85 - 115				
MS	Sample ID: HS19030298-01MS	Units: mg/L		Analysis Date: 08-Mar-2019 11:50						
Client ID: LH18/24-SP650_030619		Run ID: UV-2450_334240		SeqNo: 4980071	PrepDate:	DF: 10				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	2.83	0.250	2.5	0.48	94.0	80 - 120				
MSD	Sample ID: HS19030298-01MSD	Units: mg/L		Analysis Date: 08-Mar-2019 11:50						
Client ID: LH18/24-SP650_030619		Run ID: UV-2450_334240		SeqNo: 4980072	PrepDate:	DF: 10				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Phosphorus, Total Orthophosphate (As P)	2.92	0.250	2.5	0.48	97.6	80 - 120	2.83	3.13	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030298

QC BATCH REPORT NEW

Batch ID: R334781 (0)		Instrument: WetChem_HS		Method: AMMONIA AS N BY E350.3(ISE)						
MBLK	Sample ID: MBLK-R334781	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 4995674			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	0.20	0.20							U	
LCS	Sample ID: LCS-R334781	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 5007379			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	9.398	0.20	10	0	94.0	80 - 120				
MS	Sample ID: HS19030689-05MS	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 4995676			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	10.4	0.20	10	0.1809	102	80 - 120				
MSD	Sample ID: HS19030689-05MSD	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 4995675			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	10.38	0.20	10	0.1809	102	80 - 120	10.4	0.192	20	

The following samples were analyzed in this batch: HS19030298-01

ALS Houston, US

Date: 28-Mar-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19030298	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

ALS Houston, US

Date: 28-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030298

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19030298-01	LH18/24-SP650_030619	Login	3/7/2019 11:27:32 AM	NDR	WET176
HS19030298-01	LH18/24-SP650_030619	Login	3/7/2019 11:27:32 AM	NDR	WET176
HS19030298-01	LH18/24-SP650_030619	Login	3/7/2019 11:27:32 AM	NDR	Sub
HS19030298-02	LH18/24-SP650_030619_BIX	Login	3/7/2019 11:27:32 AM	NDR	Sub

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030298

Date/Time Received: **07-Mar-2019 08:51**
 Received by: **NDR**

Checklist completed by: Nilesh D. Ranchod 7-Mar-2019
 eSignature | Date

Reviewed by: RJ Modashia 7-Mar-2019
 eSignature | Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 0.2C/0.2C UC/C | IR11
 Cooler(s)/Kit(s): 43601
 Date/Time sample(s) sent to storage: 03/07/2019 13:00

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:


Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:


Corrective Action:

CHAIN OF CUSTODY


Name Of Lab Shipping To: ALS 10450 Stancliff Rd. Suite 210 Houston, TX. 77099 (281) 530-5656 ATTN: R.J Modshia

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001		Analyses										HS19030298 Bhate Environmental Associates, Inc. 1/18/24 Longhorn GW Treatment Plant Weekly Sample								
Job: GROUNDWATER TREATMENT PLANT WEEKLY SAMPLES					MS / MSD	No. OF CONTAINERS	AMMONIA-N	TOTAL ORGANIC CARBON	ORTHO-PHOSPHATE	PERCHLORATE											Remarks (Preservatives, etc.)	Lab I.D.#	
Prepared By: Scott Beesinger			P.O. Number																				
Field Sample I.D.	Sample Matrix	Date / Time																					
LH18/24-SP650_030619	Water	03/06/19 / 13:00		2	X	X															H2SO4		
LH18/24-SP650_030619	Water	03/06/19 / 13:00		1				X														NONE	
LH18/24-SP650_030619_BIX	Water	03/06/19 / 13:00		1						X												NONE	
Additional Remarks: Standard TAT on all parameters																							
Relinquished By: <i>Scott Beesinger</i>		Date 03/06/19	Time 13:30	Received By: N/A		Date 3/6/19	Time 08:51	Relinquished By:		Date	Time	Received By:		Date	Time								
Received At Lab By:		Date	Time	Airbill No.	For Lab Use Only Opened By:		Date	Time	Temp of Container	Seal No.	Condition												
Remarks:				43601		1R x 11		Temp at 0.2															

(Word) S:\1-ces\Forms\Chain of Custody - BiWeekly

 ALS 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CUSTOMER	
	Date: 3/6/19	
	Name: Scott	
	Company: SHER	

ODY SEAL	
Time: 13:30	Seal Broken By:
2665-NGIC	Date:

 TRK# 0221 4380 9530 9478	THU - 07 MAR 10:30A PRIORITY OVERNIGHT
AB SGRA	77099 TX-US IAH
	
FTD 5121968 @0MAR19 60GA 563C1/46D3/8C8A	



ALS Environmental
ALS Group USA, Corp
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Kelso, WA 98626
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www.alsglobal.com

March 27, 2019

Analytical Report for Service Request No: K1902047

RJ Modashia
ALS Laboratory Group
10450 Stancliff Road
Suite 210
Houston, TX 77099-4338

RE: HS19030298

Dear RJ,

Enclosed are the results of the sample(s) submitted to our laboratory March 08, 2019
For your reference, these analyses have been assigned our service request number **K1902047**.

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. The test results meet requirements of the current NELAP standards, where applicable, and except as noted in the laboratory case narrative provided. For a specific list of NELAP-accredited analytes, refer to the certifications section at www.alsglobal.com. All results are intended to be considered in their entirety, and ALS Group USA Corp. dba ALS Environmental (ALS) is not responsible for use of less than the complete report. Results apply only to the items submitted to the laboratory for analysis and individual items (samples) analyzed, as listed in the report.

Please contact me if you have any questions. My extension is 3350. You may also contact me via email at Kelley.Lovejoy@alsglobal.com.

Respectfully submitted,

ALS Group USA, Corp. dba ALS Environmental

Kelley Lovejoy
Project Manager



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Table of Contents

Acronyms

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 General Chemistry

Acronyms

ASTM	American Society for Testing and Materials
A2LA	American Association for Laboratory Accreditation
CARB	California Air Resources Board
CAS Number	Chemical Abstract Service registry Number
CFC	Chlorofluorocarbon
CFU	Colony-Forming Unit
DEC	Department of Environmental Conservation
DEQ	Department of Environmental Quality
DHS	Department of Health Services
DOE	Department of Ecology
DOH	Department of Health
EPA	U. S. Environmental Protection Agency
ELAP	Environmental Laboratory Accreditation Program
GC	Gas Chromatography
GC/MS	Gas Chromatography/Mass Spectrometry
LOD	Limit of Detection
LOQ	Limit of Quantitation
LUFT	Leaking Underground Fuel Tank
M	Modified
MCL	Maximum Contaminant Level is the highest permissible concentration of a substance allowed in drinking water as established by the USEPA.
MDL	Method Detection Limit
MPN	Most Probable Number
MRL	Method Reporting Limit
NA	Not Applicable
NC	Not Calculated
NCASI	National Council of the Paper Industry for Air and Stream Improvement
ND	Not Detected
NIOSH	National Institute for Occupational Safety and Health
PQL	Practical Quantitation Limit
RCRA	Resource Conservation and Recovery Act
SIM	Selected Ion Monitoring
TPH	Total Petroleum Hydrocarbons
tr	Trace level is the concentration of an analyte that is less than the PQL but greater than or equal to the MDL.

Inorganic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- E The result is an estimate amount because the value exceeded the instrument calibration range.
- J The result is an estimated value.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.
- H The holding time for this test is immediately following sample collection. The samples were analyzed as soon as possible after receipt by the laboratory.

Metals Data Qualifiers

- # The control limit criteria is not applicable. See case narrative.
- J The result is an estimated value.
- E The percent difference for the serial dilution was greater than 10%, indicating a possible matrix interference in the sample.
- M The duplicate injection precision was not met.
- N The Matrix Spike sample recovery is not within control limits. See case narrative.
- S The reported value was determined by the Method of Standard Additions (MSA).
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- W The post-digestion spike for furnace AA analysis is out of control limits, while sample absorbance is less than 50% of spike absorbance.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- + The correlation coefficient for the MSA is less than 0.995.
- Q See case narrative. One or more quality control criteria was outside the limits.

Organic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- A A tentatively identified compound, a suspected aldol-condensation product.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- C The analyte was qualitatively confirmed using GC/MS techniques, pattern recognition, or by comparing to historical data.
- D The reported result is from a dilution.
- E The result is an estimated value.
- J The result is an estimated value.
- N The result is presumptive. The analyte was tentatively identified, but a confirmation analysis was not performed.
- P The GC or HPLC confirmation criteria was exceeded. The relative percent difference is greater than 40% between the two analytical results.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a chromatographic interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.

Additional Petroleum Hydrocarbon Specific Qualifiers

- F The chromatographic fingerprint of the sample matches the elution pattern of the calibration standard.
- L The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of lighter molecular weight constituents than the calibration standard.
- H The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of heavier molecular weight constituents than the calibration standard.
- O The chromatographic fingerprint of the sample resembles an oil, but does not match the calibration standard.
- Y The chromatographic fingerprint of the sample resembles a petroleum product eluting in approximately the correct carbon range, but the elution pattern does not match the calibration standard.
- Z The chromatographic fingerprint does not resemble a petroleum product.

**ALS Group USA Corp. dba ALS Environmental (ALS) - Kelso
State Certifications, Accreditations, and Licenses**

Agency	Web Site	Number
Alaska DEH	http://dec.alaska.gov/eh/lab/cs/csapproval.htm	UST-040
Arizona DHS	http://www.azdhs.gov/lab/license/env.htm	AZ0339
Arkansas - DEQ	http://www.adeq.state.ar.us/techsvs/labcert.htm	88-0637
California DHS (ELAP)	http://www.cdph.ca.gov/certlic/labs/Pages/ELAP.aspx	2795
DOD ELAP	http://www.denix.osd.mil/edqw/Accreditation/AccreditedLabs.cfm	L16-58-R4
Florida DOH	http://www.doh.state.fl.us/lab/EnvLabCert/WaterCert.htm	E87412
Hawaii DOH	http://health.hawaii.gov/	-
ISO 17025	http://www.pjllabs.com/	L16-57
Louisiana DEQ	http://www.deq.louisiana.gov/page/la-lab-accreditation	03016
Maine DHS	http://www.maine.gov/dhhs/	WA01276
Minnesota DOH	http://www.health.state.mn.us/accreditation	053-999-457
Nevada DEP	http://ndep.nv.gov/bsdw/labservice.htm	WA01276
New Jersey DEP	http://www.nj.gov/dep/enforcement/oqa.html	WA005
New York - DOH	https://www.wadsworth.org/regulatory/elap	12060
North Carolina DEQ	https://deq.nc.gov/about/divisions/water-resources/water-resources-data/water-sciences-home-page/laboratory-certification-branch/non-field-lab-certification	605
Oklahoma DEQ	http://www.deq.state.ok.us/CSDnew/labcert.htm	9801
Oregon – DEQ (NELAP)	http://public.health.oregon.gov/LaboratoryServices/EnvironmentalLaboratoryAccreditation/Pages/index.aspx	WA100010
South Carolina DHEC	http://www.scdhec.gov/environment/EnvironmentalLabCertification/	61002
Texas CEQ	http://www.tceq.texas.gov/field/qa/env_lab_accreditation.html	T104704427
Washington DOE	http://www.ecy.wa.gov/programs/eap/labs/lab-accreditation.html	C544
Wyoming (EPA Region 8)	https://www.epa.gov/region8-waterops/epa-region-8-certified-drinking-water	-
Kelso Laboratory Website	www.alsglobal.com	NA

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. A complete listing of specific NELAP-certified analytes, can be found in the certification section at www.ALSGlobal.com or at the accreditation bodies web site.

Please refer to the certification and/or accreditation body's web site if samples are submitted for compliance purposes. The states highlighted above, require the analysis be listed on the state certification if used for compliance purposes and if the method/analyte is offered by that state.



Case Narrative

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com



Client: ALS Environmental - US
Project: HS19030298
Sample Matrix: Water

Service Request: K1902047
Date Received: 03/08/2019

CASE NARRATIVE

All analyses were performed consistent with the quality assurance program of ALS Environmental. This report contains analytical results for samples designated for Tier IV validation deliverables including summary forms and all of the associated raw data for each of the analyses. When appropriate to the method, method blank results have been reported with each analytical test.

Sample Receipt:

One water sample was received for analysis at ALS Environmental on 03/08/2019. The sample was received in good condition and consistent with the accompanying chain of custody form. The sample was stored in a refrigerator at 4°C upon receipt at the laboratory.

General Chemistry:

No significant anomalies were noted with this analysis.

Approved by Kelley Avejoy

Date 03/27/2019



Chain of Custody

ALS Environmental—Kelso Laboratory
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Phone (360)577- 7222 Fax (360)636-1 068
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K1902047



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 Houston, TX 77099
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www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10881

SUBCONTRACT TO:

ALS Environmental Kelso
 1317 S. 13th Avenue
 Kelso, WA 98626

Phone: +1 360 501 3312

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com



INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030298
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030298-01	LH18/24-SP650_031619	Water	06 Mar 2019 13:00
TOC Analysis for DOD Level IV			21 Mar 2019

Comments: Please analyze for the analysis listed above.
 Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: 
 Received By: 
 Cooler ID(s): _____

Date/Time: 3/7/19 1800
 Date/Time: 3/8/19 0940
 Temperature(s): _____

RIGHT SOLUTIONS | RIGHT PARTNER



PC KV

Cooler Receipt and Preservation Form

Client ALS-Houston Service Request K19 07047
 Received: 3/8/19 Opened: 3/8/19 By: BR Unloaded: 3/8/19 By: BR

1. Samples were received via? USPS Fed Ex UPS DHL PDX Courier Hand Delivered
 2. Samples were received in: (circle) Cooler Box Envelope Other NA
 3. Were custody seals on coolers? NA Y N If yes, how many and where? 2 front
 If present, were custody seals intact? Y N If present, were they signed and dated? Y N

Raw Cooler Temp	Corrected Cooler Temp	Raw Temp Blank	Corrected Temp Blank	Corr. Factor	Thermometer ID	Cooler/COC ID	Tracking Number	NA	Filed
-0.1	-0.3	1.9	1.7	-0.2	350	<u>NA</u>	480978314288		

4. Packing material: Inserts Baggies Bubble Wrap Gel Packs Wet Ice Dry Ice Sleeves
 5. Were custody papers properly filled out (ink, signed, etc.)? NA Y N
 6. Were samples received in good condition (temperature, unbroken)? Indicate in the table below. NA Y N
 If applicable, tissue samples were received: Frozen Partially Thawed Thawed
 7. Were all sample labels complete (i.e analysis, preservation, etc.)? NA Y N
 8. Did all sample labels and tags agree with custody papers? Indicate major discrepancies in the table on page 2. NA Y N
 9. Were appropriate bottles/containers and volumes received for the tests indicated? NA Y N
 10. Were the pH-preserved bottles (see SMO GEN SOP) received at the appropriate pH? Indicate in the table below NA Y N
 11. Were VOA vials received without headspace? Indicate in the table below. NA Y N
 12. Was C12/Res negative? NA Y N

Sample ID on Bottle	Sample ID on COC	Identified by:

Sample ID	Bottle Count	Bottle Type	Out of Temp	Head-space	Broke	pH	Reagent	Volume added	Reagent Lot Number	Initials	Time

Notes, Discrepancies, & Resolutions: _____



General Chemistry

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Analytical Report

Client: ALS Environmental - US
Project: HS19030298
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1902047
Date Collected: 03/6/19
Date Received: 03/8/19
Units: mg/L
Basis: NA

Carbon, Total Organic

Sample Name	Lab Code	Result	LOQ	LOD	MDL	Dil.	Date Analyzed	Q
LH18/24-SP650_030619	K1902047-001	2.94	0.50	0.20	0.07	1	03/21/19 23:03	
Method Blank	K1902047-MB	ND U	0.50	0.20	0.07	1	03/22/19 04:44	

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030298
Sample Matrix: Water

Service Request: K1902047
Date Collected: N/A
Date Received: N/A
Date Analyzed: 03/22/19
Date Extracted: NA

Matrix Spike Summary
Carbon, Total Organic

Sample Name: Batch QC
Lab Code: K1902389-001
Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA

Matrix Spike
K1902389-001MS

Analyte Name	Sample Result	Result	Spike Amount	% Rec	% Rec Limits
Carbon, Total Organic	3.34	31.0	25.0	111	83-117

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030298
Sample Matrix: Water

Service Request: K1902047
Date Analyzed: 03/22/19
Date Extracted: NA

Lab Control Sample Summary
Carbon, Total Organic

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA
Analysis Lot: 629144

Sample Name	Lab Code	Result	Spike Amount	% Rec	% Rec Limits
Lab Control Sample	K1902047-LCS	26.9	25.0	107	83-117

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030298

Service Request: K1902047

Continuing Calibration Verification (CCV) Summary

Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis		Date	True	Measured	Percent	Acceptance
	Lot	Lab Code	Analyzed	Value	Value	Recovery	Limits
CCV1	629144	KQ1903793-06	03/21/19 19:42	25.0	24.5	98	90-110
CCV2	629144	KQ1903793-07	03/21/19 23:31	25.0	24.5	98	90-110
CCV3	629144	KQ1903793-08	03/22/19 04:15	25.0	24.9	100	90-110
CCV4	629144	KQ1903793-09	03/22/19 08:58	25.0	24.4	98	90-110
CCV5	629144	KQ1903793-10	03/22/19 14:09	25.0	24.2	97	90-110

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030298

Service Request: K1902047

Continuing Calibration Blank (CCB) Summary
Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis Lot	Lab Code	Date Analyzed	LOQ	LOD	MDL	Result	Q
CCB1	629144	KQ1903793-01	03/21/19 19:57	0.50	0.20	0.07	ND	U
CCB2	629144	KQ1903793-02	03/21/19 23:46	0.50	0.20	0.07	ND	U
CCB3	629144	KQ1903793-03	03/22/19 04:30	0.50	0.20	0.07	ND	U
CCB4	629144	KQ1903793-04	03/22/19 09:13	0.50	0.20	0.07	ND	U
CCB5	629144	KQ1903793-05	03/22/19 14:23	0.50	0.20	0.07	ND	U



Raw Data

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General Chemistry

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Work Request # (Original) K1902074, 2162, 1927, 2047, 2102, 2121, 2127, 2147, 2153, 2168, 2346, 2389, 2121, 2169, 2330, 2232
 Tier: II, II, I, IV, II, I, II, II, II, II, IV, IV, I, II, II
 Date Analyzed: 3/2/19 DOC: 629142
TOC: 629144
629140
629144
 Analyst: BCD Run # _____
 Analysis: TOC/DOC

**DATA QUALITY REPORT
INORGANICS**

Explain any "no" responses to questions below, and any corrective actions in the comments section below.

1. Is the method name and number correct and appropriate? yes/no/NA
2. Holding times met for all analyses and for all samples? yes/no/NA
3. Are calculations correct? yes/no/NA
4. Is the reporting basis correct? (Dry Weight) yes/no/NA
5. All quality control criteria met? yes/no
6. Is the calibration curve correlation coefficient ≥ 0.995 ? yes/no/NA
7. MBs, CCVs, CCBs, LCSs, Dups, and Spikes, analyzed at proper frequency? yes/no/NA
8. Are ICVs, CCVs, and CCBs all within acceptance limits? yes/no/NA
9. Are results for methods blanks all ND? yes/no/NA
10. Are all QC samples within acceptance criteria? (LCS % rec, MS/DMS % rec, DUP or MS/DMS RPDs, etc.) yes/no/NA
11. Are all exceptions explained? yes/no/NA
12. Have all applicable service requests been reviewed? yes/no/NA
13. Are all samples labeled correctly? yes/no/NA
14. Have all instructions on the service request been followed? (e.g. Special MRLs, QC on a specific sample, Form V) yes/no/NA
15. Are detection limits and units reported correctly? yes/no/NA
16. Is the unused space on the benchsheet crossed out? yes/no/NA
17. Was analysis turned in by the due date? (n-2) (If not record SR#) yes/no/NA

COMMENTS: R 1902232-2/3, R1902121-1/2, K1902435-1/2/3/4/5/6/7, K1902254-1/2/3
 Sent for reanalysis due to CCB above the MRL.
 K1902102-1, K1902074-2/1 sent for reanalysis due to MB requiring a dilution.
 K1902102-1 for sent reanalysis due to requiring a dilution.
 K1902147-1 sent for reanalysis because it is overloaded.
 K1902155-1/dups have a high PRSD due to dirty, non-homogeneous sample.
 K1902168-2/2d report a high %RSD, but these samples are less than 5x the MRL.

Final Approved by: Houyer Date: 03/25/19 DQREPORT

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629142 Method/Testcode: SM 5310 C/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902074-001	Carbon, Dissolved Organic N/A (DOC)			Effluent	1.64 mg/L	10 ml	1.64 mg/L	1	0.07	0.50			3/21/19 20:56	N	II
K1902074-002	Carbon, Dissolved Organic N/A (DOC)			Water	1.65 mg/L	10 ml	1.65 mg/L	1	0.07	0.50			3/21/19 21:24	N	II
K1902102-001	Carbon, Dissolved Organic N/A (DOC)			Water	76.61 mg/L	10 ml	76.6 mg/L	1	0.07	0.50			3/21/19 21:52	N	II
KQ1903792-01	Carbon, Dissolved Organic CCB (DOC)			Effluent	0.01 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 19:57	N	II
KQ1903792-02	Carbon, Dissolved Organic CCB (DOC)			Effluent	0.03 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 23:46	N	II
KQ1903792-03	Carbon, Dissolved Organic CCV (DOC)			Effluent	24.51 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 19:42	N	II
KQ1903792-04	Carbon, Dissolved Organic CCV (DOC)			Effluent	24.47 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 23:31	N	II
KQ1903792-05	Carbon, Dissolved Organic MB (DOC)			Effluent	-0.04 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 20:12	N	II
KQ1903792-06	Carbon, Dissolved Organic LCS (DOC)			Effluent	26.54 mg/L	10 ml	26.5 mg/L	1	0.07	0.50	106		3/21/19 20:26	N	II
KQ1903792-07	Carbon, Dissolved Organic MS (DOC)		K1902102-001	Water	99.38 mg/L	10 ml	99.4 mg/L	1	0.07	0.50	91		3/21/19 22:20	N	II
KQ1903792-08	Carbon, Dissolved Organic DUP (DOC)		K1902102-001	Water	78.20 mg/L	10 ml	78.2 mg/L	1	0.07	0.50		2	3/21/19 21:52	N	II
KQ1903792-09	Carbon, Dissolved Organic DUP (DOC)		K1902074-002	Water	1.68 mg/L	10 ml	1.68 mg/L	1	0.07	0.50		2	3/21/19 21:24	N	II
KQ1903792-10	Carbon, Dissolved Organic DUP (DOC)		K1902074-001	Effluent	1.62 mg/L	10 ml	1.62 mg/L	1	0.07	0.50		1	3/21/19 20:56	N	II

03/25/19
Free up

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629144 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1901927-005	Carbon, Total Organic	N/A		Ground Water	45.71 mg/L	10 ml	45.7 mg/L	1	0.07	0.50			3/22/19 00:01	N	I
K1902047-001	Carbon, Total Organic	N/A		Water	2.94 mg/L	10 ml	2.94 mg/L	1	0.07	0.50			3/21/19 23:03	N	IV
K1902102-001	Carbon, Total Organic	N/A		Water	83.22 mg/L	10 ml	83.2 mg/L	1	0.07	0.50			3/22/19 05:42	N	II
K1902121-002	Carbon, Total Organic	N/A		Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 20:19	N	I
K1902127-001	Carbon, Total Organic	N/A		Water	15.73 mg/L	10 ml	15.7 mg/L	1	0.07	0.50			3/22/19 06:10	N	II
K1902127-002	Carbon, Total Organic	N/A		Water	7.06 mg/L	10 ml	28.2 mg/L	4	0.3	2.0			3/22/19 06:38	N	II
K1902127-003	Carbon, Total Organic	N/A		Water	7.65 mg/L	10 ml	7.65 mg/L	1	0.07	0.50			3/22/19 07:06	N	II
K1902127-004	Carbon, Total Organic	N/A		Water	4.54 mg/L	10 ml	18.1 mg/L	4	0.3	2.0			3/22/19 07:34	N	II
K1902147-001	Carbon, Total Organic	N/A		Water	0.23 mg/L	10 ml	50 mg/L U	100	7	50			3/22/19 08:02	N	II
K1902147-002	Carbon, Total Organic	N/A		Water	0.95 mg/L	10 ml	0.95 mg/L	1	0.07	0.50			3/22/19 08:30	N	II
K1902153-001	Carbon, Total Organic	N/A		Water	2.23 mg/L	10 ml	223 mg/L	100	7	50			3/22/19 02:50	N	II
K1902153-002	Carbon, Total Organic	N/A		Water	3.16 mg/L	10 ml	316 mg/L	100	7	50			3/22/19 03:19	N	II
K1902153-003	Carbon, Total Organic	N/A		Water	1.20 mg/L	10 ml	120 mg/L	100	7	50			3/22/19 03:47	N	II
K1902168-001	Carbon, Total Organic	N/A		Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50			3/22/19 09:28	N	II
K1902168-002	Carbon, Total Organic	N/A		Water	0.87 mg/L	10 ml	0.87 mg/L	1	0.07	0.50			3/22/19 09:56	N	II
K1902346-001	Carbon, Total Organic	N/A		Water	2.96 mg/L	10 ml	2.96 mg/L	1	0.07	0.50			3/22/19 05:14	N	IV
K1902389-001	Carbon, Total Organic	N/A		Water	3.34 mg/L	10 ml	3.34 mg/L	1	0.07	0.50			3/22/19 00:57	Y	IV
K1902389-002	Carbon, Total Organic	N/A		Water	27.06 mg/L	10 ml	27.1 mg/L	1	0.07	0.50			3/22/19 01:54	N	IV
K1902389-003	Carbon, Total Organic	N/A		Water	27.93 mg/L	10 ml	27.9 mg/L	1	0.07	0.50			3/22/19 02:22	N	IV
KQ1903793-01	Carbon, Total Organic	CCB		Water	9.45000000000001E	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 19:57	N	IV
KQ1903793-02	Carbon, Total Organic	CCB		Water	0.03 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 23:46	N	IV
KQ1903793-03	Carbon, Total Organic	CCB		Water	0.05 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 04:30	N	IV
KQ1903793-04	Carbon, Total Organic	CCB		Water	-0.17 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 09:13	N	IV
KQ1903793-05	Carbon, Total Organic	CCB		Water	-0.19 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 14:23	N	IV
KQ1903793-06	Carbon, Total Organic	CCV		Water	24.51 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 19:42	N	IV
KQ1903793-07	Carbon, Total Organic	CCV		Water	24.47 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 23:31	N	IV
KQ1903793-08	Carbon, Total Organic	CCV		Water	24.91 mg/L	10 ml	24.9 mg/L	1			100		3/22/19 04:15	N	IV
KQ1903793-09	Carbon, Total Organic	CCV		Water	24.39 mg/L	10 ml	24.4 mg/L	1			98		3/22/19 08:58	N	IV
KQ1903793-10	Carbon, Total Organic	CCV		Water	24.24 mg/L	10 ml	24.2 mg/L	1			97		3/22/19 14:09	N	IV
KQ1903793-11	Carbon, Total Organic	MB		Water	1.65000000000001E	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 04:44	N	IV
KQ1903793-12	Carbon, Total Organic	LCS		Water	26.86 mg/L	10 ml	26.9 mg/L	1	0.07	0.50	107		3/22/19 04:59	N	IV
KQ1903793-13	Carbon, Total Organic	MS	K1902389-001	Water	30.99 mg/L	10 ml	31.0 mg/L	1	0.07	0.50	111		3/22/19 01:25	N	IV
KQ1903793-14	Carbon, Total Organic	DUP	K1902047-001	Water	2.93 mg/L	10 ml	2.93 mg/L	1	0.07	0.50		<1	3/21/19 23:03	N	IV
KQ1903793-15	Carbon, Total Organic	DUP	K1901927-005	Ground Water	46.15 mg/L	10 ml	46.1 mg/L	1	0.07	0.50		<1	3/22/19 00:01	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

03/25/19
The sample

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629144 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1903793-16	Carbon, Total Organic	DUP	K1902389-001	Water	3.34 mg/L	10 ml	3.34 mg/L	1	0.07	0.50		<1	3/22/19 00:57	N	IV
KQ1903793-17	Carbon, Total Organic	DUP	K1902389-002	Water	27.77 mg/L	10 ml	27.8 mg/L	1	0.07	0.50		3	3/22/19 01:54	N	IV
KQ1903793-18	Carbon, Total Organic	DUP	K1902389-003	Water	27.82 mg/L	10 ml	27.8 mg/L	1	0.07	0.50		<1	3/22/19 02:22	N	IV
KQ1903793-19	Carbon, Total Organic	DUP	K1902153-001	Water	1.67 mg/L	10 ml	167 mg/L	100	7	50		29*	3/22/19 02:50	N	II
KQ1903793-20	Carbon, Total Organic	DUP	K1902153-002	Water	3.14 mg/L	10 ml	314 mg/L	100	7	50		<1	3/22/19 03:19	N	II
KQ1903793-21	Carbon, Total Organic	DUP	K1902153-003	Water	1.13 mg/L	10 ml	113 mg/L	100	7	50		6	3/22/19 03:47	N	II
KQ1903793-22	Carbon, Total Organic	DUP	K1902346-001	Water	2.84 mg/L	10 ml	2.84 mg/L	1	0.07	0.50		4	3/22/19 05:14	N	IV
KQ1903793-23	Carbon, Total Organic	DUP	K1902102-001	Water	83.37 mg/L	10 ml	83.4 mg/L	1	0.07	0.50		<1	3/22/19 05:42	N	II
KQ1903793-24	Carbon, Total Organic	DUP	K1902121-002	Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L	U	1	0.07	0.50	NC	3/22/19 20:19	N	I
KQ1903793-25	Carbon, Total Organic	DUP	K1902127-001	Water	15.57 mg/L	10 ml	15.6 mg/L	1	0.07	0.50		<1	3/22/19 06:10	N	II
KQ1903793-26	Carbon, Total Organic	DUP	K1902127-002	Water	6.91 mg/L	10 ml	27.6 mg/L	4	0.3	2.0		2	3/22/19 06:38	N	II
KQ1903793-27	Carbon, Total Organic	DUP	K1902127-003	Water	7.52 mg/L	10 ml	7.52 mg/L	1	0.07	0.50		2	3/22/19 07:06	N	II
KQ1903793-28	Carbon, Total Organic	DUP	K1902127-004	Water	4.39 mg/L	10 ml	17.6 mg/L	4	0.3	2.0		3	3/22/19 07:34	N	II
KQ1903793-29	Carbon, Total Organic	DUP	K1902147-001	Water	0.22 mg/L	10 ml	22 mg/L	J	100	7	50	NC	3/22/19 08:02	N	II
KQ1903793-30	Carbon, Total Organic	DUP	K1902147-002	Water	0.91 mg/L	10 ml	0.91 mg/L	1	0.07	0.50		4	3/22/19 08:30	N	II
KQ1903793-31	Carbon, Total Organic	DUP	K1902168-001	Water	1.78 mg/L	10 ml	1.78 mg/L	1	0.07	0.50		4	3/22/19 09:28	N	II
KQ1903793-32	Carbon, Total Organic	DUP	K1902168-002	Water	0.76 mg/L	10 ml	0.76 mg/L	1	0.07	0.50		14*	3/22/19 09:56	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629146 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902121-001	AA Carbon, Total Organic	N/A		Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 19:51	N	I
K1902169-001	Carbon, Total Organic	N/A		Water	14.29 mg/L	10 ml	14.3 mg/L	1	0.07	0.50			3/22/19 10:24	N	II
K1902169-002	Carbon, Total Organic	N/A		Water	17.52 mg/L	10 ml	17.5 mg/L	1	0.07	0.50			3/22/19 10:52	N	II
K1902169-003	Carbon, Total Organic	N/A		Water	18.67 mg/L	10 ml	18.7 mg/L	1	0.07	0.50			3/22/19 11:20	N	II
K1902169-004	Carbon, Total Organic	N/A		Water	9.18 mg/L	10 ml	9.18 mg/L	1	0.07	0.50			3/22/19 11:48	N	II
K1902169-005	Carbon, Total Organic	N/A		Water	11.61 mg/L	10 ml	11.6 mg/L	1	0.07	0.50			3/22/19 12:16	N	II
K1902169-006	Carbon, Total Organic	N/A		Water	11.71 mg/L	10 ml	11.7 mg/L	1	0.07	0.50			3/22/19 12:44	N	II
K1902169-007	Carbon, Total Organic	N/A		Water	12.46 mg/L	10 ml	12.5 mg/L	1	0.07	0.50			3/22/19 13:12	N	II
K1902169-008	Carbon, Total Organic	N/A		Water	15.31 mg/L	10 ml	15.3 mg/L	1	0.07	0.50			3/22/19 13:41	N	II
K1902230-001	Carbon, Total Organic	N/A		Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50			3/22/19 15:07	N	II
K1902230-002	Carbon, Total Organic	N/A		Water	0.70 mg/L	10 ml	0.70 mg/L	1	0.07	0.50			3/22/19 15:35	N	II
K1902230-003	Carbon, Total Organic	N/A		Water	0.80 mg/L	10 ml	0.80 mg/L	1	0.07	0.50			3/22/19 16:03	N	II
K1902230-004	Carbon, Total Organic	N/A		Water	1.23 mg/L	10 ml	1.23 mg/L	1	0.07	0.50			3/22/19 16:32	N	II
K1902230-005	Carbon, Total Organic	N/A		Water	0.32 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 17:00	N	II
K1902232-001	Carbon, Total Organic	N/A		Water	5.72 mg/L	10 ml	5.72 mg/L	1	0.07	0.50			3/22/19 17:28	N	II
K1902232-002	RA Carbon, Total Organic	N/A		Water	9.75 mg/L	10 ml	9.75 mg/L	1	0.07	0.50			3/22/19 18:55	N	II
K1902232-003	RA Carbon, Total Organic	N/A		Water	13.90 mg/L	10 ml	13.9 mg/L	1	0.07	0.50			3/22/19 19:23	N	II
KQ1903794-01	Carbon, Total Organic	CCB		Water	-0.17 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 09:13	N	II
KQ1903794-02	Carbon, Total Organic	CCB		Water	-0.19 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 14:23	N	II
KQ1903794-03	Carbon, Total Organic	CCB		Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 18:40	N	II
KQ1903794-04	Carbon, Total Organic	CCB (High)		Water	0.89 mg/L	10 ml	0.89 mg/L	1	0.07	0.50			3/22/19 23:50	N	II
KQ1903794-05	Carbon, Total Organic	CCV		Water	24.39 mg/L	10 ml	24.4 mg/L	1					3/22/19 08:58	N	II
KQ1903794-06	Carbon, Total Organic	CCV		Water	24.24 mg/L	10 ml	24.2 mg/L	1					3/22/19 14:09	N	II
KQ1903794-07	Carbon, Total Organic	CCV		Water	23.91 mg/L	10 ml	23.9 mg/L	1					3/22/19 18:25	N	II
KQ1903794-08	Carbon, Total Organic	CCV		Water	27.39 mg/L	10 ml	27.4 mg/L	1					3/22/19 23:35	N	II
KQ1903794-09	Carbon, Total Organic	MB		Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 14:38	N	II
KQ1903794-10	Carbon, Total Organic	LCS		Water	26.10 mg/L	10 ml	26.1 mg/L	1	0.07	0.50	104		3/22/19 14:53	N	II
KQ1903794-11	Carbon, Total Organic	MS	K1902232-001	Water	33.31 mg/L	10 ml	33.3 mg/L	1	0.07	0.50	110		3/22/19 17:56	N	II
KQ1903794-12	Carbon, Total Organic	DUP	K1902169-001	Water	14.17 mg/L	10 ml	14.2 mg/L	1	0.07	0.50		<1	3/22/19 10:24	N	II
KQ1903794-13	Carbon, Total Organic	DUP	K1902169-002	Water	17.70 mg/L	10 ml	17.7 mg/L	1	0.07	0.50		1	3/22/19 10:52	N	II
KQ1903794-14	Carbon, Total Organic	DUP	K1902169-003	Water	18.63 mg/L	10 ml	18.6 mg/L	1	0.07	0.50		<1	3/22/19 11:20	N	II
KQ1903794-15	Carbon, Total Organic	DUP	K1902169-004	Water	9.05 mg/L	10 ml	9.05 mg/L	1	0.07	0.50		1	3/22/19 11:48	N	II
KQ1903794-16	Carbon, Total Organic	DUP	K1902169-005	Water	11.53 mg/L	10 ml	11.5 mg/L	1	0.07	0.50		<1	3/22/19 12:16	N	II
KQ1903794-17	Carbon, Total Organic	DUP	K1902169-006	Water	11.48 mg/L	10 ml	11.5 mg/L	1	0.07	0.50		2	3/22/19 12:44	N	II
KQ1903794-18	Carbon, Total Organic	DUP	K1902169-007	Water	12.85 mg/L	10 ml	12.9 mg/L	1	0.07	0.50		3	3/22/19 13:12	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

03/25/19
[Handwritten Signature]

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629146 Method/Testcode: SM 5310 C/TOC T

<u>Lab Code</u>	<u>Target Analytes</u>	<u>QC</u>	<u>Parent Sample</u>	<u>Matrix</u>	<u>Raw Result</u>	<u>Sample Amt.</u>	<u>Final Result</u>	<u>Dil</u>	<u>MDL</u>	<u>PQL</u>	<u>% Rec</u>	<u>% RSD</u>	<u>Date Analyzed</u>	<u>QC?</u>	<u>Tier</u>
KQ1903794-19	Carbon, Total Organic	DUP	K1902169-008	Water	15.02 mg/L	10 ml	15.0 mg/L	1	0.07	0.50		2	3/22/19 13:41	N	II
KQ1903794-20	Carbon, Total Organic	DUP	K1902230-001	Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50		<1	3/22/19 15:07	N	II
KQ1903794-21	Carbon, Total Organic	DUP	K1902230-002	Water	0.64 mg/L	10 ml	0.64 mg/L	1	0.07	0.50		8	3/22/19 15:35	N	II
KQ1903794-22	Carbon, Total Organic	DUP	K1902230-003	Water	0.79 mg/L	10 ml	0.79 mg/L	1	0.07	0.50		<1	3/22/19 16:03	N	II
KQ1903794-23	Carbon, Total Organic	DUP	K1902230-004	Water	1.14 mg/L	10 ml	1.14 mg/L	1	0.07	0.50		7	3/22/19 16:32	N	II
KQ1903794-24	Carbon, Total Organic	DUP	K1902230-005	Water	0.30 mg/L	10 ml	0.30 mg/L	J 1	0.07	0.50		NC	3/22/19 17:00	N	II
KQ1903794-25	Carbon, Total Organic	DUP	K1902232-001	Water	5.79 mg/L	10 ml	5.79 mg/L	1	0.07	0.50		1	3/22/19 17:28	N	II
KQ1903794-26	Carbon, Total Organic	DUP	K1902232-002	Water	9.86 mg/L	10 ml	9.86 mg/L	1	0.07	0.50		1	3/22/19 18:55	N	II
KQ1903794-27	Carbon, Total Organic	DUP	K1902232-003	Water	13.87 mg/L	10 ml	13.9 mg/L	1	0.07	0.50		<1	3/22/19 19:23	N	II
KQ1903794-28	Carbon, Total Organic	DUP	K1902121-001	Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50		NC	3/22/19 19:51	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629149 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902254-001	Carbon, Total Organic	N/A		Water	6.54 mg/L	10 ml	6.54 mg/L	1	0.07	0.50			3/23/19 01:02	N	II
K1902254-002	Carbon, Total Organic	N/A		Water	18.37 mg/L	10 ml	18.4 mg/L	1	0.07	0.50			3/23/19 01:30	N	II
K1902254-003	Carbon, Total Organic	N/A		Water	22.01 mg/L	10 ml	22.0 mg/L	1	0.07	0.50			3/23/19 01:58	N	II
K1902435-001	Carbon, Total Organic	N/A		Ground Water	119.91 mg/L	10 ml	120 mg/L	1	0.07	0.50			3/22/19 20:47	N	IV
K1902435-002	Carbon, Total Organic	N/A		Ground Water	5.07 mg/L	10 ml	5.07 mg/L	1	0.07	0.50			3/22/19 21:15	N	IV
K1902435-003	Carbon, Total Organic	N/A		Ground Water	3.20 mg/L	10 ml	3.20 mg/L	1	0.07	0.50			3/22/19 21:43	N	IV
K1902435-004	Carbon, Total Organic	N/A		Ground Water	5.08 mg/L	10 ml	5.08 mg/L	1	0.07	0.50			3/22/19 22:11	N	IV
K1902435-005	Carbon, Total Organic	N/A		Ground Water	137.01 mg/L	10 ml	137 mg/L	1	0.07	0.50			3/22/19 22:39	N	IV
K1902435-006	Carbon, Total Organic	N/A		Ground Water	118.27 mg/L	10 ml	118 mg/L	1	0.07	0.50			3/22/19 23:07	N	IV
K1902435-007	Carbon, Total Organic	N/A		Ground Water	12.50 mg/L	10 ml	12.5 mg/L	1	0.07	0.50			3/23/19 00:05	Y	IV
KQ1903795-01	Carbon, Total Organic	CCB		Ground Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 18:40	N	IV
KQ1903795-02	Carbon, Total Organic	CCB		Ground Water	0.89 mg/L	10 ml	0.89 mg/L	1	0.07	0.50			3/22/19 23:50	N	IV
KQ1903795-03	Carbon, Total Organic	CCB		Ground Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/23/19 03:09	N	IV
KQ1903795-04	Carbon, Total Organic	CCV		Ground Water	23.91 mg/L	10 ml	23.9 mg/L	1			96		3/22/19 18:25	N	IV
KQ1903795-05	Carbon, Total Organic	CCV		Ground Water	27.39 mg/L	10 ml	27.4 mg/L	1			110		3/22/19 23:35	N	IV
KQ1903795-06	Carbon, Total Organic	CCV		Ground Water	23.81 mg/L	10 ml	23.8 mg/L	1			95		3/23/19 02:55	N	IV
KQ1903795-09	Carbon, Total Organic	MS	K1902435-007	Ground Water	33.24 mg/L	10 ml	33.2 mg/L	1	0.07	0.50	83		3/23/19 00:33	N	IV
KQ1903795-10	Carbon, Total Organic	DUP	K1902435-001	Ground Water	112.05 mg/L	10 ml	112 mg/L	1	0.07	0.50		7	3/22/19 20:47	N	IV
KQ1903795-11	Carbon, Total Organic	DUP	K1902435-002	Ground Water	3.05 mg/L	10 ml	3.05 mg/L	1	0.07	0.50		50*	3/22/19 21:15	N	IV
KQ1903795-12	Carbon, Total Organic	DUP	K1902435-003	Ground Water	2.88 mg/L	10 ml	2.88 mg/L	1	0.07	0.50		11*	3/22/19 21:43	N	IV
KQ1903795-13	Carbon, Total Organic	DUP	K1902435-004	Ground Water	4.70 mg/L	10 ml	4.70 mg/L	1	0.07	0.50		8	3/22/19 22:11	N	IV
KQ1903795-14	Carbon, Total Organic	DUP	K1902435-005	Ground Water	127.76 mg/L	10 ml	128 mg/L	1	0.07	0.50		7	3/22/19 22:39	N	IV
KQ1903795-15	Carbon, Total Organic	DUP	K1902435-006	Ground Water	127.20 mg/L	10 ml	127 mg/L	1	0.07	0.50		7	3/22/19 23:07	N	IV
KQ1903795-16	Carbon, Total Organic	DUP	K1902435-007	Ground Water	12.30 mg/L	10 ml	12.3 mg/L	1	0.07	0.50		2	3/23/19 00:05	N	IV
KQ1903795-17	Carbon, Total Organic	DUP	K1902254-001	Water	6.38 mg/L	10 ml	6.38 mg/L	1	0.07	0.50		3	3/23/19 01:02	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Printed 3/23/19 15:44

Results Summary

03/25/19
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Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629149 Method/Testcode: SM 5310 C/TOC T

<u>Lab Code</u>	<u>Target Analytes</u>	<u>QC</u>	<u>Parent Sample</u>	<u>Matrix</u>	<u>Raw Result</u>	<u>Sample Amt.</u>	<u>Final Result</u>	<u>Dil</u>	<u>MDL</u>	<u>PQL</u>	<u>% Rec</u>	<u>% RSD</u>	<u>Date Analyzed</u>	<u>QC?</u>	<u>Tier</u>
KQ1903795-18	Carbon, Total Organic	DUP	K1902254-002	Water	18.17 mg/L	10 ml	18.2 mg/L	1	0.07	0.50		1	3/23/19 01:30	N	II
KQ1903795-19	Carbon, Total Organic	DUP	K1902254-003	Water	21.93 mg/L	10 ml	21.9 mg/L	1	0.07	0.50		<1	3/23/19 01:58	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

DOC: 629142
 TOC: 629144,
 629146,
 629149

Schedule: 03212019

Version: 4

Instrument: Fusion1

Last Saved by: Fusion1 (Fusion1)

Last Saved on: 2019/03/21 18:03 - Thursday

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Blank)	Blank	Reagent/Acid Blank		1	True	Ready
D	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
1	Sample	MB1	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
2	Sample	ICS	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
3	Sample	K1902074-001.03 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
4	Sample	K1902074-002.03 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
5	Sample	K1902102-001.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
6	Sample	K1902102-001.01 ms doc	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
7	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
8	Sample	FB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
9	Sample	K1902047-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
10	Sample	K1901927-005.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
11	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
12	Sample	K1902389-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
13	Sample	K1902389-001.01 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
14	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
15	Sample	K1902389-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
16	Sample	K1902389-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
17	Sample	K1902153-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
18	Sample	K1902153-002.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
19	Sample	K1902153-003.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
20	Sample	MB2	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
21	Sample	K1902346-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
22	Sample	K1902102-001.02	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
23	Sample	K1902127-001.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
24	Sample	K1902127-002.12 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
25	Sample	K1902127-003.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
26	Sample	K1902127-004.12 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
27	Sample	K1902147-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
28	Sample	K1902147-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
29	Sample	K1902168-001.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
30	Sample	K1902168-002.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
31	Sample	K1902169-001.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
32	Sample	K1902169-002.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
33	Sample	K1902169-003.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
34	Sample	K1902169-004.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
35	Sample	K1902169-005.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
36	Sample	K1902169-006.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
37	Sample	K1902169-007.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
38	Sample	K1902169-008.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready

Printed on: March 23, 2019 11:38:49

Schedule: 03212019

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
39	Sample	MB3	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
40	Sample	K1902230-001.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
41	Sample	K1902230-002.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
42	Sample	K1902230-003.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
43	Sample	K1902230-004.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
44	Sample	K1902230-005.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
45	Sample	K1902232-001.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
46	Sample	K1902232-001.12 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
47	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
48	Sample	K1902232-002.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
49	Sample	K1902232-003.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
50	Sample	K1902121-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
51	Sample	K1902121-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
52	Sample	K1902435-001.02	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
53	Sample	K1902435-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
54	Sample	K1902435-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
55	Sample	K1902435-004.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
56	Sample	K1902435-005.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
57	Sample	K1902435-006.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
58	Sample	MB4	CAS_salt_010711 (CAS_salt_010711)	1	False	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	False	Ready
59	Sample	K1902435-007.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
60	Sample	K1902435-007.01 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
61	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
62	Sample	K1902254-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
63	Sample	K1902254-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
64	Sample	K1902254-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
65	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
					False	

Fusion Report - 03212019

Thursday, March 21, 2019 04:58 PM

(View - Repts, Unused Repts, Meta-Data, Signature, History)
Printed on 2019/03/23 11:38 - Saturday

Report Summary Information

Company Location: Gen Chem Lab
 Schedule Name: 03212019
 Instrument Name: Fusion1
 Report Version: 1 of 1
 Report Creation by Operators (schedule version): Fusion1 (Fusion1) (v2)
 Fusion1 (Fusion1) (v3)
 Fusion1 (Fusion1) (v4)
 Comment:

Engine Version: 1.1.5.1
 Firmware Version: 1.2.0696
 Connection: RS232 COM1

Report Results

03/25/19
[Signature]

Sample Type: Clean From Schedule Version 2

Pos	Analysis Type	Sample ID	Start Time
♦ (clean)		Clean	2019/03/21 16:58

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	13.69	16.58	2.89	49.93	07:59
2	TC Clean	20.38	23.31	2.93	49.73	07:19
3	TC Clean	2.42	5.55	3.13	49.79	07:01
4	TC Clean	1.97	4.69	2.71	49.80	07:01

Sample Type: Clean From Schedule Version 3

Pos	Analysis Type	Sample ID	Start Time
♦ (clean)		Clean	2019/03/21 17:32

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	10.80	13.84	3.04	49.90	08:01
2	TC Clean	4.74	7.49	2.75	49.73	07:16
3	TC Clean	1.91	4.68	2.77	49.81	06:59
4	TC Clean	1.32	4.10	2.77	49.81	07:04

Sample Type: Clean							From Schedule Version 4
Pos	Analysis Type	Sample ID			Start Time		
◆ (clean)		Clean			2019/03/21 18:06		
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	IC Clean	1.03	3.77	2.74	49.95	08:00	
2	TC Clean	4.31	6.95	2.64	49.79	07:16	
3	TC Clean	1.72	4.56	2.84	49.85	07:02	
4	TC Clean	1.74	4.48	2.74	49.84	07:03	

Sample Type: Blank (Creating v1238)							From Schedule Version 4
Pos	Analysis Type	Sample ID			Start Time		
◆ (blank)		Reagent/Acid Blank			2019/03/21 18:41		
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	IC Clean	1.00	3.60	2.60	49.84	05:29	
2	TC Clean	4.32	7.12	2.80	49.86	07:17	
3	TC Clean	2.15	5.14	2.99	49.89	07:02	
4	TC Clean	2.00	4.74	2.74	49.91	07:02	
5	Reagent Blank	8.64	11.55	2.91	49.90	08:11	
6	Acid Blank	1.76	4.51	2.75	49.59	05:29	

Sample Type: Sample							From Schedule Version 4	
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◆ D	TOC	RB	0.6753 ppm	0.0000 ppm	0.0000%	2019/03/21 19:27		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6753	6.7532	13.68	16.57	2.90	50.05	10:32
Dilution		Blank Contribution		Method		Calibration		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Sample Type: Check Standard --> CCV 25 ppm							From Schedule Version 4
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Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
* B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.7429 ppm (PASS)	0.0000 ppm	0%	2019/03/21 19:42

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.7429	247.4293	177.42	180.64	3.22	50.08	10:34

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
* D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.2462 ppm (PASS)	0.0000 ppm	0%	2019/03/21 19:57

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.2462	2.4625	11.13	14.12	2.99	50.11	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
* 1	TOC	MB1	0.1945 ppm	0.0000 ppm	0.0000%	2019/03/21 20:12

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1945	1.9447	10.41	13.45	3.03	50.13	10:31

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
* C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.7752 ppm (PASS)	0.0000 ppm	0%	2019/03/21 20:26

Pos	Base Analysis	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
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Type										
C	TOC	25.0 ppm	1	26.7752	267.7521	191.21	194.15	2.94	50.12	10:31

Completion State **Success Action** **Method** **Calibration** **STD Conc - Pos C**
 Success - Criteria met. Do Nothing CAS_salt_010711 (v4) CAS_salt_010711 (v30) 25 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
2	TOC	ICS	0.5996 ppm	0.0000 ppm	0.0000%	2019/03/21 20:41

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5996	5.9960	13.16	16.20	3.04	50.14	10:30

Dilution **Blank Contribution** **Method** **Calibration**
 1:10 (TC) 9.0920 (IC) (v1238) CAS_salt_010711 (v4) CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
3	TOC	K1902074-001.03 doc	1.8667 ppm	0.0140 ppm	0.7500%	2019/03/21 20:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.8766	18.7657	21.83	24.73	2.90	50.18	10:27
2	TOC	1.8568	18.5683	21.70	24.80	3.10	50.17	10:27

Dilution **Blank Contribution** **Method** **Calibration**
 1:10 (TC) 9.0920 (IC) (v1238) CAS_salt_010711 (v4) CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
4	TOC	K1902074-002.03 doc	1.9020 ppm	0.0253 ppm	1.3300%	2019/03/21 21:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.8841	18.8408	21.88	24.73	2.85	50.15	10:26
2	TOC	1.9199	19.1988	22.12	24.96	2.84	50.11	10:30

Dilution **Blank Contribution** **Method** **Calibration**
 1:10 (TC) 9.0920 (IC) (v1238) CAS_salt_010711 (v4) CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
5	TOC	K1902102-001.01 doc	77.6461 ppm	1.1244 ppm	1.4500%	2019/03/21 21:52

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	76.8510	768.5100	530.75	533.65	2.90	50.09	10:28
2	TOC	78.4412	784.4117	541.55	544.57	3.03	50.08	10:32

Dilution **Blank Contribution** **Method** **Calibration**
 1:10 (TC) 9.0920 (IC) CAS_salt_010711 CAS_salt_010711

(v1238) (v4) (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
6	TOC	K1902102-001.01 ms doc	99.6155 ppm	0.0000 ppm	0.0000%	2019/03/21 22:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	99.6155	996.1549	685.28	688.33	3.06	50.06	10:33

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
7	TOC	RB	0.8853 ppm	0.0000 ppm	0.0000%	2019/03/21 22:34

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.8853	8.8525	15.10	18.27	3.17	50.02	10:33

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
8	TOC	FB	0.3991 ppm	0.0000 ppm	0.0000%	2019/03/21 22:49

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3991	3.9909	11.80	14.63	2.83	50.08	10:32

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
9	TOC	K1902047-001.01	3.1747 ppm	0.0064 ppm	0.2000%	2019/03/21 23:03

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.1792	31.7917	30.67	33.69	3.02	50.09	10:26
2	TOC	3.1702	31.7018	30.61	33.43	2.82	50.08	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.7049 ppm (PASS)	0.0000 ppm	0%	2019/03/21 23:31

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.7049	247.0492	177.16	180.15	2.99	50.08	10:32
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>		<u>Calibration</u>		<u>STD Conc - Pos B</u>		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		50 ppmC		

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
◊	D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.2679 ppm (PASS)	0.0000 ppm	0%	2019/03/21 23:46
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.2679	2.6790	11.28	14.33	3.04	50.09	10:32
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>		<u>Calibration</u>		<u>STD Conc - Pos D</u>		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		0 ppmC		

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◊	10	TOC	K1901927-005.03	46.1675 ppm	0.3093 ppm	0.6700%	2019/03/22 00:01	
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	45.9488	459.4884	320.99	323.90	2.91	50.06	10:30
2	TOC	46.3862	463.8624	323.96	326.76	2.80	50.07	10:28
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◊	11	TOC	RB	0.6373 ppm	0.1548 ppm	24.2900%	2019/03/22 00:29	
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7468	7.4677	14.16	17.16	3.00	50.07	10:28
2	TOC	0.5279	5.2785	12.68	15.49	2.81	50.05	10:25
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◊	12	TOC	K1902389-001.01	3.5762 ppm	0.0031 ppm	0.0900%	2019/03/22 00:57	

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.5740	35.7399	33.35	36.29	2.94	50.02	10:24
2	TOC	3.5784	35.7841	33.38	36.25	2.87	50.02	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
13	TOC	K1902389-001.01 ms	31.2217 ppm	0.0000 ppm	0.0000%	2019/03/22 01:25

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	31.2217	312.2170	221.02	223.80	2.78	50.00	10:31

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
14	TOC	RB	0.3757 ppm	0.0000 ppm	0.0000%	2019/03/22 01:40

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3757	3.7567	11.64	14.47	2.82	50.01	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
15	TOC	K1902389-002.01	27.6512 ppm	0.5067 ppm	1.8300%	2019/03/22 01:54

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	27.2930	272.9296	194.36	197.15	2.80	50.00	10:27
2	TOC	28.0095	280.0953	199.22	201.90	2.69	49.98	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
16	TOC	K1902389-003.01	28.1127 ppm	0.0750 ppm	0.2700%	2019/03/22 02:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	28.1657	281.6569	200.28	203.19	2.91	49.97	10:27
2	TOC	28.0596	280.5962	199.56	202.41	2.85	49.98	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Analysis	Std. Dev.

Pos	Type	Sample ID	Result (ppmC)	(ppmC)	RSD	Start Time
17	TOC	K1902153-001.01 100x	2.1834 ppm	0.3944 ppm	18.0600%	2019/03/22 02:50

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.4623	24.6231	25.81	28.60	2.80	49.95	10:30
2	TOC	1.9046	19.0456	22.02	24.86	2.84	49.96	10:28

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
18	TOC	K1902153-002.01 100x	3.3888 ppm	0.0121 ppm	0.3600%	2019/03/22 03:19

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.3974	33.9735	32.15	35.09	2.93	50.01	10:28
2	TOC	3.3803	33.8026	32.04	34.91	2.87	49.96	10:29

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
19	TOC	K1902153-003.01 100x	1.4009 ppm	0.0500 ppm	3.5700%	2019/03/22 03:47

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.4362	14.3623	18.84	21.63	2.79	49.95	10:28
2	TOC	1.3655	13.6551	18.36	21.10	2.74	49.93	10:26

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.1491 ppm (PASS)	0.0000 ppm	0%	2019/03/22 04:15

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.1491	251.4909	180.17	182.80	2.63	49.90	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.1491 ppm (PASS)	0.0000 ppm	0%	2019/03/22 04:15

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.1491	251.4909	180.17	182.80	2.63	49.90	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.2848 ppm (PASS)	0.0000 ppm	0%	2019/03/22 04:30

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.2848	2.8484	11.40	14.09	2.70	49.89	10:31

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 20	TOC	MB2	0.2384 ppm	0.0000 ppm	0.0000%	2019/03/22 04:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2384	2.3837	10.71	13.39	2.68	49.91	10:32

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	27.0967 ppm (PASS)	0.0000 ppm	0%	2019/03/22 04:59

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	27.0967	270.9666	193.39	196.26	2.87	49.90	10:29

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos C 25 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 21	TOC	K1902346-001.01	3.1362 ppm	0.0803 ppm	2.5600%	2019/03/22 05:14

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.1930	31.9302	30.77	33.56	2.79	49.89	10:30

2	TOC	3.0794	30.7944	30.00	32.92	2.92	49.86	10:26
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
22	TOC	K1902102-001.02	83.5303 ppm	0.1033 ppm	0.1200%	2019/03/22 05:42		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	83.4573	834.5727	575.59	578.46	2.87	49.88	10:28
2	TOC	83.6034	836.0342	576.59	579.49	2.90	49.93	10:28
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
23	TOC	K1902127-001.12	15.8880 ppm	0.1089 ppm	0.6900%	2019/03/22 06:10		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	15.9649	159.6493	117.46	120.43	2.96	49.89	10:27
2	TOC	15.8110	158.1098	116.42	119.03	2.61	49.79	10:30
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
24	TOC	K1902127-002.12 4x	7.2203 ppm	0.1077 ppm	1.4900%	2019/03/22 06:38		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.2965	72.9647	58.62	61.29	2.67	49.75	10:31
2	TOC	7.1441	71.4414	57.59	60.28	2.69	49.70	10:28
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
25	TOC	K1902127-003.12	7.8209 ppm	0.0934 ppm	1.1900%	2019/03/22 07:06		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.8869	78.8693	62.63	65.27	2.64	49.66	10:27
2	TOC	7.7548	77.5479	61.73	64.50	2.77	49.65	10:30
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		

♦	26	TOC	K1902127-004.12 4x	4.6989 ppm	0.1032 ppm	2.2000%	2019/03/22 07:34		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	4.7718	47.7185	41.48	44.27	2.79	49.59	10:29	
2	TOC	4.6259	46.2585	40.49	43.16	2.66	49.58	10:28	
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>				
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)				
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
♦	27	TOC	K1902147-001.01 100x	0.4629 ppm	0.0021 ppm	0.4500%	2019/03/22 08:02		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	0.4644	4.6436	12.24	14.96	2.71	49.56	10:28	
2	TOC	0.4614	4.6141	12.22	14.96	2.74	49.50	10:30	
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>				
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)				
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
♦	28	TOC	K1902147-002.01	1.1631 ppm	0.0267 ppm	2.2900%	2019/03/22 08:30		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	1.1820	11.8195	17.11	19.97	2.86	49.64	10:31	
2	TOC	1.1442	11.4424	16.86	19.60	2.74	49.76	10:28	
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>				
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)				

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
♦	B	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.6255 ppm (PASS)	0.0000 ppm	0%	2019/03/22 08:58	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.6255	246.2551	176.62	179.28	2.66	49.91	10:30
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>	<u>Calibration</u>		<u>STD Conc - Pos B</u>			
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)		50 ppmC			

Sample Type: Check Standard --> CCB From Schedule Version 4

Concentration	Min / Max

Pos	BAT	(ppm)	Dil	Sample ID	(% dev)	Result	Std. Dev.	RSD	Start Time	
♦	D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0686 ppm (PASS)	0.0000 ppm	0%	2019/03/22 09:13

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0686	0.6858	9.93	12.79	2.86	49.93	10:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	29	TOC	K1902168-001.04	2.0571 ppm	0.0518 ppm	2.5200%	2019/03/22 09:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.0937	20.9372	23.30	25.96	2.66	49.96	10:30
2	TOC	2.0205	20.2050	22.81	25.66	2.86	50.00	10:27

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	30	TOC	K1902168-002.04	1.0525 ppm	0.0826 ppm	7.8500%	2019/03/22 09:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.1109	11.1094	16.63	19.46	2.83	50.01	10:25
2	TOC	0.9941	9.9412	15.84	18.81	2.97	50.01	10:28

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	31	TOC	K1902169-001.24	14.4701 ppm	0.0860 ppm	0.5900%	2019/03/22 10:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	14.5309	145.3092	107.73	110.45	2.72	49.97	10:29
2	TOC	14.4092	144.0923	106.90	109.89	2.99	49.97	10:30

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	32	TOC	K1902169-002.24	17.8450 ppm	0.1318 ppm	0.7400%	2019/03/22 10:52

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	17.7518	177.5178	129.59	132.45	2.86	49.97	10:26
2	TOC	17.9381	179.3814	130.85	133.67	2.81	49.97	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
33	TOC	K1902169-003.24	18.8828 ppm	0.0274 ppm	0.1500%	2019/03/22 11:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	18.9022	189.0220	137.40	140.21	2.81	49.95	10:32
2	TOC	18.8635	188.6346	137.14	140.09	2.96	49.96	10:25

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
34	TOC	K1902169-004.24	9.3550 ppm	0.0898 ppm	0.9600%	2019/03/22 11:48

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	9.4185	94.1847	73.02	75.99	2.97	49.97	10:27
2	TOC	9.2915	92.9148	72.16	75.02	2.86	49.97	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
35	TOC	K1902169-005.24	11.8106 ppm	0.0557 ppm	0.4700%	2019/03/22 12:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	11.8500	118.4999	89.53	92.49	2.96	49.98	10:28
2	TOC	11.7712	117.7117	88.99	92.00	3.01	49.97	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
36	TOC	K1902169-006.24	11.8325 ppm	0.1654 ppm	1.4000%	2019/03/22 12:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	11.9494	119.4943	90.20	93.00	2.79	49.99	10:29
2	TOC	11.7155	117.1549	88.62	91.50	2.88	49.96	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC)
Method CAS_salt_010711
Calibration CAS_salt_010711

		(v1238)	(v4)	(v30)				
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
37	TOC	K1902169-007.24	12.8921 ppm	0.2771 ppm	2.1500%	2019/03/22 13:12		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	12.6962	126.9620	95.27	98.09	2.82	50.01	10:30
2	TOC	13.0881	130.8807	97.93	100.84	2.91	50.03	10:27
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
38	TOC	K1902169-008.24	15.4016 ppm	0.2088 ppm	1.3600%	2019/03/22 13:41		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	15.5492	155.4920	114.64	117.52	2.88	50.05	10:30
2	TOC	15.2540	152.5397	112.64	115.67	3.04	50.06	10:25
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.4810 ppm (PASS)	0.0000 ppm	0%	2019/03/22 14:09	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.4810	244.8099	175.64	178.60	2.96	50.07	10:31
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos B</u>				
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC				

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0485 ppm (PASS)	0.0000 ppm	0%	2019/03/22 14:23	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0485	0.4854	9.79	12.80	3.01	50.09	10:33

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos D</u>
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
39	TOC	MB3	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/22 14:38

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.49	11.48	2.99	50.09	10:29

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.3372 ppm (PASS)	0.0000 ppm	0%	2019/03/22 14:53

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.3372	263.3722	188.24	191.16	2.92	50.10	10:32

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos C</u>
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	25 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
40	TOC	K1902230-001.05	2.0965 ppm	0.0063 ppm	0.3000%	2019/03/22 15:07

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.1009	21.0094	23.35	26.30	2.94	50.08	10:27
2	TOC	2.0921	20.9210	23.29	26.37	3.08	50.08	10:27

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
41	TOC	K1902230-002.05	0.9090 ppm	0.0394 ppm	4.3300%	2019/03/22 15:35

Rep	Base	ppm	µg	Adjusted	NDIR (Abs)	Baseline	Pressure	Run

#	Analysis Type			(Abs)		(Abs)	(psig)	Time
1	TOC	0.9368	9.3681	15.45	18.19	2.74	50.11	10:24
2	TOC	0.8811	8.8113	15.07	17.96	2.89	50.10	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
42	TOC	K1902230-003.05	1.0304 ppm	0.0033 ppm	0.3200%	2019/03/22 16:03

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.0327	10.3272	16.10	18.96	2.86	50.11	10:29
2	TOC	1.0280	10.2800	16.07	18.99	2.92	50.10	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
43	TOC	K1902230-004.05	1.4208 ppm	0.0626 ppm	4.4100%	2019/03/22 16:32

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.4651	14.6510	19.04	22.03	2.99	50.09	10:29
2	TOC	1.3766	13.7656	18.44	21.48	3.04	50.06	10:23

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
44	TOC	K1902230-005.05	0.5452 ppm	0.0150 ppm	2.7500%	2019/03/22 17:00

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5558	5.5584	12.86	15.72	2.85	50.11	10:28
2	TOC	0.5346	5.3463	12.72	15.64	2.92	50.10	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
45	TOC	K1902232-001.12	5.9922 ppm	0.0482 ppm	0.8000%	2019/03/22 17:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.9581	59.5807	49.54	52.48	2.94	50.09	10:26
2	TOC	6.0263	60.2628	50.00	53.07	3.07	50.11	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	33.5471	335.4714	236.81	239.79	2.98	50.10	10:32

Dilution 1:10 **Blank Contribution** (TC) 9.0920 (IC) (v1238) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
47	TOC	RB	0.0443 ppm	0.0000 ppm	0.0000%	2019/03/22 18:11

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0443	0.4435	9.39	12.26	2.87	50.11	10:29

Dilution 1:10 **Blank Contribution** (TC) 9.0920 (IC) (v1238) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.1426 ppm (PASS)	0.0000 ppm	0%	2019/03/22 18:25

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.1426	241.4260	173.34	176.18	2.84	50.17	10:30

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30) **STD Conc - Pos B** 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/22 18:40

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	8.99	12.03	3.03	50.10	10:32

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30) **STD Conc - Pos D** 0 ppmC

Sample Type: Sample							From Schedule Version 4		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
48	TOC	K1902232-002.12	10.0441 ppm	0.0777 ppm	0.7700%	2019/03/22 18:55			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	9.9892	99.8919	76.90	79.97	3.07	50.16	10:29	
2	TOC	10.0991	100.9909	77.64	80.42	2.78	50.05	10:27	
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>				
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)				
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
49	TOC	K1902232-003.12	14.1220 ppm	0.0201 ppm	0.1400%	2019/03/22 19:23			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	14.1362	141.3625	105.05	108.04	3.00	50.06	10:26	
2	TOC	14.1078	141.0782	104.86	107.79	2.93	50.04	10:23	
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>				
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)				
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
50	TOC	K1902121-001.01	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/22 19:51			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	0.0000	0.0000	8.86	11.75	2.89	50.04	10:25	
2	TOC	0.0000	0.0000	9.03	12.04	3.01	50.03	10:29	
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>				
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)				
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
51	TOC	K1902121-002.01	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/22 20:19			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	0.0000	0.0000	8.20	11.22	3.02	50.00	10:31	
2	TOC	0.0000	0.0000	8.53	11.57	3.04	50.01	10:27	
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>				
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)				
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
52	TOC	K1902435-001.02	116.2162 ppm	5.5539 ppm	4.7800%	2019/03/22 20:47			

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	120.1434	1201.4336	824.62	827.61	2.99	49.98	10:28
2	TOC	112.2890	1122.8899	771.30	774.91	3.61	49.98	10:24

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
53	TOC	K1902435-002.01	4.2973 ppm	1.4335 ppm	33.3600%	2019/03/22 21:15

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.3109	53.1089	45.14	48.45	3.31	49.96	10:25
2	TOC	3.2836	32.8362	31.38	34.42	3.04	49.97	10:25

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
54	TOC	K1902435-003.01	3.2785 ppm	0.2293 ppm	6.9900%	2019/03/22 21:43

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.4407	34.4066	32.45	35.43	2.98	49.95	10:30
2	TOC	3.1164	31.1641	30.25	33.19	2.94	49.96	10:29

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
55	TOC	K1902435-004.01	5.1276 ppm	0.2636 ppm	5.1400%	2019/03/22 22:11

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.3140	53.1399	45.16	48.09	2.93	49.95	10:26
2	TOC	4.9413	49.4127	42.63	45.55	2.91	49.94	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
56	TOC	K1902435-005.01	132.6232 ppm	6.5383 ppm	4.9300%	2019/03/22 22:39

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	137.2465	1372.4649	940.71	943.70	2.99	49.92	10:25
2	TOC	127.9999	1279.9994	877.95	881.72	3.77	49.99	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC)
Method CAS_salt_010711
Calibration CAS_salt_010711

		(v1238)	(v4)	(v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
57	TOC	K1902435-006.01	122.9733 ppm	6.3170 ppm	5.1400%	2019/03/22 23:07

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	118.5065	1185.0649	813.51	817.08	3.57	50.07	10:26
2	TOC	127.4401	1274.4013	874.15	878.10	3.95	50.09	10:27

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	27.6282 ppm (PASS)	0.0000 ppm	0%	2019/03/22 23:35

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	27.6282	276.2819	197.00	200.68	3.68	50.18	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	1.1283 ppm (PASS)	0.0000 ppm	0%	2019/03/22 23:50

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	1.1283	11.2825	17.12	20.21	3.09	50.20	10:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
59	TOC	K1902435-007.01	12.6387 ppm	0.1397 ppm	1.1100%	2019/03/23 00:05

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	12.7374	127.3745	95.55	98.46	2.91	50.20	10:30
2	TOC	12.5399	125.3989	94.21	97.27	3.06	50.20	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
60	TOC	K1902435-007.01 ms	33.4785 ppm	0.0000 ppm	0.0000%	2019/03/23 00:33

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	33.4785	334.7849	236.34	239.34	3.00	50.22	10:33

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
61	TOC	RB	0.6159 ppm	0.0000 ppm	0.0000%	2019/03/23 00:48

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6159	6.1595	13.27	16.42	3.14	50.22	10:33

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
62	TOC	K1902254-001.01	6.6998 ppm	0.1143 ppm	1.7100%	2019/03/23 01:02

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.7806	67.8056	55.12	58.04	2.92	50.23	10:28
2	TOC	6.6189	66.1895	54.02	57.06	3.04	50.22	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
63	TOC	K1902254-002.01	18.5047 ppm	0.1440 ppm	0.7800%	2019/03/23 01:30

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	18.6065	186.0653	135.39	138.31	2.92	50.23	10:30
2	TOC	18.4029	184.0293	134.01	137.12	3.11	50.22	10:29

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Analysis	Std. Dev.

Pos	Type	Sample ID	Result (ppmC)	(ppmC)	RSD	Start Time
64	TOC	K1902254-003.01	22.2028 ppm	0.0577 ppm	0.2600%	2019/03/23 01:58

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	22.2436	222.4357	160.08	163.10	3.02	50.23	10:28
2	TOC	22.1620	221.6195	159.53	162.65	3.12	50.23	10:25

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
65	TOC	RB	0.0278 ppm	0.0393 ppm	141.4200%	2019/03/23 02:27

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0555	0.5554	9.47	12.55	3.08	50.21	10:29
2	TOC	0.0000	0.0000	8.79	11.75	2.96	50.21	10:26

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.0426 ppm (PASS)	0.0000 ppm	0%	2019/03/23 02:55

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.0426	240.4257	172.66	175.53	2.87	50.20	10:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/23 03:09

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	9.20	12.29	3.09	50.20	10:29

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria	Do Nothing	CAS_salt_010711	CAS_salt_010711	0 ppmC

met.

(v4)

(v30)

Meta Data Used in this Report

Blanks

Version	Reagent (Abs)	Acid (Abs)	DI IC (Abs)	DI TC (Abs)	DI TOC (Abs)	Save Time	Operator
v1237	1.0520	0.7490	0.0000	0.0000	0.0000	2019/03/19 18:04	Fusion1 (Fusion1)
v1238	2.8787	1.7600	0.0000	0.0000	0.0000	2019/03/21 19:27	Fusion1 (Fusion1)

Calibrations

Name: CAS_salt_010711 (TOC)

Version: v30
 Calibration curve formula: TOC: $y = 6.788x + 9.463$
 Ver Creation: 2019/03/05 17:42
 r² value: TOC: $r^2 = 0.99963$
 Comment:
 Operator: Fusion1 (Fusion1)
 Basic Analysis Type: TOC

Basic Analysis Type: TOC

Sample ID	Y Raw Value	X Expected	Message	End Time
DI Water	7.8970	0.0000		2019/03/05 16:15
0.500 ppm	11.5280	0.5000		2019/03/05 16:29
1.0 ppm	14.9760	1.0000		2019/03/05 16:44
5.0 ppm	43.6500	5.0000		2019/03/05 16:58
10 ppm	79.6020	10.0000		2019/03/05 17:12
25 ppm	183.3580	25.0000		2019/03/05 17:26
50 ppm	346.3230	50.0000		2019/03/05 17:40

Methods

Name: CAS_salt_010711 (TOC)

Version: v4
 Operator: Fusion1 (Fusion1)
 Ver Creation: 2019/02/21 17:57
 Comment:

Parameter	Value	Advanced Parameter	Value
SampleVolume	10.0 mL	NeedleRinseVolume	5.0 ml
Dilution	1:10	VialPrimeVolume	2.0 ml
AcidVolume	0.5 ml	ICSamplePrimeVolume	2.0 ml

ReagentVolume	2.0 ml	ICSpurgeRinseVolume	12.0 ml
UVReactorPrerinse	Off	BaselineStabilizeTime	0.70 min
UVReactorPrerinseVolume	5.0	DetectorPressureFlow	150 ml/min
NumberOfUVReactorPrerinses	1	SyringeSpeedWaste	10
ICSpurgeTime	1.00 mins	SyringeSpeedAcid	7
DetectorSweepFlow	500 ml/min	SyringeSpeedReagent	7
PreSpurgeTime	2.00 mins	SyringeSpeedDIWater	7
SystemFlow	500 ml/min	NDIRPressurization	60 psig
		SyringeSpeedSampleDispense	5
		SyringeSpeedSampleAspirate	4
		SyringeSpeedUVDispense	5
		SyringeSpeedUVAspirate	5
		SyringeSpeedICDispense	5
		SyringeSpeedICAspirate	5
		NDIRPressureStabilize	1.75 min
		SampleMixing	Off
		SampleMixingCycles	1
		SampleMixingVolume	10.0
		LowLevelFilterNDIR	Off

Acceptance / Approval

Electronic Signatures

Report Version	User Name	Acceptance	Reason	Date
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Report History

Report History

Report Version	User Name	System Reason	User Reason	Date
1	Fusion1 (Fusion1)	Schedule completed	Schedule completed	2019/03/23 03:26

StarLIMS Run: 629142, 629144, 629629146, 629149
 Analysis: TOC/DOC
 Method: 9060, 415.1, SM 5310 C, 9060A

CCV: 11-GEN-05-76B 50 ppm LCS: 11-GEN-05-74L 25.0 ppm

ICAL Date: 3/6/19

ICAL ID: 11-GEN-05-76H

ICS ID: 11-GEN-05-74A

ICS TV: 25.0 ppm ICS % R = 2

Spike ID: 11-GEN-05-700 0.05 ml of 5000 ppm stock ---> 10.0 ml = 25.0 ppm x dilution factor

Sodium Persulfate: 11-GEN-05-77B

21 % H3PO4: 11-GEN-05-77A

Equipment ID: K-TOC-03

PIPETTE ID: 124276B, 129001F, N11314F, Marge

FILTER ID: NA

Analyzed By: <i>BCD</i>	Date Analyzed: ²¹⁰ <i>3/23/19</i>
Reviewed By: <i>Hampel</i>	Date Reviewed: <i>03/25/19</i>

BCD 3/23/19



Case Narrative

Method: 6850
Analysis: Perchlorate
Analysis SOP: LC-MS-CLO4
ALS WO ID(s): 1906774

Client: ALS Laboratories (Houston, TX)
Matrix: Water
ELMS Batch (HBN): 2227 (234852)

General Set Information: There was one field sample in this Work Order. The sample was analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: NA

Method QC data: The method blank (LMB 644168) was less than 1/2 the CRDL. The recovery for the LCS (644169) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1906774001 (Client ID's: LH18/24-SP650_030619_BIX). 4.0 μ l of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4. μ g/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in μ g/L. Results were calculated in μ g/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (μ g/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 644166) is reported from the analysis of the Laboratory Control Sample (LCS – 644169) at a level of 4.0 μ g/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch March 21, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 21, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1906774**

Project ID: HS19030298

Purchase Order: HS19030298

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP650_030619_BIX	1906774001	03/06/19	03/08/19	



ANALYTICAL REPORT

Workorder: 34-1906774

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP650_030619_BIX	Sampling Site: NA	Collected: 03/06/2019				
Lab ID: 1906774001	Media: 125 mL Nalgene	Received: 03/08/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2227 (HBN: 234852) Analyzed: 03/19/2019 12:35	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/20/2019 13:02	/S/ Stephen Brose 03/21/2019 11:52

Laboratory Contact Information

ALS Environmental
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Salt Lake City, Utah 84123Phone: (801) 266-7700
Email: als.lt.lab@ALSGlobal.com
Web: www.alsslc.com



ANALYTICAL REPORT

Workorder: 34-1906774

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935047

Analysis Information

Workorder: 1906774

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2227 (HBN: 234852)
Analyzed By: Thomas Bosch

Blank

LMB: 644168 Analyzed: 03/19/2019 12:22 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 644169 Analyzed: 03/19/2019 11:55 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.09	4.00	102	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1906774001 Analyzed: 03/19/2019 12:35 Dilution: 1 Units: ug/L			MS: 644170 Analyzed: 03/19/2019 12:48 Dilution: 1 Units: ug/L			MSD: 644171 Analyzed: 03/19/2019 13:02 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	ND	4.83	4	121	78.8 123.8	4.38	110	9.67	0.0 20.0

Continuing Calibration Verification

CCV: 644165 Analyzed: 03/19/2019 11:35 Units: ug/L Criteria: ± 15%				CCV: 644172 Analyzed: 03/19/2019 13:18 Units: ug/L Criteria: ± 15%		
Analyte	Result	Target	% Rec.	Result	Target	% Rec.
Perchlorate	25.0	25.0	99.8	25.0	25.0	100

Interference Check Sample

ICSA: 644167 Analyzed: 03/19/2019 12:09 Units: ug/L Criteria: ± 30%			
Analyte	Result	Target	% Rec.
Perchlorate	3.88	4.00	97.1



Quality Control Sample Batch Report

00935048

Analysis Information

Workorder: 1906774

Limits: Client SOW/Contract Specified

Preparation: NA

Analysis: EPA 6850, DoD QSM

Basis: DoD QSM

Batch: NA

Batch: ELMS/2227 (HBN: 234852)

Prepared By: NA

Analyzed By: Thomas Bosch

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/20/2019 13:14	/S/ Stephen Brose 03/21/2019 11:52

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



1906774



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Subcontract Chain of Custody

18698/#2

COC ID: 10880

SUBCONTRACT TO:

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1906774

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INVOICE INFORMATION:

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Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030298
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030298-02	LH18/24-SP650_030619_BIX	Water	06 Mar 2019 13:00
SUB_Perch-6850			21 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: [Signature]
Received By: [Signature]
Cooler ID(s): 9224

Date/Time: 3/7/19 18:00
Date/Time: 3/8/19 19:35
Temperature(s): 2

RIGHT SOLUTIONS | RIGHT PARTNER

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: ALS Houston Project/Task/Site: 1906774
 Date/Time of Receipt: 3/8/15 Number of Coolers Received: 1

Condition of Coolers: Acceptable/Unacceptable Temperature Control: Present/Not Included
 Cooler Custody Seals: Present/Absent/NA Location Temp Taken: Control/Between Samples
 Container Custody Seals: Present/Absent/NA Intact/Broken/NA
 Intact/Broken/NA
 Ice Present: Yes/No/NA Are all temperatures within project specific guidelines? Yes/No/NA
Frozen/Melted/NA VOA Headspace Present? Yes/No/NA

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9224</u>	<u>2</u> °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: Meredith Signature Printed Name Date: 3/8/15

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Part # 159469-434 R1T2 EXP 11/19

ORIGIN ID:SGRA (281) 530-5656
SHIPPING DEPT
ALS LABORATORY GROUP
10450 STANCLIFF RD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

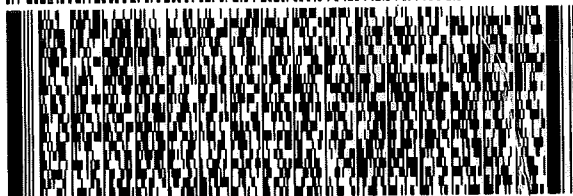
SHIP DATE: 07MAR19
ACTWGT: 13.60 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY...

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(801) 266-7700

REF: HS19030298 - RJ



FedEx
Express



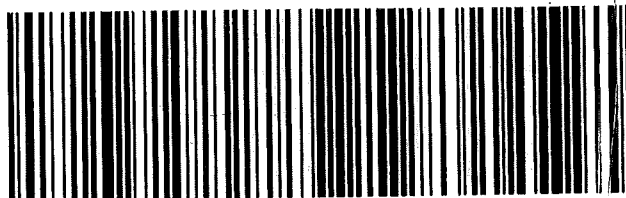
0109090811181J

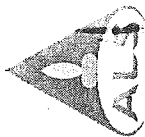
TRK# 4809 7831 4299
0201

FRI - 08 MAR 3:00P
STANDARD OVERNIGHT

AX BTFA

84123
UT-US SLC





Batch Worklist

HBN: 234852



Instrument:

Status: WP

Created: 3/19/2019 11:27

Analyst: T. Bosch

Batch: ELMS/ 2227

Rule: EPA 6850, DoD QSM Water

Workorder: 1906774 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	644165	CCV for HBN 234852 [ELMS/2227]				CCV	3		E685041C3Q	5311		3/21/2019	
2	644166	RLYS for HBN 234852 [ELMS/2227]				RLYS	3		E685041C3Q	5311		3/21/2019	
3	644167	ICS for HBN 234852 [ELMS/2227]				ICS	3		E6850.D3Q	5311		3/21/2019	
4	644168	LMB for HBN 234852 [ELMS/2227]				LMB	3		E6850Q413Q	5311		3/21/2019	
5	644169	LCS for HBN 234852 [ELMS/2227]				LCS	3		E6850Q413Q	5311		3/21/2019	
6	1906774001	LH18/24-SP650_030619_BIX				SAMPLE	3	1906774001-A	E6850Q41.3	5480	4/3/2019		3/21/2019
7	644170	LH18/24-SP650...(1906774001MS)				MS	3		E6850Q413Q	5311			3/21/2019
8	644171	LH18/24-SP65...(1906774001MSD)				MSD	3		E6850Q413Q	5311			3/21/2019
9	644172	CCV for HBN 234852 [ELMS/2227]				CCV	3		E685041C3Q	5311			3/21/2019



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1906774 (001)
 ELMS Batch/HBN ID: 2227 (234852)
 Prep Date: 03/19/2019 Analysis Date: 03/19/2019 Analyst: T. Bosch
 Analyte: ~~Perchlorate~~ Matrix: Water Method: 6850
 Sequence: \\HPCHEM\SEQUENCE\CLO4\2019\MAR\19MAR19D.s
 Reported DL: 1.0µg/L Reported LOD: 2.0µg/L Reported LOQ: 4.0µg/L

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by TNB. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 644169; Target = 4.0µg/L. ASTM type II water was used for LMB 644168.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1906774001 (Client ID's: LH18/24-SP650_030619_BIX). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alslts013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\234852-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATA\REVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 644166) is reported from the analysis of the Laboratory Control Sample (LCS – 644169) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
<u>Batch(es)/SDG: E LMS: 2227 HBN: 234852</u>		
<u>Sample Set IDs if Applicable: 1906774</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850.WKG Std 100.ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK		Description - 6850 Stock AccStd 1,000ug/mL	
Standard: 43659		Created By: Thomas Bosch	
MFG: AccuStandard		Create Date: 09/17/2018 09:09AM	
MFG Lot: 218065075		Amount: 100 mL	
Part ID: IC-PER-10X-1		Expires: 07/25/2020	
		Usable: No	
		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT			Description - 6850 Intermdt AccStd 10.ug/mL		
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK			Description - 6850 QC WKG STD 100ug/L		
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT			Description - 6850 QC Intmdt Std-QC 10ug/mL		
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK			Description: 6850 QC Stock STD 1,000ug/mL
Standard: 36748		Created By: Thomas Bosch	
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	
MFG Lot: CP-0860		Amount: 100 mL	
Part ID: ICC-013		Expires: 03/31/2020	
		Usable: Yes	
		Lab Lot: CLO4 QC STOCK	
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK			Description - Perchlorate ISTD Wrk 1,000ug/L		
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

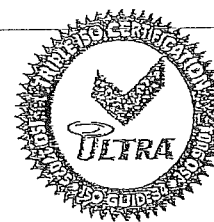
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK			Description - Perchlorate ISTD Stock
Standard: 43729		Created By: Thomas Bosch	Amount: 1 mL
MFG: Cambridge Isotope		Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026
MFG Lot: SDFF-012A		Verified By: Thomas Bosch	Usable: Yes
Part ID: OLM-7310-S		Verify Date:	Lab Lot: CLO4ISTDSTK
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:
This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:
Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:
The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:
This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:
This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:
Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

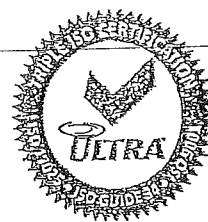
Hazards:
Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:
The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis

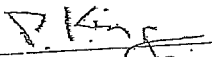


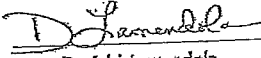
ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:
The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QA/RA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Melgan O'Leary

Melgan O'Leary, Inorganic QC Manager

For use in routine laboratory analysis.

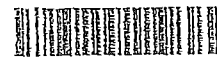


Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:

ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: NaCl^*O_4

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Ebersley

Timothy J. Ebersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 $\mu\text{g/mL}$
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 \pm 2.8 $\mu\text{g/mL}$ (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount	
*	644165	CCV@25	Vial 71	1	Control	1	1.94296e6	8.846	24.95399
*	644169	QC@4.0	Vial 72	1	Control	2	2.95659e5	8.943	4.09149
*	644167	ICS@4.0	Vial 73	1	Control	3	2.65691e5	8.627	3.88435
*	644168	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1906774001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	644170	67741MS	Vial 76	1	Sample	6	3.98555e5	8.368	4.82506
*	644171	67741SD	Vial 77	1	Sample	7	3.62343e5	8.358	4.37993
*	644172	CCV@25	Vial 71	1	Control	8	2.11382e6	8.746	24.99412

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount	
*	644165	CCV@25	Vial 71	1	Control	1	5.76297e5	8.860	24.93587
*	644169	QC@4.0	Vial 72	1	Control	2	9.68807e4	8.956	4.36015
*	644167	ICS@4.0	Vial 73	1	Control	3	8.63749e4	8.649	4.09819
*	644168	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1906774001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	644170	67741MS	Vial 76	1	Sample	6	1.30024e5	8.381	5.15091
*	644171	67741SD	Vial 77	1	Sample	7	1.14519e5	8.376	4.51535
*	644172	CCV@25	Vial 71	1	Control	8	6.19581e5	8.761	24.69614

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount	
*	644165	CCV@25	Vial 71	1	Control	1	2.36912e5	8.874	5.00000
*	644169	QC@4.0	Vial 72	1	Control	2	2.37845e5	8.962	5.00000
*	644167	ICS@4.0	Vial 73	1	Control	3	2.25720e5	8.647	5.00000
*	644168	LMB	Vial 74	1	Control	4	2.83967e5	8.871	5.00000
*	1906774001		Vial 75	1	Sample	5	2.69232e5	8.436	5.00000
*	644170	67741MS	Vial 76	1	Sample	6	2.69789e5	8.388	5.00000
*	644171	67741SD	Vial 77	1	Sample	7	2.71402e5	8.381	5.00000
*	644172	CCV@25	Vial 71	1	Control	8	2.57308e5	8.763	5.00000

*** End of Report ***

Sequence: C:\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\19MAR19D.S

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
1	Vial 71	644165	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	644169	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	644167	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	644168	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1906774001		CLO4-AQN	1	Sample	
6	Vial 76	644170	67741MS	CLO4-AQN	1	Sample	
7	Vial 77	644171	67741SD	CLO4-AQN	1	Sample	
8	Vial 71	644172	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD01.D

Sample Name: 644165 CCV@25

Injection Date: 3/19/2019 11:35:36

Seq Line: 1

Sample Name: 644165 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

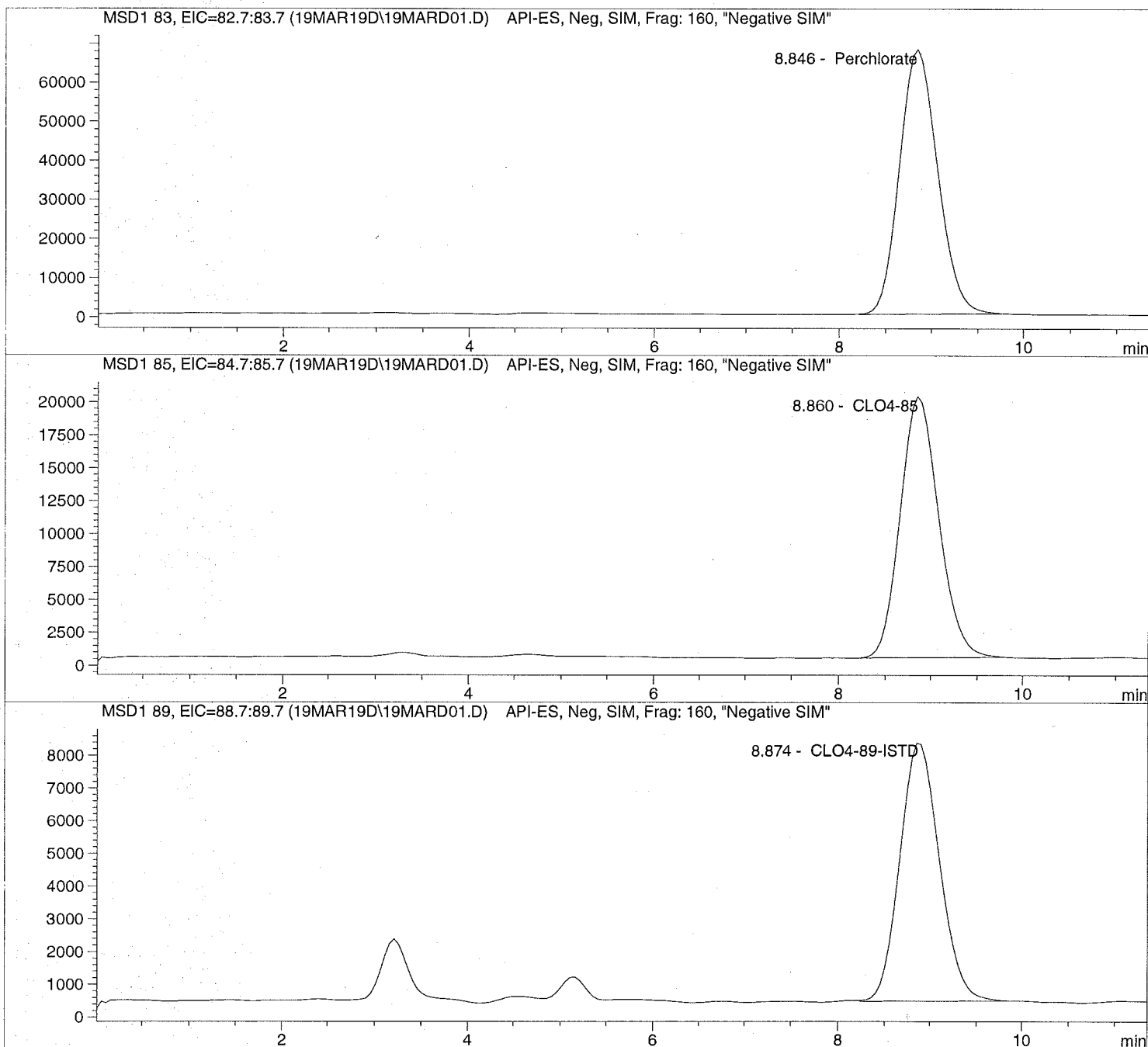
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD01.D Sample Name: 644165 CCV@25

```

=====
Injection Date: 3/19/2019 11:35:36      Seq Line: 1
Sample Name: 644165      CCV@25      Location: Vial 71
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25

```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.846	PBA	1942956.7	24.9540	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.860	PBA	576297.2	24.9359	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.874	BBA	236912.0	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD02.D

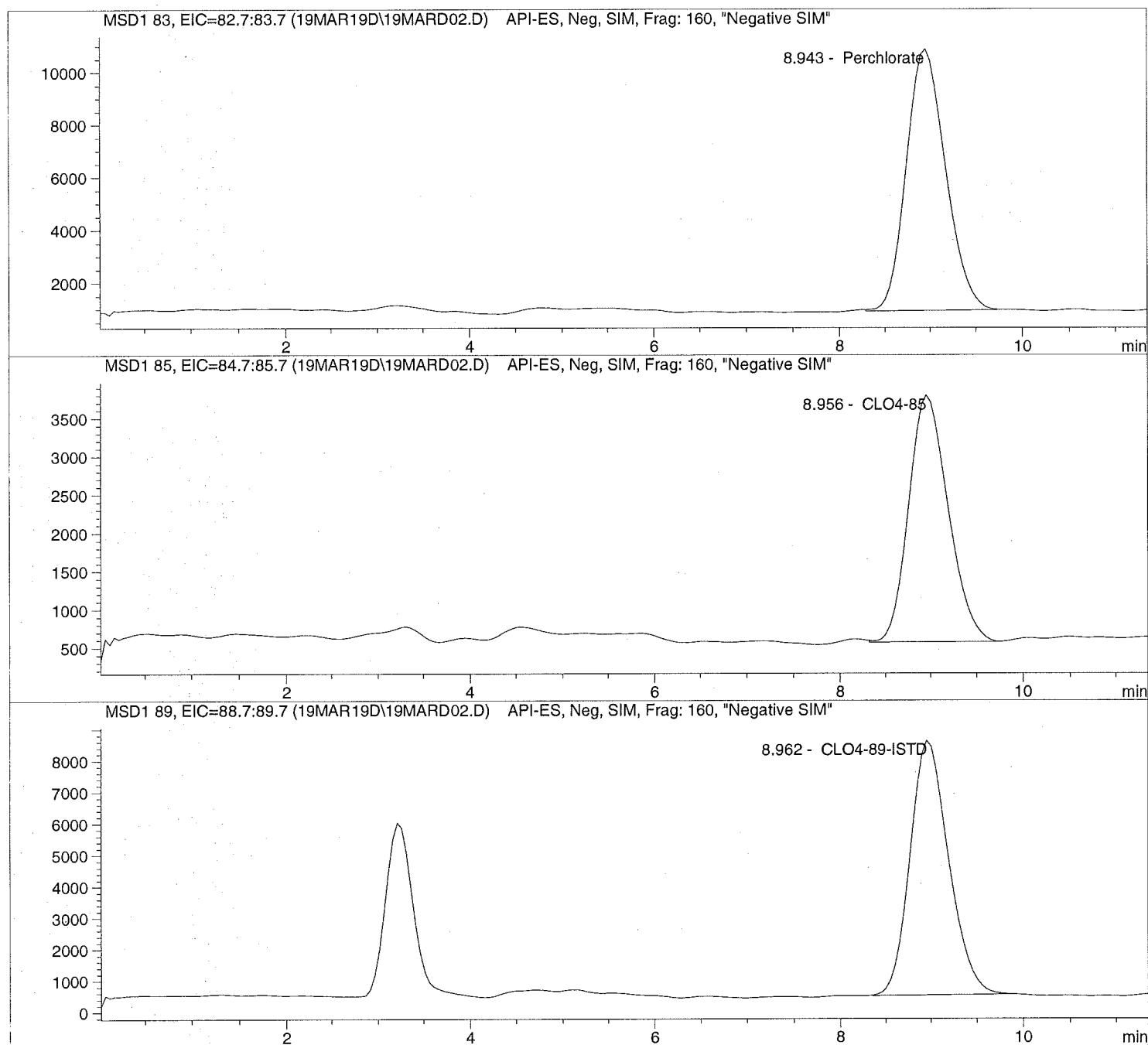
Sample Name: 644169 QC@4.0

Injection Date: 3/19/2019 11:55:42
Sample Name: 644169 QC@4.0
Acq Operator: TNB

Seq Line: 2
Location: Vial 72
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD02.D Sample Name: 644169 QC@4.0

```

=====
Injection Date: 3/19/2019 11:55:42      Seq Line: 2
Sample Name:    644169 QC@4.0           Location:  Vial 72
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  4.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.943	BBA	295659.5	4.0915	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.956	BBA	96880.7	4.3601	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.962	BBA	237845.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

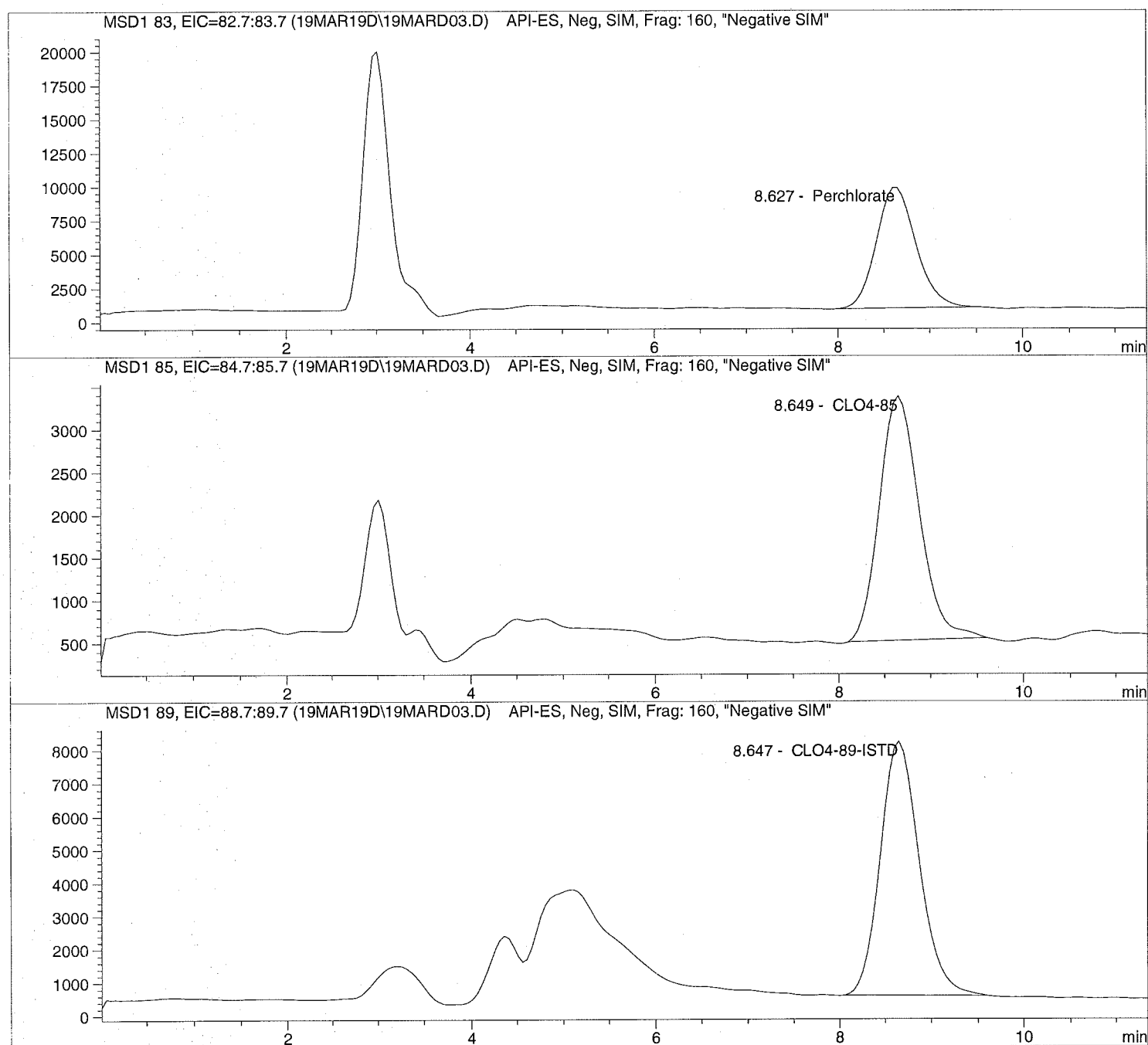
```

Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD03.D Sample Name: 644167 ICS@4.0

```
=====
Injection Date: 3/19/2019 12:09:00 Seq Line: 3
Sample Name: 644167 ICS@4.0 Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD03.D Sample Name: 644167 ICS@4.0

```
=====
Injection Date:  3/19/2019  12:09:00      Seq Line:           3
Sample Name:    644167    ICS@4.0        Location:           Vial 73
Acq Operator:   TNB                               Inj. No.:          1
                                           Inj. Vol.:         30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:38:25
```

Perchlorate analysis

=====

Sample Information

=====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  4.000
```

=====

LCMS Results

=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.627	PBA	265691.1	3.8844	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.649	PBA	86374.9	4.0982	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.647	PBA	225720.1	5.0000	CLO4-89-ISTD

=====

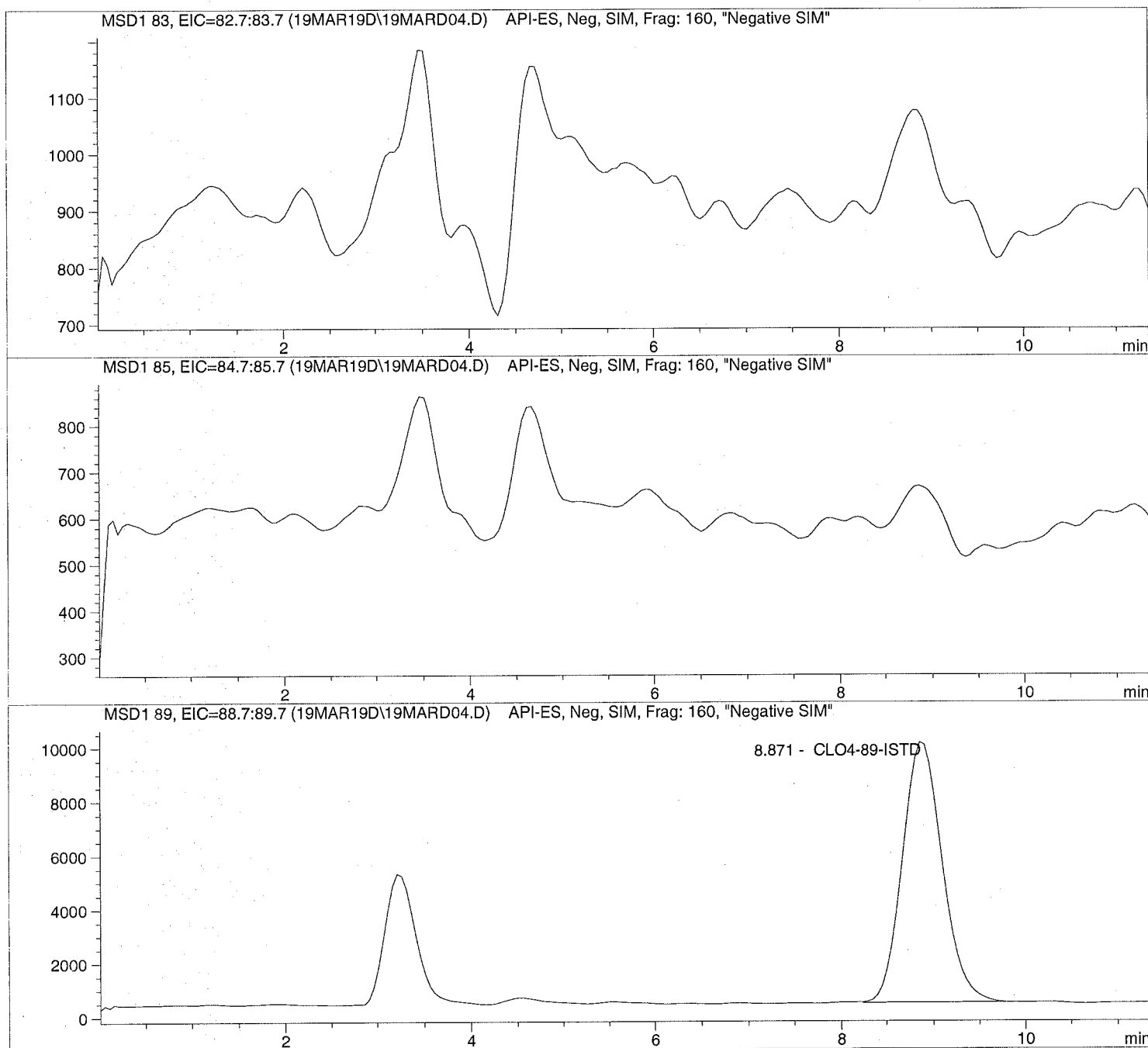
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD04.D Sample Name: 644168 LMB

```
=====
Injection Date: 3/19/2019 12:22:20      Seq Line: 4
Sample Name: 644168 LMB                 Location: Vial 74
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD05.D

Sample Name: 1906774001

Injection Date: 3/19/2019 12:35:37

Seq Line: 5

Sample Name: 1906774001

Location: Vial 75

Acq Operator: TNB

Inj. No.: 1

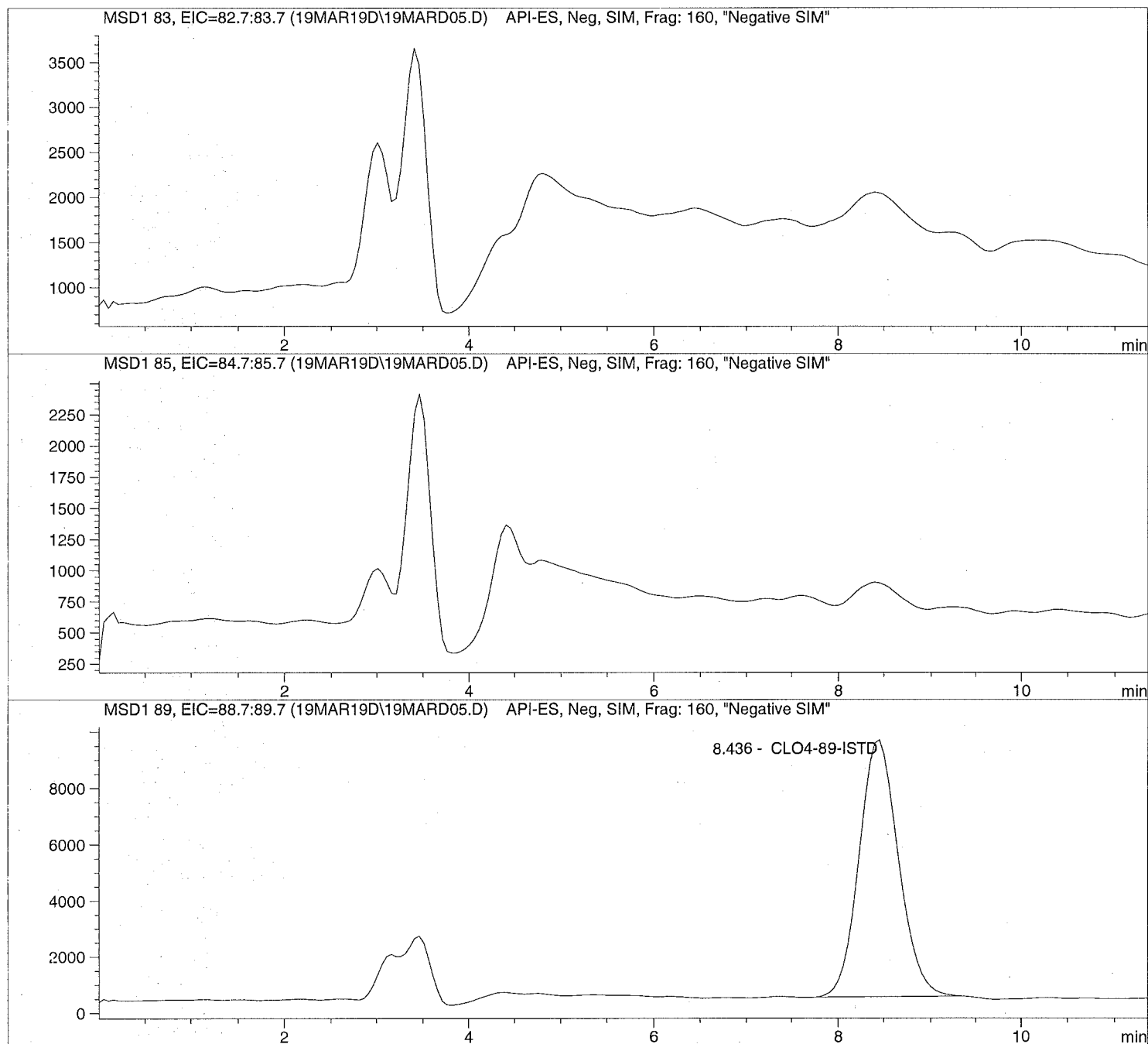
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD05.D

Sample Name: 1906774001

```

=====
Injection Date: 3/19/2019 12:35:37      Seq Line:      5
Sample Name:    1906774001              Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.436	PBA	269231.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD06.D

Sample Name: 644170 67741MS

Injection Date: 3/19/2019 12:48:55

Seq Line: 6

Sample Name: 644170 67741MS

Location: Vial 76

Acq Operator: TNB

Inj. No.: 1

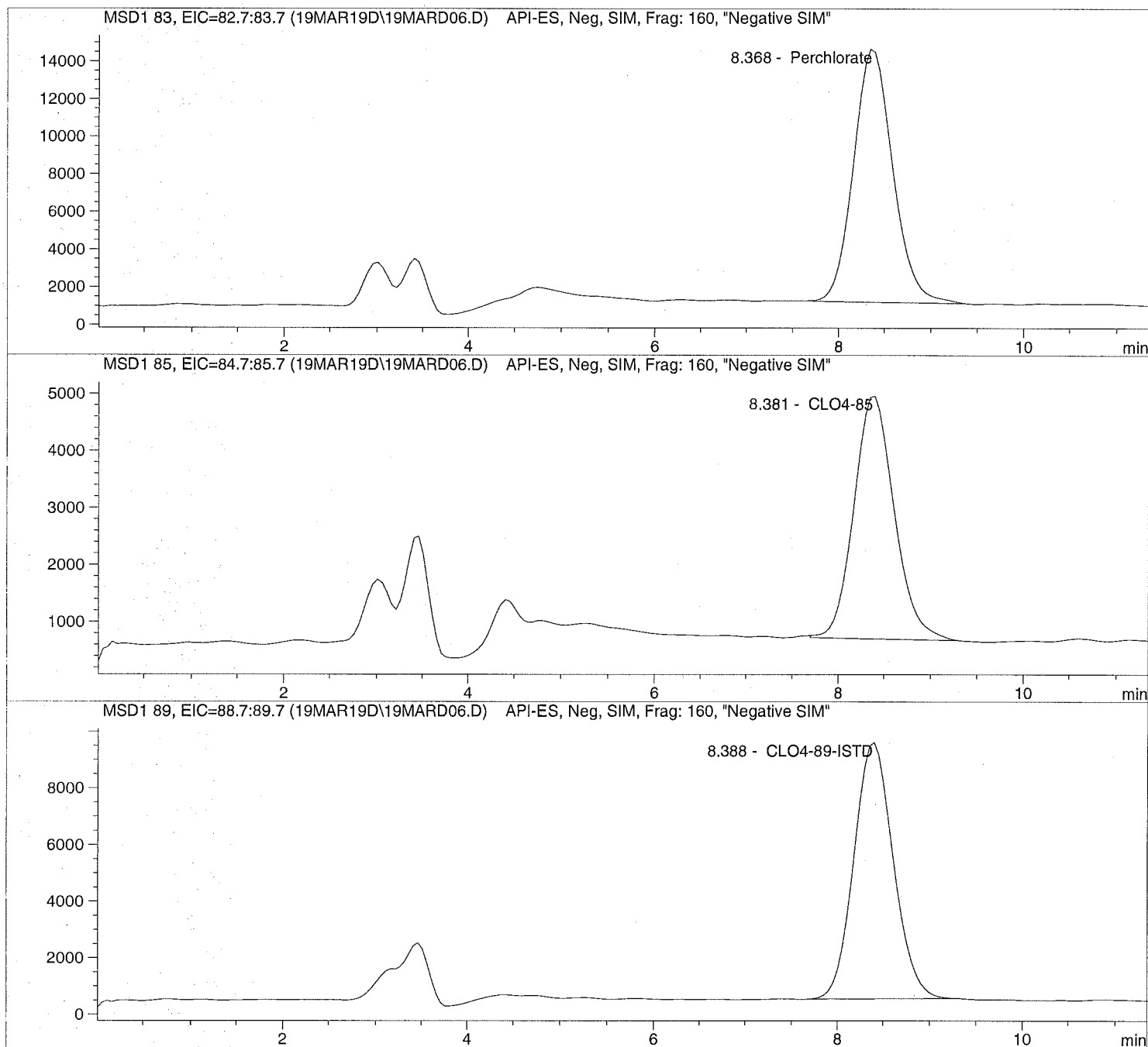
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD06.D Sample Name: 644170 67741MS

```

=====
Injection Date: 3/19/2019 12:48:55      Seq Line: 6
Sample Name:    644170 67741MS          Location:  Vial 76
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.368	BBA	398555.1	4.8251	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.381	BBA	130023.9	5.1509	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.388	BBA	269788.8	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

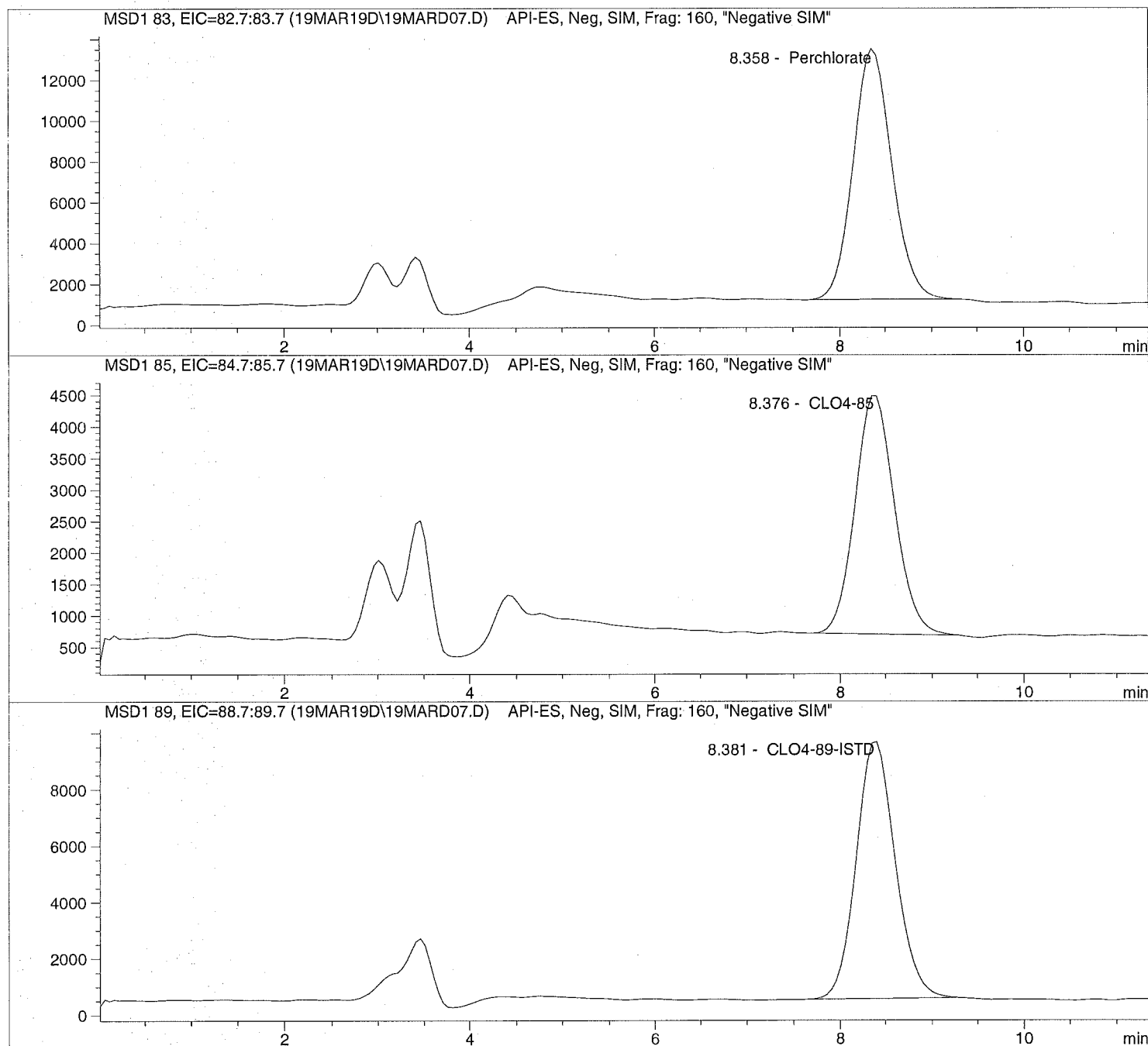
```

Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD07.D Sample Name: 644171 67741SD

```
=====
Injection Date: 3/19/2019 13:02:13      Seq Line: 7
Sample Name:    644171 67741SD          Location: Vial 77
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD07.D Sample Name: 644171 67741SD

```

=====
Injection Date: 3/19/2019 13:02:13      Seq Line:            7
Sample Name:    644171    67741SD            Location:            Vial 77
Acq Operator:   TNB                        Inj. No.:            1
                                              Inj. Vol.:            30 µl
  
```

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
 Last Changed: 3/19/2019 14:38:25

Perchlorate analysis

Sample Information

```

=====
Sorted By:            Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:          1.000000
Dilution:            1.000000
Sample Amount:        0.000
  
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.358	PBA	362342.5	4.3799	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.376	PBA	114518.6	4.5153	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.381	BBA	271402.1	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19D\19MARD08.D

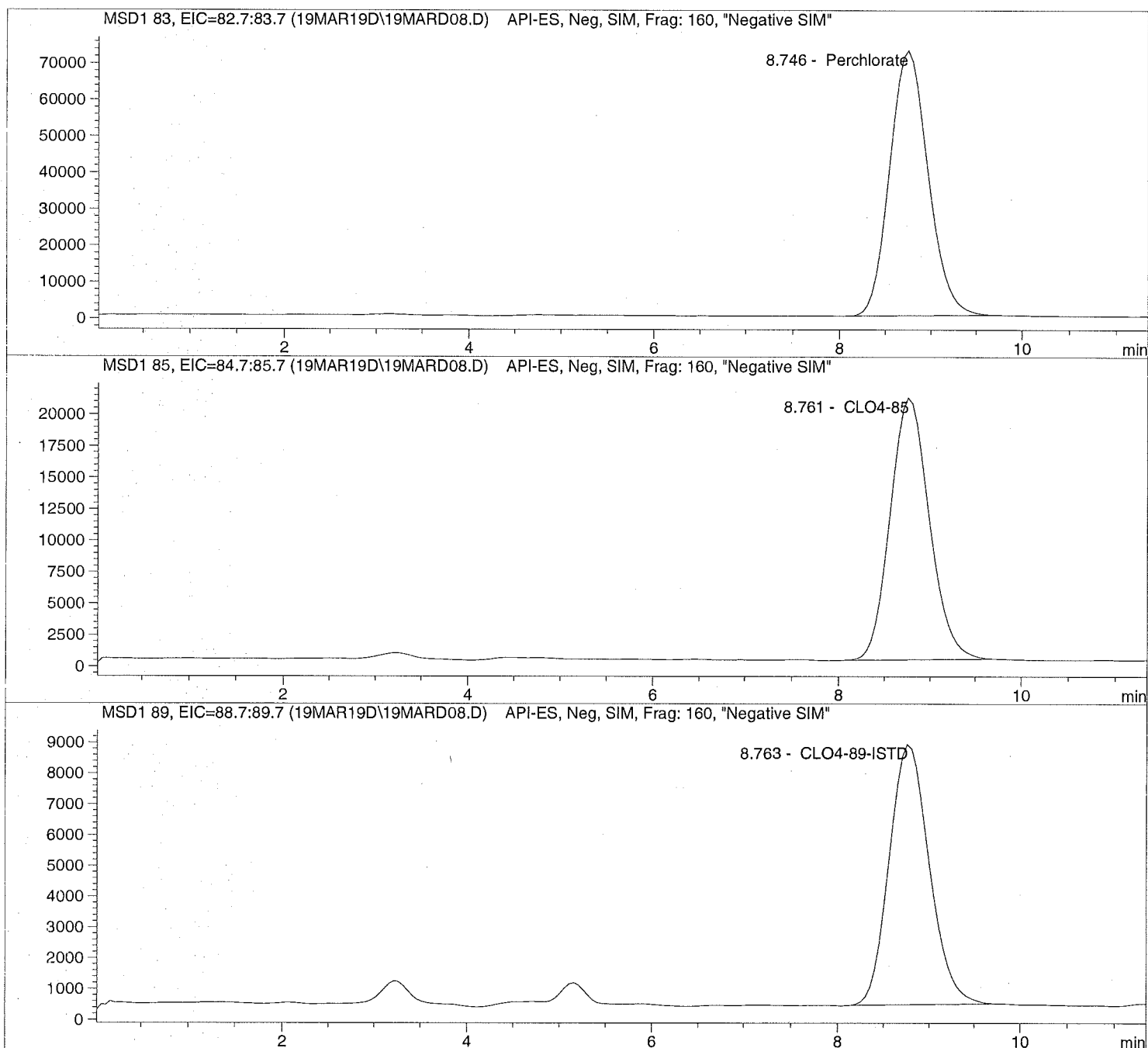
Sample Name: 644172 CCV@25

Injection Date: 3/19/2019 13:18:38
Sample Name: 644172 CCV@25
Acq Operator: TNB

Seq Line: 8
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25

Perchlorate analysis





ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
 Calibration Table
 =====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
 Based on : Peak Area

Rel. Reference Window : 20.000 %
 Abs. Reference Window : 0.000 min
 Rel. Non-ref. Window : 20.000 %
 Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
 Uncalibrated Peaks : not reported
 Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
 Origin : Ignored (some peaks differ, see below)
 Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
 Average Response : Average all calibrations
 Average Retention Time: Floating Average New 75%

Calibration Report Options :
 Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
 If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
 Signal 2: MSD1 85, EIC=84.7:85.7
 Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl	Amount	Area	Amt/Area	Ref Grp	Name
8.744	1	1	1.00000	7.76074e4	1.28854e-5	1 Perchlorate
		2	2.00000	1.35273e5	1.47849e-5	
		3	5.00000	3.37764e5	1.48033e-5	
		4	10.00000	6.83454e5	1.46316e-5	
		5	25.00000	2.08433e6	1.19943e-5	
		6	50.00000	4.13334e6	1.20968e-5	
		7	75.00000	5.99313e6	1.25143e-5	
8.755	2	1	1.00000	2.36780e4	4.22333e-5	1 CLO4-85
		2	2.00000	4.69486e4	4.25998e-5	
		3	5.00000	1.06124e5	4.71147e-5	
		4	10.00000	2.13523e5	4.68335e-5	
		5	25.00000	6.14295e5	4.06971e-5	
		6	50.00000	1.19814e6	4.17315e-5	
		7	75.00000	1.78355e6	4.20509e-5	
8.766	3	1	5.00000	2.73208e5	1.83011e-5	+I1 CLO4-89-ISTD
		2	5.00000	2.24886e5	2.22335e-5	
		3	5.00000	2.33196e5	2.14412e-5	
		4	5.00000	2.34454e5	2.13262e-5	
		5	5.00000	2.50568e5	1.99547e-5	
		6	5.00000	2.30977e5	2.16472e-5	

RetTime [min]	Lvl Sig	Amount	Area	Amt/Area	Ref Grp Name
7		5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
 Curve Type : Quadratic
 Origin : Ignored
 Calibration Level Weights:/
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333

Compound: CLO4-85

Time Window : From 6.650 min To 12.505 min
 Curve Type : Quadratic
 Origin : Ignored
 Calibration Level Weights:/
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333

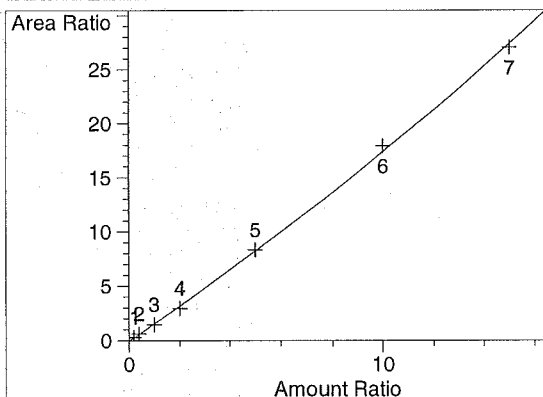
Compound: CLO4-89-ISTD

Time Window : From 6.659 min To 12.466 min
 Curve Type : Linear
 Origin : Included
 Calibration Level Weights:/
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

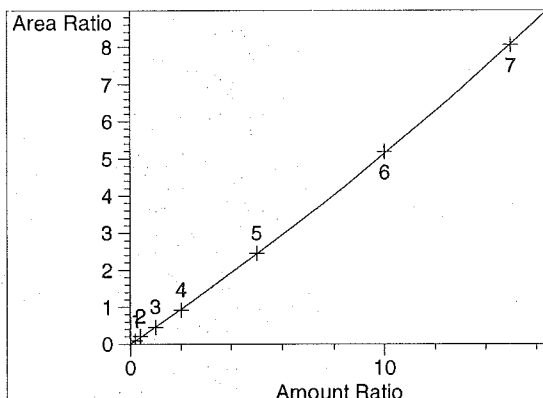
=====
 Peak Sum Table
 =====

No Entries in table

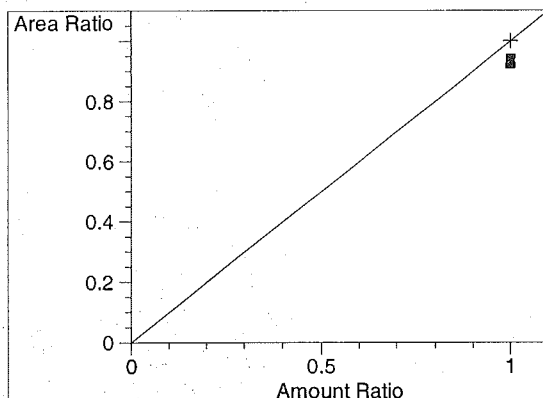
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

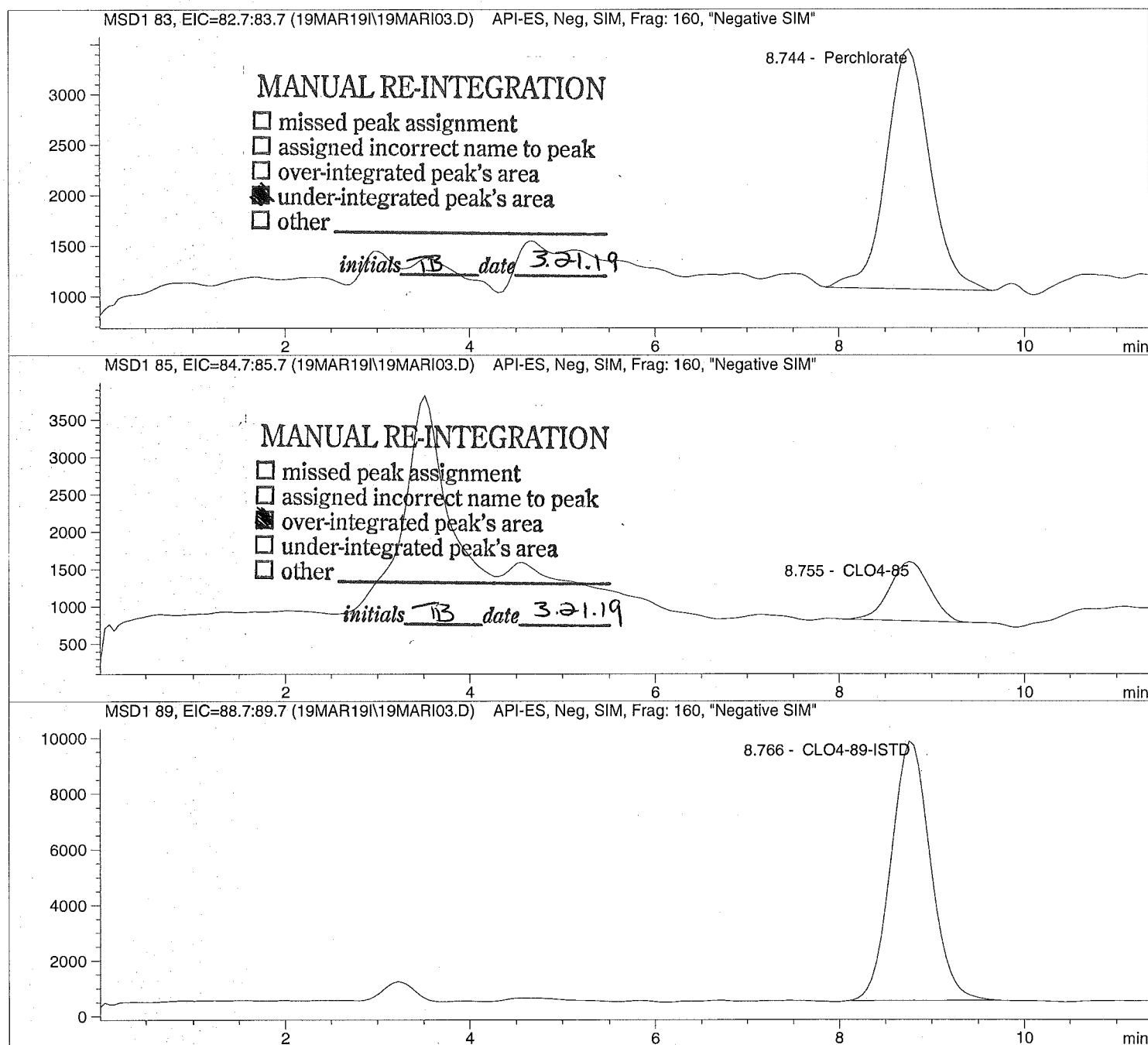
Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:          3
Sample Name:    CLO4@ 1.0ug/L           Location:          Vial 73
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D

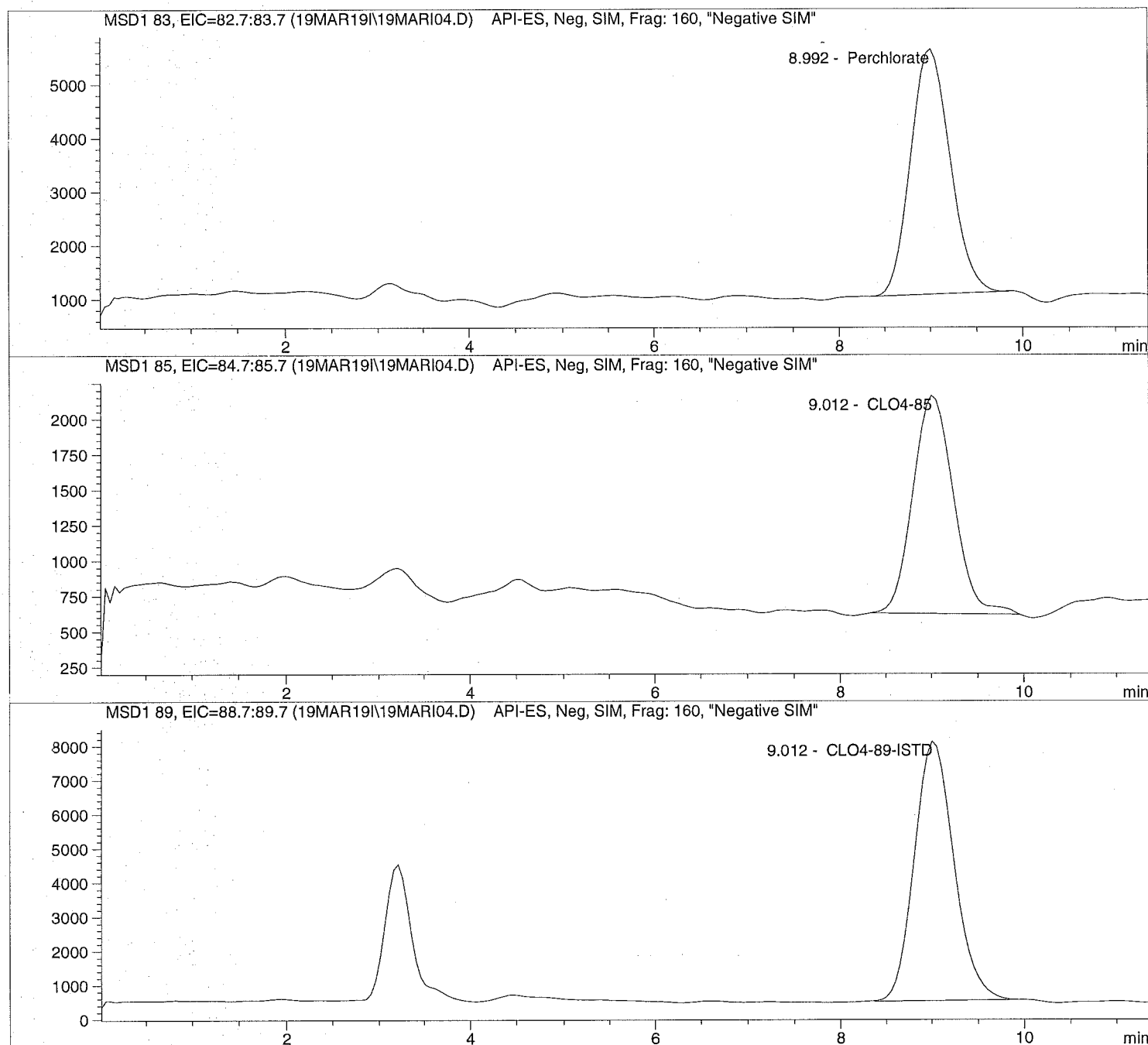
Sample Name: CLO4@ 2.0ug/L

=====
Injection Date: 3/19/2019 09:53:00
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line: 4
Sample Name:    CLO4@ 2.0ug/L          Location:  Vial 74
Acq Operator:  TNB                    Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI05.D

Sample Name: CLO4@ 5.0ug/L

Injection Date: 3/19/2019 10:06:16

Seq Line: 5

Sample Name: CLO4@ 5.0ug/L

Location: Vial 75

Acq Operator: TNB

Inj. No.: 1

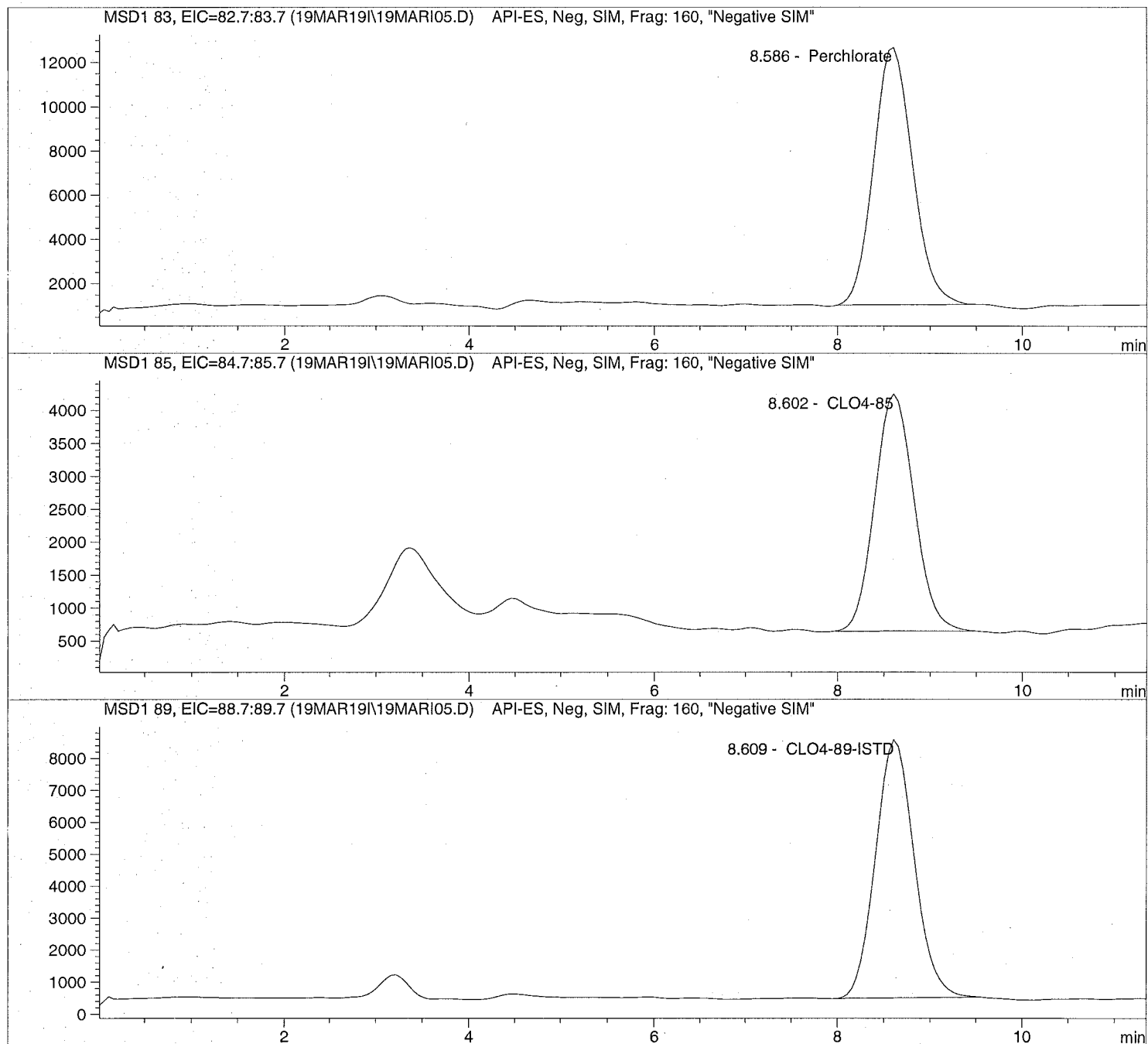
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line: 5
Sample Name: CLO4@ 5.0ug/L      Location: Vial 75
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 5.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI06.D

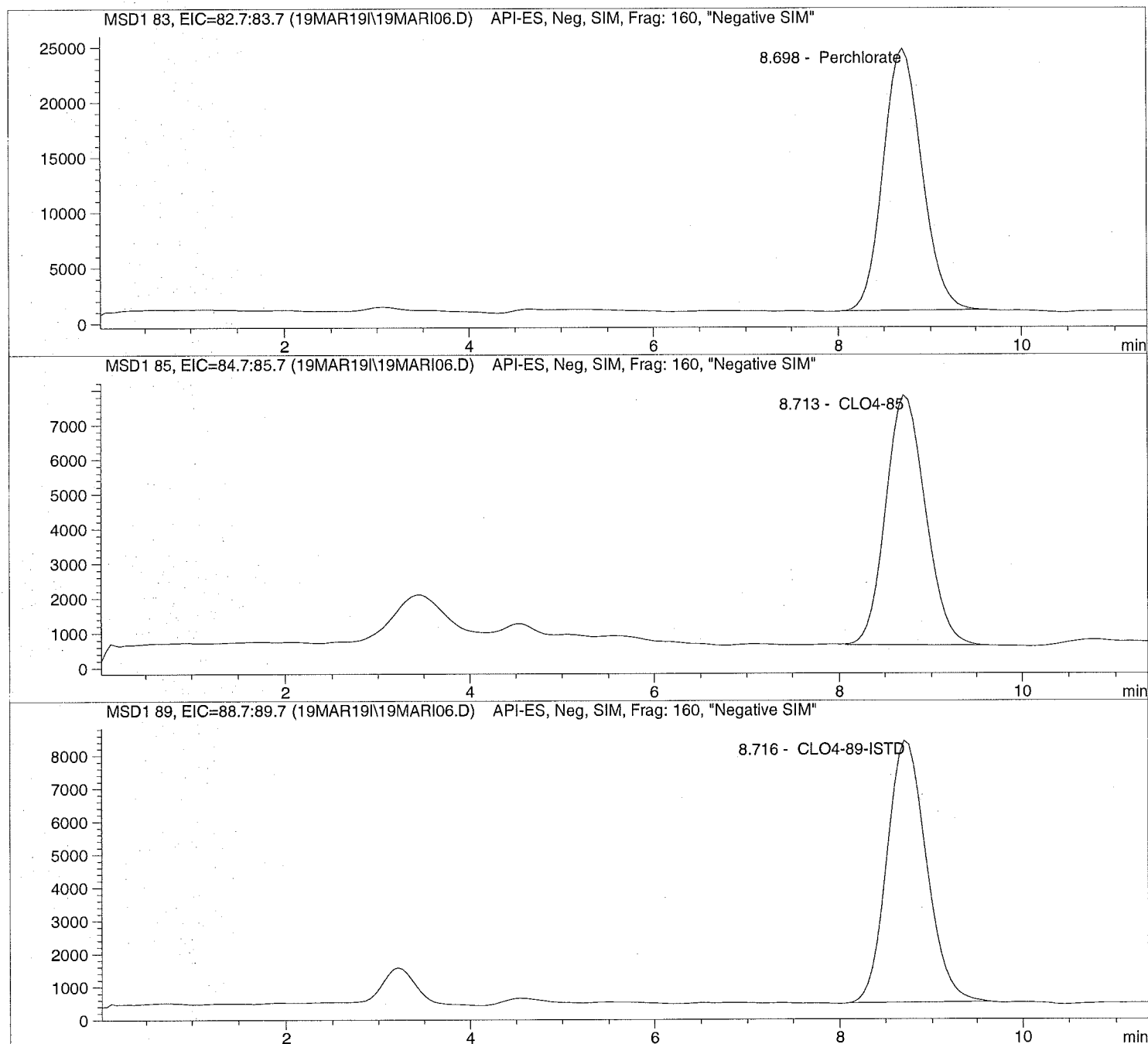
Sample Name: CLO4@ 10.ug/L

Injection Date: 3/19/2019 10:19:32
Sample Name: CLO4@ 10.ug/L
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line:            6
Sample Name:    CLO4@ 10.ug/L            Location:            Vial 76
Acq Operator:   TNB                      Inj. No.:            1
                                          Inj. Vol.:            30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:            Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:          1.000000
Dilution:            1.000000
Sample Amount:        10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI07.D

Sample Name: CLO4@ 25.ug/L

Injection Date: 3/19/2019 10:32:49

Seq Line: 7

Sample Name: CLO4@ 25.ug/L

Location: Vial 77

Acq Operator: TNB

Inj. No.: 1

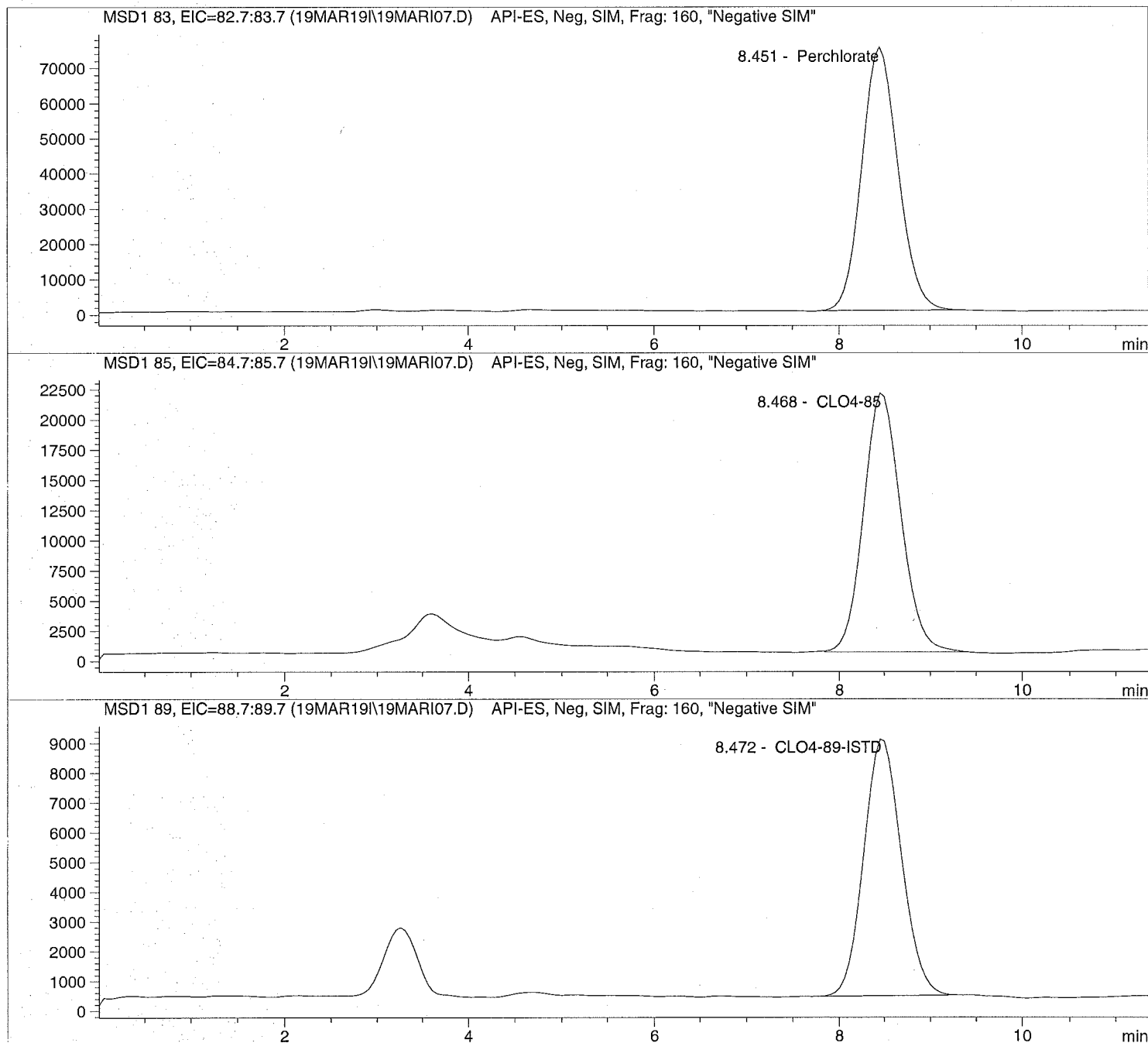
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 3/19/2019 10:32:49      Seq Line: 7
Sample Name:    CLO4@ 25.ug/L          Location:  Vial 77
Acq Operator:  TNB                    Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D

Sample Name: CLO4@ 50.ug/L

Injection Date: 3/19/2019 10:46:05

Seq Line: 8

Sample Name: CLO4@ 50.ug/L

Location: Vial 78

Acq Operator: TNB

Inj. No.: 1

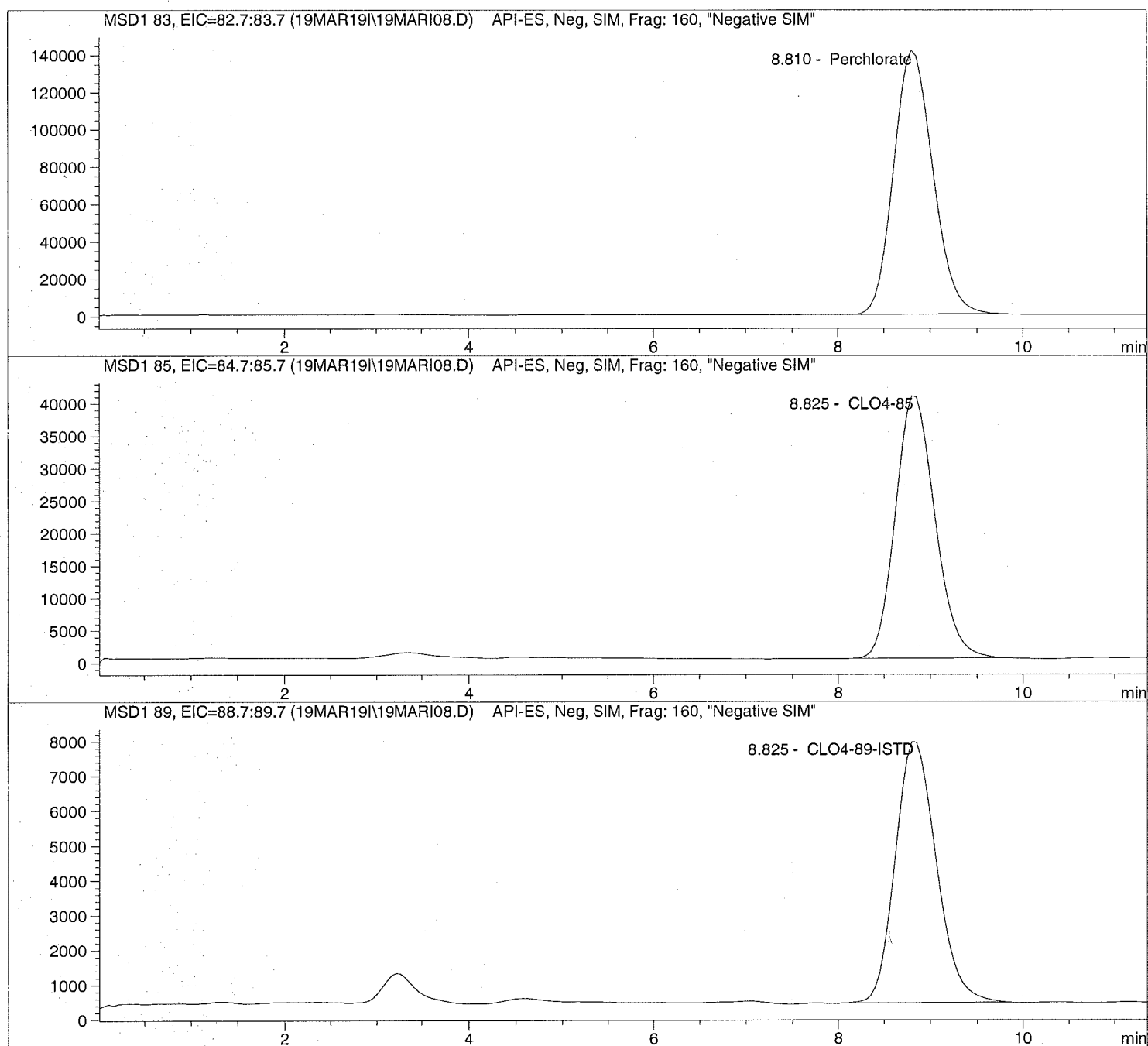
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  50.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 3/19/2019 10:59:22

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

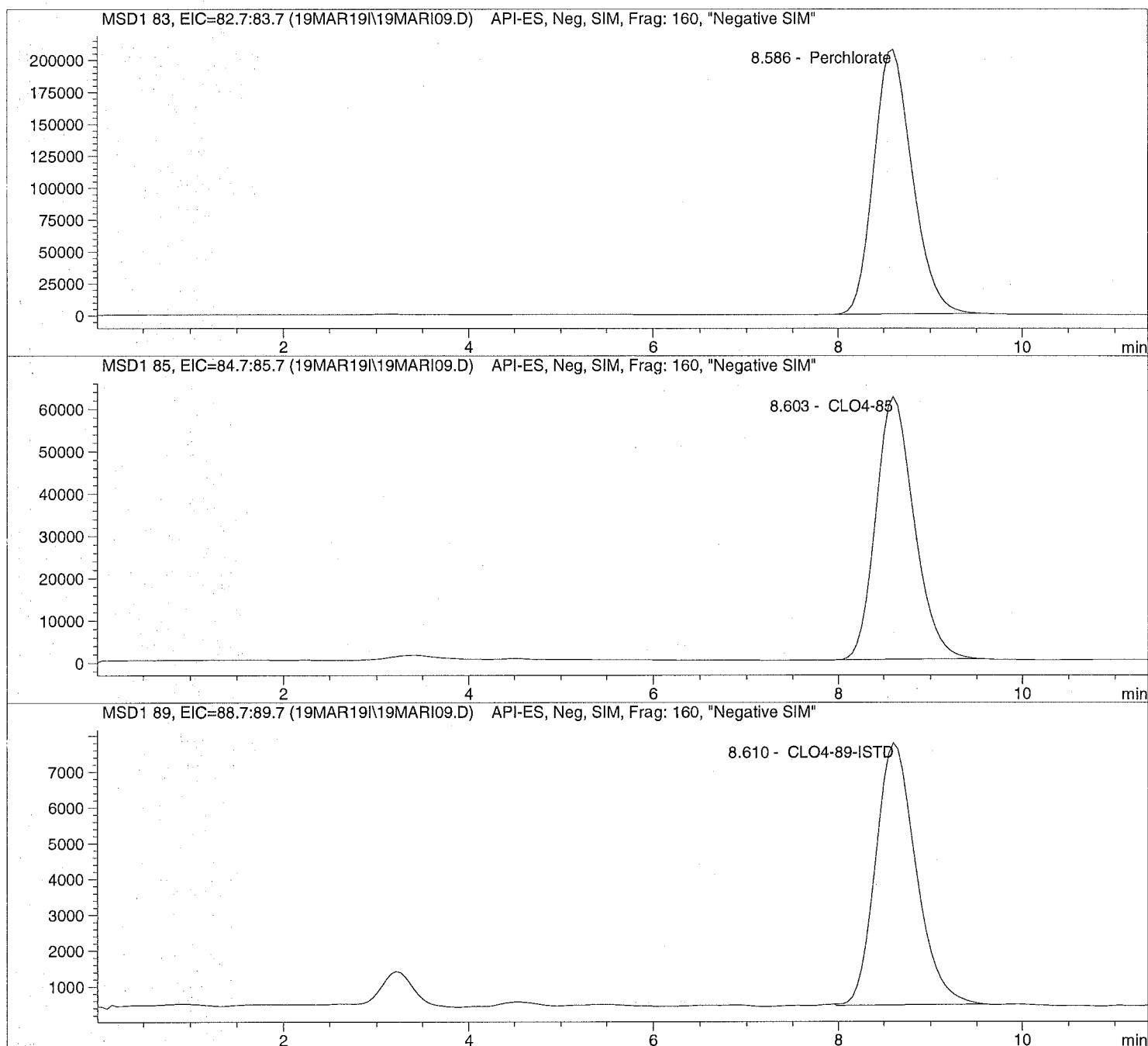
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date: 3/19/2019 10:59:22      Seq Line:          9
Sample Name:    CLO4@ 75.ug/L           Location:          Vial 79
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:       30 µl
  
```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
  
```

Perchlorate analysis

Sample Information

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  75.000
  
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D

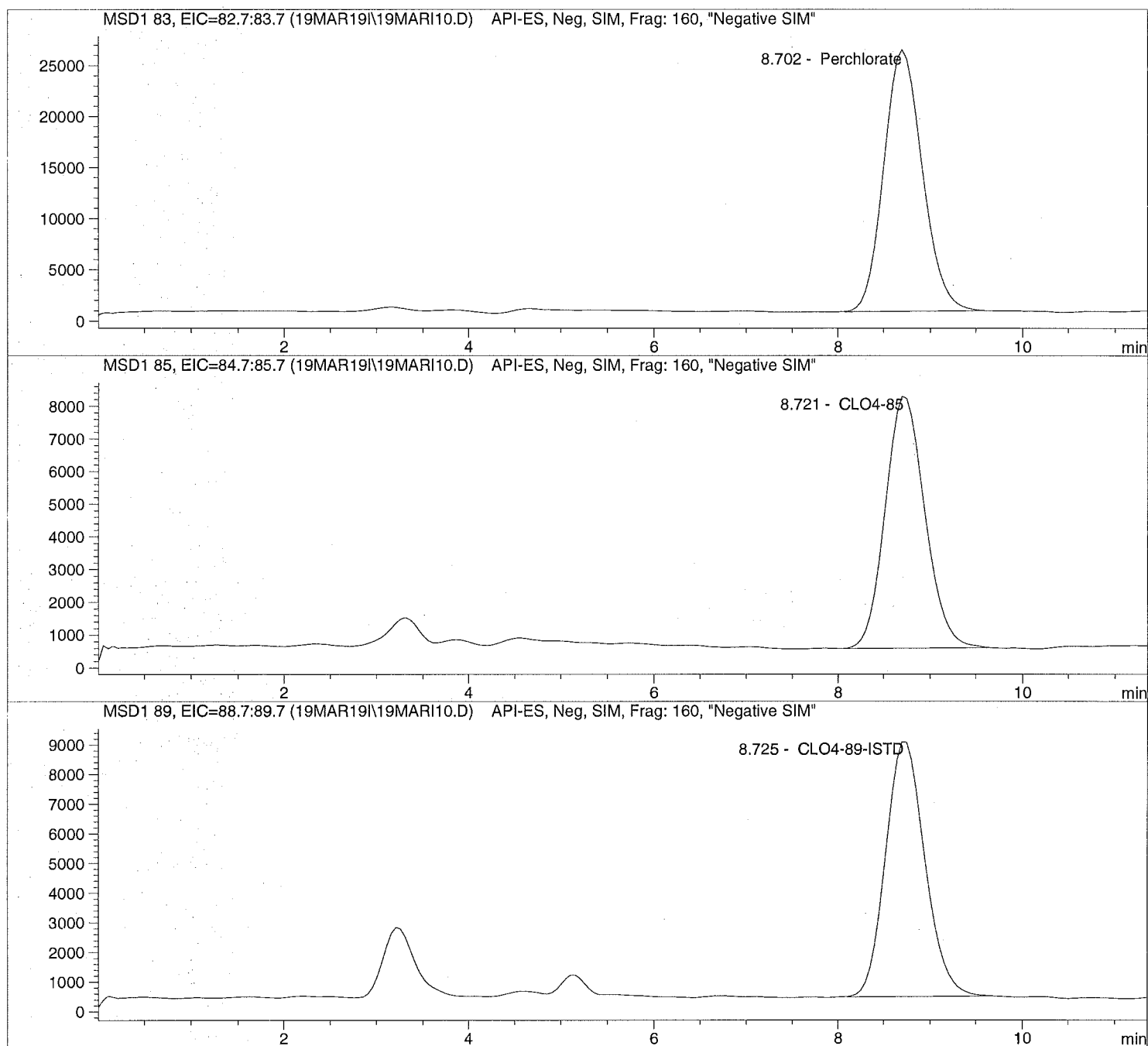
Sample Name: ICAL Verf@10ug/L

=====
Injection Date: 3/19/2019 11:12:42
Sample Name: ICAL Verf@10ug/L
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 3/19/2019 11:12:42      Seq Line:      10
Sample Name:   ICAL Verf@10ug/L        Location:      Vial 80
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

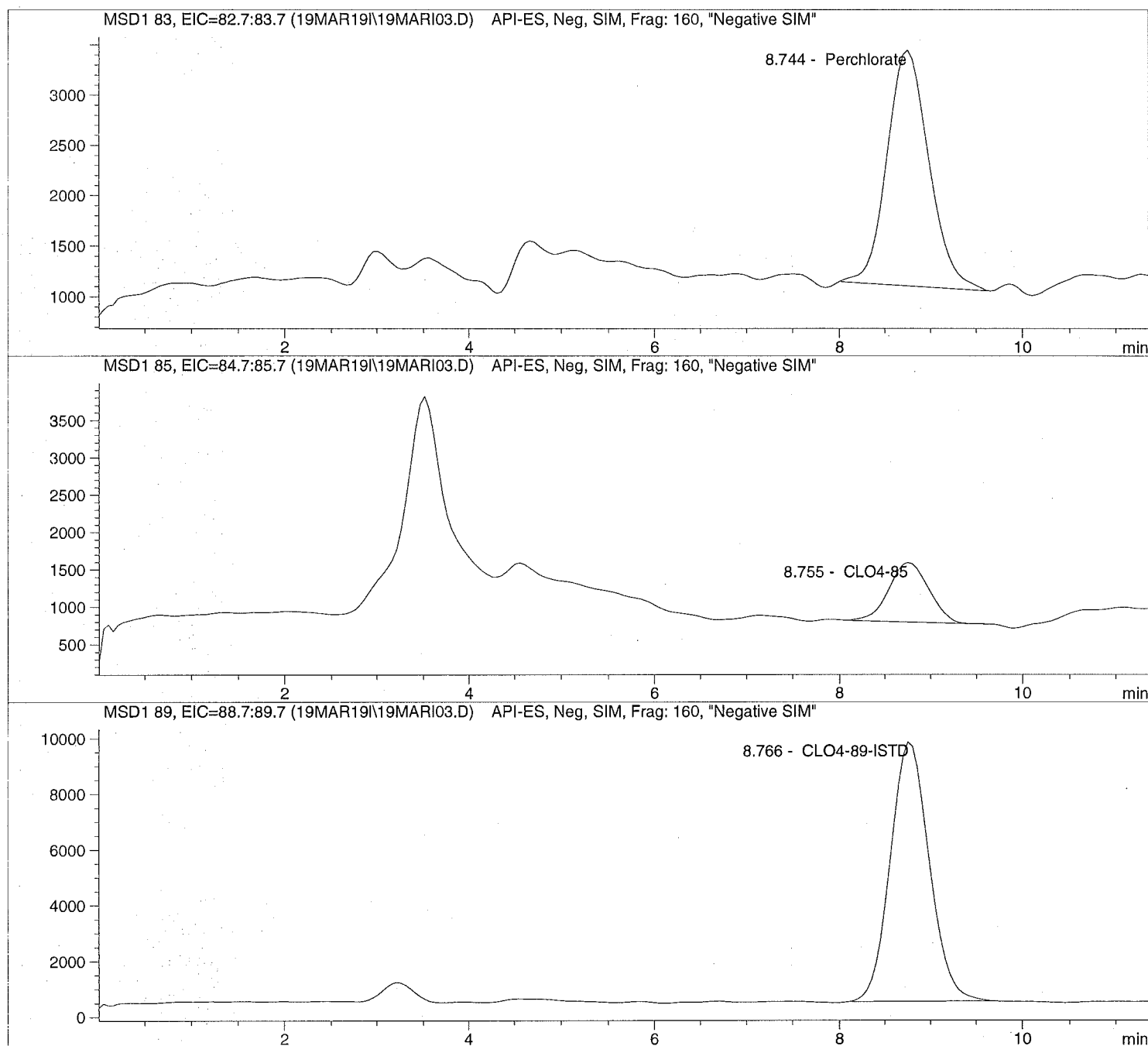
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L          Location:  Vial 73
Acq Operator:  TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 1.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
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March 11, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030301**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 07, 2019 for the analysis presented in the following report.

This is a REVISED REPORT. Please see the Case Narrative for discussion concerning this revision.

Regards,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a hand-drawn oval.

Generated By: RJ.MODASHIA
RJ Modashia
Project Manager

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030301

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030301-01	LH18/24-SP650_030619	Water		06-Mar-2019 13:00	07-Mar-2019 08:51	<input type="checkbox"/>
HS19030301-02	TRIP BLANK	Water	ALS-020119-60	06-Mar-2019 00:00	07-Mar-2019 08:51	<input type="checkbox"/>

Revision:1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
 - Updated the Sample ID to the correct name: LH18/24-SP650_030619
-

GCMS Volatiles by Method SW8260**Batch ID: R334270****Sample ID: HS19030163-01MS**

- MS and MSD are for an unrelated sample

Sample ID: VLCSW-190308

- 1,2,3-Trichlorobenzene exceeded QC limits for LCS. CCV is OK. Samples are ND for this compound.
-

WetChemistry by Method SW9056**Batch ID: R334216**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_030619
 Collection Date: 06-Mar-2019 13:00

ANALYTICAL REPORT
 WorkOrder:HS19030301
 Lab ID:HS19030301-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	08-Mar-2019 12:53	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	08-Mar-2019 12:53	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	08-Mar-2019 12:53	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	08-Mar-2019 12:53	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	08-Mar-2019 12:53	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_030619
 Collection Date: 06-Mar-2019 13:00

ANALYTICAL REPORT
 WorkOrder:HS19030301
 Lab ID:HS19030301-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
cis-1,2-Dichloroethene	1.6		0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	08-Mar-2019 12:53	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	08-Mar-2019 12:53	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	08-Mar-2019 12:53	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Trichloroethene	0.74	J	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:53	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>83.4</i>			0	<i>81-118</i>	%REC	1	08-Mar-2019 12:53	
<i>Surr: 4-Bromofluorobenzene</i>	<i>101</i>			0	<i>85-114</i>	%REC	1	08-Mar-2019 12:53	
<i>Surr: Dibromofluoromethane</i>	<i>84.5</i>			0	<i>80-119</i>	%REC	1	08-Mar-2019 12:53	
<i>Surr: Toluene-d8</i>	<i>108</i>			0	<i>89-112</i>	%REC	1	08-Mar-2019 12:53	
ANIONS BY SW9056A		Method:SW9056						Analyst: KMU	
Chloride	312		2.00	5.00	5.00	mg/L	10	08-Mar-2019 00:17	
Sulfate	13.9		2.00	5.00	5.00	mg/L	10	08-Mar-2019 00:17	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: TRIP BLANK
 Collection Date: 06-Mar-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19030301
 Lab ID:HS19030301-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	08-Mar-2019 12:29	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	08-Mar-2019 12:29	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	08-Mar-2019 12:29	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	08-Mar-2019 12:29	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	08-Mar-2019 12:29	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: TRIP BLANK
 Collection Date: 06-Mar-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19030301
 Lab ID:HS19030301-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD		Method:SW8260							Analyst: PC
8260C									
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	08-Mar-2019 12:29	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	08-Mar-2019 12:29	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	08-Mar-2019 12:29	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Trichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	08-Mar-2019 12:29	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>82.5</i>			<i>0</i>	<i>81-118</i>	<i>%REC</i>	<i>1</i>	<i>08-Mar-2019 12:29</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>101</i>			<i>0</i>	<i>85-114</i>	<i>%REC</i>	<i>1</i>	<i>08-Mar-2019 12:29</i>	
<i>Surr: Dibromofluoromethane</i>	<i>83.8</i>			<i>0</i>	<i>80-119</i>	<i>%REC</i>	<i>1</i>	<i>08-Mar-2019 12:29</i>	
<i>Surr: Toluene-d8</i>	<i>108</i>			<i>0</i>	<i>89-112</i>	<i>%REC</i>	<i>1</i>	<i>08-Mar-2019 12:29</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R334216	Test Name : ANIONS BY SW9056A		Matrix: Water			
HS19030301-01	LH18/24-SP650_030619	06 Mar 2019 13:00			08 Mar 2019 00:17	10
Batch ID R334270	Test Name : VOLATILES ORGANICS BY METHOD 8260C		Matrix: Water			
HS19030301-01	LH18/24-SP650_030619	06 Mar 2019 13:00			08 Mar 2019 12:53	1
HS19030301-02	TRIP BLANK	06 Mar 2019 00:00			08 Mar 2019 12:29	1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
MBLK	Sample ID: VBLKW-190308	Units: UG/L			Analysis Date: 08-Mar-2019 12:05					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984642	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	0.50	1.0								U
1,1,1-Trichloroethane	0.50	1.0								U
1,1,2,2-Tetrachloroethane	0.50	1.0								U
1,1,2-Trichloroethane	0.50	1.0								U
1,1-Dichloroethane	0.50	1.0								U
1,1-Dichloroethene	0.50	1.0								U
1,1-Dichloropropene	0.50	1.0								U
1,2,3-Trichlorobenzene	0.50	1.0								U
1,2,3-Trichloropropane	0.50	1.0								U
1,2,4-Trichlorobenzene	0.50	1.0								U
1,2,4-Trimethylbenzene	0.50	1.0								U
1,2-Dibromo-3-chloropropane	0.50	1.0								U
1,2-Dibromoethane	0.50	1.0								U
1,2-Dichlorobenzene	0.50	1.0								U
1,2-Dichloroethane	0.50	1.0								U
1,2-Dichloropropane	0.50	1.0								U
1,3,5-Trimethylbenzene	0.50	1.0								U
1,3-Dichlorobenzene	0.50	1.0								U
1,3-Dichloropropane	0.50	1.0								U
1,4-Dichlorobenzene	0.50	1.0								U
2,2-Dichloropropane	0.50	1.0								U
2-Butanone	1.0	2.0								U
2-Chlorotoluene	0.50	1.0								U
2-Hexanone	1.0	2.0								U
4-Chlorotoluene	0.50	1.0								U
4-Isopropyltoluene	0.50	1.0								U
4-Methyl-2-pentanone	1.0	2.0								U
Acetone	2.0	2.0								U
Benzene	0.50	1.0								U
Bromobenzene	0.50	1.0								U
Bromochloromethane	0.50	1.0								U
Bromodichloromethane	0.50	1.0								U
Bromoform	0.50	1.0								U
Bromomethane	0.50	1.0								U

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ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
MBLK	Sample ID: VBLKW-190308	Units: UG/L			Analysis Date: 08-Mar-2019 12:05					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984642	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	41.37	1.0	50	0	82.7	81 - 118				
Surr: 4-Bromofluorobenzene	50.23	1.0	50	0	100	85 - 114				
Surr: Dibromofluoromethane	40.97	1.0	50	0	81.9	80 - 119				
Surr: Toluene-d8	53.9	1.0	50	0	108	89 - 112				

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ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
LCS	Sample ID: VLCSW-190308	Units: UG/L			Analysis Date: 08-Mar-2019 11:17					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984641	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	21.24	1.0	20	0	106	78 - 124				
1,1,1-Trichloroethane	20.25	1.0	20	0	101	74 - 131				
1,1,2,2-Tetrachloroethane	21.25	1.0	20	0	106	71 - 121				
1,1,2-Trichloroethane	21.41	1.0	20	0	107	80 - 119				
1,1-Dichloroethane	19.78	1.0	20	0	98.9	77 - 125				
1,1-Dichloroethene	19.2	1.0	20	0	96.0	71 - 131				
1,1-Dichloropropene	19.85	1.0	20	0	99.2	78 - 125				
1,2,3-Trichlorobenzene	26.35	1.0	20	0	132	69 - 129				S
1,2,3-Trichloropropane	20.6	1.0	20	0	103	73 - 122				
1,2,4-Trichlorobenzene	23.18	1.0	20	0	116	69 - 130				
1,2,4-Trimethylbenzene	19.42	1.0	20	0	97.1	76 - 124				
1,2-Dibromo-3-chloropropane	23.86	1.0	20	0	119	62 - 128				
1,2-Dibromoethane	22.06	1.0	20	0	110	77 - 121				
1,2-Dichlorobenzene	20.01	1.0	20	0	100	80 - 119				
1,2-Dichloroethane	21.49	1.0	20	0	107	73 - 128				
1,2-Dichloropropane	21.18	1.0	20	0	106	78 - 122				
1,3,5-Trimethylbenzene	19.3	1.0	20	0	96.5	75 - 124				
1,3-Dichlorobenzene	19.63	1.0	20	0	98.2	80 - 119				
1,3-Dichloropropane	21.34	1.0	20	0	107	80 - 119				
1,4-Dichlorobenzene	19.59	1.0	20	0	97.9	79 - 118				
2,2-Dichloropropane	20.16	1.0	20	0	101	60 - 139				
2-Butanone	47.93	2.0	40	0	120	56 - 143				
2-Chlorotoluene	19.09	1.0	20	0	95.4	79 - 122				
2-Hexanone	46.31	2.0	40	0	116	57 - 139				
4-Chlorotoluene	19.42	1.0	20	0	97.1	78 - 122				
4-Isopropyltoluene	19.19	1.0	20	0	95.9	77 - 127				
4-Methyl-2-pentanone	44.94	2.0	40	0	112	67 - 130				
Acetone	48.14	2.0	40	0	120	39 - 160				
Benzene	20.74	1.0	20	0	104	79 - 120				
Bromobenzene	19.89	1.0	20	0	99.5	80 - 120				
Bromochloromethane	20.61	1.0	20	0	103	78 - 123				
Bromodichloromethane	21.23	1.0	20	0	106	79 - 125				
Bromoform	22.88	1.0	20	0	114	66 - 130				
Bromomethane	24.1	1.0	20	0	120	53 - 141				

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Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
LCS	Sample ID: VLCSW-190308	Units: UG/L			Analysis Date: 08-Mar-2019 11:17					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984641		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	40.69	2.0	40	0	102	64 - 133				
Carbon tetrachloride	21.12	1.0	20	0	106	72 - 136				
Chlorobenzene	20.68	1.0	20	0	103	82 - 118				
Chloroethane	19.46	1.0	20	0	97.3	60 - 138				
Chloroform	20.35	1.0	20	0	102	79 - 124				
Chloromethane	20.05	1.0	20	0	100	50 - 139				
cis-1,2-Dichloroethene	20.09	1.0	20	0	100	78 - 123				
cis-1,3-Dichloropropene	21.64	1.0	20	0	108	75 - 124				
Dibromochloromethane	22.28	1.0	20	0	111	74 - 126				
Dibromomethane	21.51	1.0	20	0	108	79 - 123				
Dichlorodifluoromethane	19.85	1.0	20	0	99.3	32 - 152				
Ethylbenzene	20.06	1.0	20	0	100	79 - 121				
Hexachlorobutadiene	22.92	1.0	20	0	115	66 - 134				
Isopropylbenzene	19.83	1.0	20	0	99.1	72 - 131				
m,p-Xylene	40.07	2.0	40	0	100	80 - 121				
Methylene chloride	20.39	2.0	20	0	102	74 - 124				
Naphthalene	24.65	1.0	20	0	123	61 - 128				
n-Butylbenzene	19.03	1.0	20	0	95.1	75 - 128				
n-Propylbenzene	19.08	1.0	20	0	95.4	76 - 126				
o-Xylene	20.48	1.0	20	0	102	78 - 122				
sec-Butylbenzene	18.75	1.0	20	0	93.8	77 - 126				
Styrene	21.13	1.0	20	0	106	78 - 123				
tert-Butylbenzene	18.73	1.0	20	0	93.7	78 - 124				
Tetrachloroethene	20.12	1.0	20	0	101	74 - 129				
Toluene	20.87	1.0	20	0	104	80 - 121				
trans-1,2-Dichloroethene	20.1	1.0	20	0	101	75 - 124				
trans-1,3-Dichloropropene	21.95	1.0	20	0	110	73 - 127				
Trichloroethene	21.02	1.0	20	0	105	79 - 123				
Trichlorofluoromethane	19.19	1.0	20	0	95.9	65 - 141				
Vinyl chloride	18.48	1.0	20	0	92.4	58 - 137				
Surr: 1,2-Dichloroethane-d4	51.89	1.0	50	0	104	81 - 118				
Surr: 4-Bromofluorobenzene	53.23	1.0	50	0	106	85 - 114				
Surr: Dibromofluoromethane	51.27	1.0	50	0	103	80 - 119				
Surr: Toluene-d8	47.66	1.0	50	0	95.3	89 - 112				

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ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
MS	Sample ID: HS19030163-01MS	Units: UG/L			Analysis Date: 08-Mar-2019 16:29					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984650	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	20.64	1.0	20	0	103	78 - 124				
1,1,1-Trichloroethane	17.28	1.0	20	0	86.4	74 - 131				
1,1,2,2-Tetrachloroethane	23.34	1.0	20	0	117	71 - 121				
1,1,2-Trichloroethane	21.51	1.0	20	0	108	80 - 119				
1,1-Dichloroethane	15.49	1.0	20	0	77.5	77 - 125				
1,1-Dichloroethene	16.33	1.0	20	0	81.6	71 - 131				
1,1-Dichloropropene	18.4	1.0	20	0	92.0	78 - 125				
1,2,3-Trichlorobenzene	31.17	1.0	20	0	156	69 - 129				S
1,2,3-Trichloropropane	32.3	1.0	20	0	161	73 - 122				S
1,2,4-Trichlorobenzene	27.02	1.0	20	0	135	69 - 130				S
1,2,4-Trimethylbenzene	21.71	1.0	20	0	109	76 - 124				
1,2-Dibromo-3-chloropropane	27.4	1.0	20	0	137	62 - 128				S
1,2-Dibromoethane	22.13	1.0	20	0	111	77 - 121				
1,2-Dichlorobenzene	21.24	1.0	20	0	106	80 - 119				
1,2-Dichloroethane	18.17	1.0	20	0	90.9	73 - 128				
1,2-Dichloropropane	18.26	1.0	20	0	91.3	78 - 122				
1,3,5-Trimethylbenzene	22.06	1.0	20	0	110	75 - 124				
1,3-Dichlorobenzene	21.05	1.0	20	0	105	80 - 119				
1,3-Dichloropropane	21.22	1.0	20	0	106	80 - 119				
1,4-Dichlorobenzene	20.65	1.0	20	0	103	79 - 118				
2,2-Dichloropropane	16.44	1.0	20	0	82.2	60 - 139				
2-Butanone	41.77	2.0	40	0	104	56 - 143				
2-Chlorotoluene	20.76	1.0	20	0	104	79 - 122				
2-Hexanone	50.33	2.0	40	0	126	57 - 139				
4-Chlorotoluene	21.05	1.0	20	0	105	78 - 122				
4-Isopropyltoluene	22.51	1.0	20	0	113	77 - 127				
4-Methyl-2-pentanone	48.85	2.0	40	0	122	67 - 130				
Acetone	40.25	2.0	40	0	101	39 - 160				
Benzene	18.1	1.0	20	0	90.5	79 - 120				
Bromobenzene	20.7	1.0	20	0	103	80 - 120				
Bromochloromethane	15.3	1.0	20	0	76.5	78 - 123				S
Bromodichloromethane	18.48	1.0	20	0	92.4	79 - 125				
Bromoform	22.63	1.0	20	0	113	66 - 130				
Bromomethane	20.71	1.0	20	0	104	53 - 141				

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ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
MS	Sample ID: HS19030163-01MS	Units: UG/L			Analysis Date: 08-Mar-2019 16:29					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984650	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	30.91	2.0	40	0	77.3	64 - 133				
Carbon tetrachloride	21.19	1.0	20	0	106	72 - 136				
Chlorobenzene	20.56	1.0	20	0	103	82 - 118				
Chloroethane	15.55	1.0	20	0	77.7	60 - 138				
Chloroform	15.88	1.0	20	0	79.4	79 - 124				
Chloromethane	11.94	1.0	20	0	59.7	50 - 139				
cis-1,2-Dichloroethene	16	1.0	20	0	80.0	78 - 123				
cis-1,3-Dichloropropene	18.11	1.0	20	0	90.6	75 - 124				
Dibromochloromethane	21.65	1.0	20	0	108	74 - 126				
Dibromomethane	18.93	1.0	20	0	94.6	79 - 123				
Dichlorodifluoromethane	9.761	1.0	20	0	48.8	32 - 152				
Ethylbenzene	21.29	1.0	20	0	106	79 - 121				
Hexachlorobutadiene	25.33	1.0	20	0	127	66 - 134				
Isopropylbenzene	23.53	1.0	20	1.608	110	72 - 131				
m,p-Xylene	42.68	2.0	40	0	107	80 - 121				
Methylene chloride	16.05	2.0	20	0	80.3	74 - 124				
Naphthalene	29.28	1.0	20	0	146	61 - 128				S
n-Butylbenzene	22.33	1.0	20	0	112	75 - 128				
n-Propylbenzene	22	1.0	20	0	110	76 - 126				
o-Xylene	21.3	1.0	20	0	106	78 - 122				
sec-Butylbenzene	22.53	1.0	20	0	113	77 - 126				
Styrene	20.92	1.0	20	0	105	78 - 123				
tert-Butylbenzene	22.24	1.0	20	0	111	78 - 124				
Tetrachloroethene	22.8	1.0	20	0	114	74 - 129				
Toluene	21.05	1.0	20	0	105	80 - 121				
trans-1,2-Dichloroethene	16.49	1.0	20	0	82.5	75 - 124				
trans-1,3-Dichloropropene	18.72	1.0	20	0	93.6	73 - 127				
Trichloroethene	19.06	1.0	20	0	95.3	79 - 123				
Trichlorofluoromethane	15.35	1.0	20	0	76.7	65 - 141				
Vinyl chloride	13.26	1.0	20	0	66.3	58 - 137				
Surr: 1,2-Dichloroethane-d4	42.6	1.0	50	0	85.2	81 - 118				
Surr: 4-Bromofluorobenzene	50.07	1.0	50	0	100	85 - 114				
Surr: Dibromofluoromethane	42.23	1.0	50	0	84.5	80 - 119				
Surr: Toluene-d8	54.26	1.0	50	0	109	89 - 112				

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
MSD	Sample ID: HS19030163-01MSD	Units: UG/L			Analysis Date: 08-Mar-2019 16:53					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984651	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	20.78	1.0	20	0	104	78 - 124	20.64	0.661	20	
1,1,1-Trichloroethane	18.25	1.0	20	0	91.2	74 - 131	17.28	5.44	20	
1,1,2,2-Tetrachloroethane	20.8	1.0	20	0	104	71 - 121	23.34	11.5	20	
1,1,2-Trichloroethane	20.81	1.0	20	0	104	80 - 119	21.51	3.32	20	
1,1-Dichloroethane	16.04	1.0	20	0	80.2	77 - 125	15.49	3.46	20	
1,1-Dichloroethene	17.15	1.0	20	0	85.7	71 - 131	16.33	4.89	20	
1,1-Dichloropropene	20.56	1.0	20	0	103	78 - 125	18.4	11.1	20	
1,2,3-Trichlorobenzene	31	1.0	20	0	155	69 - 129	31.17	0.541	20	S
1,2,3-Trichloropropane	28.78	1.0	20	0	144	73 - 122	32.3	11.5	20	S
1,2,4-Trichlorobenzene	26.83	1.0	20	0	134	69 - 130	27.02	0.679	20	S
1,2,4-Trimethylbenzene	21.09	1.0	20	0	105	76 - 124	21.71	2.9	20	
1,2-Dibromo-3-chloropropane	24.34	1.0	20	0	122	62 - 128	27.4	11.9	20	
1,2-Dibromoethane	21.34	1.0	20	0	107	77 - 121	22.13	3.67	20	
1,2-Dichlorobenzene	19.82	1.0	20	0	99.1	80 - 119	21.24	6.9	20	
1,2-Dichloroethane	19.09	1.0	20	0	95.4	73 - 128	18.17	4.9	20	
1,2-Dichloropropane	18.64	1.0	20	0	93.2	78 - 122	18.26	2.07	20	
1,3,5-Trimethylbenzene	21.5	1.0	20	0	108	75 - 124	22.06	2.55	20	
1,3-Dichlorobenzene	20.17	1.0	20	0	101	80 - 119	21.05	4.24	20	
1,3-Dichloropropane	20.54	1.0	20	0	103	80 - 119	21.22	3.28	20	
1,4-Dichlorobenzene	19.56	1.0	20	0	97.8	79 - 118	20.65	5.41	20	
2,2-Dichloropropane	17.08	1.0	20	0	85.4	60 - 139	16.44	3.82	20	
2-Butanone	40.18	2.0	40	0	100	56 - 143	41.77	3.89	20	
2-Chlorotoluene	20.11	1.0	20	0	101	79 - 122	20.76	3.17	20	
2-Hexanone	46.6	2.0	40	0	117	57 - 139	50.33	7.68	20	
4-Chlorotoluene	20.2	1.0	20	0	101	78 - 122	21.05	4.14	20	
4-Isopropyltoluene	22.99	1.0	20	0	115	77 - 127	22.51	2.1	20	
4-Methyl-2-pentanone	45.37	2.0	40	0	113	67 - 130	48.85	7.39	20	
Acetone	37.06	2.0	40	0	92.7	39 - 160	40.25	8.26	20	
Benzene	19.25	1.0	20	0	96.2	79 - 120	18.1	6.17	20	
Bromobenzene	19.33	1.0	20	0	96.7	80 - 120	20.7	6.82	20	
Bromochloromethane	15.9	1.0	20	0	79.5	78 - 123	15.3	3.8	20	
Bromodichloromethane	19.03	1.0	20	0	95.2	79 - 125	18.48	2.92	20	
Bromoform	21.66	1.0	20	0	108	66 - 130	22.63	4.36	20	
Bromomethane	19.94	1.0	20	0	99.7	53 - 141	20.71	3.76	20	

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270		Instrument: VOA6		Method: SW8260						
MSD	Sample ID: HS19030163-01MSD	Units: UG/L			Analysis Date: 08-Mar-2019 16:53					
Client ID:	Run ID: VOA6_334270	SeqNo: 4984651	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	33.15	2.0	40	0	82.9	64 - 133	30.91	7	20	
Carbon tetrachloride	22.13	1.0	20	0	111	72 - 136	21.19	4.31	20	
Chlorobenzene	20.65	1.0	20	0	103	82 - 118	20.56	0.397	20	
Chloroethane	17.3	1.0	20	0	86.5	60 - 138	15.55	10.7	20	
Chloroform	16.33	1.0	20	0	81.7	79 - 124	15.88	2.83	20	
Chloromethane	12.62	1.0	20	0	63.1	50 - 139	11.94	5.51	20	
cis-1,2-Dichloroethene	16.36	1.0	20	0	81.8	78 - 123	16	2.18	20	
cis-1,3-Dichloropropene	18.69	1.0	20	0	93.5	75 - 124	18.11	3.17	20	
Dibromochloromethane	21.6	1.0	20	0	108	74 - 126	21.65	0.221	20	
Dibromomethane	19.3	1.0	20	0	96.5	79 - 123	18.93	1.97	20	
Dichlorodifluoromethane	10.51	1.0	20	0	52.6	32 - 152	9.761	7.43	20	
Ethylbenzene	21.74	1.0	20	0	109	79 - 121	21.29	2.07	20	
Hexachlorobutadiene	27.51	1.0	20	0	138	66 - 134	25.33	8.26	20	S
Isopropylbenzene	24.58	1.0	20	1.608	115	72 - 131	23.53	4.35	20	
m,p-Xylene	43.51	2.0	40	0	109	80 - 121	42.68	1.94	20	
Methylene chloride	16.16	2.0	20	0	80.8	74 - 124	16.05	0.673	20	
Naphthalene	27.74	1.0	20	0	139	61 - 128	29.28	5.41	20	S
n-Butylbenzene	23.43	1.0	20	0	117	75 - 128	22.33	4.82	20	
n-Propylbenzene	21.88	1.0	20	0	109	76 - 126	22	0.545	20	
o-Xylene	21.5	1.0	20	0	108	78 - 122	21.3	0.948	20	
sec-Butylbenzene	23.02	1.0	20	0	115	77 - 126	22.53	2.13	20	
Styrene	21.25	1.0	20	0	106	78 - 123	20.92	1.56	20	
tert-Butylbenzene	22.27	1.0	20	0	111	78 - 124	22.24	0.114	20	
Tetrachloroethene	24.06	1.0	20	0	120	74 - 129	22.8	5.36	20	
Toluene	21.57	1.0	20	0	108	80 - 121	21.05	2.46	20	
trans-1,2-Dichloroethene	17.04	1.0	20	0	85.2	75 - 124	16.49	3.29	20	
trans-1,3-Dichloropropene	19.24	1.0	20	0	96.2	73 - 127	18.72	2.72	20	
Trichloroethene	20.31	1.0	20	0	102	79 - 123	19.06	6.33	20	
Trichlorofluoromethane	16.52	1.0	20	0	82.6	65 - 141	15.35	7.37	20	
Vinyl chloride	13.85	1.0	20	0	69.3	58 - 137	13.26	4.37	20	
Surr: 1,2-Dichloroethane-d4	41.92	1.0	50	0	83.8	81 - 118	42.6	1.62	20	
Surr: 4-Bromofluorobenzene	50.79	1.0	50	0	102	85 - 114	50.07	1.42	20	
Surr: Dibromofluoromethane	41.48	1.0	50	0	83.0	80 - 119	42.23	1.8	20	
Surr: Toluene-d8	52.71	1.0	50	0	105	89 - 112	54.26	2.9	20	

Revision: 1

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334270	Instrument: VOA6	Method: SW8260
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The following samples were analyzed in this batch:

HS19030301-01	HS19030301-02
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ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334216		Instrument: ICS2100		Method: SW9056						
MBLK	Sample ID: WBLKW1-030719	Units: mg/L			Analysis Date: 07-Mar-2019 16:22					
Client ID:	Run ID: ICS2100_334216	SeqNo: 4979622		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	0.500	0.500							U	
Sulfate	0.500	0.500							U	
LCS	Sample ID: WLCSW1-030719	Units: mg/L			Analysis Date: 07-Mar-2019 16:37					
Client ID:	Run ID: ICS2100_334216	SeqNo: 4979623		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	21.43	0.500	20	0	107	80 - 120				
Sulfate	20.79	0.500	20	0	104	80 - 120				
LCSD	Sample ID: WLCSDW1-030719	Units: mg/L			Analysis Date: 07-Mar-2019 16:51					
Client ID:	Run ID: ICS2100_334216	SeqNo: 4979624		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	20.62	0.500	20	0	103	80 - 120	21.43	3.84	20	
Sulfate	20	0.500	20	0	100.0	80 - 120	20.79	3.91	20	
MS	Sample ID: HS19030301-01MS	Units: mg/L			Analysis Date: 08-Mar-2019 00:31					
Client ID: LH18/24-SP650_030619	Run ID: ICS2100_334216	SeqNo: 4979646		PrepDate:			DF: 10			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	412.7	5.00	100	312	101	80 - 120				
Sulfate	107.6	5.00	100	13.91	93.7	80 - 120				
MS	Sample ID: HS19030293-01MS	Units: mg/L			Analysis Date: 07-Mar-2019 19:04					
Client ID:	Run ID: ICS2100_334216	SeqNo: 4979627		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	23.07	0.500	10	13.61	94.6	80 - 120				
Sulfate	33.83	0.500	10	24.5	93.3	80 - 120				

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030301

QC BATCH REPORT

Batch ID: R334216 **Instrument:** ICS2100 **Method:** SW9056

MSD		Sample ID: HS19030301-01MSD			Units: mg/L		Analysis Date: 08-Mar-2019 00:46			
Client ID: LH18/24-SP650_030619		Run ID: ICS2100_334216			SeqNo: 4979647		PrepDate:		DF: 10	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chloride	396.8	5.00	100	312	84.8	80 - 120	412.7	3.93	20	
Sulfate	103	5.00	100	13.91	89.1	80 - 120	107.6	4.34	20	

MSD		Sample ID: HS19030293-01MSD			Units: mg/L		Analysis Date: 07-Mar-2019 19:19			
Client ID:		Run ID: ICS2100_334216			SeqNo: 4979628		PrepDate:		DF: 1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chloride	23.2	0.500	10	13.61	95.9	80 - 120	23.07	0.553	20	
Sulfate	33.85	0.500	10	24.5	93.5	80 - 120	33.83	0.0621	20	

The following samples were analyzed in this batch: HS19030301-01

Revision: 1

ALS Houston, US

Date: 11-Mar-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19030301	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

Unit Reported	Description
mg/L	Milligrams per Liter

CERTIFICATIONS, ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

ALS Houston, US

Date: 11-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030301

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19030301-01	LH18/24-SP650_030619	Login	3/7/2019 11:41:15 AM	NDR	WET311
HS19030301-01	LH18/24-SP650_030619	Login	3/7/2019 11:41:15 AM	NDR	WET311
HS19030301-01	LH18/24-SP650_030619	Login	3/7/2019 11:41:15 AM	NDR	Sub

Revision:1

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030301

Date/Time Received: **07-Mar-2019 08:51**
 Received by: **NDR**

Checklist completed by: Nilesh D. Ranchod 7-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 7-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 0.2C/0.2C UC/C IR11
 Cooler(s)/Kit(s): 43601
 Date/Time sample(s) sent to storage: 03/07/2019 13:00

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A


pH adjusted by:

Login Notes:


Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

 <p>ALS 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887</p>	CUSTOMER	
	Date:	3/6/19
	Name:	See #
	Company:	B.H.P.

ODY SEAL	
Time:	1330
Beck NGel	
Seal Broken By:	[Signature]
Date:	[Signature]

FedEx	THU - 07 MAR 10:30A
TRK# 0227 4380 9530 9478	PRIORITY OVERNIGHT
AB SGRA	77099
	TX-US IAH
	
F10 5121956 06MAR19 0000 553C1/4603/8CBA	



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 31, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030749**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 15, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval outline.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030749

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030749-01	LH18/24-SP650_031419	Water		14-Mar-2019 14:00	15-Mar-2019 08:38	<input type="checkbox"/>
HS19030749-02	LH18/24-SP650_031419_BIX	Water		14-Mar-2019 14:00	15-Mar-2019 08:38	<input type="checkbox"/>

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
 - The analysis for TOC was subcontracted to ALS Kelso, WA. Final report attached.
-

WetChemistry by Method E350.3**Batch ID: R334781**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

WetChemistry by Method E365.3**Batch ID: R334753**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_031419
 Collection Date: 14-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030749
 Lab ID:HS19030749-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
AMMONIA AS N BY E350.3(ISE)								Analyst: KVL
Nitrogen, Ammonia (As N)	18		0.20	0.20	0.20	mg/L	1	18-Mar-2019 12:30
ORTHO PHOSPHATE (PO4) AS P BY E365.3								Analyst: MZD
Phosphorus, Total Orthophosphate (As P)	4.65		0.100	0.250	0.250	mg/L	10	15-Mar-2019 14:30
SUBCONTRACT ANALYSIS - TOC ANALYSIS								Analyst: SUBK
Subcontract Analysis	See Attached		0	0		NA	1	29-Mar-2019 15:52

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_031419_BIX
 Collection Date: 14-Mar-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19030749
 Lab ID:HS19030749-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	31-Mar-2019 12:13

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030749

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R334753	Test Name : ORTHO PHOSPHATE (PO4) AS P BY E365.3		Matrix: Water			
HS19030749-01	LH18/24-SP650_031419	14 Mar 2019 14:00			15 Mar 2019 14:30	10
Batch ID R334781	Test Name : AMMONIA AS N BY E350.3(ISE)		Matrix: Water			
HS19030749-01	LH18/24-SP650_031419	14 Mar 2019 14:00			18 Mar 2019 12:30	1
Batch ID R335613	Test Name : SUBCONTRACT ANALYSIS - TOC ANALYSIS		Matrix: Water			
HS19030749-01	LH18/24-SP650_031419	14 Mar 2019 14:00			29 Mar 2019 15:52	1
Batch ID R335660	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19030749-02	LH18/24-SP650_031419_BIX	14 Mar 2019 14:00			31 Mar 2019 12:13	1

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030749

QC BATCH REPORT NEW

Batch ID:	R334753 (0)	Instrument:	UV-2450	Method:	ORTHO PHOSPHATE (PO4) AS P BY E365.3					
MBLK	Sample ID: MBLK-334753	Units:	mg/L	Analysis Date:	15-Mar-2019 14:30					
Client ID:		Run ID:	UV-2450_334753	SeqNo:	4995078	PrepDate:		DF:	1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	0.0250	0.0250								U
LCS	Sample ID: LCS-334753	Units:	mg/L	Analysis Date:	15-Mar-2019 14:30					
Client ID:		Run ID:	UV-2450_334753	SeqNo:	4995079	PrepDate:		DF:	1	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	0.253	0.0250	0.25	0	101	85 - 115				
MS	Sample ID: HS19030749-01MS	Units:	mg/L	Analysis Date:	15-Mar-2019 14:30					
Client ID:	LH18/24-SP650_031419	Run ID:	UV-2450_334753	SeqNo:	4995081	PrepDate:		DF:	10	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	7.48	0.250	2.5	4.65	113	80 - 120				
MSD	Sample ID: HS19030749-01MSD	Units:	mg/L	Analysis Date:	15-Mar-2019 14:30					
Client ID:	LH18/24-SP650_031419	Run ID:	UV-2450_334753	SeqNo:	4995082	PrepDate:		DF:	10	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	7.46	0.250	2.5	4.65	112	80 - 120	7.48	0.268	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19030749

QC BATCH REPORT NEW

Batch ID: R334781 (0)		Instrument: WetChem_HS		Method: AMMONIA AS N BY E350.3(ISE)						
MBLK	Sample ID: MBLK-R334781	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 4995674			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	0.20	0.20							U	
LCS	Sample ID: LCS-R334781	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 5007379			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	9.398	0.20	10	0	94.0	80 - 120				
MS	Sample ID: HS19030689-05MS	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 4995676			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	10.4	0.20	10	0.1809	102	80 - 120				
MSD	Sample ID: HS19030689-05MSD	Units: mg/L			Analysis Date: 18-Mar-2019 12:30					
Client ID:	Run ID: WetChem_HS_334781	SeqNo: 4995675			PrepDate:		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	10.38	0.20	10	0.1809	102	80 - 120	10.4	0.192	20	

The following samples were analyzed in this batch: HS19030749-01

ALS Houston, US

Date: 31-Mar-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19030749	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19030749

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19030749-01	LH18/24-SP650_031419	Login	3/15/2019 9:47:50 AM	JRM	WET311
HS19030749-01	LH18/24-SP650_031419	Login	3/15/2019 9:47:50 AM	JRM	WET311
HS19030749-01	LH18/24-SP650_031419	Login	3/15/2019 9:47:50 AM	JRM	Sub
HS19030749-02	LH18/24-SP650_031419_BIX	Login	3/15/2019 9:47:50 AM	JRM	Sub

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030749

Date/Time Received: **15-Mar-2019 08:38**
 Received by: **JRM**

Checklist completed by: Jared R. Makan 15-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 15-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **ALS Courier**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s):

1.3c/1.3c UC/C	IR25
----------------	------

Cooler(s)/Kit(s):

25780

Date/Time sample(s) sent to storage:

03/15/2019 10:55

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

--

Login Notes:

Client Contacted: _____ Date Contacted: _____ Person Contacted: _____
 Contacted By: _____ Regarding: _____

Comments:

--

Corrective Action:

--

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd. Suite 210 Houston, TX. 77099 (281) 530-5656 ATTN: R.J Modshia

Page 1 of 1

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001		Analyses										Remarks (Preservatives, etc.)	Lab I.D.#	
Job: GROUNDWATER TREATMENT PLANT WEEKLY SAMPLES			MS / MSD	No. OF CONTAINERS	AMMONIA-N	TOTAL ORGANIC CARBON	ORTHO-PHOSPHATE	PERCHLORATE									
Prepared By: Scott Beesinger	P.O. Number														Field Sample I.D.	Sample Matrix	
LH18/24-SP650_031419	Water	03/14/19 / 14:00	2	X	X											H2SO4	
LH18/24-SP650_031419	Water	03/14/19 / 14:00	1			X										NONE	
LH18/24-SP650_031419_BIX	Water	03/14/19 / 14:00	1				X									NONE	

Additional Remarks: **Standard TAT on all parameters**

Relinquished By: <i>Scott Beesinger</i>	Date: 03/14/19	Time: 14:30	Received By:	Date:	Time:	Relinquished By:	Date:	Time:	Received By:	Date:	Time:
---	----------------	-------------	--------------	-------	-------	------------------	-------	-------	--------------	-------	-------

For Lab Use Only									
Received At Lab By:	Date	Time	Airbill No.	Opened By:	Date	Time	Temp of Container	Seal No.	Condition
<i>J. Wynn</i>	3/15/19	08:28							

Remarks: *Coder 25780 11225
Temp 1.3 CFO.0*

HS19030749

Bhate Environmental Associates, Inc.
18/24 Longhorn GW Treatment Plant Weekly Sampl



(Word) S:\1-ces\Forms\Chain of Custody - BiWeekly

	ALS 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 6666 Fax. +1 281 530 5887	CU: Date: <u>3/14/19</u> Name: <u>Scott</u> Company: <u>BIT</u>
---	--	---

STUDY SEAL Time: <u>1430</u> <u>Scott</u>	Seal Broken By: <u>SM</u> Date: <u>3/15/19</u>
--	---


 TRK#
 0221 4380 9530 9490

FRI - 15 MAR 10:30A
 PRIORITY OVERNIGHT

AB SGRA

77099
 TX-US
 IAH



FTD 3586291 14MAR19 0GGA 563C1/4603/41CA



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1907867; 1907869; 1907871;
1908794

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2230 (235529)

General Set Information: There were four field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Method QC data: The method blank (LMB 645533) was less than 1/2 the CRDL. The recovery for the LCS (645534) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0 μ l of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4. μ g/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in μ g/L. Results were calculated in μ g/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (μ g/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level of 4.0 μ g/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch March 29, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 29, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1907871**

Project ID: HS19030749

Purchase Order: HS19030749

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP650_031419_BIX	1907871001	03/14/19	03/19/19	



ANALYTICAL REPORT

Workorder: 34-1907871

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP650_031419_BIX	Sampling Site: NA	Collected: 03/14/2019				
Lab ID: 1907871001	Media: 125 mL Nalgene	Received: 03/19/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2230 (HBN: 235529) Analyzed: 03/28/2019 10:21	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 235529)

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/28/2019 14:08	/S/ Stephen Brose 03/29/2019 12:58

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1907871

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935159

Analysis Information

Workorder: 1907871

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2230 (HBN: 235529)
Analyzed By: Thomas Bosch

Blank

LMB: 645533 Analyzed: 03/28/2019 09:14 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 645534 Analyzed: 03/28/2019 08:47 Dilution: 1 Units: ug/L					
Analyte	Result	Target	% Rec	QC Limits	
Perchlorate	4.00	4.00	99.9	78.8	123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1907869001 Analyzed: 03/28/2019 09:41 Dilution: 1 Units: ug/L		MS: 645535 Analyzed: 03/28/2019 09:54 Dilution: 1 Units: ug/L				MSD: 645536 Analyzed: 03/28/2019 10:08 Dilution: 1 Units: ug/L					
Analyte	Result	Result	Target	% Rec	QC Limits		Result	% Rec	RPD	QC Limits	
Perchlorate	ND	4.66	4	117	78.8	123.8	4.05	101	14	0.0	20.0

Comments

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/28/2019 14:23	/S/ Stephen Brose 03/29/2019 12:58

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



1907871

18698/#2

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10915

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

1907871

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030749
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030749-02	LH18/24-SP650_031419_BIX	Water	14 Mar 2019 14:00
	SUB_Perch-6850		29 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. [Signature]
Received By: Jumoke M. Lawal
Cooler ID(s): _____

Date/Time: 3/18/19 18:00
Date/Time: 03-19-19 9:50
Temperature(s): _____

RIGHT SOLUTIONS | RIGHT PARTNER



ALS Environmental
CHAIN-OF-CUSTODY

Project / Job / Task: HS19030749		Split:	Workorder ID: 1907871	Level: ENV_LVL4	Requested Analysis	
Client: ALS Environmental (Houston)		Account: 8101		Type: 125Poly		
Comments:						
Collect Date/Time	Sample ID	Lab ID	QC	Matrix	ID(s)	Count
1 03/14/2019 14:00	LH18/24-SP650_031419_BIX	1907871001		Water	A	1
2						
3						
4						
5						
6						
7						
8						
9						
10						

EPA 6950, DDD GSM

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY				SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY			
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location	Sample Prep / Analysis for:	Lab Notebook No.:	Prepared / Analyzed by:	Date / Time:
Warath, Julie	03/19/2019 09:50	ALS Sample Receiving	Sample Login				
<i>Julie Warath</i>	3/21/19 11:45	<i>ISC</i>	<i>Storage</i>				
R-33-1	3-27-19 11:45	<i>T.B. ...</i>	<i>6850</i>				

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: <u>ALS Houston</u>		Project/Task/Site: <u>1907871</u>						
Date/Time of Receipt: <u>03-19-19 9:50</u>		Number of Coolers Received: <u>1</u>						
Condition of Coolers: <u>Acceptable/Unacceptable</u>		Temperature Control: <u>Present/Not Included</u>						
Cooler Custody Seals: <u>Present/Absent/NA</u>		Location Temp Taken: <u>Control/Between Samples</u>						
Container Custody Seals: <u>Present/Absent/NA</u>		Are all temperatures within project specific guidelines? <u>Yes/No/NA</u>						
Ice Present: <u>Frozen/Melted/NA</u>		VOA Headspace Present? <u>Yes/No/NA</u>						
pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA		
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA		
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA		
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA		
Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9050</u>	<u>1</u> °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C
Taken By: <u>Tam Vantassel</u>		Signature		Printed Name		Date		<u>03-19-19</u>

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Part # 169469-434 RIT2 EXP 11/19

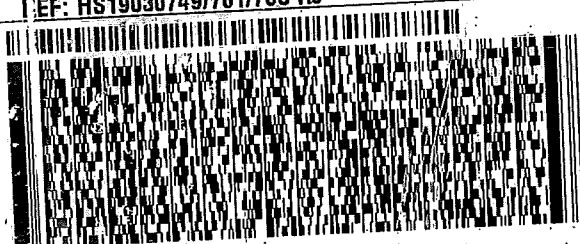
ORIGIN ID:SGRA (281) 530-5656
SHIPPING DEPT
ALS LABORATORY GROUP
10450 STANCLIFF RD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

SHIP DATE: 18MAR19
ACTWTG: 9.95 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(01) 288-7700
REF: HS19030749/761/763 RJ



FedEx
Express



11811116060606060606


TRK# 4809 7831 7725
0201

TUE - 19 MAR 3:00P
STANDARD OVERNIGHT

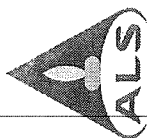
AX BTFA

84123
UT-US SLC




ALS
 10450 Stancliff Rd, Suite 210
 Houston, Texas 77099
 Tel. +1 281 530 5656
 Fax. +1 281 530 5987

Date
Name
Company



Batch Worklist

Batch: ELMS/ 2230 **Created:** 3/28/2019 07:45 **Instrument:** HBN: 235529
Rule: EPA 6850, DoD QSM Water **Analyst:** T. Bosch **Status:** WP



Workorder: 1907867 [ENV_LVL4]
Workorder: 1907869 [ENV_LVL4]
Workorder: 1907871 [ENV_LVL4]
Workorder: 1908794 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	64530	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	
2	64531	RLV'S for HBN 235529 [ELMS/2230]				RLV'S	3		E685041C3Q	5311		4/1/2019	
3	64532	ICS for HBN 235529 [ELMS/2230]				ICS	3		E6850.D3Q	5311		4/1/2019	
4	64533	LMB for HBN 235529 [ELMS/2230]				LMB	3		E6850Q413Q	5311		4/1/2019	
5	64534	LCS for HBN 235529 [ELMS/2230]				LCS	3		E6850Q413Q	5311		4/1/2019	
6	1907867001	LH18/24-SP140_031419				SAMPLE	3	1907867001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
7	1907869001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907869001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
8	64535	LH18/24-SP650...(1907869001MS)				MS	3		E6850Q413Q	5311		4/1/2019	
9	64536	LH18/24-SP65...(1907869001MSD)				MSD	3		E6850Q413Q	5311		4/1/2019	
10	1907871001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907871001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
11	1908794001	LH18/24-SP650_032119_BIX				SAMPLE	3	1908794001-A	E6850Q41.3	5480	4/18/2019	4/8/2019	
12	64537	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #'s: 1907867 (001); 1907869 (001); 1907871 (001); 1908794 (001)

ELMS Batch/HBN ID: 2230 (235529)

Prep Date: 03/27/2019 Analysis Date: 03/28/2019 Analyst: T. Bosch

Analyte: **Perchlorate** Matrix: **Water** Method: **6850**

Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\28MAR19D.s

Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by **TNB**. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 25µL
Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 645534; Target = 4.0µg/L. ASTM type II water was used for LMB 645533.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\235529-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
<u>Batch(es)/SDG: ELMS: 2230 HBN: 235529</u>		
<u>Sample Set IDs if Applicable: 1907867/1907869/1907871/1908794</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850 WKG Std 100 ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK			Description - 6850 Stock AccStd 1,000ug/mL
Standard: 43659		Created By: Thomas Bosch	
MFG: AccuStandard		Create Date: 09/17/2018 09:09AM	
MFG Lot: 218065075		Amount: 100 mL	
Part ID: IC-PER-10X-1		Expires: 07/25/2020	
		Usable: No	
		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT			Description - 6850 Intermdt AccStd 10.ug/mL		
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK			Description: 6850 QC WKG STD 100ug/L		
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK			Description - 6850 QC Stock STD 1,000ug/mL
Standard: 36748		Created By: Thomas Bosch	
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	
MFG Lot: CP-0860		Amount: 100 mL	
Part ID: ICC-013		Expires: 03/31/2020	
		Usable: Yes	
		Lab Lot: CLO4 QC STOCK	
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730		Created By: Thomas Bosch		Amount: 25 mL	
MFG: ALS/SLC		Create Date: 09/20/2018 09:09AM		Expires: 09/20/2019	
MFG Lot: TNB: 05/09/2018		Verified By: Thomas Bosch		Usable: Yes	
Pipette ID: Not Provided		Verify Date:		Lab Lot: CLO4ISTDWRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

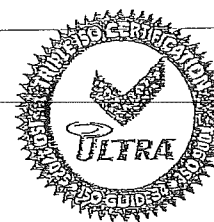
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:
This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

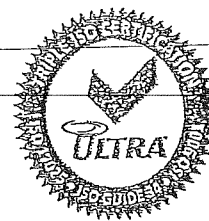
Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis

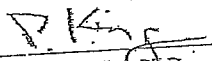


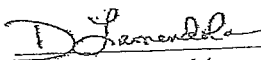
ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:
The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Larnendola
Director of QA/RA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

For use in routine laboratory analysis.

Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:

ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: NaCl*O4

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration data.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 µg/mL
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 ± 2.8 µg/mL (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
*	645530	CCV@25	Vial 71	1	Control	1	1.81499e6	24.13647
*	645534	QC@4.0	Vial 72	1	Control	2	2.89937e5	3.99582
*	645532	ICS@4.0	Vial 73	1	Control	3	2.38655e5	3.65624
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	7.55115e5	8344.77963
*	1907869001		Vial 76	1	Sample	6	0.00000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	2.77046e5	4.66163
*	645536	78691SD	Vial 78	1	Sample	8	2.89384e5	4.05130
*	1907871001		Vial 79	1	Sample	9	0.00000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	1.79709e6	24.49119

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
*	645530	CCV@25	Vial 71	1	Control	1	5.33395e5	23.90219
*	645534	QC@4.0	Vial 72	1	Control	2	9.52963e4	4.26699
*	645532	ICS@4.0	Vial 73	1	Control	3	7.88339e4	3.90865
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.38827e5	8762.35701
*	1907869001		Vial 76	1	Sample	6	0.00000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	8.93399e4	4.91371
*	645536	78691SD	Vial 78	1	Sample	8	9.85690e4	4.48457
*	1907871001		Vial 79	1	Sample	9	0.00000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	5.35802e5	24.59111

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
*	645530	CCV@25	Vial 71	1	Control	1	2.29255e5	5.00000
*	645534	QC@4.0	Vial 72	1	Control	2	2.39106e5	5.00000
*	645532	ICS@4.0	Vial 73	1	Control	3	2.16080e5	5.00000
*	645533	LMB	Vial 74	1	Control	4	2.42742e5	5.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.89136e5	5000.00000
*	1907869001		Vial 76	1	Sample	6	1.89925e5	5.00000
*	645535	78691MS	Vial 77	1	Sample	7	1.94413e5	5.00000
*	645536	78691SD	Vial 78	1	Sample	8	2.35220e5	5.00000
*	1907871001		Vial 79	1	Sample	9	1.89609e5	5.00000
*	1908794001		Vial 80	1	Sample	10	1.92453e5	5.00000
*	645537	CCV@25	Vial 71	1	Control	11	2.23514e5	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	645530	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	645534	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	645532	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	645533	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1907867001	1K	CLO4-AQN	1	Sample	
6	Vial 76	1907869001		CLO4-AQN	1	Sample	
7	Vial 77	645535	78691MS	CLO4-AQN	1	Sample	
8	Vial 78	645536	78691SD	CLO4-AQN	1	Sample	
9	Vial 79	1907871001		CLO4-AQN	1	Sample	
10	Vial 80	1908794001		CLO4-AQN	1	Sample	
11	Vial 71	645537	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D

Sample Name: 645530 CCV@25

Injection Date: 3/28/2019 08:30:53

Seq Line: 1

Sample Name: 645530 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

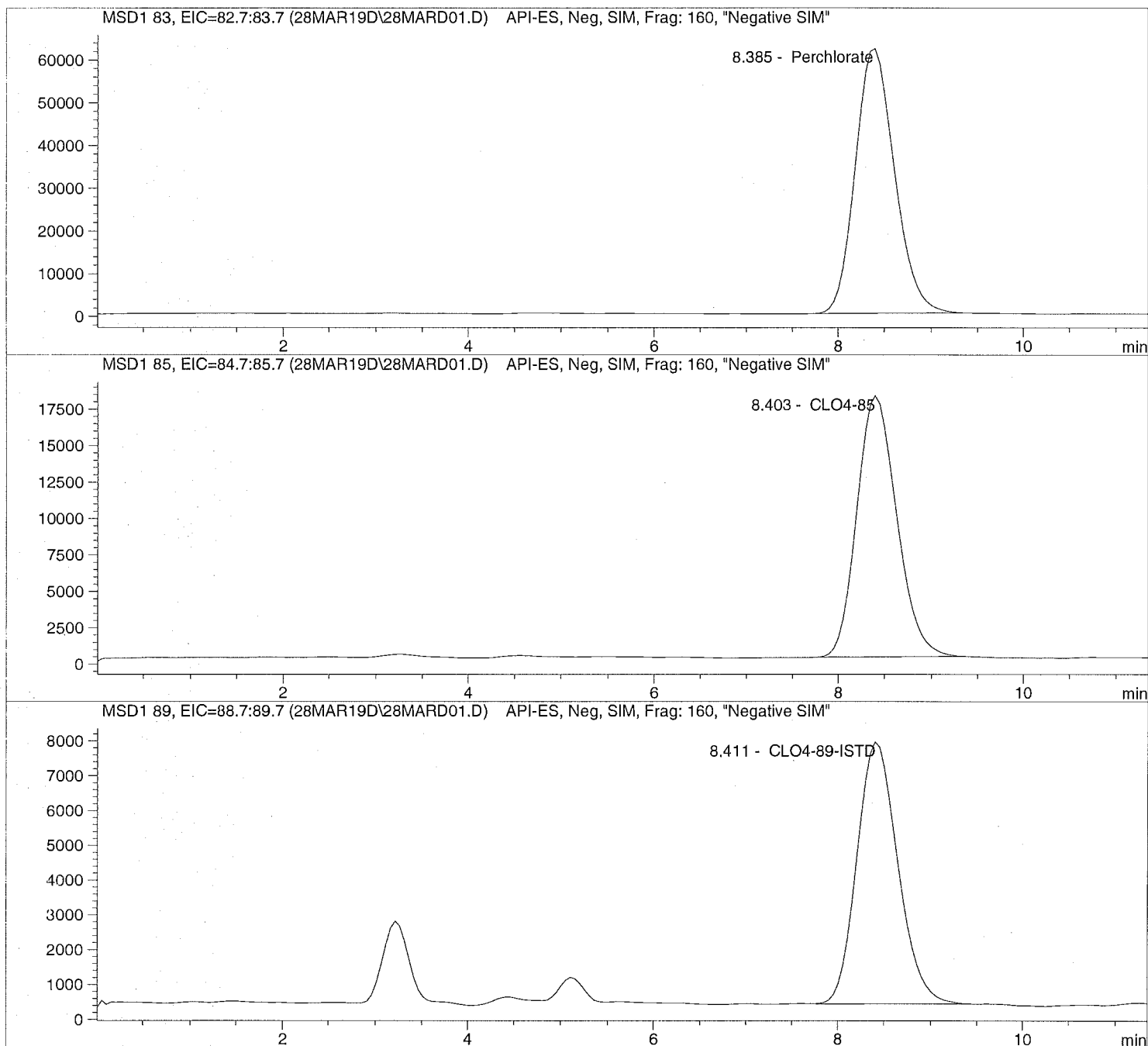
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D Sample Name: 645530 CCV@25

```

=====
Injection Date: 3/28/2019 08:30:53      Seq Line: 1
Sample Name:    645530 CCV@25          Location: Vial 71
Acq Operator:   TNB                    Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis

Sample Information

```

=====
Sorted By:          Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:        1.000000
Dilution:          1.000000
Sample Amount:     25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.385	PBA	1814992.7	24.1365	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.403	PBA	533395.5	23.9022	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.411	BBA	229255.1	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D

Sample Name: 645534 QC@4.0

Injection Date: 3/28/2019 08:47:42

Seq Line: 2

Sample Name: 645534 QC@4.0

Location: Vial 72

Acq Operator: TNB

Inj. No.: 1

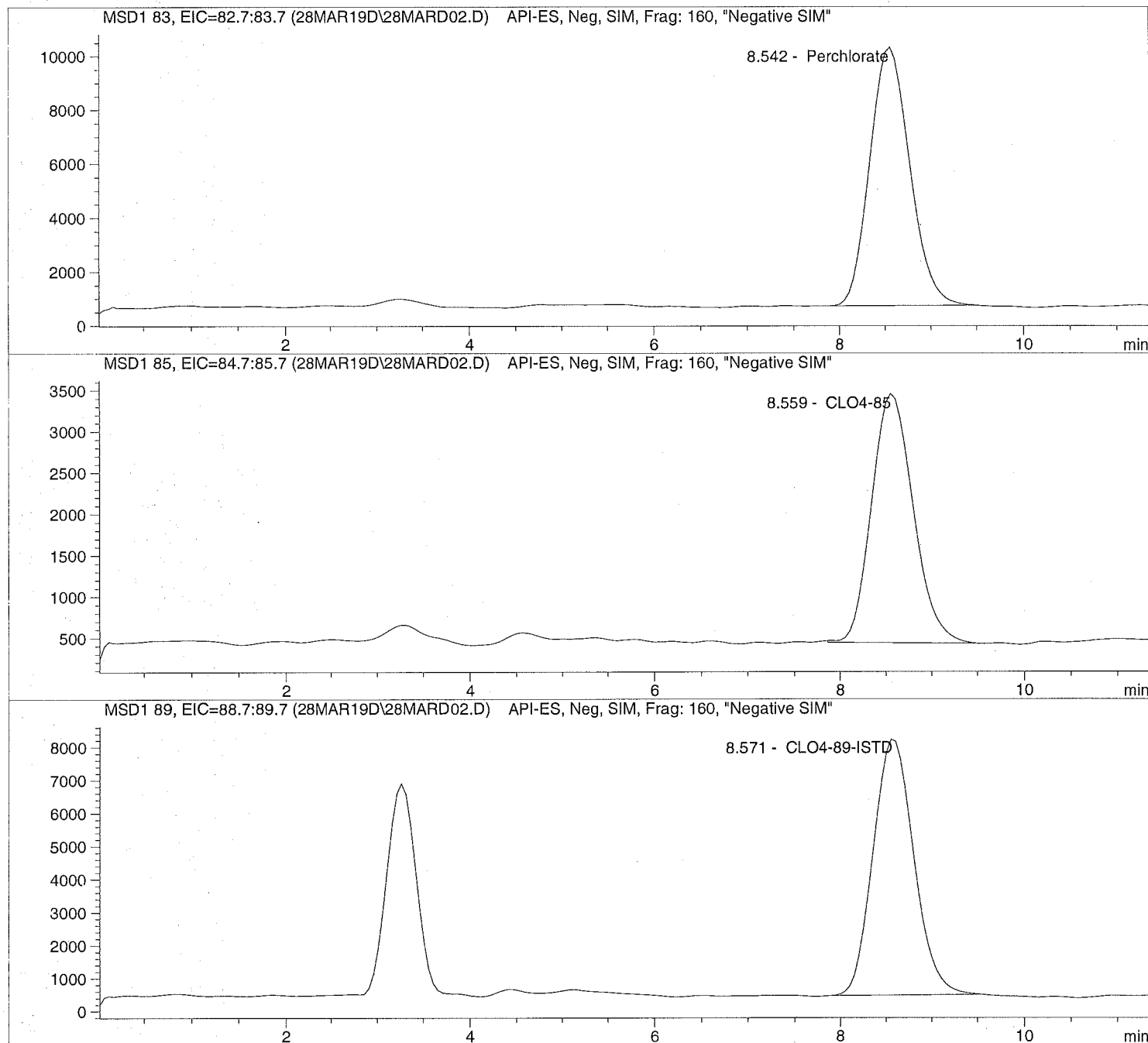
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D Sample Name: 645534 QC@4.0

```

=====
Injection Date: 3/28/2019 08:47:42      Seq Line: 2
Sample Name: 645534    QC@4.0      Location: Vial 72
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 25 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.542	PBA	289937.1	3.9958	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.559	BBA	95296.3	4.2670	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.571	PBA	239105.9	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD03.D

Sample Name: 645532 ICS@4.0

Injection Date: 3/28/2019 09:00:54
Sample Name: 645532 ICS@4.0
Acq Operator: TNB

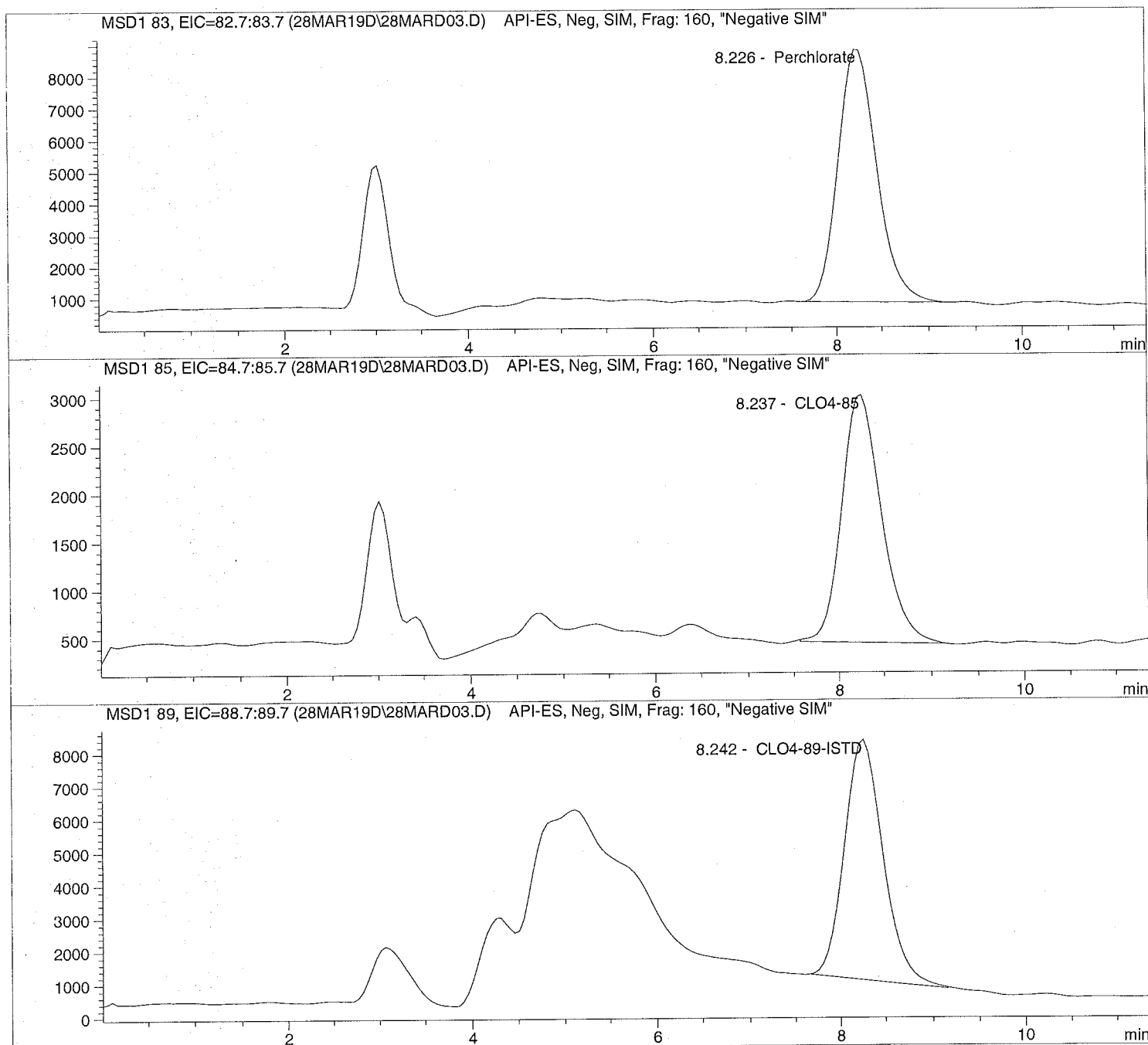
Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD03.D Sample Name: 645532 ICS@4.0

```

=====
Injection Date: 3/28/2019 09:00:54      Seq Line: 3
Sample Name: 645532 ICS@4.0            Location: Vial 73
Acq Operator: TNB                        Inj. No.: 1
                                          Inj. Vol.: 25 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

=====
Sample Information
=====

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000

```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.226	BBA	238654.8	3.6562	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.237	BBA	78833.9	3.9087	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.242	PBA	216079.7	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D

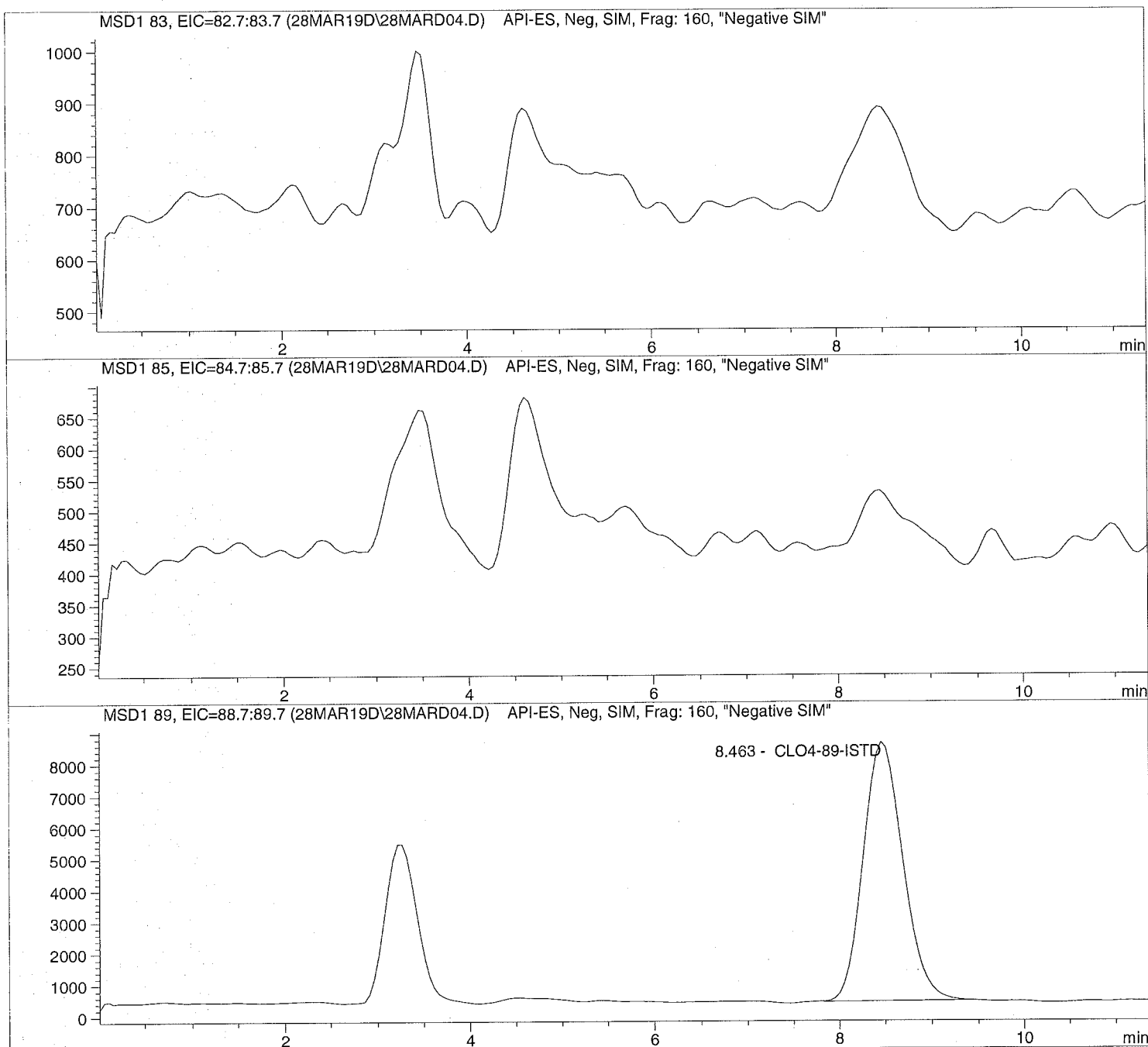
Sample Name: 645533 LMB

Injection Date: 3/28/2019 09:14:08
Sample Name: 645533 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D Sample Name: 645533 LMB

```

=====
Injection Date: 3/28/2019 09:14:08      Seq Line: 4
Sample Name: 645533 LMB                  Location: Vial 74
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.463	BBA	242742.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

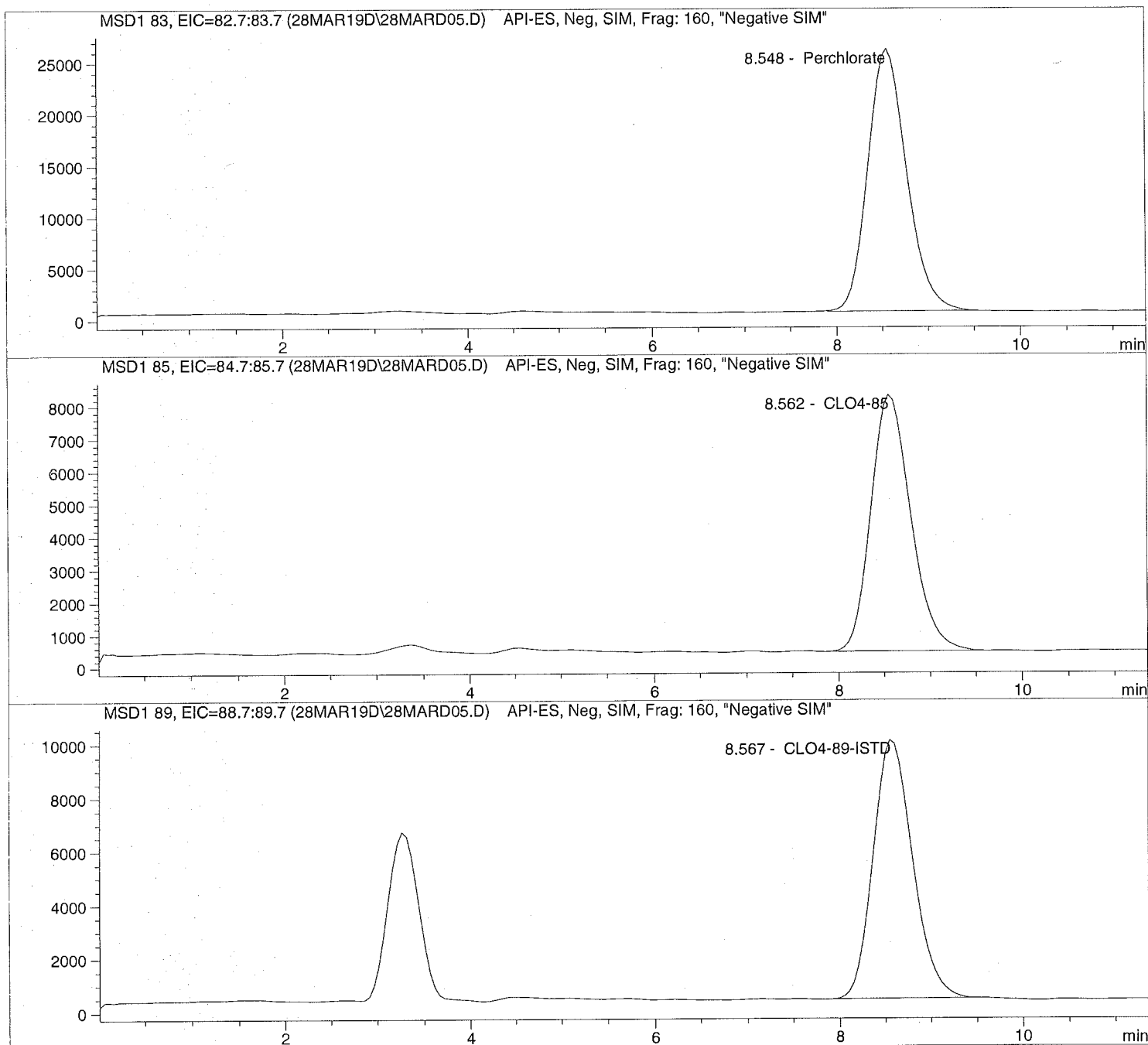
```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```
=====
Injection Date: 3/28/2019 09:28:25      Seq Line:      5
Sample Name:    1907867001 1K           Location:      Vial 75
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```

=====
Injection Date: 3/28/2019 09:28:25      Seq Line:      5
Sample Name:    1907867001 1K          Location:      Vial 75
Acq Operator:   TNB                    Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1000.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.548	BBA	755115.1	8344.7796	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.562	PBA	238826.8	8762.3570	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.567	PBA	289135.6	5000.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D

Sample Name: 1907869001

=====
Injection Date: 3/28/2019 09:41:37
Sample Name: 1907869001
Acq Operator: TNB

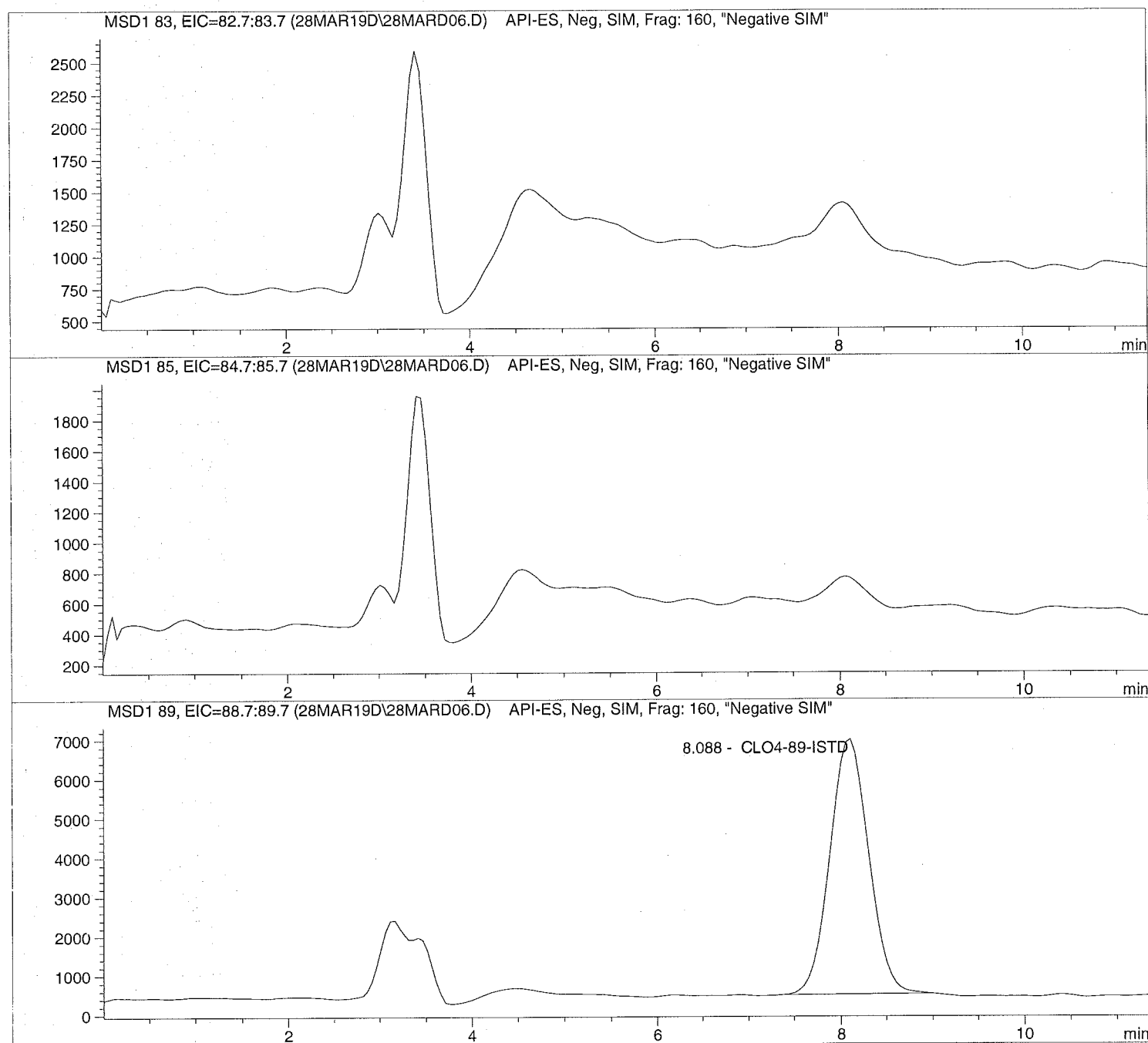
Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D

Sample Name: 1907869001

```

=====
Injection Date:  3/28/2019  09:41:37      Seq Line:      6
Sample Name:    1907869001      Location:      Vial 76
Acq Operator:   TNB              Inj. No.:     1
                                      Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.088	BBA	189925.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

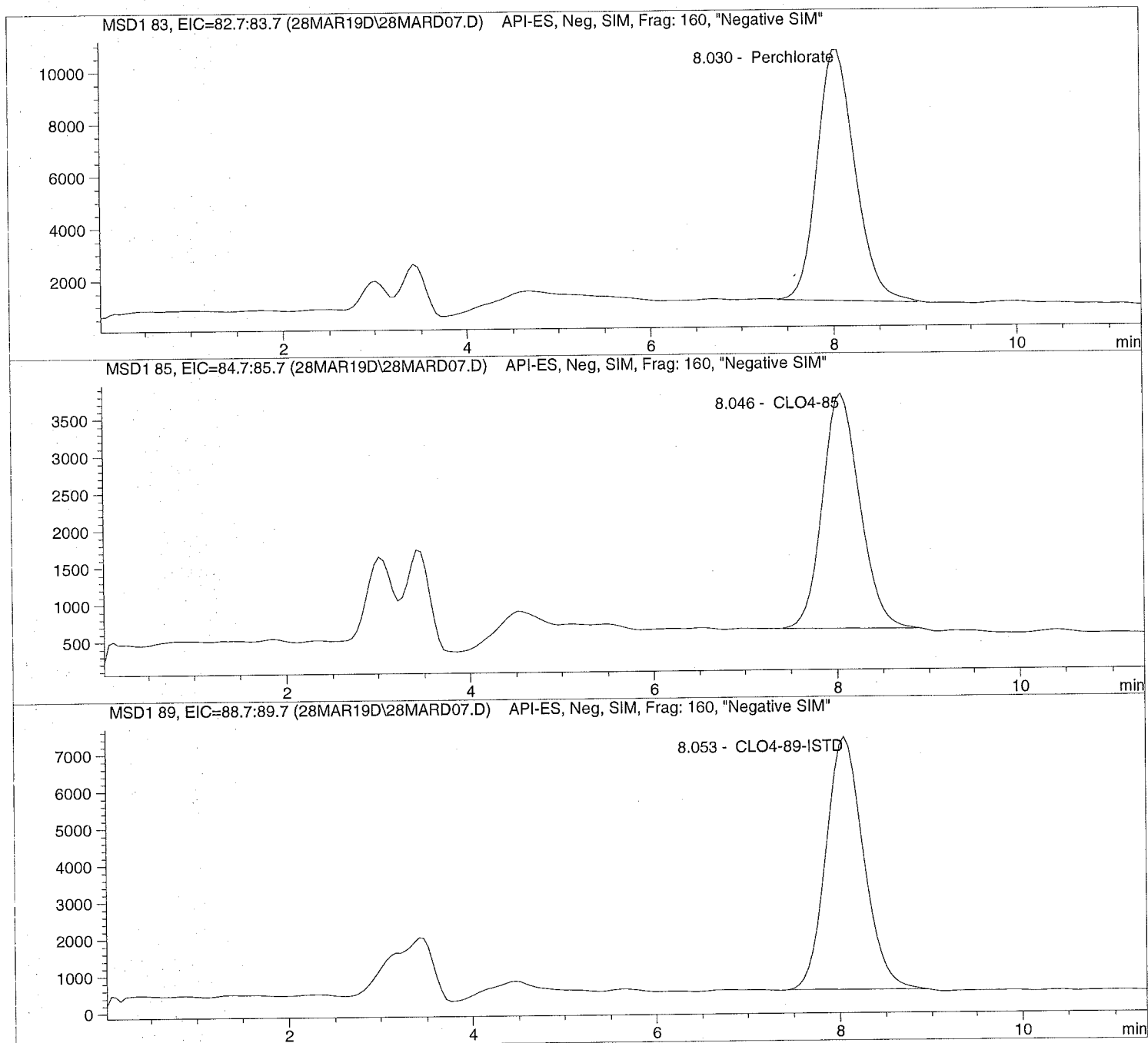
```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D Sample Name: 645535 78691MS

```
=====
Injection Date: 3/28/2019 09:54:47      Seq Line: 7
Sample Name:    645535 78691MS          Location:  Vial 77
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:  CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D Sample Name: 645535 78691MS

```

=====
Injection Date: 3/28/2019 09:54:47      Seq Line: 7
Sample Name:    645535 78691MS          Location:  Vial 77
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.030	BBA	277046.3	4.6616	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.046	BBA	89339.9	4.9137	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	194412.9	5.0000	CLO4-89-ISTD

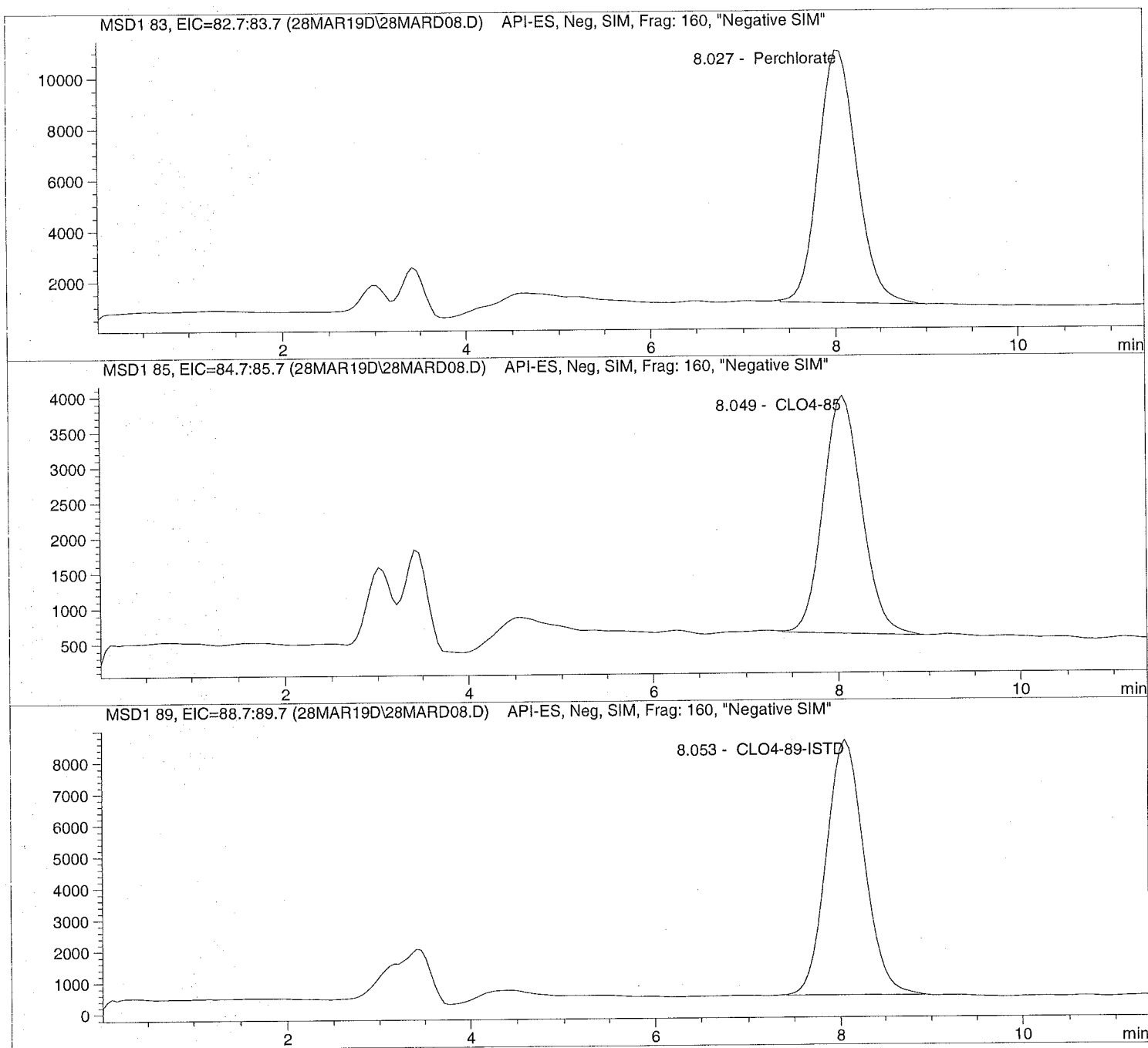
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D Sample Name: 645536 78691SD

```
=====
Injection Date: 3/28/2019 10:08:05 Seq Line: 8
Sample Name: 645536 78691SD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D Sample Name: 645536 78691SD

```

=====
Injection Date: 3/28/2019 10:08:05 Seq Line: 8
Sample Name: 645536 78691SD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.027	BBA	289383.7	4.0513	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.049	BBA	98569.0	4.4846	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	235219.8	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D

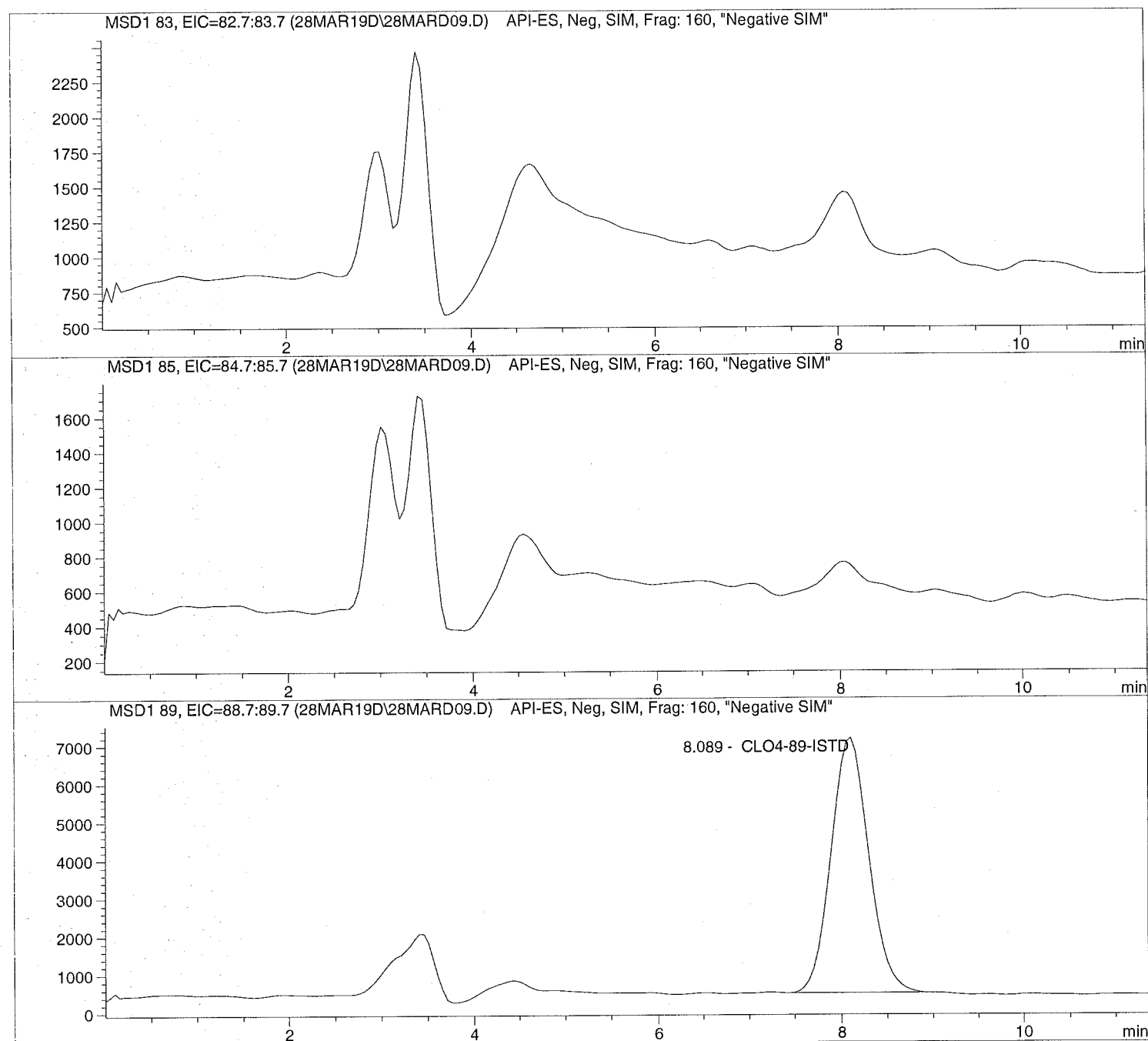
Sample Name: 1907871001

=====
Injection Date: 3/28/2019 10:21:14
Sample Name: 1907871001
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D

Sample Name: 1907871001

```

=====
Injection Date:  3/28/2019  10:21:14      Seq Line:      9
Sample Name:    1907871001              Location:      Vial 79
Acq Operator:   TNB                      Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified:  Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.089	PBA	189608.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD10.D

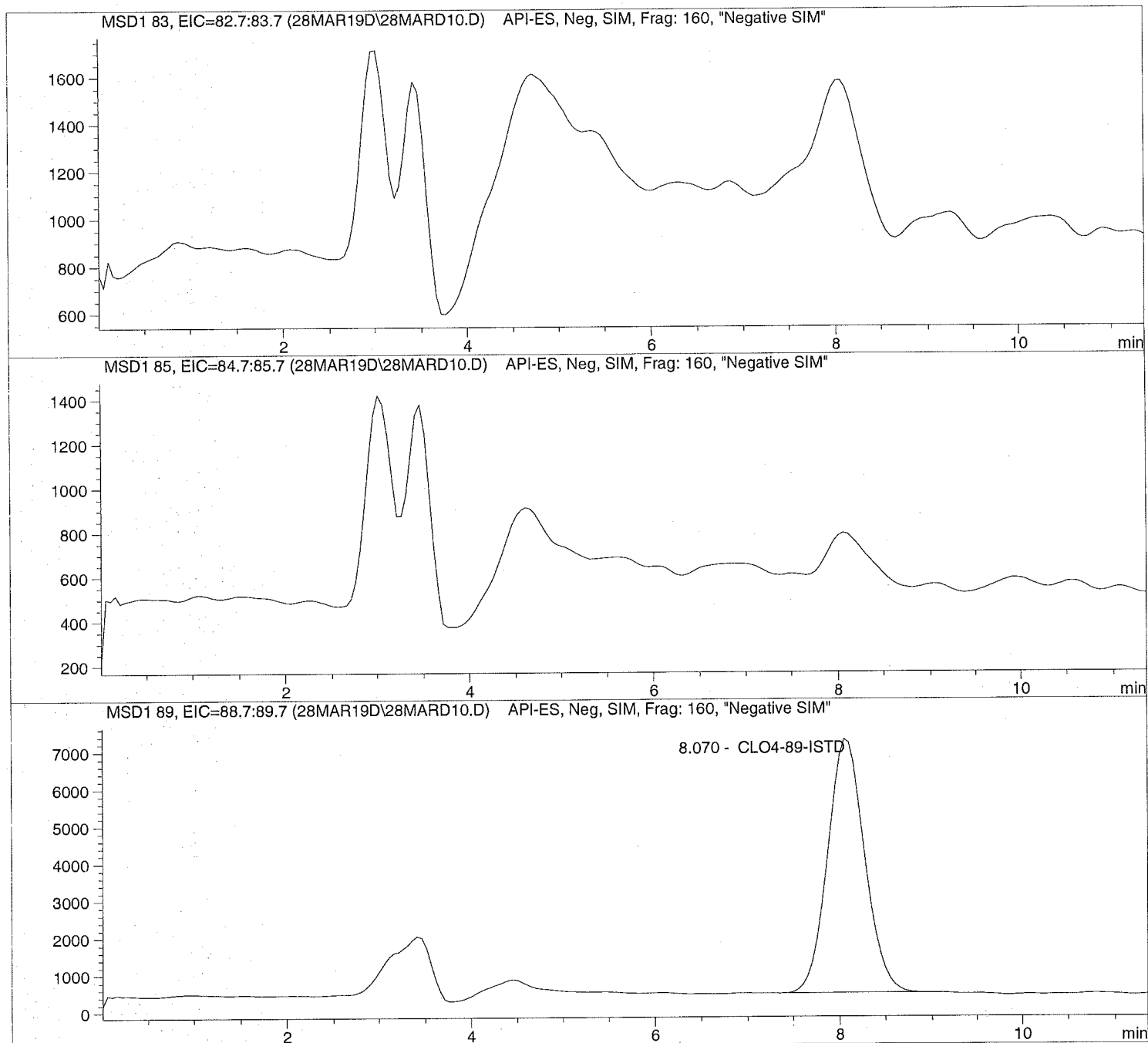
Sample Name: 1908794001

Injection Date: 3/28/2019 10:34:27
Sample Name: 1908794001
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD10.D Sample Name: 1908794001

```

=====
Injection Date: 3/28/2019 10:34:27      Seq Line: 10
Sample Name: 1908794001                Location: Vial 80
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.070	PBA	192453.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D

Sample Name: 645537 CCV@25

Injection Date: 3/28/2019 10:49:41

Seq Line: 11

Sample Name: 645537 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

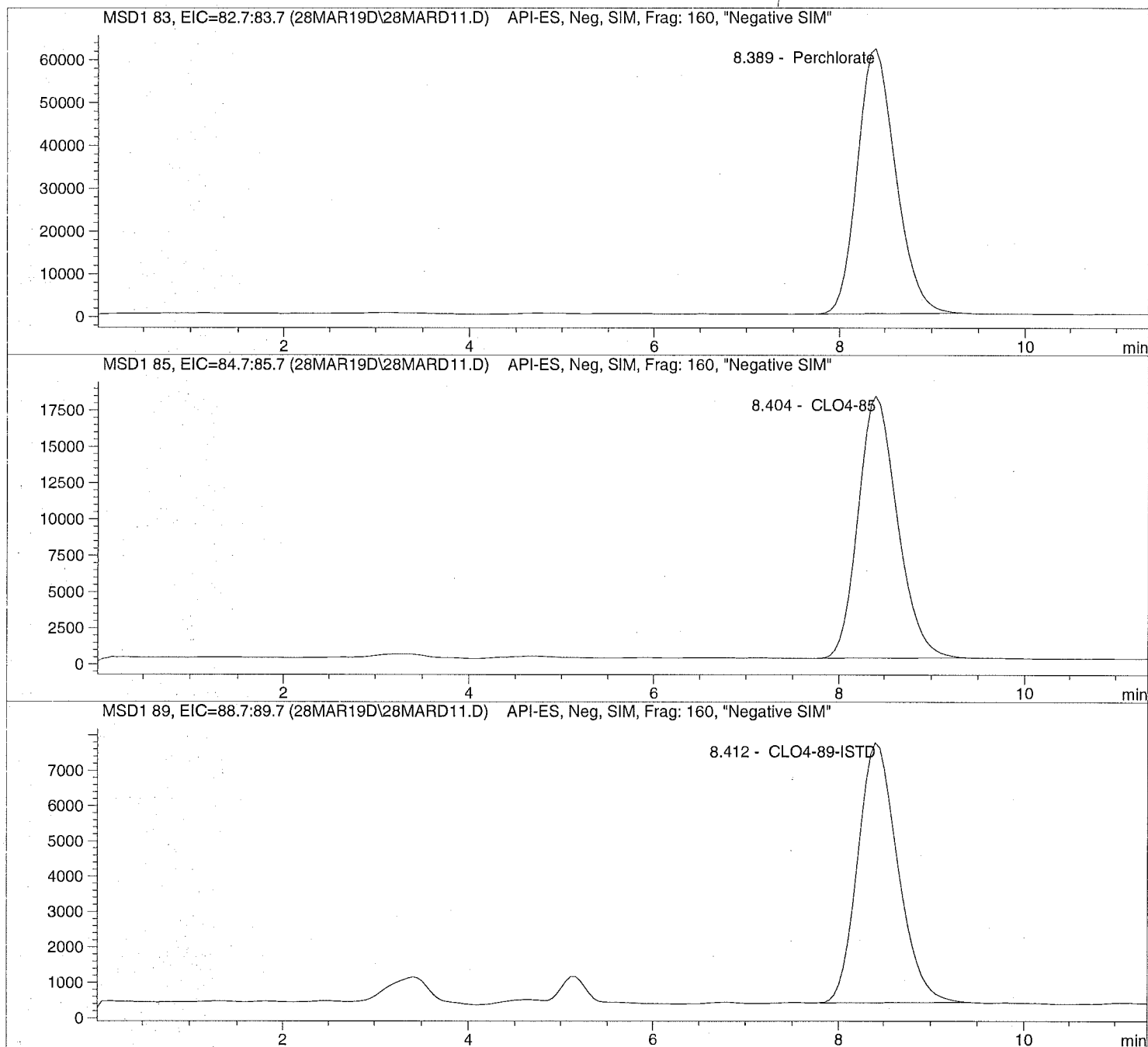
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D Sample Name: 645537 CCV@25

```

=====
Injection Date:  3/28/2019  10:49:41                    Seq Line:                    11
Sample Name:     645537    CCV@25                      Location:                    Vial 71
Acq Operator:    TNB                                    Inj. No.:                    1
                                                          Inj. Vol.:                    25 µl
=====

```

```

Acq. Method:     CLO4-AQN.M
Analysis Method:  C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:    3/19/2019  14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:                    Signal
Calib. Data Modified:        Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                  1.000000
Dilution:                    1.000000
Sample Amount:                25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.389	PBA	1797085.1	24.4912	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.404	PBA	535801.9	24.5911	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.412	BBA	223514.2	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
 Calibration Table
 =====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
 Based on : Peak Area

Rel. Reference Window : 20.000 %
 Abs. Reference Window : 0.000 min
 Rel. Non-ref. Window : 20.000 %
 Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
 Uncalibrated Peaks : not reported
 Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
 Origin : Ignored (some peaks differ, see below)
 Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
 Average Response : Average all calibrations
 Average Retention Time: Floating Average New 75%

Calibration Report Options :
 Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
 If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
 Signal 2: MSD1 85, EIC=84.7:85.7
 Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl	Amount	Area	Amt/Area	Ref	Grp	Name
8.744	1	1.00000	7.76074e4	1.28854e-5	1		Perchlorate
	2	2.00000	1.35273e5	1.47849e-5			
	3	5.00000	3.37764e5	1.48033e-5			
	4	10.00000	6.83454e5	1.46316e-5			
	5	25.00000	2.08433e6	1.19943e-5			
	6	50.00000	4.13334e6	1.20968e-5			
	7	75.00000	5.99313e6	1.25143e-5			
8.755	2	1.00000	2.36780e4	4.22333e-5	1		CLO4-85
	2	2.00000	4.69486e4	4.25998e-5			
	3	5.00000	1.06124e5	4.71147e-5			
	4	10.00000	2.13523e5	4.68335e-5			
	5	25.00000	6.14295e5	4.06971e-5			
	6	50.00000	1.19814e6	4.17315e-5			
	7	75.00000	1.78355e6	4.20509e-5			
8.766	3	5.00000	2.73208e5	1.83011e-5	+I1		CLO4-89-ISTD
	2	5.00000	2.24886e5	2.22335e-5			
	3	5.00000	2.33196e5	2.14412e-5			
	4	5.00000	2.34454e5	2.13262e-5			
	5	5.00000	2.50568e5	1.99547e-5			
	6	5.00000	2.30977e5	2.16472e-5			

RetTime	Lvl	Amount	Area	Amt/Area	Ref Grp Name
[min]	Sig				
7		5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-85

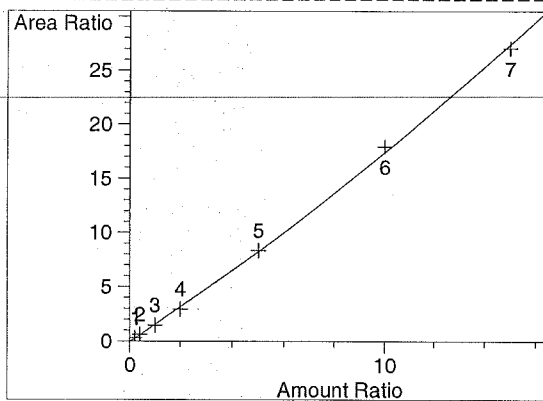
Time Window : From 6.650 min To 12.505 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-89-ISTD

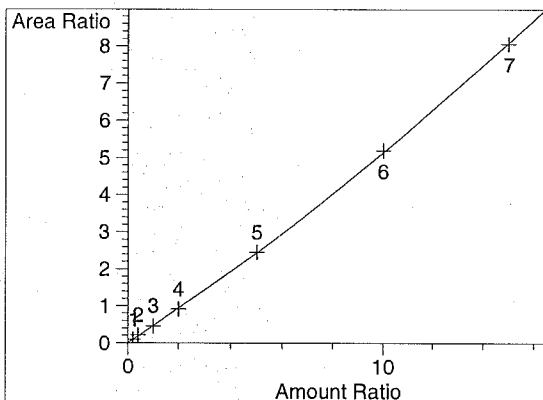
Time Window : From 6.659 min To 12.466 min
Curve Type : Linear
Origin : Included
Calibration Level Weights:/
Level 1 : 1
Level 2 : 1
Level 3 : 1
Level 4 : 1
Level 5 : 1
Level 6 : 1
Level 7 : 1

=====
Peak Sum Table
=====***No Entries in table***
=====

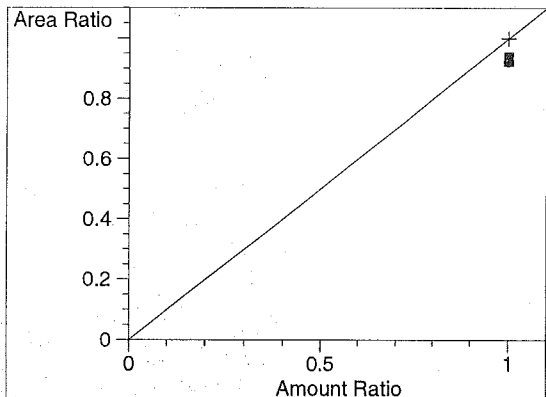
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

##	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
##	---	---	---	---	---	---	---	---
##	CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
##	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
##	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
##	CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
##	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
##	CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
##	CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
##	ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

##	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
##	---	---	---	---	---	---	---	---
##	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
##	CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
##	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
##	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
##	CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
##	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
##	CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
##	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

##	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
##	---	---	---	---	---	---	---	---
##	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
##	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
##	CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
##	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
##	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
##	CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
##	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
##	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

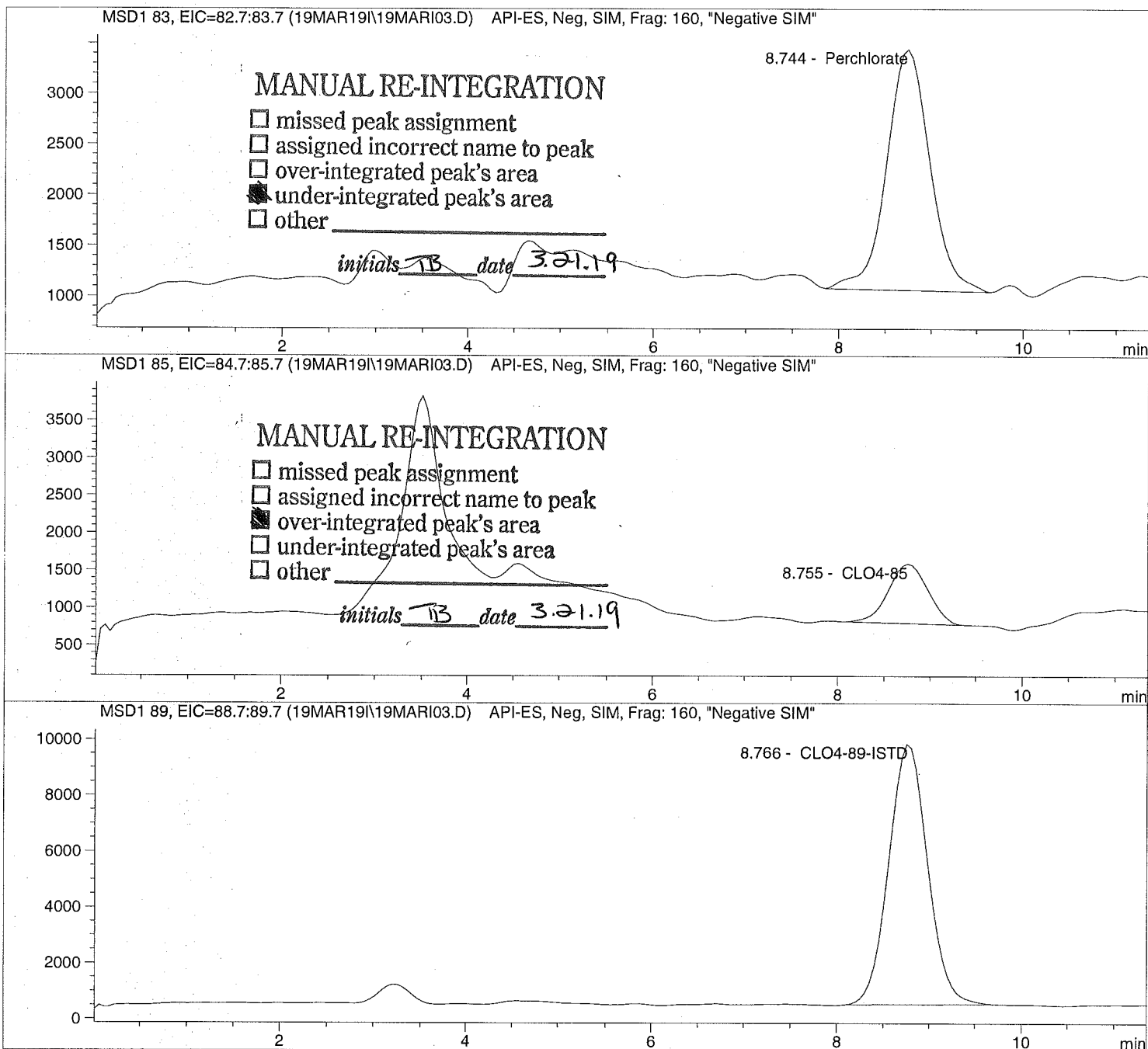
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:      3
Sample Name:    CLO4@ 1.0ug/L           Location:      Vial 73
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D

Sample Name: CLO4@ 2.0ug/L

Injection Date: 3/19/2019 09:53:00

Seq Line: 4

Sample Name: CLO4@ 2.0ug/L

Location: Vial 74

Acq Operator: TNB

Inj. No.: 1

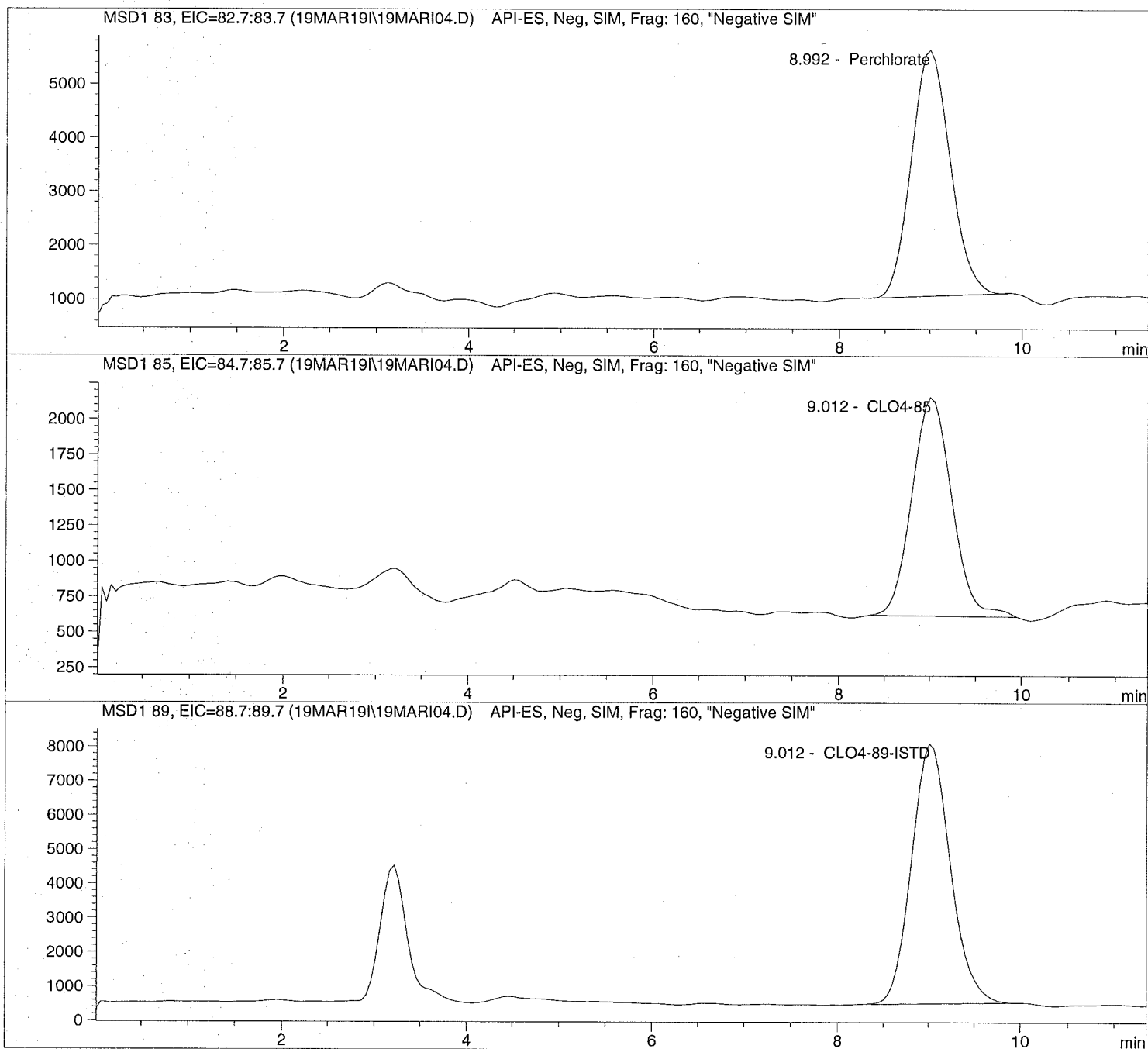
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line:      4
Sample Name:    CLO4@ 2.0ug/L          Location:      Vial 74
Acq Operator:   TNB                    Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D

Sample Name: CLO4@ 5.0ug/L

Injection Date: 3/19/2019 10:06:16

Seq Line: 5

Sample Name: CLO4@ 5.0ug/L

Location: Vial 75

Acq Operator: TNB

Inj. No.: 1

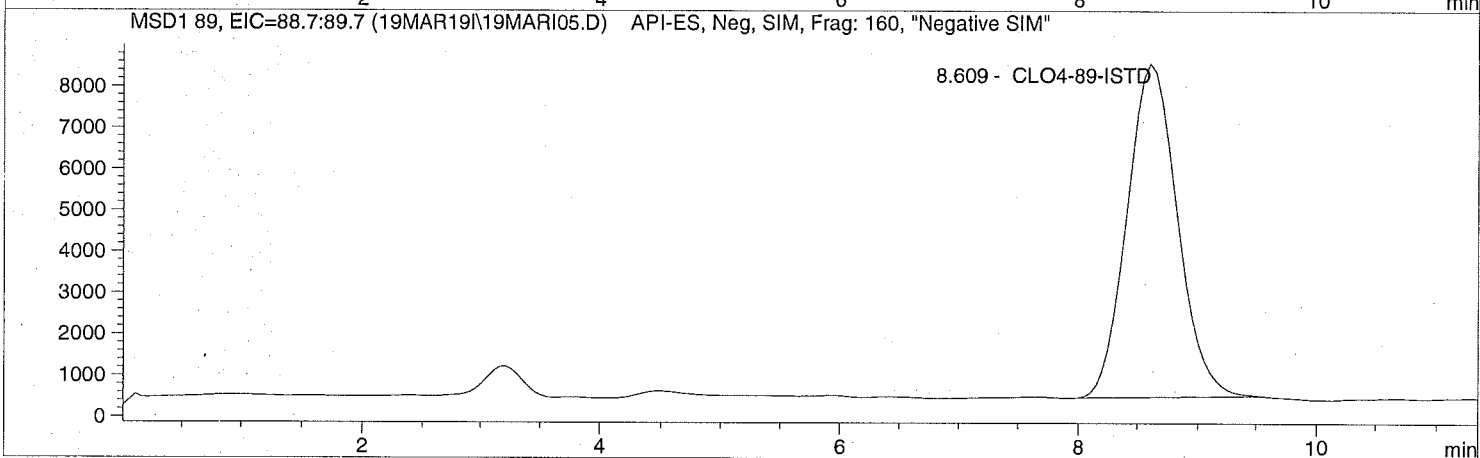
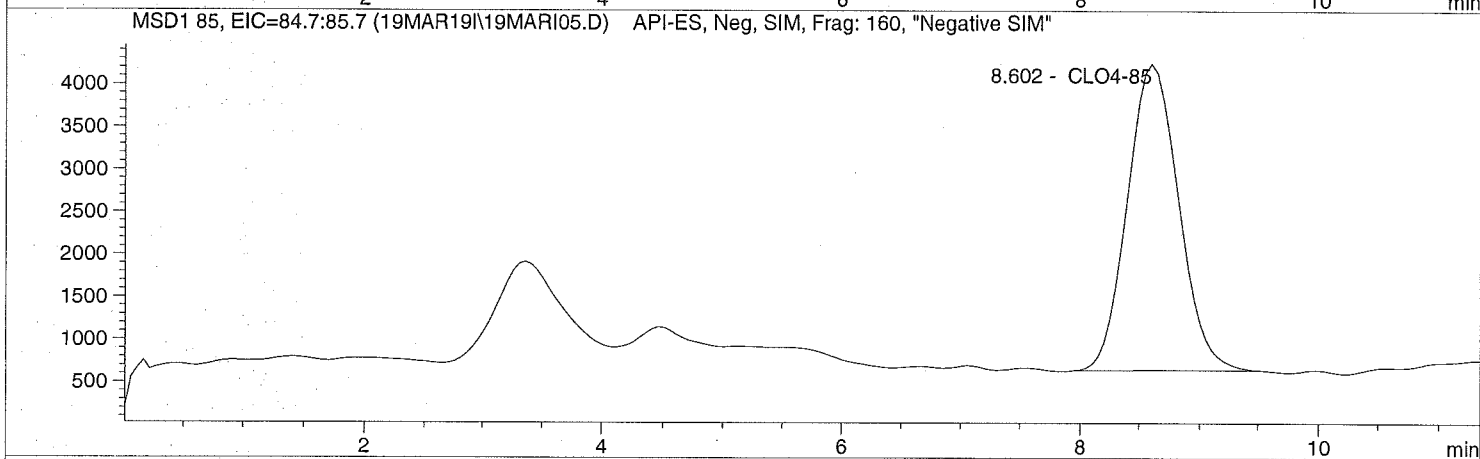
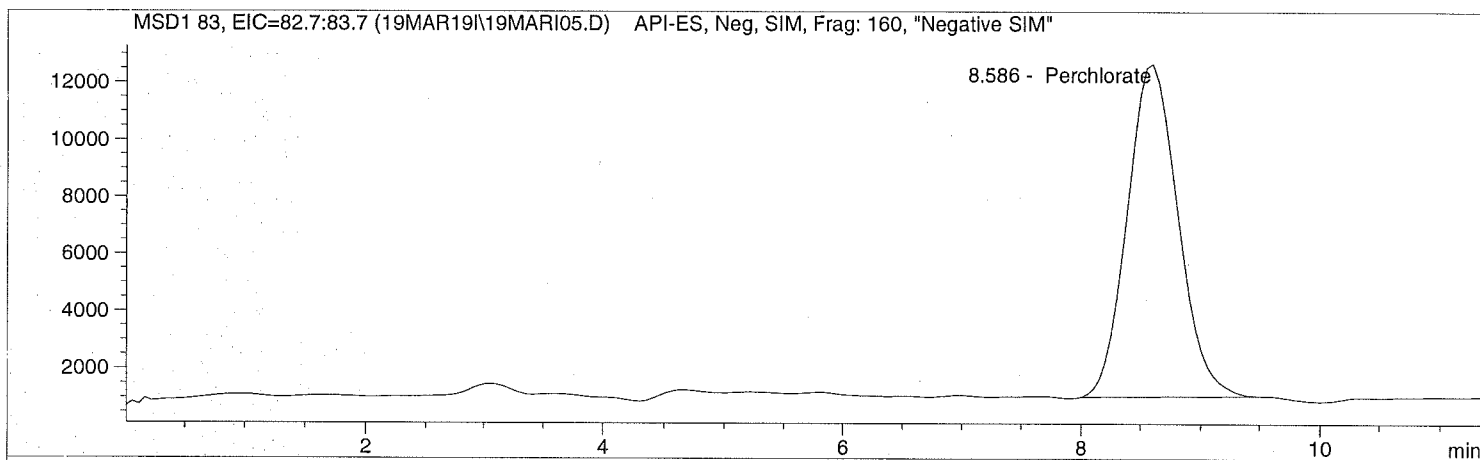
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line:      5
Sample Name:    CLO4@ 5.0ug/L           Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D

Sample Name: CLO4@ 10.ug/L

Injection Date: 3/19/2019 10:19:32

Seq Line: 6

Sample Name: CLO4@ 10.ug/L

Location: Vial 76

Acq Operator: TNB

Inj. No.: 1

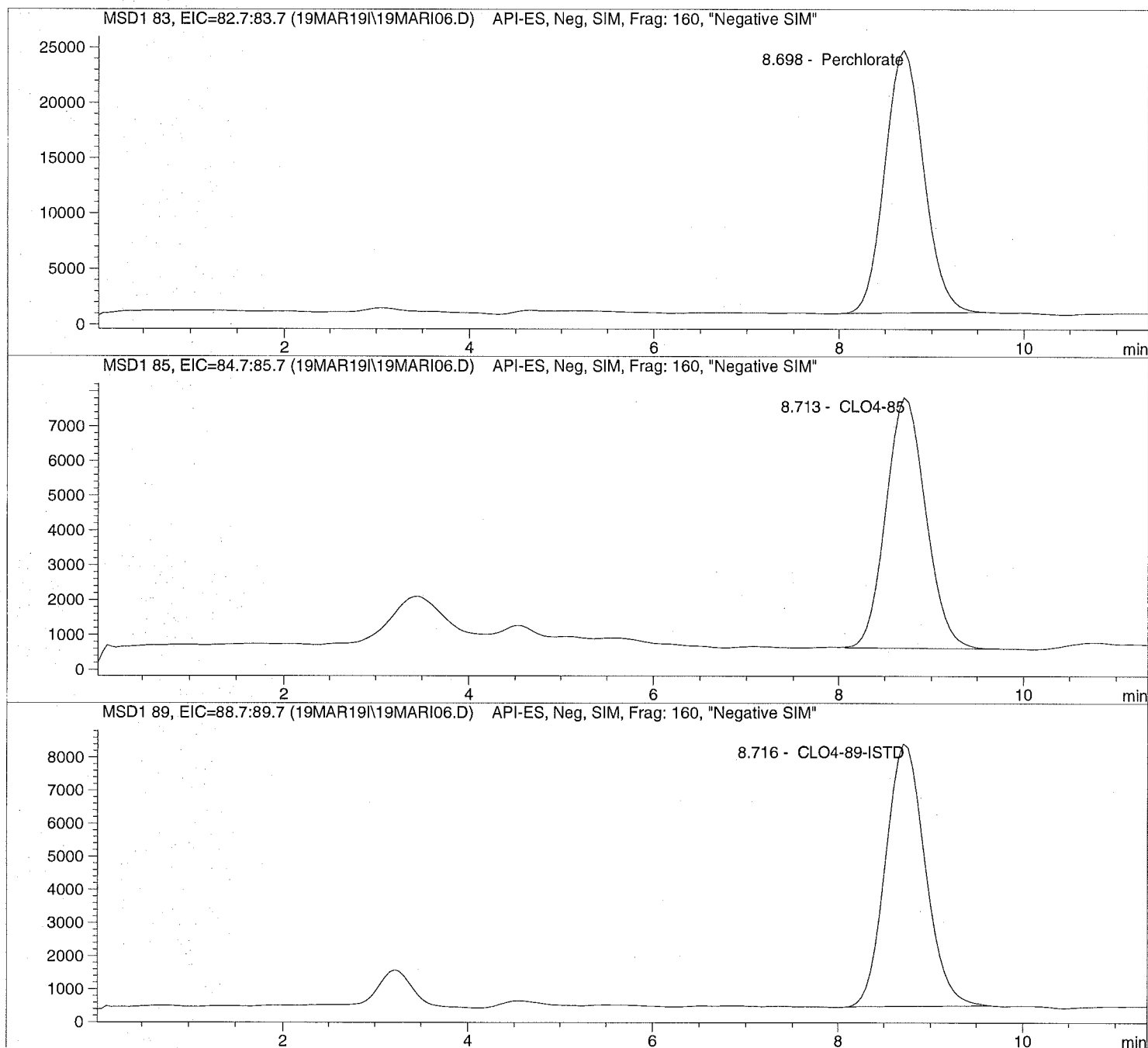
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line:            6
Sample Name:    CLO4@ 10.ug/L            Location:            Vial 76
Acq Operator:   TNB                      Inj. No.:            1
                                         Inj. Vol.:            30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22

```

Perchlorate analysis

Sample Information

```

Sorted By:            Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:          1.000000
Dilution:            1.000000
Sample Amount:        10.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D

Sample Name: CLO4@ 25.ug/L

Injection Date: 3/19/2019 10:32:49

Seq Line: 7

Sample Name: CLO4@ 25.ug/L

Location: Vial 77

Acq Operator: TNB

Inj. No.: 1

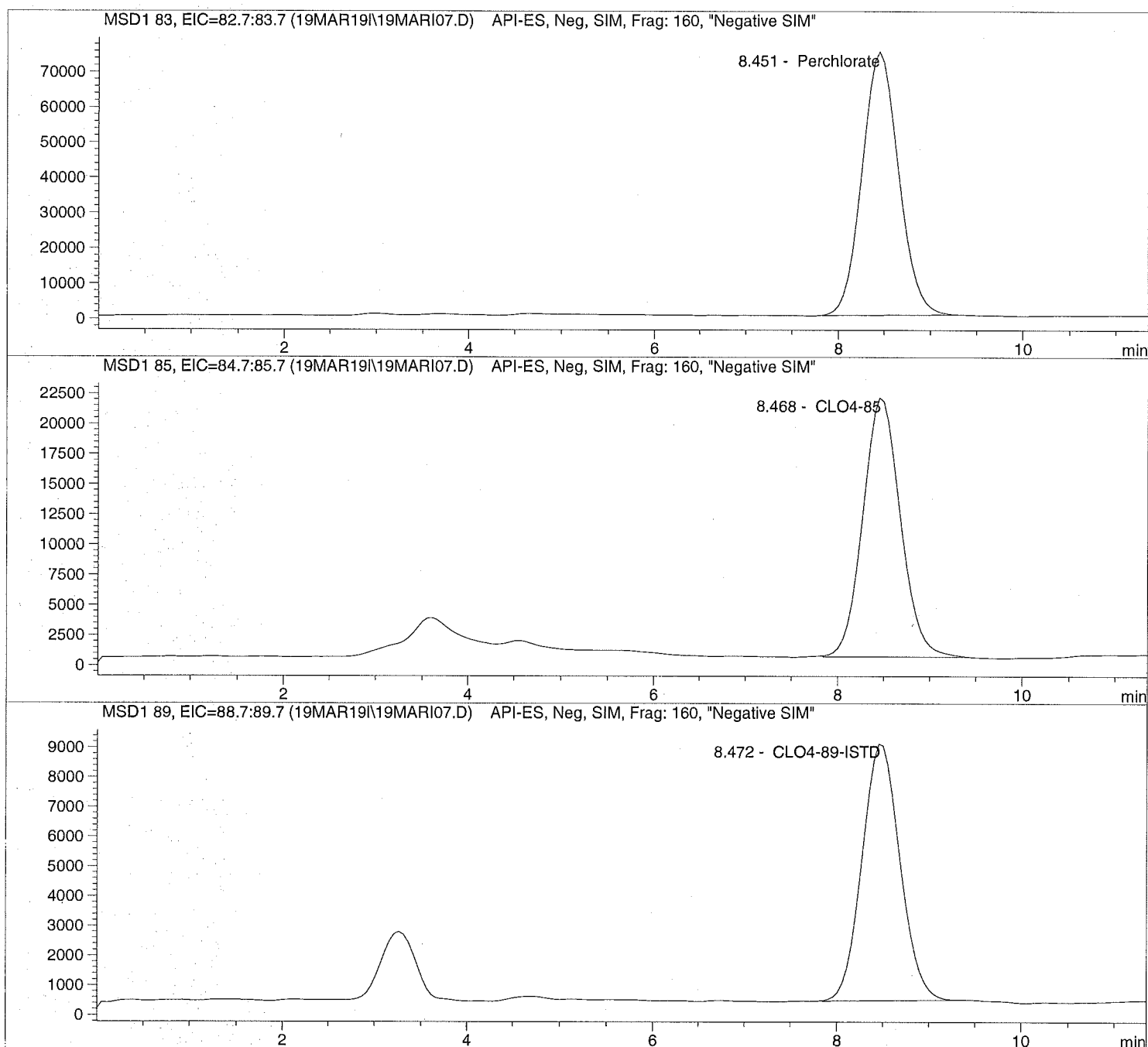
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```
=====
Injection Date:  3/19/2019  10:32:49                    Seq Line:                    7
Sample Name:     CLO4@ 25.ug/L                         Location:                    Vial 77
Acq Operator:    TNB                                    Inj. No.:                    1
                                                         Inj. Vol.:                    30 µl
=====
```

```
Acq. Method:     CLO4-AQN.M
Analysis Method:  C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:    3/19/2019  14:35:22
```

Perchlorate analysis

=====

Sample Information

=====

```
Sorted By:                    Signal
Calib. Data Modified:        Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                  1.000000
Dilution:                    1.000000
Sample Amount:                25.000
```

=====

LCMS Results

=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

=====

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D

Sample Name: CLO4@ 50.ug/L

Injection Date: 3/19/2019 10:46:05

Seq Line: 8

Sample Name: CLO4@ 50.ug/L

Location: Vial 78

Acq Operator: TNB

Inj. No.: 1

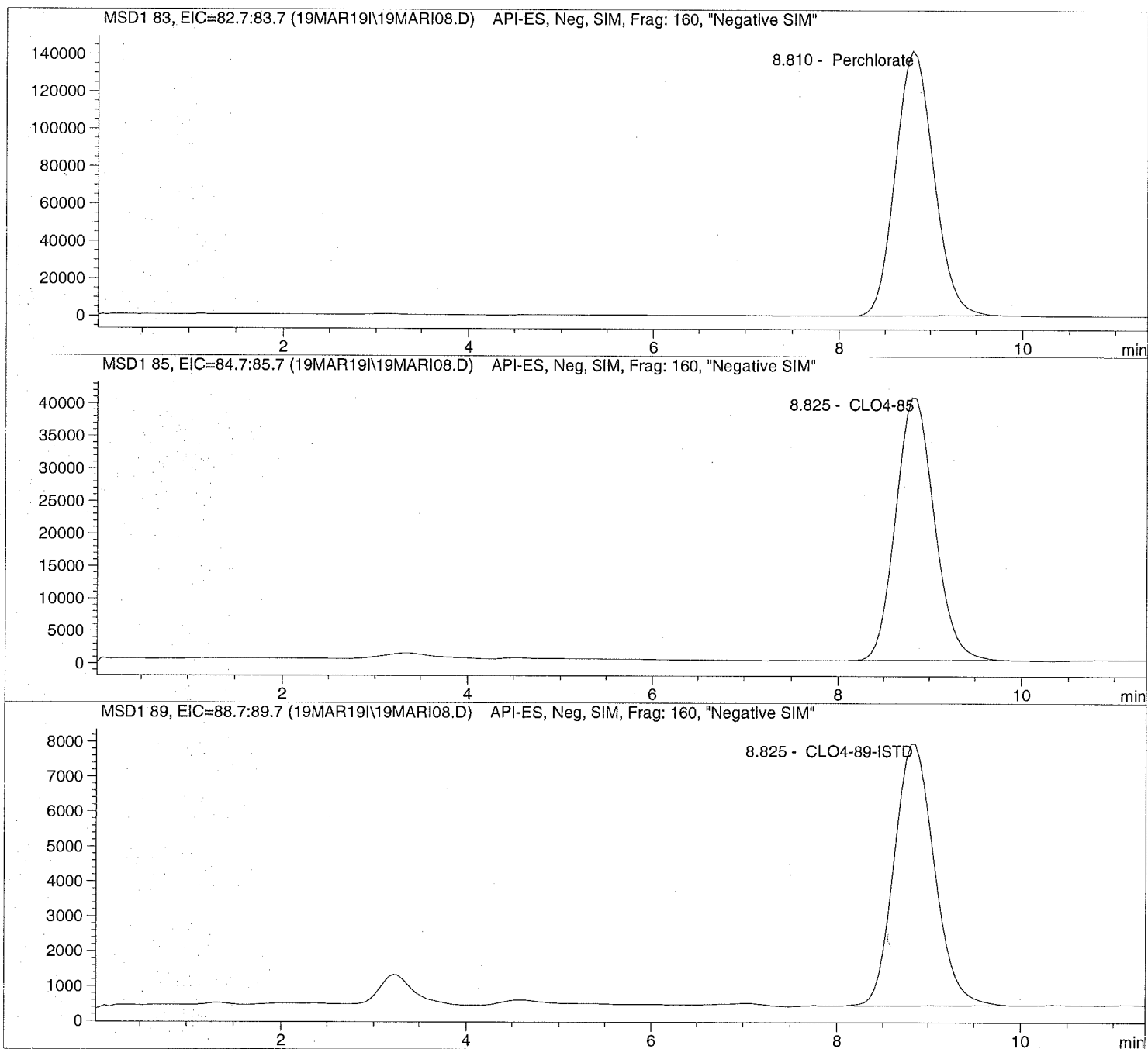
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 3/19/2019 10:46:05      Seq Line:                8
Sample Name:    CLO4@ 50.ug/L            Location:                Vial 78
Acq Operator:   TNB                      Inj. No.:                1
                                         Inj. Vol.:                30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:                                Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                               1.000000
Dilution:                                 1.000000
Sample Amount:                            50.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 3/19/2019 10:59:22

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

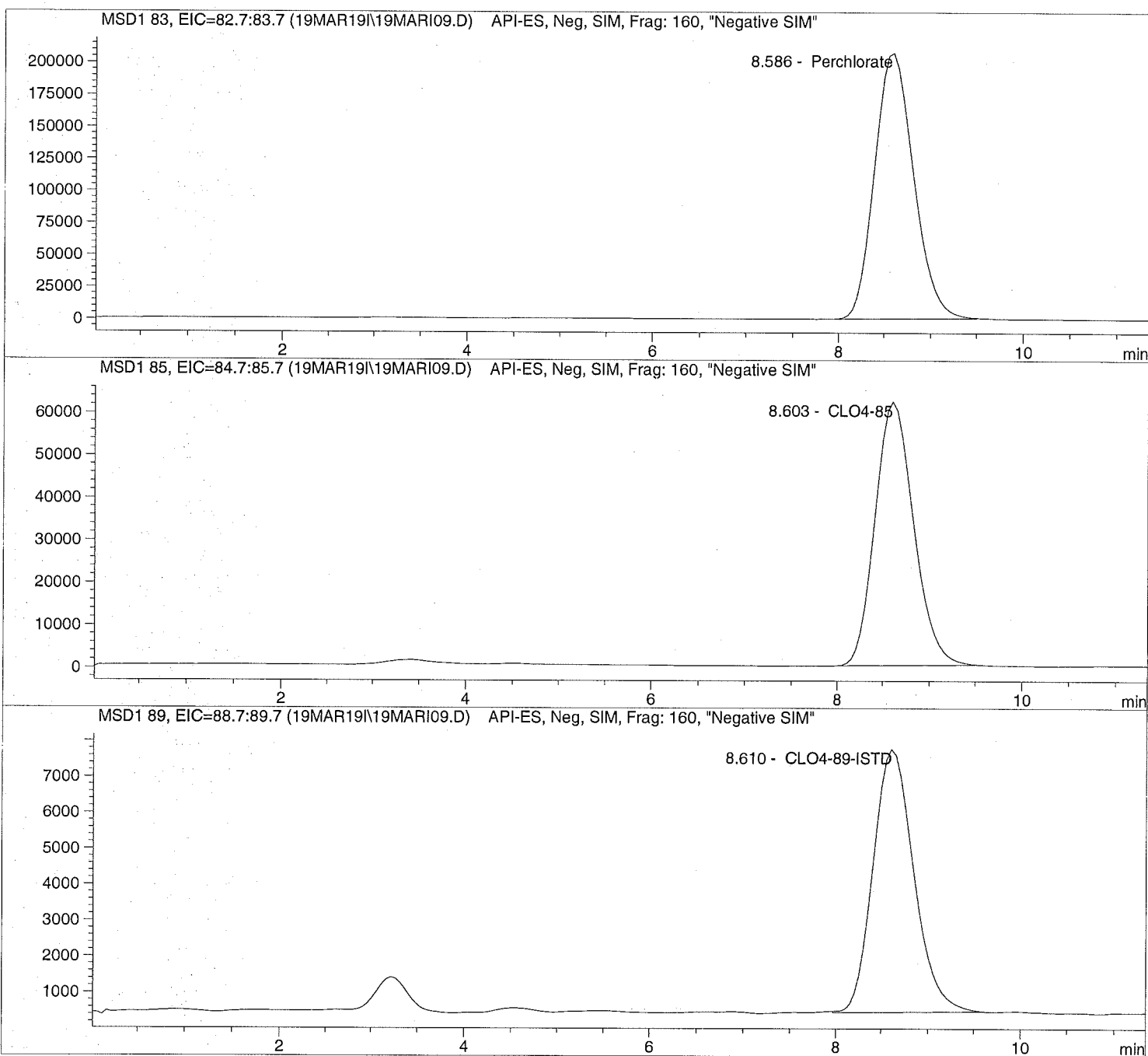
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date:  3/19/2019  10:59:22          Seq Line:           9
Sample Name:    CLO4@ 75.ug/L              Location:           Vial 79
Acq Operator:   TNB                        Inj. No.:          1
                                           Inj. Vol.:         30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  75.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D

Sample Name: ICAL Verf@10ug/L

Injection Date: 3/19/2019 11:12:42

Seq Line: 10

Sample Name: ICAL, Verf@10ug/L

Location: Vial 80

Acq Operator: TNB

Inj. No.: 1

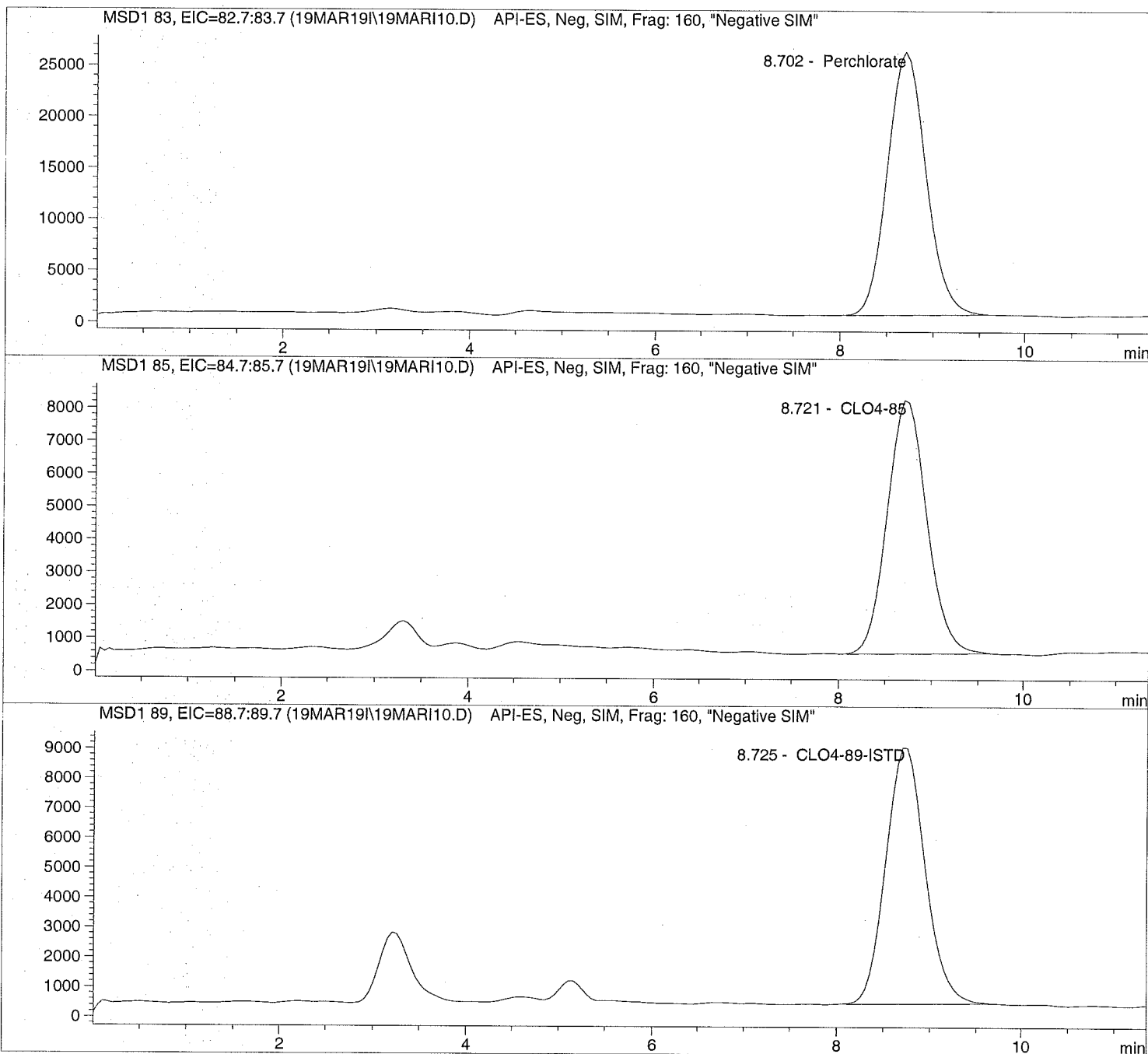
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 3/19/2019 11:12:42      Seq Line:      10
Sample Name:    ICAL Verf@10ug/L        Location:      Vial 80
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

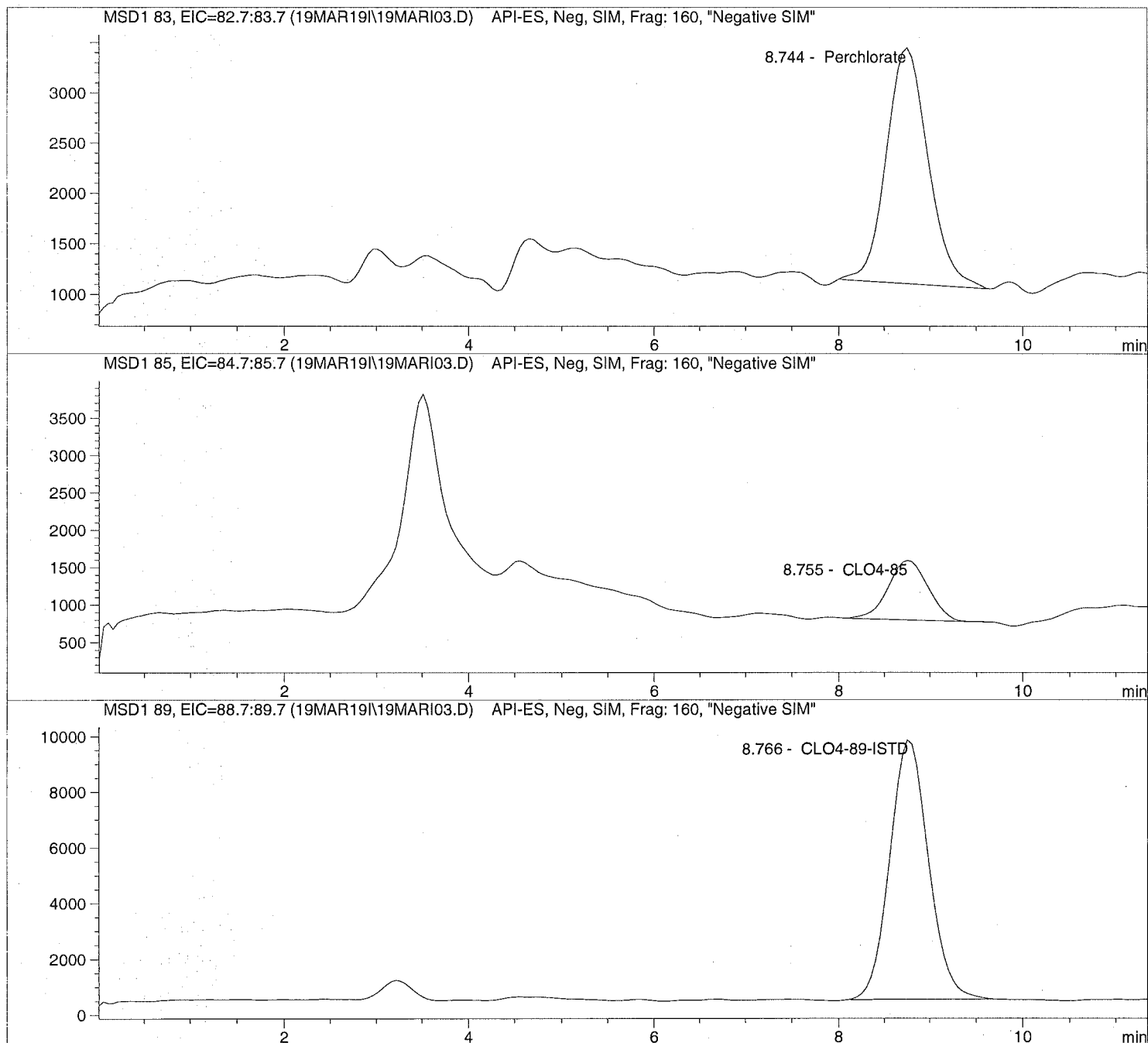
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:      3
Sample Name:    CLO4@ 1.0ug/L           Location:      Vial 73
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



ALS Environmental
ALS Group USA, Corp
1317 South 13th Avenue
Kelso, WA 98626
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www.alsglobal.com

March 29, 2019

Analytical Report for Service Request No: K1902346

RJ Modashia
ALS Laboratory Group
10450 Stancliff Road
Suite 210
Houston, TX 77099-4338

RE: HS19030749

Dear RJ,

Enclosed are the results of the sample(s) submitted to our laboratory March 19, 2019
For your reference, these analyses have been assigned our service request number **K1902346**.

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. The test results meet requirements of the current NELAP standards, where applicable, and except as noted in the laboratory case narrative provided. For a specific list of NELAP-accredited analytes, refer to the certifications section at www.alsglobal.com. All results are intended to be considered in their entirety, and ALS Group USA Corp. dba ALS Environmental (ALS) is not responsible for use of less than the complete report. Results apply only to the items submitted to the laboratory for analysis and individual items (samples) analyzed, as listed in the report.

Please contact me if you have any questions. My extension is 3350. You may also contact me via email at Kelley.Lovejoy@alsglobal.com.

Respectfully submitted,

ALS Group USA, Corp. dba ALS Environmental

for Kelley Lovejoy
Project Manager



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Acronyms

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 General Chemistry

Acronyms

ASTM	American Society for Testing and Materials
A2LA	American Association for Laboratory Accreditation
CARB	California Air Resources Board
CAS Number	Chemical Abstract Service registry Number
CFC	Chlorofluorocarbon
CFU	Colony-Forming Unit
DEC	Department of Environmental Conservation
DEQ	Department of Environmental Quality
DHS	Department of Health Services
DOE	Department of Ecology
DOH	Department of Health
EPA	U. S. Environmental Protection Agency
ELAP	Environmental Laboratory Accreditation Program
GC	Gas Chromatography
GC/MS	Gas Chromatography/Mass Spectrometry
LOD	Limit of Detection
LOQ	Limit of Quantitation
LUFT	Leaking Underground Fuel Tank
M	Modified
MCL	Maximum Contaminant Level is the highest permissible concentration of a substance allowed in drinking water as established by the USEPA.
MDL	Method Detection Limit
MPN	Most Probable Number
MRL	Method Reporting Limit
NA	Not Applicable
NC	Not Calculated
NCASI	National Council of the Paper Industry for Air and Stream Improvement
ND	Not Detected
NIOSH	National Institute for Occupational Safety and Health
PQL	Practical Quantitation Limit
RCRA	Resource Conservation and Recovery Act
SIM	Selected Ion Monitoring
TPH	Total Petroleum Hydrocarbons
tr	Trace level is the concentration of an analyte that is less than the PQL but greater than or equal to the MDL.

Inorganic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- E The result is an estimate amount because the value exceeded the instrument calibration range.
- J The result is an estimated value.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.
- H The holding time for this test is immediately following sample collection. The samples were analyzed as soon as possible after receipt by the laboratory.

Metals Data Qualifiers

- # The control limit criteria is not applicable. See case narrative.
- J The result is an estimated value.
- E The percent difference for the serial dilution was greater than 10%, indicating a possible matrix interference in the sample.
- M The duplicate injection precision was not met.
- N The Matrix Spike sample recovery is not within control limits. See case narrative.
- S The reported value was determined by the Method of Standard Additions (MSA).
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- W The post-digestion spike for furnace AA analysis is out of control limits, while sample absorbance is less than 50% of spike absorbance.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- + The correlation coefficient for the MSA is less than 0.995.
- Q See case narrative. One or more quality control criteria was outside the limits.

Organic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- A A tentatively identified compound, a suspected aldol-condensation product.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- C The analyte was qualitatively confirmed using GC/MS techniques, pattern recognition, or by comparing to historical data.
- D The reported result is from a dilution.
- E The result is an estimated value.
- J The result is an estimated value.
- N The result is presumptive. The analyte was tentatively identified, but a confirmation analysis was not performed.
- P The GC or HPLC confirmation criteria was exceeded. The relative percent difference is greater than 40% between the two analytical results.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a chromatographic interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.

Additional Petroleum Hydrocarbon Specific Qualifiers

- F The chromatographic fingerprint of the sample matches the elution pattern of the calibration standard.
- L The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of lighter molecular weight constituents than the calibration standard.
- H The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of heavier molecular weight constituents than the calibration standard.
- O The chromatographic fingerprint of the sample resembles an oil, but does not match the calibration standard.
- Y The chromatographic fingerprint of the sample resembles a petroleum product eluting in approximately the correct carbon range, but the elution pattern does not match the calibration standard.
- Z The chromatographic fingerprint does not resemble a petroleum product.

**ALS Group USA Corp. dba ALS Environmental (ALS) - Kelso
State Certifications, Accreditations, and Licenses**

Agency	Web Site	Number
Alaska DEH	http://dec.alaska.gov/eh/lab/cs/csapproval.htm	UST-040
Arizona DHS	http://www.azdhs.gov/lab/license/env.htm	AZ0339
Arkansas - DEQ	http://www.adeq.state.ar.us/techsvs/labcert.htm	88-0637
California DHS (ELAP)	http://www.cdph.ca.gov/certlic/labs/Pages/ELAP.aspx	2795
DOD ELAP	http://www.denix.osd.mil/edqw/Accreditation/AccreditedLabs.cfm	L16-58-R4
Florida DOH	http://www.doh.state.fl.us/lab/EnvLabCert/WaterCert.htm	E87412
Hawaii DOH	http://health.hawaii.gov/	-
ISO 17025	http://www.pjlabs.com/	L16-57
Louisiana DEQ	http://www.deq.louisiana.gov/page/la-lab-accreditation	03016
Maine DHS	http://www.maine.gov/dhhs/	WA01276
Minnesota DOH	http://www.health.state.mn.us/accreditation	053-999-457
Nevada DEP	http://ndep.nv.gov/bsdw/labservice.htm	WA01276
New Jersey DEP	http://www.nj.gov/dep/enforcement/oqa.html	WA005
New York - DOH	https://www.wadsworth.org/regulatory/elap	12060
North Carolina DEQ	https://deq.nc.gov/about/divisions/water-resources/water-resources-data/water-sciences-home-page/laboratory-certification-branch/non-field-lab-certification	605
Oklahoma DEQ	http://www.deq.state.ok.us/CSDnew/labcert.htm	9801
Oregon – DEQ (NELAP)	http://public.health.oregon.gov/LaboratoryServices/EnvironmentalLaboratoryAccreditation/Pages/index.aspx	WA100010
South Carolina DHEC	http://www.scdhec.gov/environment/EnvironmentalLabCertification/	61002
Texas CEQ	http://www.tceq.texas.gov/field/qa/env_lab_accreditation.html	T104704427
Washington DOE	http://www.ecy.wa.gov/programs/eap/labs/lab-accreditation.html	C544
Wyoming (EPA Region 8)	https://www.epa.gov/region8-waterops/epa-region-8-certified-drinking-water	-
Kelso Laboratory Website	www.alsglobal.com	NA

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. A complete listing of specific NELAP-certified analytes, can be found in the certification section at www.ALSGlobal.com or at the accreditation bodies web site.

Please refer to the certification and/or accreditation body's web site if samples are submitted for compliance purposes. The states highlighted above, require the analysis be listed on the state certification if used for compliance purposes and if the method/analyte is offered by that state.



Case Narrative

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com



Client: ALS Environmental - US
Project: HS19030749
Sample Matrix: Water

Service Request: K1902346
Date Received: 03/19/2019

CASE NARRATIVE

All analyses were performed consistent with the quality assurance program of ALS Group. This report contains analytical results and quality control parameters for samples designated for Tier IV validation deliverables, which includes summary forms and all of the associated raw data for each of the analyses. When appropriate to the method, method blank results have been reported with each analytical test.

Sample Receipt:

One water sample was received for analysis at ALS Environmental on 03/19/2019. The sample was received in good condition and consistent with the accompanying chain of custody form. The sample was stored in a refrigerator at 4°C upon receipt at the laboratory.

General Chemistry:

No significant anomalies were noted with this analysis.

Approved by Noel D. O'Connell

Date 03/29/2019



Chain of Custody

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

K1902346



10450 Stancliff Rd, Ste 210
 Houston, TX 77099
 T: +1 281 530 5656
 F: +1 281 530 5887
 www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10916

SUBCONTRACT TO:

ALS Environmental Kelso
 1317 S. 13th Avenue
 Kelso, WA 98626

Phone: +1 360 501 3312

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030749
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030749-01	LH18/24-SP650_031419	Water	14 Mar 2019 14:00
TOC Analysis for DOD Level IV			29 Mar 2019

Comments: Please analyze for the analysis listed above.
 Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. MAURY Date/Time: 3/19/19 18:00
 Received By: [Signature] Date/Time: 3/19/19 09:30
 Cooler ID(s): _____ Temperature(s): _____

RIGHT SOLUTIONS | RIGHT PARTNER



PC KL

Cooler Receipt and Preservation Form

Client ALS Houston Service Request K19 03346
 Received: 3/19/19 Opened: 3/19/19 By: CB Unloaded: 3/19/19 By: CB

1. Samples were received via? USPS Fed Ex UPS DHL PDX Courier Hand Delivered
2. Samples were received in: (circle) Cooler Box Envelope Other NA
3. Were custody seals on coolers? NA Y N If yes, how many and where? 1 Left 1 Right
 If present, were custody seals intact? Y N If present, were they signed and dated? Y N

Raw Cooler Temp	Corrected Cooler Temp	Raw Temp Blank	Corrected Temp Blank	Corr. Factor	Thermometer ID	Cooler/COC ID	Tracking Number	NA	Filed
-0.3	-0.1	1.4	1.6	+0.2	379	10916	4809 7831 7714		

4. Packing material: Inserts Baggies Bubble Wrap Gel Packs Wet Ice Dry Ice Sleeves
5. Were custody papers properly filled out (ink, signed, etc.)? NA Y N
6. Were samples received in good condition (temperature, unbroken)? *Indicate in the table below.* NA Y N
 If applicable, tissue samples were received: Frozen Partially Thawed Thawed
7. Were all sample labels complete (i.e analysis, preservation, etc.)? NA Y N
8. Did all sample labels and tags agree with custody papers? *Indicate major discrepancies in the table on page 2.* NA Y N
9. Were appropriate bottles/containers and volumes received for the tests indicated? NA Y N
10. Were the pH-preserved bottles (*see SMO GEN SOP*) received at the appropriate pH? *Indicate in the table below* NA Y N
11. Were VOA vials received without headspace? *Indicate in the table below.* NA Y N
12. Was C12/Res negative? NA Y N

Sample ID on Bottle	Sample ID on COC	Identified by:

Sample ID	Bottle Count	Out of	Head-	Broke	pH	Reagent	Volume	Reagent Lot	Initials	Time
	Bottle Type	Temp	space				added	Number		

Notes, Discrepancies, & Resolutions:



General Chemistry

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Analytical Report

Client: ALS Environmental - US
Project: HS19030749
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1902346
Date Collected: 03/14/19
Date Received: 03/19/19
Units: mg/L
Basis: NA

Carbon, Total Organic

Sample Name	Lab Code	Result	LOQ	LOD	MDL	Dil.	Date Analyzed	Q
LH18/24-SP650_031419	K1902346-001	2.96	0.50	0.20	0.07	1	03/22/19 05:14	
Method Blank	K1902346-MB	ND U	0.50	0.20	0.07	1	03/22/19 04:44	

QA/QC Report

Client: ALS Environmental - US
Project: HS19030749
Sample Matrix: Water

Service Request: K1902346
Date Collected: N/A
Date Received: N/A
Date Analyzed: 03/22/19
Date Extracted: NA

Matrix Spike Summary
Carbon, Total Organic

Sample Name: Batch QC
Lab Code: K1902389-001
Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA

Matrix Spike
K1902389-001MS

<u>Analyte Name</u>	<u>Sample Result</u>	<u>Result</u>	<u>Spike Amount</u>	<u>% Rec</u>	<u>% Rec Limits</u>
Carbon, Total Organic	3.34	31.0	25.0	111	83-117

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030749
Sample Matrix: Water

Service Request: K1902346
Date Analyzed: 03/22/19
Date Extracted: NA

Lab Control Sample Summary
Carbon, Total Organic

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA
Analysis Lot: 629144

Sample Name	Lab Code	Result	Spike Amount	% Rec	% Rec Limits
Lab Control Sample	K1902346-LCS	26.9	25.0	107	83-117

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030749

Service Request: K1902346

Continuing Calibration Verification (CCV) Summary

Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis Lot	Lab Code	Date Analyzed	True Value	Measured Value	Percent Recovery	Acceptance Limits
CCV1	629144	KQ1903793-06	03/21/19 19:42	25.0	24.5	98	90-110
CCV2	629144	KQ1903793-07	03/21/19 23:31	25.0	24.5	98	90-110
CCV3	629144	KQ1903793-08	03/22/19 04:15	25.0	24.9	100	90-110
CCV4	629144	KQ1903793-09	03/22/19 08:58	25.0	24.4	98	90-110
CCV5	629144	KQ1903793-10	03/22/19 14:09	25.0	24.2	97	90-110

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19030749

Service Request: K1902346

Continuing Calibration Blank (CCB) Summary
Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis Lot	Lab Code	Date Analyzed	LOQ	LOD	MDL	Result	Q
CCB1	629144	KQ1903793-01	03/21/19 19:57	0.50	0.20	0.07	ND	U
CCB2	629144	KQ1903793-02	03/21/19 23:46	0.50	0.20	0.07	ND	U
CCB3	629144	KQ1903793-03	03/22/19 04:30	0.50	0.20	0.07	ND	U
CCB4	629144	KQ1903793-04	03/22/19 09:13	0.50	0.20	0.07	ND	U
CCB5	629144	KQ1903793-05	03/22/19 14:23	0.50	0.20	0.07	ND	U



Raw Data

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General Chemistry

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Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Work Request # (Original) K1902074, 2162, 1927, 2047, 2102, 2121, 2127, 2147, 2153, 2168, 2346, 2389, 2121, 2169, 2330, 2232
 Tier: II, II, I, IV, II, I, II, II, II, II, IV, IV, I, II, II
 Date Analyzed: 3/2/19 DOC: 629144
TOC: 629144
629144
629144
 Analyst: BCD Run # _____
 Analysis: TOC/DOC

**DATA QUALITY REPORT
INORGANICS**

Explain any "no" responses to questions below, and any corrective actions in the comments section below.

1. Is the method name and number correct and appropriate? yes/no/NA
2. Holding times met for all analyses and for all samples? yes/no/NA
3. Are calculations correct? yes/no/NA
4. Is the reporting basis correct? (Dry Weight) yes/no/NA
5. All quality control criteria met? yes/no
6. Is the calibration curve correlation coefficient ≥ 0.995 ? yes/no/NA
7. MBs, CCVs, CCBs, LCSs, Dups, and Spikes, analyzed at proper frequency? yes/no/NA
8. Are ICVs, CCVs, and CCBs all within acceptance limits? yes/no/NA
9. Are results for methods blanks all ND? yes/no/NA
10. Are all QC samples within acceptance criteria? (LCS % rec, MS/DMS % rec, DUP or MS/DMS RPDs, etc.) yes/no/NA
11. Are all exceptions explained? yes/no/NA
12. Have all applicable service requests been reviewed? yes/no/NA
13. Are all samples labeled correctly? yes/no/NA
14. Have all instructions on the service request been followed? (e.g. Special MRLs, QC on a specific sample, Form V) yes/no/NA
15. Are detection limits and units reported correctly? yes/no/NA
16. Is the unused space on the benchsheet crossed out? yes/no/NA
17. Was analysis turned in by the due date? (n-2) (If not record SR#) yes/no/NA

COMMENTS: R 1902232-2/3, R1902121-1/2, K1902435-1/2/3/4/5/6/7, K1902254-1/2/3
 Sent for reanalysis due to CCB above the MRL.
 K1902102-1, K1902074-2/1 sent for reanalysis due to MB requiring a dilution.
 K1902102-1 for sent reanalysis due to requiring a dilution.
 K1902147-1 sent for reanalysis because it is overloaded.
 K1902155-1/dups have a high PRSD due to dirty, non-homogeneous sample.
 K1902168-2/2d report a high PRSD, but these samples are less than 5x the MRL.

Final Approved by: Houyer Date: 03/25/19 DQREPORT

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629142 Method/Testcode: SM 5310 C/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902074-001	Carbon, Dissolved Organic N/A (DOC)			Effluent	1.64 mg/L	10 ml	1.64 mg/L	1	0.07	0.50			3/21/19 20:56	N	II
K1902074-002	Carbon, Dissolved Organic N/A (DOC)			Water	1.65 mg/L	10 ml	1.65 mg/L	1	0.07	0.50			3/21/19 21:24	N	II
K1902102-001	Carbon, Dissolved Organic N/A (DOC)			Water	76.61 mg/L	10 ml	76.6 mg/L	1	0.07	0.50			3/21/19 21:52	N	II
KQ1903792-01	Carbon, Dissolved Organic CCB (DOC)			Effluent	0.01 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 19:57	N	II
KQ1903792-02	Carbon, Dissolved Organic CCB (DOC)			Effluent	0.03 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 23:46	N	II
KQ1903792-03	Carbon, Dissolved Organic CCV (DOC)			Effluent	24.51 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 19:42	N	II
KQ1903792-04	Carbon, Dissolved Organic CCV (DOC)			Effluent	24.47 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 23:31	N	II
KQ1903792-05	Carbon, Dissolved Organic MB (DOC)			Effluent	-0.04 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 20:12	N	II
KQ1903792-06	Carbon, Dissolved Organic LCS (DOC)			Effluent	26.54 mg/L	10 ml	26.5 mg/L	1	0.07	0.50	106		3/21/19 20:26	N	II
KQ1903792-07	Carbon, Dissolved Organic MS (DOC)		K1902102-001	Water	99.38 mg/L	10 ml	99.4 mg/L	1	0.07	0.50	91		3/21/19 22:20	N	II
KQ1903792-08	Carbon, Dissolved Organic DUP (DOC)		K1902102-001	Water	78.20 mg/L	10 ml	78.2 mg/L	1	0.07	0.50		2	3/21/19 21:52	N	II
KQ1903792-09	Carbon, Dissolved Organic DUP (DOC)		K1902074-002	Water	1.68 mg/L	10 ml	1.68 mg/L	1	0.07	0.50		2	3/21/19 21:24	N	II
KQ1903792-10	Carbon, Dissolved Organic DUP (DOC)		K1902074-001	Effluent	1.62 mg/L	10 ml	1.62 mg/L	1	0.07	0.50		1	3/21/19 20:56	N	II

03/25/19
Free up

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

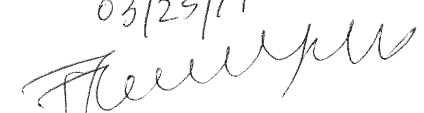
Analyst: BDITZLER

Analysis Lot: 629144

Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1901927-005	Carbon, Total Organic	N/A		Ground Water	45.71 mg/L	10 ml	45.7 mg/L	1	0.07	0.50			3/22/19 00:01	N	I
K1902047-001	Carbon, Total Organic	N/A		Water	2.94 mg/L	10 ml	2.94 mg/L	1	0.07	0.50			3/21/19 23:03	N	IV
K1902102-001	Carbon, Total Organic	N/A		Water	83.22 mg/L	10 ml	83.2 mg/L	1	0.07	0.50			3/22/19 05:42	N	II
K1902121-002	Carbon, Total Organic	N/A		Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 20:19	N	I
K1902127-001	Carbon, Total Organic	N/A		Water	15.73 mg/L	10 ml	15.7 mg/L	1	0.07	0.50			3/22/19 06:10	N	II
K1902127-002	Carbon, Total Organic	N/A		Water	7.06 mg/L	10 ml	28.2 mg/L	4	0.3	2.0			3/22/19 06:38	N	II
K1902127-003	Carbon, Total Organic	N/A		Water	7.65 mg/L	10 ml	7.65 mg/L	1	0.07	0.50			3/22/19 07:06	N	II
K1902127-004	Carbon, Total Organic	N/A		Water	4.54 mg/L	10 ml	18.1 mg/L	4	0.3	2.0			3/22/19 07:34	N	II
K1902147-001	Carbon, Total Organic	N/A		Water	0.23 mg/L	10 ml	50 mg/L U	100	7	50			3/22/19 08:02	N	II
K1902147-002	Carbon, Total Organic	N/A		Water	0.95 mg/L	10 ml	0.95 mg/L	1	0.07	0.50			3/22/19 08:30	N	II
K1902153-001	Carbon, Total Organic	N/A		Water	2.23 mg/L	10 ml	223 mg/L	100	7	50			3/22/19 02:50	N	II
K1902153-002	Carbon, Total Organic	N/A		Water	3.16 mg/L	10 ml	316 mg/L	100	7	50			3/22/19 03:19	N	II
K1902153-003	Carbon, Total Organic	N/A		Water	1.20 mg/L	10 ml	120 mg/L	100	7	50			3/22/19 03:47	N	II
K1902168-001	Carbon, Total Organic	N/A		Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50			3/22/19 09:28	N	II
K1902168-002	Carbon, Total Organic	N/A		Water	0.87 mg/L	10 ml	0.87 mg/L	1	0.07	0.50			3/22/19 09:56	N	II
K1902346-001	Carbon, Total Organic	N/A		Water	2.96 mg/L	10 ml	2.96 mg/L	1	0.07	0.50			3/22/19 05:14	N	IV
K1902389-001	Carbon, Total Organic	N/A		Water	3.34 mg/L	10 ml	3.34 mg/L	1	0.07	0.50			3/22/19 00:57	Y	IV
K1902389-002	Carbon, Total Organic	N/A		Water	27.06 mg/L	10 ml	27.1 mg/L	1	0.07	0.50			3/22/19 01:54	N	IV
K1902389-003	Carbon, Total Organic	N/A		Water	27.93 mg/L	10 ml	27.9 mg/L	1	0.07	0.50			3/22/19 02:22	N	IV
KQ1903793-01	Carbon, Total Organic	CCB		Water	9.45000000000001E	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 19:57	N	IV
KQ1903793-02	Carbon, Total Organic	CCB		Water	0.03 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/21/19 23:46	N	IV
KQ1903793-03	Carbon, Total Organic	CCB		Water	0.05 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 04:30	N	IV
KQ1903793-04	Carbon, Total Organic	CCB		Water	-0.17 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 09:13	N	IV
KQ1903793-05	Carbon, Total Organic	CCB		Water	-0.19 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 14:23	N	IV
KQ1903793-06	Carbon, Total Organic	CCV		Water	24.51 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 19:42	N	IV
KQ1903793-07	Carbon, Total Organic	CCV		Water	24.47 mg/L	10 ml	24.5 mg/L	1			98		3/21/19 23:31	N	IV
KQ1903793-08	Carbon, Total Organic	CCV		Water	24.91 mg/L	10 ml	24.9 mg/L	1			100		3/22/19 04:15	N	IV
KQ1903793-09	Carbon, Total Organic	CCV		Water	24.39 mg/L	10 ml	24.4 mg/L	1			98		3/22/19 08:58	N	IV
KQ1903793-10	Carbon, Total Organic	CCV		Water	24.24 mg/L	10 ml	24.2 mg/L	1			97		3/22/19 14:09	N	IV
KQ1903793-11	Carbon, Total Organic	MB		Water	1.65000000000001E	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 04:44	N	IV
KQ1903793-12	Carbon, Total Organic	LCS		Water	26.86 mg/L	10 ml	26.9 mg/L	1	0.07	0.50	107		3/22/19 04:59	N	IV
KQ1903793-13	Carbon, Total Organic	MS	K1902389-001	Water	30.99 mg/L	10 ml	31.0 mg/L	1	0.07	0.50	111		3/22/19 01:25	N	IV
KQ1903793-14	Carbon, Total Organic	DUP	K1902047-001	Water	2.93 mg/L	10 ml	2.93 mg/L	1	0.07	0.50		<1	3/21/19 23:03	N	IV
KQ1903793-15	Carbon, Total Organic	DUP	K1901927-005	Ground Water	46.15 mg/L	10 ml	46.1 mg/L	1	0.07	0.50		<1	3/22/19 00:01	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

03/25/19


Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629144 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1903793-16	Carbon, Total Organic	DUP	K1902389-001	Water	3.34 mg/L	10 ml	3.34 mg/L	1	0.07	0.50		<1	3/22/19 00:57	N	IV
KQ1903793-17	Carbon, Total Organic	DUP	K1902389-002	Water	27.77 mg/L	10 ml	27.8 mg/L	1	0.07	0.50		3	3/22/19 01:54	N	IV
KQ1903793-18	Carbon, Total Organic	DUP	K1902389-003	Water	27.82 mg/L	10 ml	27.8 mg/L	1	0.07	0.50		<1	3/22/19 02:22	N	IV
KQ1903793-19	Carbon, Total Organic	DUP	K1902153-001	Water	1.67 mg/L	10 ml	167 mg/L	100	7	50		29*	3/22/19 02:50	N	II
KQ1903793-20	Carbon, Total Organic	DUP	K1902153-002	Water	3.14 mg/L	10 ml	314 mg/L	100	7	50		<1	3/22/19 03:19	N	II
KQ1903793-21	Carbon, Total Organic	DUP	K1902153-003	Water	1.13 mg/L	10 ml	113 mg/L	100	7	50		6	3/22/19 03:47	N	II
KQ1903793-22	Carbon, Total Organic	DUP	K1902346-001	Water	2.84 mg/L	10 ml	2.84 mg/L	1	0.07	0.50		4	3/22/19 05:14	N	IV
KQ1903793-23	Carbon, Total Organic	DUP	K1902102-001	Water	83.37 mg/L	10 ml	83.4 mg/L	1	0.07	0.50		<1	3/22/19 05:42	N	II
KQ1903793-24	Carbon, Total Organic	DUP	K1902121-002	Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50		NC	3/22/19 20:19	N	I
KQ1903793-25	Carbon, Total Organic	DUP	K1902127-001	Water	15.57 mg/L	10 ml	15.6 mg/L	1	0.07	0.50		<1	3/22/19 06:10	N	II
KQ1903793-26	Carbon, Total Organic	DUP	K1902127-002	Water	6.91 mg/L	10 ml	27.6 mg/L	4	0.3	2.0		2	3/22/19 06:38	N	II
KQ1903793-27	Carbon, Total Organic	DUP	K1902127-003	Water	7.52 mg/L	10 ml	7.52 mg/L	1	0.07	0.50		2	3/22/19 07:06	N	II
KQ1903793-28	Carbon, Total Organic	DUP	K1902127-004	Water	4.39 mg/L	10 ml	17.6 mg/L	4	0.3	2.0		3	3/22/19 07:34	N	II
KQ1903793-29	Carbon, Total Organic	DUP	K1902147-001	Water	0.22 mg/L	10 ml	22 mg/L	J 100	7	50		NC	3/22/19 08:02	N	II
KQ1903793-30	Carbon, Total Organic	DUP	K1902147-002	Water	0.91 mg/L	10 ml	0.91 mg/L	1	0.07	0.50		4	3/22/19 08:30	N	II
KQ1903793-31	Carbon, Total Organic	DUP	K1902168-001	Water	1.78 mg/L	10 ml	1.78 mg/L	1	0.07	0.50		4	3/22/19 09:28	N	II
KQ1903793-32	Carbon, Total Organic	DUP	K1902168-002	Water	0.76 mg/L	10 ml	0.76 mg/L	1	0.07	0.50		14*	3/22/19 09:56	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629146 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902121-001	AA Carbon, Total Organic	N/A		Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 19:51	N	I
K1902169-001	Carbon, Total Organic	N/A		Water	14.29 mg/L	10 ml	14.3 mg/L	1	0.07	0.50			3/22/19 10:24	N	II
K1902169-002	Carbon, Total Organic	N/A		Water	17.52 mg/L	10 ml	17.5 mg/L	1	0.07	0.50			3/22/19 10:52	N	II
K1902169-003	Carbon, Total Organic	N/A		Water	18.67 mg/L	10 ml	18.7 mg/L	1	0.07	0.50			3/22/19 11:20	N	II
K1902169-004	Carbon, Total Organic	N/A		Water	9.18 mg/L	10 ml	9.18 mg/L	1	0.07	0.50			3/22/19 11:48	N	II
K1902169-005	Carbon, Total Organic	N/A		Water	11.61 mg/L	10 ml	11.6 mg/L	1	0.07	0.50			3/22/19 12:16	N	II
K1902169-006	Carbon, Total Organic	N/A		Water	11.71 mg/L	10 ml	11.7 mg/L	1	0.07	0.50			3/22/19 12:44	N	II
K1902169-007	Carbon, Total Organic	N/A		Water	12.46 mg/L	10 ml	12.5 mg/L	1	0.07	0.50			3/22/19 13:12	N	II
K1902169-008	Carbon, Total Organic	N/A		Water	15.31 mg/L	10 ml	15.3 mg/L	1	0.07	0.50			3/22/19 13:41	N	II
K1902230-001	Carbon, Total Organic	N/A		Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50			3/22/19 15:07	N	II
K1902230-002	Carbon, Total Organic	N/A		Water	0.70 mg/L	10 ml	0.70 mg/L	1	0.07	0.50			3/22/19 15:35	N	II
K1902230-003	Carbon, Total Organic	N/A		Water	0.80 mg/L	10 ml	0.80 mg/L	1	0.07	0.50			3/22/19 16:03	N	II
K1902230-004	Carbon, Total Organic	N/A		Water	1.23 mg/L	10 ml	1.23 mg/L	1	0.07	0.50			3/22/19 16:32	N	II
K1902230-005	Carbon, Total Organic	N/A		Water	0.32 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 17:00	N	II
K1902232-001	Carbon, Total Organic	N/A		Water	5.72 mg/L	10 ml	5.72 mg/L	1	0.07	0.50			3/22/19 17:28	N	II
K1902232-002	RA Carbon, Total Organic	N/A		Water	9.75 mg/L	10 ml	9.75 mg/L	1	0.07	0.50			3/22/19 18:55	N	II
K1902232-003	RA Carbon, Total Organic	N/A		Water	13.90 mg/L	10 ml	13.9 mg/L	1	0.07	0.50			3/22/19 19:23	N	II
KQ1903794-01	Carbon, Total Organic	CCB		Water	-0.17 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 09:13	N	II
KQ1903794-02	Carbon, Total Organic	CCB		Water	-0.19 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 14:23	N	II
KQ1903794-03	Carbon, Total Organic	CCB		Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 18:40	N	II
KQ1903794-04	Carbon, Total Organic	CCB	(High)	Water	0.89 mg/L	10 ml	0.89 mg/L	1	0.07	0.50			3/22/19 23:50	N	II
KQ1903794-05	Carbon, Total Organic	CCV		Water	24.39 mg/L	10 ml	24.4 mg/L	1					3/22/19 08:58	N	II
KQ1903794-06	Carbon, Total Organic	CCV		Water	24.24 mg/L	10 ml	24.2 mg/L	1					3/22/19 14:09	N	II
KQ1903794-07	Carbon, Total Organic	CCV		Water	23.91 mg/L	10 ml	23.9 mg/L	1					3/22/19 18:25	N	II
KQ1903794-08	Carbon, Total Organic	CCV		Water	27.39 mg/L	10 ml	27.4 mg/L	1					3/22/19 23:35	N	II
KQ1903794-09	Carbon, Total Organic	MB		Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 14:38	N	II
KQ1903794-10	Carbon, Total Organic	LCS		Water	26.10 mg/L	10 ml	26.1 mg/L	1	0.07	0.50	104		3/22/19 14:53	N	II
KQ1903794-11	Carbon, Total Organic	MS	K1902232-001	Water	33.31 mg/L	10 ml	33.3 mg/L	1	0.07	0.50	110		3/22/19 17:56	N	II
KQ1903794-12	Carbon, Total Organic	DUP	K1902169-001	Water	14.17 mg/L	10 ml	14.2 mg/L	1	0.07	0.50		<1	3/22/19 10:24	N	II
KQ1903794-13	Carbon, Total Organic	DUP	K1902169-002	Water	17.70 mg/L	10 ml	17.7 mg/L	1	0.07	0.50		1	3/22/19 10:52	N	II
KQ1903794-14	Carbon, Total Organic	DUP	K1902169-003	Water	18.63 mg/L	10 ml	18.6 mg/L	1	0.07	0.50		<1	3/22/19 11:20	N	II
KQ1903794-15	Carbon, Total Organic	DUP	K1902169-004	Water	9.05 mg/L	10 ml	9.05 mg/L	1	0.07	0.50		1	3/22/19 11:48	N	II
KQ1903794-16	Carbon, Total Organic	DUP	K1902169-005	Water	11.53 mg/L	10 ml	11.5 mg/L	1	0.07	0.50		<1	3/22/19 12:16	N	II
KQ1903794-17	Carbon, Total Organic	DUP	K1902169-006	Water	11.48 mg/L	10 ml	11.5 mg/L	1	0.07	0.50		2	3/22/19 12:44	N	II
KQ1903794-18	Carbon, Total Organic	DUP	K1902169-007	Water	12.85 mg/L	10 ml	12.9 mg/L	1	0.07	0.50		3	3/22/19 13:12	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

03/25/19
Free

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629146 Method/Testcode: SM 5310 C/TOC T

<u>Lab Code</u>	<u>Target Analytes</u>	<u>QC</u>	<u>Parent Sample</u>	<u>Matrix</u>	<u>Raw Result</u>	<u>Sample Amt.</u>	<u>Final Result</u>	<u>Dil</u>	<u>MDL</u>	<u>PQL</u>	<u>% Rec</u>	<u>% RSD</u>	<u>Date Analyzed</u>	<u>QC?</u>	<u>Tier</u>
KQ1903794-19	Carbon, Total Organic	DUP	K1902169-008	Water	15.02 mg/L	10 ml	15.0 mg/L	1	0.07	0.50		2	3/22/19 13:41	N	II
KQ1903794-20	Carbon, Total Organic	DUP	K1902230-001	Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50		<1	3/22/19 15:07	N	II
KQ1903794-21	Carbon, Total Organic	DUP	K1902230-002	Water	0.64 mg/L	10 ml	0.64 mg/L	1	0.07	0.50		8	3/22/19 15:35	N	II
KQ1903794-22	Carbon, Total Organic	DUP	K1902230-003	Water	0.79 mg/L	10 ml	0.79 mg/L	1	0.07	0.50		<1	3/22/19 16:03	N	II
KQ1903794-23	Carbon, Total Organic	DUP	K1902230-004	Water	1.14 mg/L	10 ml	1.14 mg/L	1	0.07	0.50		7	3/22/19 16:32	N	II
KQ1903794-24	Carbon, Total Organic	DUP	K1902230-005	Water	0.30 mg/L	10 ml	0.30 mg/L	J 1	0.07	0.50		NC	3/22/19 17:00	N	II
KQ1903794-25	Carbon, Total Organic	DUP	K1902232-001	Water	5.79 mg/L	10 ml	5.79 mg/L	1	0.07	0.50		1	3/22/19 17:28	N	II
KQ1903794-26	Carbon, Total Organic	DUP	K1902232-002	Water	9.86 mg/L	10 ml	9.86 mg/L	1	0.07	0.50		1	3/22/19 18:55	N	II
KQ1903794-27	Carbon, Total Organic	DUP	K1902232-003	Water	13.87 mg/L	10 ml	13.9 mg/L	1	0.07	0.50		<1	3/22/19 19:23	N	II
KQ1903794-28	Carbon, Total Organic	DUP	K1902121-001	Reagent Water	-0.24 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50		NC	3/22/19 19:51	N	I

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629149 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902254-001	Carbon, Total Organic	N/A		Water	6.54 mg/L	10 ml	6.54 mg/L	1	0.07	0.50			3/23/19 01:02	N	II
K1902254-002	Carbon, Total Organic	N/A		Water	18.37 mg/L	10 ml	18.4 mg/L	1	0.07	0.50			3/23/19 01:30	N	II
K1902254-003	Carbon, Total Organic	N/A		Water	22.01 mg/L	10 ml	22.0 mg/L	1	0.07	0.50			3/23/19 01:58	N	II
K1902435-001	Carbon, Total Organic	N/A		Ground Water	119.91 mg/L	10 ml	120 mg/L	1	0.07	0.50			3/22/19 20:47	N	IV
K1902435-002	Carbon, Total Organic	N/A		Ground Water	5.07 mg/L	10 ml	5.07 mg/L	1	0.07	0.50			3/22/19 21:15	N	IV
K1902435-003	Carbon, Total Organic	N/A		Ground Water	3.20 mg/L	10 ml	3.20 mg/L	1	0.07	0.50			3/22/19 21:43	N	IV
K1902435-004	Carbon, Total Organic	N/A		Ground Water	5.08 mg/L	10 ml	5.08 mg/L	1	0.07	0.50			3/22/19 22:11	N	IV
K1902435-005	Carbon, Total Organic	N/A		Ground Water	137.01 mg/L	10 ml	137 mg/L	1	0.07	0.50			3/22/19 22:39	N	IV
K1902435-006	Carbon, Total Organic	N/A		Ground Water	118.27 mg/L	10 ml	118 mg/L	1	0.07	0.50			3/22/19 23:07	N	IV
K1902435-007	Carbon, Total Organic	N/A		Ground Water	12.50 mg/L	10 ml	12.5 mg/L	1	0.07	0.50			3/23/19 00:05	Y	IV
KQ1903795-01	Carbon, Total Organic	CCB		Ground Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/22/19 18:40	N	IV
KQ1903795-02	Carbon, Total Organic	CCB		Ground Water	0.89 mg/L	10 ml	0.89 mg/L	1	0.07	0.50			3/22/19 23:50	N	IV
KQ1903795-03	Carbon, Total Organic	CCB		Ground Water	-0.24 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			3/23/19 03:09	N	IV
KQ1903795-04	Carbon, Total Organic	CCV		Ground Water	23.91 mg/L	10 ml	23.9 mg/L	1			96		3/22/19 18:25	N	IV
KQ1903795-05	Carbon, Total Organic	CCV		Ground Water	27.39 mg/L	10 ml	27.4 mg/L	1			110		3/22/19 23:35	N	IV
KQ1903795-06	Carbon, Total Organic	CCV		Ground Water	23.81 mg/L	10 ml	23.8 mg/L	1			95		3/23/19 02:55	N	IV
KQ1903795-09	Carbon, Total Organic	MS	K1902435-007	Ground Water	33.24 mg/L	10 ml	33.2 mg/L	1	0.07	0.50	83		3/23/19 00:33	N	IV
KQ1903795-10	Carbon, Total Organic	DUP	K1902435-001	Ground Water	112.05 mg/L	10 ml	112 mg/L	1	0.07	0.50		7	3/22/19 20:47	N	IV
KQ1903795-11	Carbon, Total Organic	DUP	K1902435-002	Ground Water	3.05 mg/L	10 ml	3.05 mg/L	1	0.07	0.50		50*	3/22/19 21:15	N	IV
KQ1903795-12	Carbon, Total Organic	DUP	K1902435-003	Ground Water	2.88 mg/L	10 ml	2.88 mg/L	1	0.07	0.50		11*	3/22/19 21:43	N	IV
KQ1903795-13	Carbon, Total Organic	DUP	K1902435-004	Ground Water	4.70 mg/L	10 ml	4.70 mg/L	1	0.07	0.50		8	3/22/19 22:11	N	IV
KQ1903795-14	Carbon, Total Organic	DUP	K1902435-005	Ground Water	127.76 mg/L	10 ml	128 mg/L	1	0.07	0.50		7	3/22/19 22:39	N	IV
KQ1903795-15	Carbon, Total Organic	DUP	K1902435-006	Ground Water	127.20 mg/L	10 ml	127 mg/L	1	0.07	0.50		7	3/22/19 23:07	N	IV
KQ1903795-16	Carbon, Total Organic	DUP	K1902435-007	Ground Water	12.30 mg/L	10 ml	12.3 mg/L	1	0.07	0.50		2	3/23/19 00:05	N	IV
KQ1903795-17	Carbon, Total Organic	DUP	K1902254-001	Water	6.38 mg/L	10 ml	6.38 mg/L	1	0.07	0.50		3	3/23/19 01:02	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Printed 3/23/19 15:44

Results Summary

03/25/19
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Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629149 Method/Testcode: SM 5310 C/TOC T

<u>Lab Code</u>	<u>Target Analytes</u>	<u>QC</u>	<u>Parent Sample</u>	<u>Matrix</u>	<u>Raw Result</u>	<u>Sample Amt.</u>	<u>Final Result</u>	<u>Dil</u>	<u>MDL</u>	<u>PQL</u>	<u>% Rec</u>	<u>% RSD</u>	<u>Date Analyzed</u>	<u>QC?</u>	<u>Tier</u>
KQ1903795-18	Carbon, Total Organic	DUP	K1902254-002	Water	18.17 mg/L	10 ml	18.2 mg/L	1	0.07	0.50		1	3/23/19 01:30	N	II
KQ1903795-19	Carbon, Total Organic	DUP	K1902254-003	Water	21.93 mg/L	10 ml	21.9 mg/L	1	0.07	0.50		<1	3/23/19 01:58	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

DOC: 629142
 TOC: 629144,
 629146,
 629149

Schedule: 03212019

Version: 4

Instrument: Fusion1

Last Saved by: Fusion1 (Fusion1)

Last Saved on: 2019/03/21 18:03 - Thursday

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Blank)	Blank	Reagent/Acid Blank		1	True	Ready
D	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
1	Sample	MB1	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
2	Sample	ICS	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
3	Sample	K1902074-001.03 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
4	Sample	K1902074-002.03 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
5	Sample	K1902102-001.01 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
6	Sample	K1902102-001.01 ms doc	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
7	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
8	Sample	FB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
9	Sample	K1902047-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
10	Sample	K1901927-005.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
11	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
12	Sample	K1902389-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
13	Sample	K1902389-001.01 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
14	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
15	Sample	K1902389-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
16	Sample	K1902389-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
17	Sample	K1902153-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
18	Sample	K1902153-002.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
19	Sample	K1902153-003.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
20	Sample	MB2	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
21	Sample	K1902346-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
22	Sample	K1902102-001.02	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
23	Sample	K1902127-001.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
24	Sample	K1902127-002.12 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
25	Sample	K1902127-003.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
26	Sample	K1902127-004.12 4x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
27	Sample	K1902147-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
28	Sample	K1902147-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
29	Sample	K1902168-001.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
30	Sample	K1902168-002.04	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
31	Sample	K1902169-001.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
32	Sample	K1902169-002.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
33	Sample	K1902169-003.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
34	Sample	K1902169-004.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
35	Sample	K1902169-005.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
36	Sample	K1902169-006.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
37	Sample	K1902169-007.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
38	Sample	K1902169-008.24	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready

Printed on: March 23, 2019 11:38:49

Schedule: 03212019

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
39	Sample	MB3	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
40	Sample	K1902230-001.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
41	Sample	K1902230-002.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
42	Sample	K1902230-003.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
43	Sample	K1902230-004.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
44	Sample	K1902230-005.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
45	Sample	K1902232-001.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
46	Sample	K1902232-001.12 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
47	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
48	Sample	K1902232-002.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
49	Sample	K1902232-003.12	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
50	Sample	K1902121-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
51	Sample	K1902121-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
52	Sample	K1902435-001.02	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
53	Sample	K1902435-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
54	Sample	K1902435-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
55	Sample	K1902435-004.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
56	Sample	K1902435-005.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
57	Sample	K1902435-006.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
58	Sample	MB4	CAS_salt_010711 (CAS_salt_010711)	1	False	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	False	Ready
59	Sample	K1902435-007.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
60	Sample	K1902435-007.01 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
61	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
62	Sample	K1902254-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
63	Sample	K1902254-002.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
64	Sample	K1902254-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
65	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
					False	

Fusion Report - 03212019

Thursday, March 21, 2019 04:58 PM

(View - Repts, Unused Repts, Meta-Data, Signature, History)
Printed on 2019/03/23 11:38 - Saturday

Report Summary Information

Company Location: Gen Chem Lab
 Schedule Name: 03212019
 Instrument Name: Fusion1
 Report Version: 1 of 1
 Report Creation by Operators (schedule version): Fusion1 (Fusion1) (v2)
 Fusion1 (Fusion1) (v3)
 Fusion1 (Fusion1) (v4)
 Comment:

Engine Version: 1.1.5.1
 Firmware Version: 1.2.0696
 Connection: RS232 COM1

Report Results

03/25/19
Fusion

Sample Type: Clean From Schedule Version 2

Pos	Analysis Type	Sample ID	Start Time
♦ (clean)		Clean	2019/03/21 16:58

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	13.69	16.58	2.89	49.93	07:59
2	TC Clean	20.38	23.31	2.93	49.73	07:19
3	TC Clean	2.42	5.55	3.13	49.79	07:01
4	TC Clean	1.97	4.69	2.71	49.80	07:01

Sample Type: Clean From Schedule Version 3

Pos	Analysis Type	Sample ID	Start Time
♦ (clean)		Clean	2019/03/21 17:32

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	10.80	13.84	3.04	49.90	08:01
2	TC Clean	4.74	7.49	2.75	49.73	07:16
3	TC Clean	1.91	4.68	2.77	49.81	06:59
4	TC Clean	1.32	4.10	2.77	49.81	07:04

Sample Type: Clean			From Schedule Version 4			
Pos	Analysis Type	Sample ID			Start Time	
♦ (clean)		Clean			2019/03/21 18:06	
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	1.03	3.77	2.74	49.95	08:00
2	TC Clean	4.31	6.95	2.64	49.79	07:16
3	TC Clean	1.72	4.56	2.84	49.85	07:02
4	TC Clean	1.74	4.48	2.74	49.84	07:03

Sample Type: Blank (Creating v1238)			From Schedule Version 4			
Pos	Analysis Type	Sample ID			Start Time	
♦ (blank)		Reagent/Acid Blank			2019/03/21 18:41	
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	1.00	3.60	2.60	49.84	05:29
2	TC Clean	4.32	7.12	2.80	49.86	07:17
3	TC Clean	2.15	5.14	2.99	49.89	07:02
4	TC Clean	2.00	4.74	2.74	49.91	07:02
5	Reagent Blank	8.64	11.55	2.91	49.90	08:11
6	Acid Blank	1.76	4.51	2.75	49.59	05:29

Sample Type: Sample			From Schedule Version 4					
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
♦ D	TOC	RB	0.6753 ppm	0.0000 ppm	0.0000%	2019/03/21 19:27		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6753	6.7532	13.68	16.57	2.90	50.05	10:32
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.7429 ppm (PASS)	0.0000 ppm	0%	2019/03/21 19:42

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.7429	247.4293	177.42	180.64	3.22	50.08	10:34

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.2462 ppm (PASS)	0.0000 ppm	0%	2019/03/21 19:57

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.2462	2.4625	11.13	14.12	2.99	50.11	10:31

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 1	TOC	MB1	0.1945 ppm	0.0000 ppm	0.0000%	2019/03/21 20:12

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1945	1.9447	10.41	13.45	3.03	50.13	10:31

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.7752 ppm (PASS)	0.0000 ppm	0%	2019/03/21 20:26

Pos	Base Analysis	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time

Type	Completion State	Success Action	Method	Calibration	STD Conc - Pos C
C TOC 25.0 ppm 1 26.7752 267.7521 191.21 194.15 2.94 50.12 10:31	Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	25 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
2	TOC	ICS	0.5996 ppm	0.0000 ppm	0.0000%	2019/03/21 20:41

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5996	5.9960	13.16	16.20	3.04	50.14	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
3	TOC	K1902074-001.03 doc	1.8667 ppm	0.0140 ppm	0.7500%	2019/03/21 20:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.8766	18.7657	21.83	24.73	2.90	50.18	10:27
2	TOC	1.8568	18.5683	21.70	24.80	3.10	50.17	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
4	TOC	K1902074-002.03 doc	1.9020 ppm	0.0253 ppm	1.3300%	2019/03/21 21:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.8841	18.8408	21.88	24.73	2.85	50.15	10:26
2	TOC	1.9199	19.1988	22.12	24.96	2.84	50.11	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
5	TOC	K1902102-001.01 doc	77.6461 ppm	1.1244 ppm	1.4500%	2019/03/21 21:52

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	76.8510	768.5100	530.75	533.65	2.90	50.09	10:28
2	TOC	78.4412	784.4117	541.55	544.57	3.03	50.08	10:32

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC)
Method CAS_salt_010711
Calibration CAS_salt_010711

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
6	TOC	K1902102-001.01 ms doc	99.6155 ppm	0.0000 ppm	0.0000%	2019/03/21 22:20		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	99.6155	996.1549	685.28	688.33	3.06	50.06	10:33
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
7	TOC	RB	0.8853 ppm	0.0000 ppm	0.0000%	2019/03/21 22:34		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.8853	8.8525	15.10	18.27	3.17	50.02	10:33
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
8	TOC	FB	0.3991 ppm	0.0000 ppm	0.0000%	2019/03/21 22:49		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3991	3.9909	11.80	14.63	2.83	50.08	10:32
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
9	TOC	K1902047-001.01	3.1747 ppm	0.0064 ppm	0.2000%	2019/03/21 23:03		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.1792	31.7917	30.67	33.69	3.02	50.09	10:26
2	TOC	3.1702	31.7018	30.61	33.43	2.82	50.08	10:26
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.7049 ppm (PASS)	0.0000 ppm	0%	2019/03/21 23:31

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.7049	247.0492	177.16	180.15	2.99	50.08	10:32
Completion State		Success Action		Method		Calibration		STD Conc - Pos B		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		50 ppmC		

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
◊	D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.2679 ppm (PASS)	0.0000 ppm	0%	2019/03/21 23:46

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.2679	2.6790	11.28	14.33	3.04	50.09	10:32
Completion State		Success Action		Method		Calibration		STD Conc - Pos D		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		0 ppmC		

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
◊	10	TOC	K1901927-005.03	46.1675 ppm	0.3093 ppm	0.6700%	2019/03/22 00:01

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	45.9488	459.4884	320.99	323.90	2.91	50.06	10:30
2	TOC	46.3862	463.8624	323.96	326.76	2.80	50.07	10:28
Dilution		Blank Contribution		Method		Calibration		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
◊	11	TOC	RB	0.6373 ppm	0.1548 ppm	24.2900%	2019/03/22 00:29

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.7468	7.4677	14.16	17.16	3.00	50.07	10:28
2	TOC	0.5279	5.2785	12.68	15.49	2.81	50.05	10:25
Dilution		Blank Contribution		Method		Calibration		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
◊	12	TOC	K1902389-001.01	3.5762 ppm	0.0031 ppm	0.0900%	2019/03/22 00:57

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.5740	35.7399	33.35	36.29	2.94	50.02	10:24
2	TOC	3.5784	35.7841	33.38	36.25	2.87	50.02	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
13	TOC	K1902389-001.01 ms	31.2217 ppm	0.0000 ppm	0.0000%	2019/03/22 01:25

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	31.2217	312.2170	221.02	223.80	2.78	50.00	10:31

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
14	TOC	RB	0.3757 ppm	0.0000 ppm	0.0000%	2019/03/22 01:40

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3757	3.7567	11.64	14.47	2.82	50.01	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
15	TOC	K1902389-002.01	27.6512 ppm	0.5067 ppm	1.8300%	2019/03/22 01:54

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	27.2930	272.9296	194.36	197.15	2.80	50.00	10:27
2	TOC	28.0095	280.0953	199.22	201.90	2.69	49.98	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
16	TOC	K1902389-003.01	28.1127 ppm	0.0750 ppm	0.2700%	2019/03/22 02:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	28.1657	281.6569	200.28	203.19	2.91	49.97	10:27
2	TOC	28.0596	280.5962	199.56	202.41	2.85	49.98	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Analysis	Std. Dev.

Pos	Type	Sample ID	Result (ppmC)	(ppmC)	RSD	Start Time
17	TOC	K1902153-001.01 100x	2.1834 ppm	0.3944 ppm	18.0600%	2019/03/22 02:50

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.4623	24.6231	25.81	28.60	2.80	49.95	10:30
2	TOC	1.9046	19.0456	22.02	24.86	2.84	49.96	10:28

Dilution 1:10 **Blank Contribution** (TC) 9.0920 (IC) (v1238) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
18	TOC	K1902153-002.01 100x	3.3888 ppm	0.0121 ppm	0.3600%	2019/03/22 03:19

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.3974	33.9735	32.15	35.09	2.93	50.01	10:28
2	TOC	3.3803	33.8026	32.04	34.91	2.87	49.96	10:29

Dilution 1:10 **Blank Contribution** (TC) 9.0920 (IC) (v1238) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
19	TOC	K1902153-003.01 100x	1.4009 ppm	0.0500 ppm	3.5700%	2019/03/22 03:47

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.4362	14.3623	18.84	21.63	2.79	49.95	10:28
2	TOC	1.3655	13.6551	18.36	21.10	2.74	49.93	10:26

Dilution 1:10 **Blank Contribution** (TC) 9.0920 (IC) (v1238) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.1491 ppm (PASS)	0.0000 ppm	0%	2019/03/22 04:15

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.1491	251.4909	180.17	182.80	2.63	49.90	10:31

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30) **STD Conc - Pos B** 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.2848 ppm (PASS)	0.0000 ppm	0%	2019/03/22 04:30

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.2848	2.8484	11.40	14.09	2.70	49.89	10:31

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 20	TOC	MB2	0.2384 ppm	0.0000 ppm	0.0000%	2019/03/22 04:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2384	2.3837	10.71	13.39	2.68	49.91	10:32

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	27.0967 ppm (PASS)	0.0000 ppm	0%	2019/03/22 04:59

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	27.0967	270.9666	193.39	196.26	2.87	49.90	10:29

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos C 25 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 21	TOC	K1902346-001.01	3.1362 ppm	0.0803 ppm	2.5600%	2019/03/22 05:14

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.1930	31.9302	30.77	33.56	2.79	49.89	10:30

2	TOC	3.0794	30.7944	30.00	32.92	2.92	49.86	10:26
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
22	TOC	K1902102-001.02	83.5303 ppm	0.1033 ppm	0.1200%	2019/03/22 05:42		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	83.4573	834.5727	575.59	578.46	2.87	49.88	10:28
2	TOC	83.6034	836.0342	576.59	579.49	2.90	49.93	10:28
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
23	TOC	K1902127-001.12	15.8880 ppm	0.1089 ppm	0.6900%	2019/03/22 06:10		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	15.9649	159.6493	117.46	120.43	2.96	49.89	10:27
2	TOC	15.8110	158.1098	116.42	119.03	2.61	49.79	10:30
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
24	TOC	K1902127-002.12 4x	7.2203 ppm	0.1077 ppm	1.4900%	2019/03/22 06:38		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.2965	72.9647	58.62	61.29	2.67	49.75	10:31
2	TOC	7.1441	71.4414	57.59	60.28	2.69	49.70	10:28
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
25	TOC	K1902127-003.12	7.8209 ppm	0.0934 ppm	1.1900%	2019/03/22 07:06		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.8869	78.8693	62.63	65.27	2.64	49.66	10:27
2	TOC	7.7548	77.5479	61.73	64.50	2.77	49.65	10:30
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		

26	TOC	K1902127-004.12 4x	4.6989 ppm	0.1032 ppm	2.2000%	2019/03/22 07:34		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	4.7718	47.7185	41.48	44.27	2.79	49.59	10:29
2	TOC	4.6259	46.2585	40.49	43.16	2.66	49.58	10:28
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
27	TOC	K1902147-001.01 100x	0.4629 ppm	0.0021 ppm	0.4500%	2019/03/22 08:02		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4644	4.6436	12.24	14.96	2.71	49.56	10:28
2	TOC	0.4614	4.6141	12.22	14.96	2.74	49.50	10:30
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
28	TOC	K1902147-002.01	1.1631 ppm	0.0267 ppm	2.2900%	2019/03/22 08:30		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.1820	11.8195	17.11	19.97	2.86	49.64	10:31
2	TOC	1.1442	11.4424	16.86	19.60	2.74	49.76	10:28
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.6255 ppm (PASS)	0.0000 ppm	0%	2019/03/22 08:58

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.6255	246.2551	176.62	179.28	2.66	49.91	10:30

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos B</u>
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Concentration	Min / Max

Pos	BAT	(ppm)	Dil	Sample ID	(% dev)	Result	Std. Dev.	RSD	Start Time	
♦	D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0686 ppm (PASS)	0.0000 ppm	0%	2019/03/22 09:13

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0686	0.6858	9.93	12.79	2.86	49.93	10:30

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos D</u>
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	29	TOC	K1902168-001.04	2.0571 ppm	0.0518 ppm	2.5200%	2019/03/22 09:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.0937	20.9372	23.30	25.96	2.66	49.96	10:30
2	TOC	2.0205	20.2050	22.81	25.66	2.86	50.00	10:27

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	30	TOC	K1902168-002.04	1.0525 ppm	0.0826 ppm	7.8500%	2019/03/22 09:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.1109	11.1094	16.63	19.46	2.83	50.01	10:25
2	TOC	0.9941	9.9412	15.84	18.81	2.97	50.01	10:28

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	31	TOC	K1902169-001.24	14.4701 ppm	0.0860 ppm	0.5900%	2019/03/22 10:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	14.5309	145.3092	107.73	110.45	2.72	49.97	10:29
2	TOC	14.4092	144.0923	106.90	109.89	2.99	49.97	10:30

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time	
♦	32	TOC	K1902169-002.24	17.8450 ppm	0.1318 ppm	0.7400%	2019/03/22 10:52

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	17.7518	177.5178	129.59	132.45	2.86	49.97	10:26
2	TOC	17.9381	179.3814	130.85	133.67	2.81	49.97	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
33	TOC	K1902169-003.24	18.8828 ppm	0.0274 ppm	0.1500%	2019/03/22 11:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	18.9022	189.0220	137.40	140.21	2.81	49.95	10:32
2	TOC	18.8635	188.6346	137.14	140.09	2.96	49.96	10:25

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
34	TOC	K1902169-004.24	9.3550 ppm	0.0898 ppm	0.9600%	2019/03/22 11:48

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	9.4185	94.1847	73.02	75.99	2.97	49.97	10:27
2	TOC	9.2915	92.9148	72.16	75.02	2.86	49.97	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
35	TOC	K1902169-005.24	11.8106 ppm	0.0557 ppm	0.4700%	2019/03/22 12:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	11.8500	118.4999	89.53	92.49	2.96	49.98	10:28
2	TOC	11.7712	117.7117	88.99	92.00	3.01	49.97	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
36	TOC	K1902169-006.24	11.8325 ppm	0.1654 ppm	1.4000%	2019/03/22 12:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	11.9494	119.4943	90.20	93.00	2.79	49.99	10:29
2	TOC	11.7155	117.1549	88.62	91.50	2.88	49.96	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC)
Method CAS_salt_010711
Calibration CAS_salt_010711

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
37	TOC	K1902169-007.24	12.8921 ppm	0.2771 ppm	2.1500%	2019/03/22 13:12

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	12.6962	126.9620	95.27	98.09	2.82	50.01	10:30
2	TOC	13.0881	130.8807	97.93	100.84	2.91	50.03	10:27

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
38	TOC	K1902169-008.24	15.4016 ppm	0.2088 ppm	1.3600%	2019/03/22 13:41

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	15.5492	155.4920	114.64	117.52	2.88	50.05	10:30
2	TOC	15.2540	152.5397	112.64	115.67	3.04	50.06	10:25

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.4810 ppm (PASS)	0.0000 ppm	0%	2019/03/22 14:09

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.4810	244.8099	175.64	178.60	2.96	50.07	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0485 ppm (PASS)	0.0000 ppm	0%	2019/03/22 14:23

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0485	0.4854	9.79	12.80	3.01	50.09	10:33

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos D</u>
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
39	TOC	MB3	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/22 14:38

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.49	11.48	2.99	50.09	10:29

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.3372 ppm (PASS)	0.0000 ppm	0%	2019/03/22 14:53

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.3372	263.3722	188.24	191.16	2.92	50.10	10:32

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos C</u>
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	25 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
40	TOC	K1902230-001.05	2.0965 ppm	0.0063 ppm	0.3000%	2019/03/22 15:07

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.1009	21.0094	23.35	26.30	2.94	50.08	10:27
2	TOC	2.0921	20.9210	23.29	26.37	3.08	50.08	10:27

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
41	TOC	K1902230-002.05	0.9090 ppm	0.0394 ppm	4.3300%	2019/03/22 15:35

Rep	Base	ppm	µg	Adjusted	NDIR (Abs)	Baseline	Pressure	Run
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#	Analysis Type			(Abs)		(Abs)	(psig)	Time
1	TOC	0.9368	9.3681	15.45	18.19	2.74	50.11	10:24
2	TOC	0.8811	8.8113	15.07	17.96	2.89	50.10	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
42	TOC	K1902230-003.05	1.0304 ppm	0.0033 ppm	0.3200%	2019/03/22 16:03

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.0327	10.3272	16.10	18.96	2.86	50.11	10:29
2	TOC	1.0280	10.2800	16.07	18.99	2.92	50.10	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
43	TOC	K1902230-004.05	1.4208 ppm	0.0626 ppm	4.4100%	2019/03/22 16:32

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.4651	14.6510	19.04	22.03	2.99	50.09	10:29
2	TOC	1.3766	13.7656	18.44	21.48	3.04	50.06	10:23

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
44	TOC	K1902230-005.05	0.5452 ppm	0.0150 ppm	2.7500%	2019/03/22 17:00

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5558	5.5584	12.86	15.72	2.85	50.11	10:28
2	TOC	0.5346	5.3463	12.72	15.64	2.92	50.10	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
45	TOC	K1902232-001.12	5.9922 ppm	0.0482 ppm	0.8000%	2019/03/22 17:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.9581	59.5807	49.54	52.48	2.94	50.09	10:26
2	TOC	6.0263	60.2628	50.00	53.07	3.07	50.11	10:27

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	33.5471	335.4714	236.81	239.79	2.98	50.10	10:32
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
47	TOC	RB	0.0443 ppm	0.0000 ppm	0.0000%	2019/03/22 18:11		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0443	0.4435	9.39	12.26	2.87	50.11	10:29
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>	<u>Calibration</u>			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)			

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
47	B	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.1426 ppm (PASS)	0.0000 ppm	0%	2019/03/22 18:25	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.1426	241.4260	173.34	176.18	2.84	50.17	10:30
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos B</u>				
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC				

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
47	D	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/22 18:40	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	8.99	12.03	3.03	50.10	10:32
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos D</u>				
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC				

Sample Type: Sample							From Schedule Version 4		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
48	TOC	K1902232-002.12	10.0441 ppm	0.0777 ppm	0.7700%	2019/03/22 18:55			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	9.9892	99.8919	76.90	79.97	3.07	50.16	10:29	
2	TOC	10.0991	100.9909	77.64	80.42	2.78	50.05	10:27	
Dilution		Blank Contribution		Method		Calibration			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
49	TOC	K1902232-003.12	14.1220 ppm	0.0201 ppm	0.1400%	2019/03/22 19:23			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	14.1362	141.3625	105.05	108.04	3.00	50.06	10:26	
2	TOC	14.1078	141.0782	104.86	107.79	2.93	50.04	10:23	
Dilution		Blank Contribution		Method		Calibration			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
50	TOC	K1902121-001.01	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/22 19:51			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	0.0000	0.0000	8.86	11.75	2.89	50.04	10:25	
2	TOC	0.0000	0.0000	9.03	12.04	3.01	50.03	10:29	
Dilution		Blank Contribution		Method		Calibration			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
51	TOC	K1902121-002.01	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/22 20:19			
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time	
1	TOC	0.0000	0.0000	8.20	11.22	3.02	50.00	10:31	
2	TOC	0.0000	0.0000	8.53	11.57	3.04	50.01	10:27	
Dilution		Blank Contribution		Method		Calibration			
1:10		(TC) 9.0920 (IC) (v1238)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)			
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time			
52	TOC	K1902435-001.02	116.2162 ppm	5.5539 ppm	4.7800%	2019/03/22 20:47			

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	120.1434	1201.4336	824.62	827.61	2.99	49.98	10:28
2	TOC	112.2890	1122.8899	771.30	774.91	3.61	49.98	10:24

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
53	TOC	K1902435-002.01	4.2973 ppm	1.4335 ppm	33.3600%	2019/03/22 21:15

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.3109	53.1089	45.14	48.45	3.31	49.96	10:25
2	TOC	3.2836	32.8362	31.38	34.42	3.04	49.97	10:25

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
54	TOC	K1902435-003.01	3.2785 ppm	0.2293 ppm	6.9900%	2019/03/22 21:43

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.4407	34.4066	32.45	35.43	2.98	49.95	10:30
2	TOC	3.1164	31.1641	30.25	33.19	2.94	49.96	10:29

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
55	TOC	K1902435-004.01	5.1276 ppm	0.2636 ppm	5.1400%	2019/03/22 22:11

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.3140	53.1399	45.16	48.09	2.93	49.95	10:26
2	TOC	4.9413	49.4127	42.63	45.55	2.91	49.94	10:30

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
56	TOC	K1902435-005.01	132.6232 ppm	6.5383 ppm	4.9300%	2019/03/22 22:39

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	137.2465	1372.4649	940.71	943.70	2.99	49.92	10:25
2	TOC	127.9999	1279.9994	877.95	881.72	3.77	49.99	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC)
Method CAS_salt_010711
Calibration CAS_salt_010711

		(v1238)	(v4)	(v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
57	TOC	K1902435-006.01	122.9733 ppm	6.3170 ppm	5.1400%	2019/03/22 23:07

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	118.5065	1185.0649	813.51	817.08	3.57	50.07	10:26
2	TOC	127.4401	1274.4013	874.15	878.10	3.95	50.09	10:27

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 9.0920 (IC) (v1238)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	27.6282 ppm (PASS)	0.0000 ppm	0%	2019/03/22 23:35

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	27.6282	276.2819	197.00	200.68	3.68	50.18	10:31

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	1.1283 ppm (PASS)	0.0000 ppm	0%	2019/03/22 23:50

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	1.1283	11.2825	17.12	20.21	3.09	50.20	10:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
59	TOC	K1902435-007.01	12.6387 ppm	0.1397 ppm	1.1100%	2019/03/23 00:05

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	12.7374	127.3745	95.55	98.46	2.91	50.20	10:30
2	TOC	12.5399	125.3989	94.21	97.27	3.06	50.20	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
60	TOC	K1902435-007.01 ms	33.4785 ppm	0.0000 ppm	0.0000%	2019/03/23 00:33

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	33.4785	334.7849	236.34	239.34	3.00	50.22	10:33

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
61	TOC	RB	0.6159 ppm	0.0000 ppm	0.0000%	2019/03/23 00:48

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6159	6.1595	13.27	16.42	3.14	50.22	10:33

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
62	TOC	K1902254-001.01	6.6998 ppm	0.1143 ppm	1.7100%	2019/03/23 01:02

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.7806	67.8056	55.12	58.04	2.92	50.23	10:28
2	TOC	6.6189	66.1895	54.02	57.06	3.04	50.22	10:26

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
63	TOC	K1902254-002.01	18.5047 ppm	0.1440 ppm	0.7800%	2019/03/23 01:30

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	18.6065	186.0653	135.39	138.31	2.92	50.23	10:30
2	TOC	18.4029	184.0293	134.01	137.12	3.11	50.22	10:29

Dilution 1:10
Blank Contribution (TC) 9.0920 (IC) (v1238)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Analysis	Std. Dev.

Pos	Type	Sample ID	Result (ppmC)	(ppmC)	RSD	Start Time
64	TOC	K1902254-003.01	22.2028 ppm	0.0577 ppm	0.2600%	2019/03/23 01:58

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	22.2436	222.4357	160.08	163.10	3.02	50.23	10:28
2	TOC	22.1620	221.6195	159.53	162.65	3.12	50.23	10:25

Dilution 1:10 Blank Contribution (TC) 9.0920 (IC) (v1238) Method CAS_salt_010711 (v4) Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
65	TOC	RB	0.0278 ppm	0.0393 ppm	141.4200%	2019/03/23 02:27

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0555	0.5554	9.47	12.55	3.08	50.21	10:29
2	TOC	0.0000	0.0000	8.79	11.75	2.96	50.21	10:26

Dilution 1:10 Blank Contribution (TC) 9.0920 (IC) (v1238) Method CAS_salt_010711 (v4) Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	24.0426 ppm (PASS)	0.0000 ppm	0%	2019/03/23 02:55

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	24.0426	240.4257	172.66	175.53	2.87	50.20	10:30

Completion State Success - Criteria met. Success Action Do Nothing Method CAS_salt_010711 (v4) Calibration CAS_salt_010711 (v30) STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/23 03:09

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	9.20	12.29	3.09	50.20	10:29

Completion State Success - Criteria Success Action Do Nothing Method CAS_salt_010711 Calibration CAS_salt_010711 STD Conc - Pos D 0 ppmC

met.

(v4)

(v30)

Meta Data Used in this Report

Blanks

Version	Reagent (Abs)	Acid (Abs)	DI IC (Abs)	DI TC (Abs)	DI TOC (Abs)	Save Time	Operator
v1237	1.0520	0.7490	0.0000	0.0000	0.0000	2019/03/19 18:04	Fusion1 (Fusion1)
v1238	2.8787	1.7600	0.0000	0.0000	0.0000	2019/03/21 19:27	Fusion1 (Fusion1)

Calibrations

Name: CAS_salt_010711 (TOC)

Version: v30
 Calibration curve formula: TOC: $y = 6.788x + 9.463$
 Ver Creation: 2019/03/05 17:42
 r^2 value: TOC: $r^2 = 0.99963$
 Comment:
 Operator: Fusion1 (Fusion1)
 Basic Analysis Type: TOC

Basic Analysis Type: TOC

Sample ID	Y Raw Value	X Expected	Message	End Time
DI Water	7.8970	0.0000		2019/03/05 16:15
0.500 ppm	11.5280	0.5000		2019/03/05 16:29
1.0 ppm	14.9760	1.0000		2019/03/05 16:44
5.0 ppm	43.6500	5.0000		2019/03/05 16:58
10 ppm	79.6020	10.0000		2019/03/05 17:12
25 ppm	183.3580	25.0000		2019/03/05 17:26
50 ppm	346.3230	50.0000		2019/03/05 17:40

Methods

Name: CAS_salt_010711 (TOC)

Version: v4
 Operator: Fusion1 (Fusion1)
 Ver Creation: 2019/02/21 17:57
 Comment:

Parameter	Value	Advanced Parameter	Value
SampleVolume	10.0 mL	NeedleRinseVolume	5.0 ml
Dilution	1:10	VialPrimeVolume	2.0 ml
AcidVolume	0.5 ml	ICSamplePrimeVolume	2.0 ml

ReagentVolume	2.0 ml	ICSpargeRinseVolume	12.0 ml
UVReactorPrerinse	Off	BaselineStabilizeTime	0.70 min
UVReactorPrerinseVolume	5.0	DetectorPressureFlow	150 ml/min
NumberOfUVReactorPrerinses	1	SyringeSpeedWaste	10
ICSpargeTime	1.00 mins	SyringeSpeedAcid	7
DetectorSweepFlow	500 ml/min	SyringeSpeedReagent	7
PreSpargeTime	2.00 mins	SyringeSpeedDIWater	7
SystemFlow	500 ml/min	NDIRPressurization	60 psig
		SyringeSpeedSampleDispense	5
		SyringeSpeedSampleAspirate	4
		SyringeSpeedUVDispense	5
		SyringeSpeedUVAspirate	5
		SyringeSpeedICDispense	5
		SyringeSpeedICAspirate	5
		NDIRPressureStabilize	1.75 min
		SampleMixing	Off
		SampleMixingCycles	1
		SampleMixingVolume	10.0
		LowLevelFilterNDIR	Off

Acceptance / Approval

Electronic Signatures

Report Version	User Name	Acceptance	Reason	Date
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Report History

Report History

Report Version	User Name	System Reason	User Reason	Date
1	Fusion1 (Fusion1)	Schedule completed	Schedule completed	2019/03/23 03:26

StarLIMS Run: 629142, 629144, 629629146, 629149
 Analysis: TOC/DOC
 Method: 9060, 415.1, SM 5310 C, 9060A

CCV: 11-GEN-05-76B 50 ppm LCS: 11-GEN-05-74L 25.0 ppm

ICAL Date: 3/6/19

ICAL ID: 11-GEN-05-76H

ICS ID: 11-GEN-05-74A

ICS TV: 25.0 ppm ICS % R = 2

Spike ID: 11-GEN-05-700 0.05 ml of 5000 ppm stock ---> 10.0 ml = 25.0 ppm x dilution factor

Sodium Persulfate: 11-GEN-05-77B

21 % H3PO4: 11-GEN-05-77A

Equipment ID: K-TOC-03

PIPETTE ID: 124276B, 129001F, N11314F, Marge

FILTER ID: NA

Analyzed By: <i>BCD</i>	Date Analyzed: ²¹⁰ <i>3/23/19</i>
Reviewed By: <i>Hamp</i>	Date Reviewed: <i>03/25/19</i>

BCD 3/23/19



10450 Stancliff Rd. Suite 210
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T: +1 281 530 5656
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March 31, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030761**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples**

Dear Marcia,

ALS Environmental received 3 sample(s) on Mar 15, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a hand-drawn oval.

Generated By: JUMOKE.LAWAL

RJ Modashia
Project Manager

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
Work Order: HS19030761

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030761-01	LH18/24-SP650_031419	Water		14-Mar-2019 14:00	15-Mar-2019 08:38	<input type="checkbox"/>
HS19030761-02	LH18/24-SP650_031419_BIX	Water		14-Mar-2019 14:00	15-Mar-2019 08:38	<input type="checkbox"/>
HS19030761-03	Trip Blank	Water	ALS 022719-66	14-Mar-2019 14:00	15-Mar-2019 08:38	<input type="checkbox"/>

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.

GCMS Semivolatiles by Method SW8270SIM**Batch ID: 138779****Sample ID: LCSD-138779**

- The RPD between the LCS and LCSD was outside of the control limit. Surrogate Nitrobenzene-d5

Sample ID: LH18/24-SP650_031419 (HS19030761-01)

- One or more of the method 8270 surrogates were recovered outside of the control limits. This was due to a dilution required for sample analysis.

GCMS Volatiles by Method SW8260**Batch ID: R334643****Sample ID: HS19030640-01MS**

- MS and MSD are for an unrelated sample

Metals by Method SW6020**Batch ID: 138939****Sample ID: HS19030918-01MS**

- MS and MSD are for an unrelated sample

WetChemistry by Method SW7196**Batch ID: R334752**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: LH18/24-SP650_031419
 Collection Date: 14-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030761
 Lab ID:HS19030761-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2-Dichloroethane	0.53	J	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
2-Butanone	3.5		0.50	1.0	2.0	UG/L	1	15-Mar-2019 13:16	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	15-Mar-2019 13:16	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	15-Mar-2019 13:16	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	15-Mar-2019 13:16	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	15-Mar-2019 13:16	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: LH18/24-SP650_031419
 Collection Date: 14-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030761
 Lab ID:HS19030761-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
cis-1,2-Dichloroethene	2.1		0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	15-Mar-2019 13:16	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	15-Mar-2019 13:16	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	15-Mar-2019 13:16	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Trichloroethene	0.74	J	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 13:16	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>88.8</i>			0	<i>81-118</i>	%REC	<i>1</i>	<i>15-Mar-2019 13:16</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>97.3</i>			0	<i>85-114</i>	%REC	<i>1</i>	<i>15-Mar-2019 13:16</i>	
<i>Surr: Dibromofluoromethane</i>	<i>90.5</i>			0	<i>80-119</i>	%REC	<i>1</i>	<i>15-Mar-2019 13:16</i>	
<i>Surr: Toluene-d8</i>	<i>106</i>			0	<i>89-112</i>	%REC	<i>1</i>	<i>15-Mar-2019 13:16</i>	
SEMIVOLATILES SIM		Method:SW8270SIM						Prep:SW3510 / 18-Mar-2019 Analyst: QX	
1,4-Dioxane	3.1		0.10	0.10	0.10	ug/L	10	18-Mar-2019 16:42	
<i>Surr: 2-Fluorobiphenyl</i>	<i>193</i>	S		0	<i>40-140</i>	%REC	<i>10</i>	<i>18-Mar-2019 16:42</i>	
<i>Surr: 4-Terphenyl-d14</i>	<i>62.7</i>			0	<i>40-140</i>	%REC	<i>10</i>	<i>18-Mar-2019 16:42</i>	
<i>Surr: Nitrobenzene-d5</i>	<i>103</i>			0	<i>40-140</i>	%REC	<i>10</i>	<i>18-Mar-2019 16:42</i>	
ICP-MS METALS BY SW6020A		Method:SW6020						Prep:SW3010A / 21-Mar-2019 Analyst: JCJ	
Barium	0.139		0.00190	0.00250	0.00400	mg/L	1	23-Mar-2019 01:05	
Lead	0.00100	U	0.000600	0.00100	0.00200	mg/L	1	23-Mar-2019 01:05	
Selenium	0.00250	U	0.00110	0.00250	0.00200	mg/L	1	23-Mar-2019 01:05	
Silver	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	23-Mar-2019 01:05	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 31-Mar-19

Client:	Bhate Environmental Associates, Inc.	ANALYTICAL REPORT
Project:	LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples	WorkOrder:HS19030761
Sample ID:	LH18/24-SP650_031419	Lab ID:HS19030761-01
Collection Date:	14-Mar-2019 14:00	Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
HEXAVALENT CHROMIUM BY SW7196A		Method:SW7196		Analyst: MZD				
Chromium, Hexavalent	0.0100	U	0.00600	0.0100	0.0100	mg/L	1	15-Mar-2019 12:14

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 31-Mar-19

Client:	Bhate Environmental Associates, Inc.	ANALYTICAL REPORT
Project:	LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples	WorkOrder:HS19030761
Sample ID:	LH18/24-SP650_031419_BIX	Lab ID:HS19030761-02
Collection Date:	14-Mar-2019 14:00	Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	31-Mar-2019 12:13

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: Trip Blank
 Collection Date: 14-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030761
 Lab ID:HS19030761-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	15-Mar-2019 12:52	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	15-Mar-2019 12:52	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	15-Mar-2019 12:52	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	15-Mar-2019 12:52	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	15-Mar-2019 12:52	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
 Sample ID: Trip Blank
 Collection Date: 14-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030761
 Lab ID:HS19030761-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	15-Mar-2019 12:52	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	15-Mar-2019 12:52	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	15-Mar-2019 12:52	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Trichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	15-Mar-2019 12:52	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>88.7</i>			0	<i>81-118</i>	<i>%REC</i>	<i>1</i>	<i>15-Mar-2019 12:52</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>102</i>			0	<i>85-114</i>	<i>%REC</i>	<i>1</i>	<i>15-Mar-2019 12:52</i>	
<i>Surr: Dibromofluoromethane</i>	<i>90.1</i>			0	<i>80-119</i>	<i>%REC</i>	<i>1</i>	<i>15-Mar-2019 12:52</i>	
<i>Surr: Toluene-d8</i>	<i>103</i>			0	<i>89-112</i>	<i>%REC</i>	<i>1</i>	<i>15-Mar-2019 12:52</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

WEIGHT LOG

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

Batch ID: 138779 **Method:** SEMIVOLATILES SIM **Prep:** 3510_B_SIM

SampID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19030761-01	1	980	1 (mL)	0.00102

Batch ID: 138939 **Method:** ICP-MS METALS BY SW6020A **Prep:** 3010A

SampID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19030761-01	1	10	10 (mL)	1

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID 138779	Test Name : SEMIVOLATILES SIM		Matrix: Water			
HS19030761-01	LH18/24-SP650_031419	14 Mar 2019 14:00		18 Mar 2019 12:38	18 Mar 2019 16:42	10
Batch ID 138939	Test Name : ICP-MS METALS BY SW6020A		Matrix: Water			
HS19030761-01	LH18/24-SP650_031419	14 Mar 2019 14:00		21 Mar 2019 10:00	23 Mar 2019 01:05	1
Batch ID R334643	Test Name : VOLATILES ORGANICS BY METHOD 8260C		Matrix: Water			
HS19030761-01	LH18/24-SP650_031419	14 Mar 2019 14:00			15 Mar 2019 13:16	1
HS19030761-03	Trip Blank	14 Mar 2019 14:00			15 Mar 2019 12:52	1
Batch ID R334752	Test Name : HEXAVALENT CHROMIUM BY SW7196A		Matrix: Water			
HS19030761-01	LH18/24-SP650_031419	14 Mar 2019 14:00			15 Mar 2019 12:14	1
Batch ID R335660	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19030761-02	LH18/24-SP650_031419_BIX	14 Mar 2019 14:00			31 Mar 2019 12:13	1

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: 138939 (0)		Instrument: ICPMS04		Method: ICP-MS METALS BY SW6020A						
MBLK	Sample ID: MBLK-138939	Units: mg/L			Analysis Date: 23-Mar-2019 00:40					
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003309		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.00250	0.00400								U
Lead	0.00100	0.00200								U
Selenium	0.00250	0.00200								U
Silver	0.000500	0.00200								U
LCS	Sample ID: LCS-138939	Units: mg/L			Analysis Date: 23-Mar-2019 00:42					
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003310		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.05067	0.00400	0.05	0	101	80 - 120				
Lead	0.05015	0.00200	0.05	0	100	80 - 120				
Selenium	0.05143	0.00200	0.05	0	103	80 - 120				
Silver	0.05011	0.00200	0.05	0	100	80 - 120				
MS	Sample ID: HS19030918-01MS	Units: mg/L			Analysis Date: 23-Mar-2019 00:49					
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003313		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.4055	0.00400	0.05	0.3415	128	80 - 120				SO
Lead	0.04827	0.00200	0.05	0.000163	96.2	80 - 120				
Selenium	0.05006	0.00200	0.05	-0.000569	101	80 - 120				
Silver	0.04751	0.00200	0.05	-0.000012	95.0	80 - 120				
MSD	Sample ID: HS19030918-01MSD	Units: mg/L			Analysis Date: 23-Mar-2019 00:51					
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003314		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.4147	0.00400	0.05	0.3415	146	80 - 120	0.4055	2.23	20	SO
Lead	0.05045	0.00200	0.05	0.000163	101	80 - 120	0.04827	4.41	20	
Selenium	0.0556	0.00200	0.05	-0.000569	112	80 - 120	0.05006	10.5	20	
Silver	0.04885	0.00200	0.05	-0.000012	97.7	80 - 120	0.04751	2.79	20	

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: 138939 (0)		Instrument: ICPMS04		Method: ICP-MS METALS BY SW6020A						
PDS	Sample ID: HS19030918-01PDS	Units: mg/L			Analysis Date: 23-Mar-2019 00:53					
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003315		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Barium	0.4541	0.00400	0.1	0.3415	113	75 - 125				
Lead	0.09808	0.00200	0.1	0.000163	97.9	75 - 125				
Selenium	0.1029	0.00200	0.1	-0.000569	103	75 - 125				
Silver	0.08679	0.00200	0.1	-0.000012	86.8	75 - 125				
SD	Sample ID: HS19030918-01SD	Units: mg/L			Analysis Date: 23-Mar-2019 00:47					
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003312		PrepDate: 21-Mar-2019		DF: 5				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	RPD Limit	Qual
Barium	0.3292	0.0200					0.3415	3.6	10	
Lead	0.00500	0.0100					0.000163	0	10	U
Selenium	0.0125	0.0100					-0.000569	0	10	U
Silver	0.00250	0.0100					-0.000012	0	10	U

The following samples were analyzed in this batch: HS19030761-01

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: 138779 (0)		Instrument: SV-5		Method: SEMIVOLATILES SIM						
MBLK	Sample ID: MBLK-138779	Units: ug/L			Analysis Date: 18-Mar-2019 12:55					
Client ID:	Run ID: SV-5_334775	SeqNo: 4995605		PrepDate: 18-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,4-Dioxane	0.010	0.010								U
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.09739</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>122</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.08402</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>105</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.07978</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>99.7</i>	<i>40 - 140</i>				
LCS	Sample ID: LCS-138779	Units: ug/L			Analysis Date: 18-Mar-2019 13:16					
Client ID:	Run ID: SV-5_334775	SeqNo: 4995606		PrepDate: 18-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,4-Dioxane	0.08961	0.010	0.08	0	112	40 - 140				
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.1004</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>126</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.05163</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>64.5</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.07478</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>93.5</i>	<i>40 - 140</i>				
LCSD	Sample ID: LCSD-138779	Units: ug/L			Analysis Date: 18-Mar-2019 13:37					
Client ID:	Run ID: SV-5_334775	SeqNo: 4995607		PrepDate: 18-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,4-Dioxane	0.09058	0.010	0.08	0	113	40 - 140	0.08961	1.07	20	
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.1045</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>131</i>	<i>40 - 140</i>	<i>0.1004</i>	<i>4.04</i>	<i>20</i>	
<i>Surr: 4-Terphenyl-d14</i>	<i>0.05772</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>72.2</i>	<i>40 - 140</i>	<i>0.05163</i>	<i>11.1</i>	<i>20</i>	
<i>Surr: Nitrobenzene-d5</i>	<i>0.09447</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>118</i>	<i>40 - 140</i>	<i>0.07478</i>	<i>23.3</i>	<i>20</i>	R
The following samples were analyzed in this batch: <input type="text" value="HS19030761-01"/>										

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190315	Units: UG/L			Analysis Date: 15-Mar-2019 12:28					
Client ID:	Run ID: VOA6_334643	SeqNo: 4992329	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	0.50	1.0								U
1,1,1-Trichloroethane	0.50	1.0								U
1,1,2,2-Tetrachloroethane	0.50	1.0								U
1,1,2-Trichloroethane	0.50	1.0								U
1,1-Dichloroethane	0.50	1.0								U
1,1-Dichloroethene	0.50	1.0								U
1,1-Dichloropropene	0.50	1.0								U
1,2,3-Trichlorobenzene	0.50	1.0								U
1,2,3-Trichloropropane	0.50	1.0								U
1,2,4-Trichlorobenzene	0.50	1.0								U
1,2,4-Trimethylbenzene	0.50	1.0								U
1,2-Dibromo-3-chloropropane	0.50	1.0								U
1,2-Dibromoethane	0.50	1.0								U
1,2-Dichlorobenzene	0.50	1.0								U
1,2-Dichloroethane	0.50	1.0								U
1,2-Dichloropropane	0.50	1.0								U
1,3,5-Trimethylbenzene	0.50	1.0								U
1,3-Dichlorobenzene	0.50	1.0								U
1,3-Dichloropropane	0.50	1.0								U
1,4-Dichlorobenzene	0.50	1.0								U
2,2-Dichloropropane	0.50	1.0								U
2-Butanone	1.0	2.0								U
2-Chlorotoluene	0.50	1.0								U
2-Hexanone	1.0	2.0								U
4-Chlorotoluene	0.50	1.0								U
4-Isopropyltoluene	0.50	1.0								U
4-Methyl-2-pentanone	1.0	2.0								U
Acetone	2.0	2.0								U
Benzene	0.50	1.0								U
Bromobenzene	0.50	1.0								U
Bromochloromethane	0.50	1.0								U
Bromodichloromethane	0.50	1.0								U
Bromoform	0.50	1.0								U
Bromomethane	0.50	1.0								U

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190315	Units: UG/L			Analysis Date: 15-Mar-2019 12:28					
Client ID:	Run ID: VOA6_334643	SeqNo: 4992329	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	45.24	1.0	50	0	90.5	81 - 118				
Surr: 4-Bromofluorobenzene	49.92	1.0	50	0	99.8	85 - 114				
Surr: Dibromofluoromethane	45.41	1.0	50	0	90.8	80 - 119				
Surr: Toluene-d8	53.38	1.0	50	0	107	89 - 112				

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190315	Units: UG/L			Analysis Date: 15-Mar-2019 11:40					
Client ID:	Run ID: VOA6_334643	SeqNo: 4992328	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	22.16	1.0	20	0	111	78 - 124				
1,1,1-Trichloroethane	16.14	1.0	20	0	80.7	74 - 131				
1,1,2,2-Tetrachloroethane	22.71	1.0	20	0	114	71 - 121				
1,1,2-Trichloroethane	22.94	1.0	20	0	115	80 - 119				
1,1-Dichloroethane	18.71	1.0	20	0	93.6	77 - 125				
1,1-Dichloroethene	15.69	1.0	20	0	78.5	71 - 131				
1,1-Dichloropropene	16.39	1.0	20	0	81.9	78 - 125				
1,2,3-Trichlorobenzene	18.01	1.0	20	0	90.1	69 - 129				
1,2,3-Trichloropropane	22.87	1.0	20	0	114	73 - 122				
1,2,4-Trichlorobenzene	19.19	1.0	20	0	95.9	69 - 130				
1,2,4-Trimethylbenzene	19.85	1.0	20	0	99.3	76 - 124				
1,2-Dibromo-3-chloropropane	22.7	1.0	20	0	114	62 - 128				
1,2-Dibromoethane	23.61	1.0	20	0	118	77 - 121				
1,2-Dichlorobenzene	21.03	1.0	20	0	105	80 - 119				
1,2-Dichloroethane	21.58	1.0	20	0	108	73 - 128				
1,2-Dichloropropane	20.91	1.0	20	0	105	78 - 122				
1,3,5-Trimethylbenzene	19.08	1.0	20	0	95.4	75 - 124				
1,3-Dichlorobenzene	20.4	1.0	20	0	102	80 - 119				
1,3-Dichloropropane	23.02	1.0	20	0	115	80 - 119				
1,4-Dichlorobenzene	20.42	1.0	20	0	102	79 - 118				
2,2-Dichloropropane	17.11	1.0	20	0	85.5	60 - 139				
2-Butanone	41.75	2.0	40	0	104	56 - 143				
2-Chlorotoluene	19.61	1.0	20	0	98.0	79 - 122				
2-Hexanone	45.67	2.0	40	0	114	57 - 139				
4-Chlorotoluene	19.89	1.0	20	0	99.5	78 - 122				
4-Isopropyltoluene	17.81	1.0	20	0	89.1	77 - 127				
4-Methyl-2-pentanone	45.87	2.0	40	0	115	67 - 130				
Acetone	41.97	2.0	40	0	105	39 - 160				
Benzene	20.55	1.0	20	0	103	79 - 120				
Bromobenzene	21.65	1.0	20	0	108	80 - 120				
Bromochloromethane	19.78	1.0	20	0	98.9	78 - 123				
Bromodichloromethane	21.11	1.0	20	0	106	79 - 125				
Bromoform	22.6	1.0	20	0	113	66 - 130				
Bromomethane	22.21	1.0	20	0	111	53 - 141				

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190315	Units: UG/L			Analysis Date: 15-Mar-2019 11:40					
Client ID:	Run ID: VOA6_334643	SeqNo: 4992328	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	32.8	2.0	40	0	82.0	64 - 133				
Carbon tetrachloride	17.12	1.0	20	0	85.6	72 - 136				
Chlorobenzene	21.13	1.0	20	0	106	82 - 118				
Chloroethane	17.29	1.0	20	0	86.4	60 - 138				
Chloroform	19.52	1.0	20	0	97.6	79 - 124				
Chloromethane	19.05	1.0	20	0	95.3	50 - 139				
cis-1,2-Dichloroethene	19.07	1.0	20	0	95.3	78 - 123				
cis-1,3-Dichloropropene	21.65	1.0	20	0	108	75 - 124				
Dibromochloromethane	23.09	1.0	20	0	115	74 - 126				
Dibromomethane	21.72	1.0	20	0	109	79 - 123				
Dichlorodifluoromethane	14.43	1.0	20	0	72.1	32 - 152				
Ethylbenzene	19.86	1.0	20	0	99.3	79 - 121				
Hexachlorobutadiene	18.24	1.0	20	0	91.2	66 - 134				
Isopropylbenzene	18.08	1.0	20	0	90.4	72 - 131				
m,p-Xylene	39.8	2.0	40	0	99.5	80 - 121				
Methylene chloride	20.23	2.0	20	0	101	74 - 124				
Naphthalene	19.53	1.0	20	0	97.6	61 - 128				
n-Butylbenzene	17.73	1.0	20	0	88.6	75 - 128				
n-Propylbenzene	18.26	1.0	20	0	91.3	76 - 126				
o-Xylene	20.76	1.0	20	0	104	78 - 122				
sec-Butylbenzene	16.87	1.0	20	0	84.4	77 - 126				
Styrene	22.12	1.0	20	0	111	78 - 123				
tert-Butylbenzene	17.52	1.0	20	0	87.6	78 - 124				
Tetrachloroethene	18.01	1.0	20	0	90.0	74 - 129				
Toluene	20.99	1.0	20	0	105	80 - 121				
trans-1,2-Dichloroethene	18.14	1.0	20	0	90.7	75 - 124				
trans-1,3-Dichloropropene	22.11	1.0	20	0	111	73 - 127				
Trichloroethene	18.65	1.0	20	0	93.3	79 - 123				
Trichlorofluoromethane	14.72	1.0	20	0	73.6	65 - 141				
Vinyl chloride	15.41	1.0	20	0	77.1	58 - 137				
Surr: 1,2-Dichloroethane-d4	44.38	1.0	50	0	88.8	81 - 118				
Surr: 4-Bromofluorobenzene	49.83	1.0	50	0	99.7	85 - 114				
Surr: Dibromofluoromethane	45.28	1.0	50	0	90.6	80 - 119				
Surr: Toluene-d8	52.92	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19030640-01MS	Units: UG/L			Analysis Date: 15-Mar-2019 15:16					
Client ID:	Run ID: VOA6_334643	SeqNo: 4993378	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	21.87	1.0	20	0	109	78 - 124				
1,1,1-Trichloroethane	18.37	1.0	20	0	91.8	74 - 131				
1,1,2,2-Tetrachloroethane	22.01	1.0	20	0	110	71 - 121				
1,1,2-Trichloroethane	22.04	1.0	20	0	110	80 - 119				
1,1-Dichloroethane	18.38	1.0	20	0	91.9	77 - 125				
1,1-Dichloroethene	16.87	1.0	20	0	84.4	71 - 131				
1,1-Dichloropropene	19.1	1.0	20	0	95.5	78 - 125				
1,2,3-Trichlorobenzene	20.17	1.0	20	0	101	69 - 129				
1,2,3-Trichloropropane	21.55	1.0	20	0	108	73 - 122				
1,2,4-Trichlorobenzene	20.7	1.0	20	0	103	69 - 130				
1,2,4-Trimethylbenzene	21.33	1.0	20	0	107	76 - 124				
1,2-Dibromo-3-chloropropane	22.55	1.0	20	0	113	62 - 128				
1,2-Dibromoethane	22.2	1.0	20	0	111	77 - 121				
1,2-Dichlorobenzene	21.17	1.0	20	0	106	80 - 119				
1,2-Dichloroethane	20.86	1.0	20	0	104	73 - 128				
1,2-Dichloropropane	20.3	1.0	20	0	102	78 - 122				
1,3,5-Trimethylbenzene	20.92	1.0	20	0	105	75 - 124				
1,3-Dichlorobenzene	21.41	1.0	20	0	107	80 - 119				
1,3-Dichloropropane	22.07	1.0	20	0	110	80 - 119				
1,4-Dichlorobenzene	20.83	1.0	20	0	104	79 - 118				
2,2-Dichloropropane	18.57	1.0	20	0	92.9	60 - 139				
2-Butanone	40.73	2.0	40	0	102	56 - 143				
2-Chlorotoluene	21.03	1.0	20	0	105	79 - 122				
2-Hexanone	44.22	2.0	40	0	111	57 - 139				
4-Chlorotoluene	20.94	1.0	20	0	105	78 - 122				
4-Isopropyltoluene	20.89	1.0	20	0	104	77 - 127				
4-Methyl-2-pentanone	44.67	2.0	40	0	112	67 - 130				
Acetone	37.88	2.0	40	0	94.7	39 - 160				
Benzene	20.21	1.0	20	0	101	79 - 120				
Bromobenzene	21.83	1.0	20	0	109	80 - 120				
Bromochloromethane	18.73	1.0	20	0	93.6	78 - 123				
Bromodichloromethane	20.96	1.0	20	0	105	79 - 125				
Bromoform	21.92	1.0	20	0	110	66 - 130				
Bromomethane	21.2	1.0	20	0	106	53 - 141				

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19030640-01MS	Units: UG/L			Analysis Date: 15-Mar-2019 15:16					
Client ID:	Run ID: VOA6_334643	SeqNo: 4993378	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	31.5	2.0	40	0	78.8	64 - 133				
Carbon tetrachloride	20.18	1.0	20	0	101	72 - 136				
Chlorobenzene	21.21	1.0	20	0	106	82 - 118				
Chloroethane	15.49	1.0	20	0	77.5	60 - 138				
Chloroform	19.18	1.0	20	0	95.9	79 - 124				
Chloromethane	13.1	1.0	20	0	65.5	50 - 139				
cis-1,2-Dichloroethene	19.38	1.0	20	0	96.9	78 - 123				
cis-1,3-Dichloropropene	20.85	1.0	20	0	104	75 - 124				
Dibromochloromethane	22.27	1.0	20	0	111	74 - 126				
Dibromomethane	20.77	1.0	20	0	104	79 - 123				
Dichlorodifluoromethane	6.432	1.0	20	0	32.2	32 - 152				
Ethylbenzene	21.28	1.0	20	0	106	79 - 121				
Hexachlorobutadiene	20.9	1.0	20	0	104	66 - 134				
Isopropylbenzene	21.35	1.0	20	0	107	72 - 131				
m,p-Xylene	43	2.0	40	0	107	80 - 121				
Methylene chloride	19.69	2.0	20	0	98.4	74 - 124				
Naphthalene	21.03	1.0	20	0	105	61 - 128				
n-Butylbenzene	20.68	1.0	20	0	103	75 - 128				
n-Propylbenzene	21.1	1.0	20	0	105	76 - 126				
o-Xylene	21.6	1.0	20	0	108	78 - 122				
sec-Butylbenzene	20.35	1.0	20	0	102	77 - 126				
Styrene	0.6311	1.0	20	0	3.16	78 - 123				JS
tert-Butylbenzene	20.72	1.0	20	0	104	78 - 124				
Tetrachloroethene	21.75	1.0	20	0.6002	106	74 - 129				
Toluene	21.49	1.0	20	0	107	80 - 121				
trans-1,2-Dichloroethene	18.66	1.0	20	0	93.3	75 - 124				
trans-1,3-Dichloropropene	21.08	1.0	20	0	105	73 - 127				
Trichloroethene	24.07	1.0	20	0	120	79 - 123				
Trichlorofluoromethane	16.28	1.0	20	0	81.4	65 - 141				
Vinyl chloride	11.55	1.0	20	0	57.8	58 - 137				S
Surr: 1,2-Dichloroethane-d4	44.6	1.0	50	0	89.2	81 - 118				
Surr: 4-Bromofluorobenzene	50.01	1.0	50	0	100	85 - 114				
Surr: Dibromofluoromethane	45.51	1.0	50	0	91.0	80 - 119				
Surr: Toluene-d8	52.76	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19030640-01MSD	Units: UG/L			Analysis Date: 15-Mar-2019 15:41					
Client ID:	Run ID: VOA6_334643	SeqNo: 4993379		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	21.49	1.0	20	0	107	78 - 124	21.87	1.74	20	
1,1,1-Trichloroethane	19.31	1.0	20	0	96.6	74 - 131	18.37	5.02	20	
1,1,2,2-Tetrachloroethane	20.34	1.0	20	0	102	71 - 121	22.01	7.9	20	
1,1,2-Trichloroethane	21.49	1.0	20	0	107	80 - 119	22.04	2.51	20	
1,1-Dichloroethane	19.13	1.0	20	0	95.6	77 - 125	18.38	3.96	20	
1,1-Dichloroethene	18.15	1.0	20	0	90.8	71 - 131	16.87	7.33	20	
1,1-Dichloropropene	20.27	1.0	20	0	101	78 - 125	19.1	5.94	20	
1,2,3-Trichlorobenzene	20.76	1.0	20	0	104	69 - 129	20.17	2.85	20	
1,2,3-Trichloropropane	20.04	1.0	20	0	100	73 - 122	21.55	7.23	20	
1,2,4-Trichlorobenzene	21.24	1.0	20	0	106	69 - 130	20.7	2.58	20	
1,2,4-Trimethylbenzene	21	1.0	20	0	105	76 - 124	21.33	1.52	20	
1,2-Dibromo-3-chloropropane	20.98	1.0	20	0	105	62 - 128	22.55	7.21	20	
1,2-Dibromoethane	21.63	1.0	20	0	108	77 - 121	22.2	2.61	20	
1,2-Dichlorobenzene	20.55	1.0	20	0	103	80 - 119	21.17	2.97	20	
1,2-Dichloroethane	20.98	1.0	20	0	105	73 - 128	20.86	0.57	20	
1,2-Dichloropropane	20.63	1.0	20	0	103	78 - 122	20.3	1.62	20	
1,3,5-Trimethylbenzene	21.04	1.0	20	0	105	75 - 124	20.92	0.553	20	
1,3-Dichlorobenzene	20.51	1.0	20	0	103	80 - 119	21.41	4.28	20	
1,3-Dichloropropane	21.28	1.0	20	0	106	80 - 119	22.07	3.66	20	
1,4-Dichlorobenzene	20.2	1.0	20	0	101	79 - 118	20.83	3.05	20	
2,2-Dichloropropane	19.45	1.0	20	0	97.3	60 - 139	18.57	4.62	20	
2-Butanone	39.77	2.0	40	0	99.4	56 - 143	40.73	2.39	20	
2-Chlorotoluene	20.53	1.0	20	0	103	79 - 122	21.03	2.42	20	
2-Hexanone	42.25	2.0	40	0	106	57 - 139	44.22	4.55	20	
4-Chlorotoluene	20.55	1.0	20	0	103	78 - 122	20.94	1.88	20	
4-Isopropyltoluene	21.3	1.0	20	0	107	77 - 127	20.89	1.96	20	
4-Methyl-2-pentanone	42.72	2.0	40	0	107	67 - 130	44.67	4.46	20	
Acetone	37.52	2.0	40	0	93.8	39 - 160	37.88	0.939	20	
Benzene	20.89	1.0	20	0	104	79 - 120	20.21	3.31	20	
Bromobenzene	20.64	1.0	20	0	103	80 - 120	21.83	5.58	20	
Bromochloromethane	19.32	1.0	20	0	96.6	78 - 123	18.73	3.08	20	
Bromodichloromethane	21.16	1.0	20	0	106	79 - 125	20.96	0.916	20	
Bromoform	21.22	1.0	20	0	106	66 - 130	21.92	3.28	20	
Bromomethane	20.44	1.0	20	0	102	53 - 141	21.2	3.64	20	

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19030640-01MSD	Units: UG/L			Analysis Date: 15-Mar-2019 15:41					
Client ID:	Run ID: VOA6_334643	SeqNo: 4993379	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit	Qual
Carbon disulfide	33.76	2.0	40	0	84.4	64 - 133	31.5	6.9	20	
Carbon tetrachloride	21.6	1.0	20	0	108	72 - 136	20.18	6.79	20	
Chlorobenzene	21.2	1.0	20	0	106	82 - 118	21.21	0.0564	20	
Chloroethane	16.4	1.0	20	0	82.0	60 - 138	15.49	5.68	20	
Chloroform	19.69	1.0	20	0	98.4	79 - 124	19.18	2.6	20	
Chloromethane	13.63	1.0	20	0	68.1	50 - 139	13.1	4	20	
cis-1,2-Dichloroethene	19.94	1.0	20	0	99.7	78 - 123	19.38	2.81	20	
cis-1,3-Dichloropropene	21.1	1.0	20	0	105	75 - 124	20.85	1.19	20	
Dibromochloromethane	21.77	1.0	20	0	109	74 - 126	22.27	2.23	20	
Dibromomethane	21.14	1.0	20	0	106	79 - 123	20.77	1.8	20	
Dichlorodifluoromethane	7.059	1.0	20	0	35.3	32 - 152	6.432	9.3	20	
Ethylbenzene	21.68	1.0	20	0	108	79 - 121	21.28	1.84	20	
Hexachlorobutadiene	22.92	1.0	20	0	115	66 - 134	20.9	9.22	20	
Isopropylbenzene	22.06	1.0	20	0	110	72 - 131	21.35	3.29	20	
m,p-Xylene	43.42	2.0	40	0	109	80 - 121	43	0.986	20	
Methylene chloride	19.76	2.0	20	0	98.8	74 - 124	19.69	0.346	20	
Naphthalene	20.95	1.0	20	0	105	61 - 128	21.03	0.392	20	
n-Butylbenzene	21.83	1.0	20	0	109	75 - 128	20.68	5.42	20	
n-Propylbenzene	21.19	1.0	20	0	106	76 - 126	21.1	0.431	20	
o-Xylene	21.72	1.0	20	0	109	78 - 122	21.6	0.556	20	
sec-Butylbenzene	21.17	1.0	20	0	106	77 - 126	20.35	3.96	20	
Styrene	0.6366	1.0	20	0	3.18	78 - 123	0.6311	0	20	JS
tert-Butylbenzene	21.05	1.0	20	0	105	78 - 124	20.72	1.59	20	
Tetrachloroethene	22.29	1.0	20	0.6002	108	74 - 129	21.75	2.43	20	
Toluene	21.39	1.0	20	0	107	80 - 121	21.49	0.458	20	
trans-1,2-Dichloroethene	19.33	1.0	20	0	96.6	75 - 124	18.66	3.49	20	
trans-1,3-Dichloropropene	21.38	1.0	20	0	107	73 - 127	21.08	1.42	20	
Trichloroethene	22.25	1.0	20	0	111	79 - 123	24.07	7.84	20	
Trichlorofluoromethane	17.9	1.0	20	0	89.5	65 - 141	16.28	9.48	20	
Vinyl chloride	12.21	1.0	20	0	61.0	58 - 137	11.55	5.5	20	
Surr: 1,2-Dichloroethane-d4	44.83	1.0	50	0	89.7	81 - 118	44.6	0.518	20	
Surr: 4-Bromofluorobenzene	51.19	1.0	50	0	102	85 - 114	50.01	2.33	20	
Surr: Dibromofluoromethane	45.7	1.0	50	0	91.4	80 - 119	45.51	0.408	20	
Surr: Toluene-d8	50.92	1.0	50	0	102	89 - 112	52.76	3.54	20	

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334643 (0) **Instrument:** VOA6 **Method:** VOLATILES ORGANICS BY METHOD 8260C

The following samples were analyzed in this batch:

HS19030761-01	HS19030761-03
---------------	---------------

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
WorkOrder: HS19030761

QC BATCH REPORT NEW

Batch ID: R334752 (0)		Instrument: UV-2450		Method: HEXAVALENT CHROMIUM BY SW7196A						
MBLK	Sample ID: MBLK-334752	Units: mg/L		Analysis Date: 15-Mar-2019 12:14						
Client ID:	Run ID: UV-2450_334752	SeqNo: 4995036		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chromium, Hexavalent	0.0100	0.0100							U	
LCS	Sample ID: LCS-334752	Units: mg/L		Analysis Date: 15-Mar-2019 12:14						
Client ID:	Run ID: UV-2450_334752	SeqNo: 4995037		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chromium, Hexavalent	0.244	0.0100	0.25	0	97.6	80 - 120				
LCSD	Sample ID: LCSD-334752	Units: mg/L		Analysis Date: 15-Mar-2019 12:14						
Client ID:	Run ID: UV-2450_334752	SeqNo: 4995038		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chromium, Hexavalent	0.244	0.0100	0.25	0	97.6	80 - 120	0.244	0	20	

The following samples were analyzed in this batch: HS19030761-01

ALS Houston, US

Date: 31-Mar-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples	
WorkOrder:	HS19030761	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Monthly Effluent Samples
Work Order: HS19030761

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19030761-01	LH18/24-SP650_031419	Login	3/15/2019 10:54:17 AM	JRM	EXT075
HS19030761-01	LH18/24-SP650_031419	Login	3/15/2019 10:54:17 AM	JRM	WET176
HS19030761-01	LH18/24-SP650_031419	Login	3/15/2019 10:54:17 AM	JRM	MET068
HS19030761-01	LH18/24-SP650_031419	Login	3/15/2019 10:54:17 AM	JRM	VOA052
HS19030761-02	LH18/24-SP650_031419_BIX	Login	3/15/2019 10:54:17 AM	JRM	Sub
HS19030761-03	Trip Blank	Login	3/15/2019 10:54:17 AM	JRM	VOA052

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030761

Date/Time Received: **15-Mar-2019 08:38**
 Received by: **JRM**

Checklist completed by: Jared R. Makan 15-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 15-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.3c/1.3c UC/C IR25
 Cooler(s)/Kit(s): 25780
 Date/Time sample(s) sent to storage: 03/15/2019 10:58

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd., Suite 210 Houston, TX 77099 (281) 530 - 5656 ATTN: RJ Modashia

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001			Analyses										Remarks (Preservatives, etc.)	Lab I.D.#		
Job: GROUNDWATER TREATMENT PLANT MONTHLY EFFLUENT SAMPLES			MS / MSD No. OF CONTAINERS	VOLATILES SILVER, SELENIUM, LEAD, BARIUM HEXAVALENT CHROMIUM 1, 4 - DIOXANE PERCHLORATE															
Prepared By: Scott Beesinger		P.O. Number																	
Field Sample I.D.	Sample Matrix	Date / Time																	
LH18/24-SP650_031419	Water	03/14/19 / 14:00	3	X															HCL
LH18/24-SP650_031419	Water	03/14/19 / 14:00	2				X	X											NONE
LH18/24-SP650_031419_BIX	Water	03/14/19 / 14:00	1								X								NONE
LH18/24-SP650_031419	Water	03/14/19 / 14:00	1		X														HNO3
Trip Blank	Water	03/14/19	2	X															HCL

Additional Remarks: STANDARD TURN AROUND TIME


Relinquished By: <i>Scott Beesinger</i>	Date 03/14/19	Time 14:30	Received By:	Date	Time	Relinquished By: <i>J. Modashia</i>	Date 3/15/19	Time 3:03	Received By:	Date	Time
---	-------------------------	----------------------	---------------------	-------------	-------------	---	------------------------	---------------------	---------------------	-------------	-------------

For Lab Use Only									
Received At Lab By:	Date	Time	Airbill No.	Opened By:	Date	Time	Temp of Container	Seal No.	Condition
Remarks: <div style="display: flex; justify-content: space-between;"> HS19030761 <i>Coder 25780</i> <i>1425</i> </div> <div style="display: flex; justify-content: space-between;"> <i>Temp 1.3</i> <i>CF0.0</i> </div>									

Bhate Environmental Associates, Inc.
 18/24 Longhorn GW Treatment Plant Monthly Effluer



(Word) S:\-ces\Forms\Chain of Custod

	ALS 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 6858 Fax. +1 281 530 5887	CU: Date: 3/14/19 Name: Scott Company: B.H.E.
---	--	---

STUDY SEAL		Seal Broken By:
Type: P30		SM
BAIRDINGER		Date: 3/15/19

 TRK# 0221 4380 9530 9490	FRI - 15 MAR 10:30A PRIORITY OVERNIGHT
AB SGRA	77099 TX-US IAH
	
<small>F10 9586291 14MAR19 GGA 553C1/4603/PCBA</small>	



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1907867; 1907869; 1907871;
1908794

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2230 (235529)

General Set Information: There were four field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ¹⁸O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50μL of an ¹⁸O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Method QC data: The method blank (LMB 645533) was less than 1/2 the CRDL. The recovery for the LCS (645534) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.µg/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in µg/L. Results were calculated in µg/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (µg/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level of 4.0µg/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch March 29, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 29, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1907869**

Project ID: HS19030761

Purchase Order: HS19030761

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP650_031419_BIX	1907869001	03/14/19	03/19/19	



ANALYTICAL REPORT

Workorder: 34-1907869

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP650_031419_BIX	Sampling Site: NA	Collected: 03/14/2019				
Lab ID: 1907869001	Media: 125 mL Nalgene	Received: 03/19/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2230 (HBN: 235529) Analyzed: 03/28/2019 09:41	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 235529)

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/28/2019 14:08	/S/ Stephen Brose 03/29/2019 12:58

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1907869

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935320

Analysis Information

Workorder: 1907869

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2230 (HBN: 235529)
Analyzed By: Thomas Bosch

Blank

LMB: 645533 Analyzed: 03/28/2019 09:14 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 645534 Analyzed: 03/28/2019 08:47 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.00	4.00	99.9	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1907869001 Analyzed: 03/28/2019 09:41 Dilution: 1 Units: ug/L			MS: 645535 Analyzed: 03/28/2019 09:54 Dilution: 1 Units: ug/L			MSD: 645536 Analyzed: 03/28/2019 10:08 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	ND	4.66	4	117	78.8 123.8	4.05	101	14	0.0 20.0

Comments

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/28/2019 14:23	/S/ Stephen Brose 03/29/2019 12:58

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



1907869



18698/#2

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10917

1907869

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030761
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030761-02	LH18/24-SP650_031419_BIX	Water	14 Mar 2019 14:00
	SUB_Perch-6850		25 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. Whalen Date/Time: 3/12/19 18:00
 Received By: Dawn Jussell Date/Time: 03/19/19 9:50
 Cooler ID(s): _____ Temperature(s): _____

RIGHT SOLUTIONS | RIGHT PARTNER



ALS Environmental
CHAIN-OF-CUSTODY

Project / Job / Task: HS19030761		Split:	Workorder ID: 1907869	Level: ENV_LVL4	Requested Analysis	
Client: ALS Environmental (Houston)		Account: 8101		Type: 125Poly		
Comments:						
Item	Collect Date/Time	Sample ID	Lab ID	QC	Matrix	Containers
1	03/14/2019 14:00	LH18/24-SP650_031419_BIX	1907869001		Water	1 A
2						
3						
4						
5						
6						
7						
8						
9						
10						

EP-A 6850, DGD GSM

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY				SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY			
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location	Sample Prep / Analysis for:	Lab Notebook No.:	Prepared / Analyzed by:	Date / Time:
<i>Julie Wraith</i>	03/19/2019 09:50	ALS Sample Receiving	Sample Login				
<i>R.33.1</i>	<i>3/27/19/11:45</i>	<i>T. Bond</i>	<i>Storage</i>				

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: ALS Houston Project/Task/Site: 1907819
 Date/Time of Receipt: 03-19-19 9:50 Number of Coolers Received: 1

Condition of Coolers: Acceptable/Unacceptable Temperature Control: Present/Not Included
 Cooler Custody Seals: Present/Absent/NA Location Temp Taken: Control/Between Samples
 Container Custody Seals: Present/Absent/NA Are all temperatures within project specific guidelines? Yes/No/NA
 Ice Present: Frozen/Melted/NA VOA Headspace Present? Yes/No/NA

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 9750	1 °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: Tamir Vassell Signature Tamir Vassell Printed Name 03-19-19 Date

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Part # 159469-434 RIT2 EXP-11/19 **

ORIGIN ID: SGRA (281) 530-5656
SHIPPING DEPT
ALS LABORATORY GROUP
10450 STANCLIFF RD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

SHIP DATE: 18MAR19
ACTWGT: 9.95 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

3401/EGW/TJ155

TO: **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(01) 266-7700
REF: HS19030749/761/763 RJ



FedEx
Express



AN105090811181F

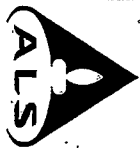
TUE - 19 MAR 3:00P
STANDARD OVERNIGHT

TRK# 4809 7831 7725
0201

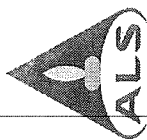
AX BTFA

84123
UT-US SLC




ALS
 10450 Stancliff Rd., Suite 210
 Houston, Texas 77099
 Tel. +1 281 530 5656
 Fax +1 281 530 5887

Date _____
 Name _____
 Company _____



Batch Worklist

Batch: ELMS/ 2230 **Created:** 3/28/2019 07:45 **Instrument:** HBN: 235529
Rule: EPA 6850, DoD QSM Water **Analyst:** T. Bosch **Status:** WP



Workorder: 1907867 [ENV_LVL4]
Workorder: 1907869 [ENV_LVL4]
Workorder: 1907871 [ENV_LVL4]
Workorder: 1908794 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	64530	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	
2	64531	RLV'S for HBN 235529 [ELMS/2230]				RLV'S	3		E685041C3Q	5311		4/1/2019	
3	64532	ICS for HBN 235529 [ELMS/2230]				ICS	3		E6850.D3Q	5311		4/1/2019	
4	64533	LMB for HBN 235529 [ELMS/2230]				LMB	3		E6850Q413Q	5311		4/1/2019	
5	64534	LCS for HBN 235529 [ELMS/2230]				LCS	3		E6850Q413Q	5311		4/1/2019	
6	1907867001	LH18/24-SP140_031419				SAMPLE	3	1907867001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
7	1907869001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907869001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
8	64535	LH18/24-SP650...(1907869001MS)				MS	3		E6850Q413Q	5311		4/1/2019	
9	64536	LH18/24-SP65...(1907869001MSD)				MSD	3		E6850Q413Q	5311		4/1/2019	
10	1907871001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907871001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
11	1908794001	LH18/24-SP650_032119_BIX				SAMPLE	3	1908794001-A	E6850Q41.3	5480	4/18/2019	4/8/2019	
12	64537	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #'s: 1907867 (001); 1907869 (001); 1907871 (001); 1908794 (001)

ELMS Batch/HBN ID: 2230 (235529)

Prep Date: 03/27/2019 Analysis Date: 03/28/2019 Analyst: T. Bosch

Analyte: **Perchlorate** Matrix: **Water** Method: **6850**

Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\28MAR19D.s

Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by **TNB**. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 25µL
Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 645534; Target = 4.0µg/L. ASTM type II water was used for LMB 645533.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\235529-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
<u>Batch(es)/SDG: ELMS: 2230 HBN: 235529</u>		
<u>Sample Set IDs if Applicable: 1907867/1907869/1907871/1908794</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850 WKG Std 100 ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK			Description - 6850 Stock AccStd 1,000ug/mL
Standard: 43659		Created By: Thomas Bosch	
MFG: AccuStandard		Create Date: 09/17/2018 09:09AM	
MFG Lot: 218065075		Amount: 100 mL	
Part ID: IC-PER-10X-1		Expires: 07/25/2020	
		Usable: No	
		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT			Description - 6850 Intermdt AccStd 10.ug/mL		
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK			Description: 6850 QC WKG STD 100ug/L		
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK			Description - 6850 QC Stock STD 1,000ug/mL
Standard: 36748		Created By: Thomas Bosch	
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	
MFG Lot: CP-0860		Amount: 100 mL	
Part ID: ICC-013		Expires: 03/31/2020	
		Usable: Yes	
		Lab Lot: CLO4 QC STOCK	
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730		Created By: Thomas Bosch		Amount: 25 mL	
MFG: ALS/SLC		Create Date: 09/20/2018 09:09AM		Expires: 09/20/2019	
MFG Lot: TNB: 05/09/2018		Verified By: Thomas Bosch		Usable: Yes	
Pipette ID: Not Provided		Verify Date:		Lab Lot: CLO4ISTDWRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

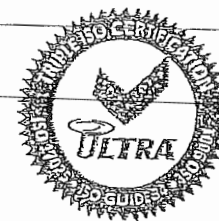
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729		Created By: Thomas Bosch	Amount: 1 mL
MFG: Cambridge Isotope		Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026
MFG Lot: SDFF-012A		Verified By: Thomas Bosch	Usable: Yes
Part ID: OLM-7310-S		Verify Date:	Lab Lot: CLO4ISTDSTK
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:

This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCCL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

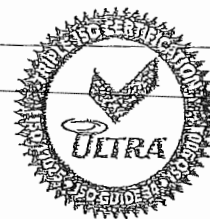
Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis



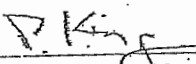
ISO Guide 34 Reference Material

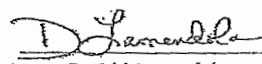
Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:

The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QAVRA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager



Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:
ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: NaCl^+O_4

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration data.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 $\mu\text{g/mL}$
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 \pm 2.8 $\mu\text{g/mL}$ (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
*	645530	CCV@25	Vial 71	1	Control	1	1.81499e6	24.13647
*	645534	QC@4.0	Vial 72	1	Control	2	2.89937e5	3.99582
*	645532	ICS@4.0	Vial 73	1	Control	3	2.38655e5	3.65624
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	7.55115e5	8344.77963
*	1907869001		Vial 76	1	Sample	6	0.00000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	2.77046e5	4.66163
*	645536	78691SD	Vial 78	1	Sample	8	2.89384e5	4.05130
*	1907871001		Vial 79	1	Sample	9	0.00000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	1.79709e6	24.49119

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
*	645530	CCV@25	Vial 71	1	Control	1	5.33395e5	23.90219
*	645534	QC@4.0	Vial 72	1	Control	2	9.52963e4	4.26699
*	645532	ICS@4.0	Vial 73	1	Control	3	7.88339e4	3.90865
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.38827e5	8762.35701
*	1907869001		Vial 76	1	Sample	6	0.00000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	8.93399e4	4.91371
*	645536	78691SD	Vial 78	1	Sample	8	9.85690e4	4.48457
*	1907871001		Vial 79	1	Sample	9	0.00000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	5.35802e5	24.59111

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
*	645530	CCV@25	Vial 71	1	Control	1	2.29255e5	5.00000
*	645534	QC@4.0	Vial 72	1	Control	2	2.39106e5	5.00000
*	645532	ICS@4.0	Vial 73	1	Control	3	2.16080e5	5.00000
*	645533	LMB	Vial 74	1	Control	4	2.42742e5	5.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.89136e5	5000.00000
*	1907869001		Vial 76	1	Sample	6	1.89925e5	5.00000
*	645535	78691MS	Vial 77	1	Sample	7	1.94413e5	5.00000
*	645536	78691SD	Vial 78	1	Sample	8	2.35220e5	5.00000
*	1907871001		Vial 79	1	Sample	9	1.89609e5	5.00000
*	1908794001		Vial 80	1	Sample	10	1.92453e5	5.00000
*	645537	CCV@25	Vial 71	1	Control	11	2.23514e5	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	645530	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	645534	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	645532	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	645533	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1907867001	1K	CLO4-AQN	1	Sample	
6	Vial 76	1907869001		CLO4-AQN	1	Sample	
7	Vial 77	645535	78691MS	CLO4-AQN	1	Sample	
8	Vial 78	645536	78691SD	CLO4-AQN	1	Sample	
9	Vial 79	1907871001		CLO4-AQN	1	Sample	
10	Vial 80	1908794001		CLO4-AQN	1	Sample	
11	Vial 71	645537	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D

Sample Name: 645530 CCV@25

Injection Date: 3/28/2019 08:30:53

Seq Line: 1

Sample Name: 645530 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

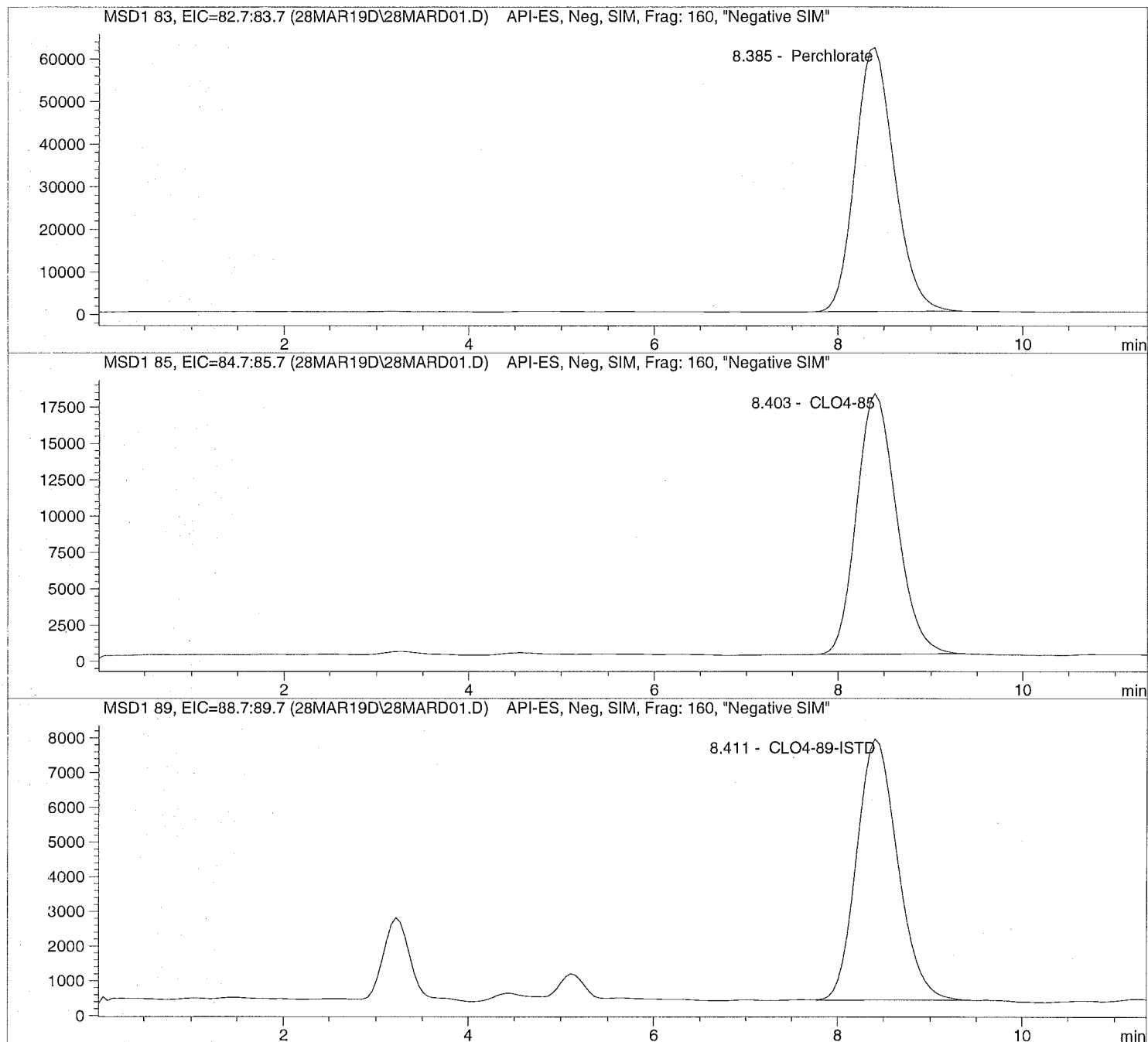
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D Sample Name: 645530 CCV@25

```

=====
Injection Date: 3/28/2019 08:30:53      Seq Line: 1
Sample Name: 645530 CCV@25             Location: Vial 71
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.385	PBA	1814992.7	24.1365	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.403	PBA	533395.5	23.9022	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.411	BBA	229255.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

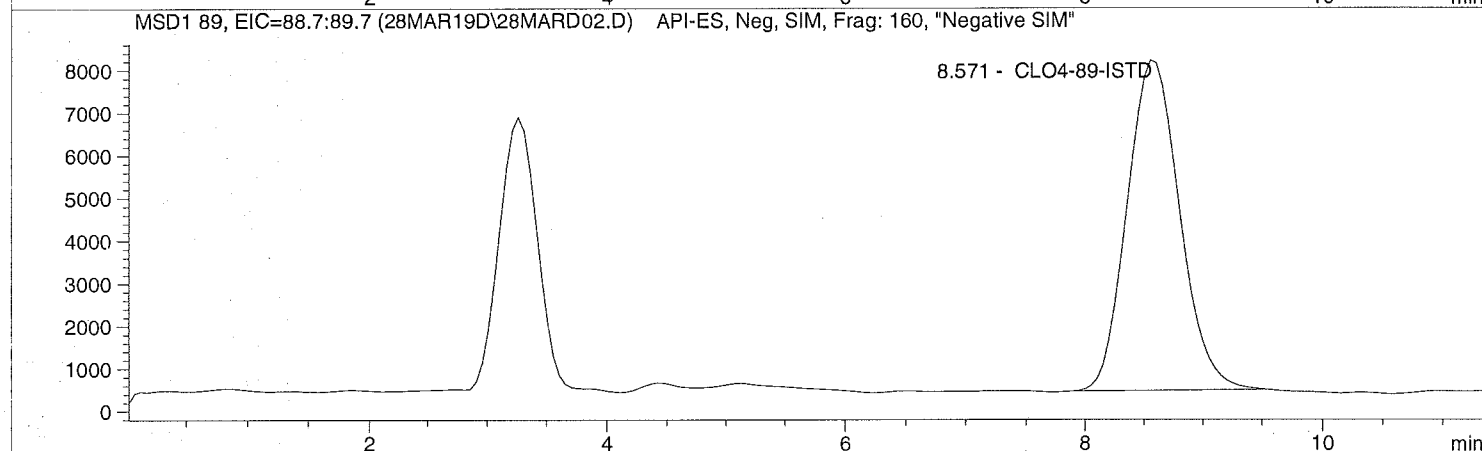
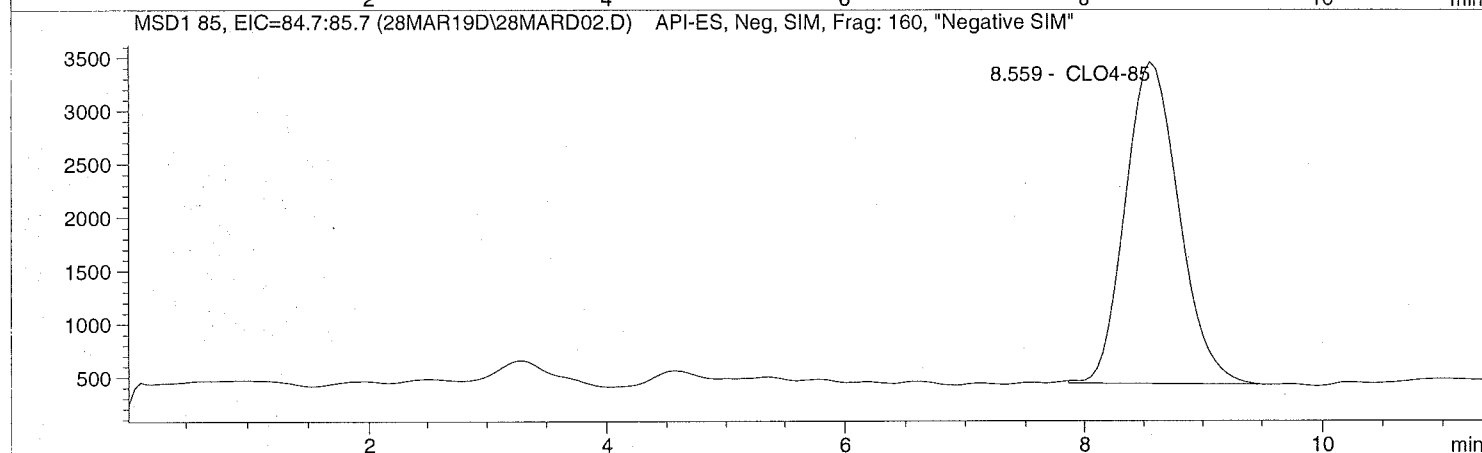
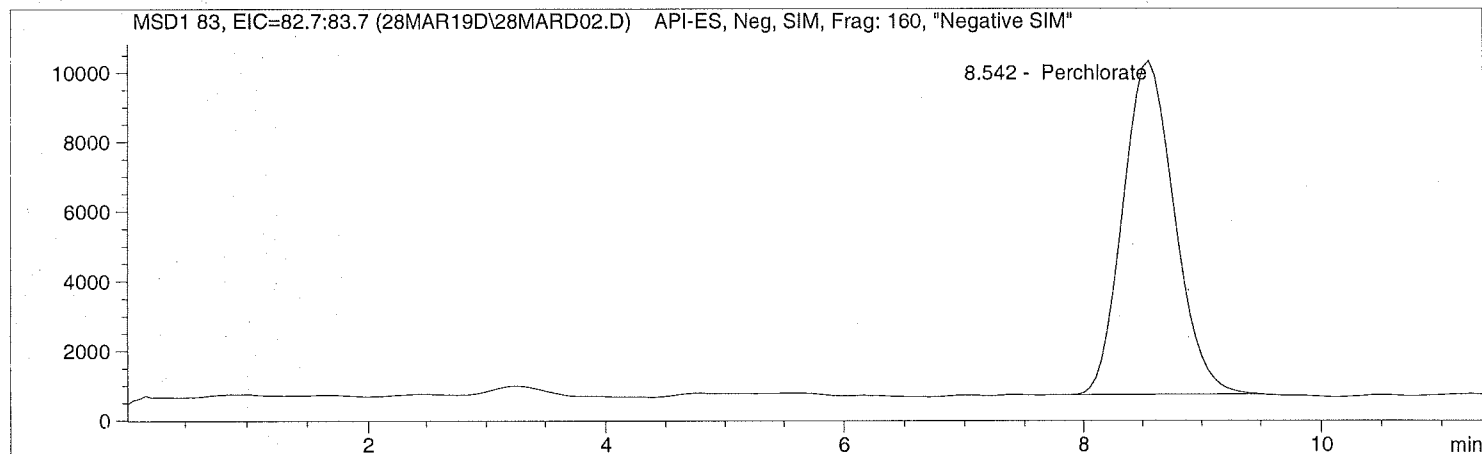
```


Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D Sample Name: 645534 QC@4.0

```
=====
Injection Date: 3/28/2019 08:47:42      Seq Line: 2
Sample Name: 645534 QC@4.0              Location: Vial 72
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D Sample Name: 645534 QC@4.0

=====
Injection Date: 3/28/2019 08:47:42 Seq Line: 2
Sample Name: 645534 QC@4.0 Location: Vial 72
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

=====
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis

=====
Sample Information
=====

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.542	PBA	289937.1	3.9958	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.559	BBA	95296.3	4.2670	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.571	PBA	239105.9	5.0000	CLO4-89-ISTD

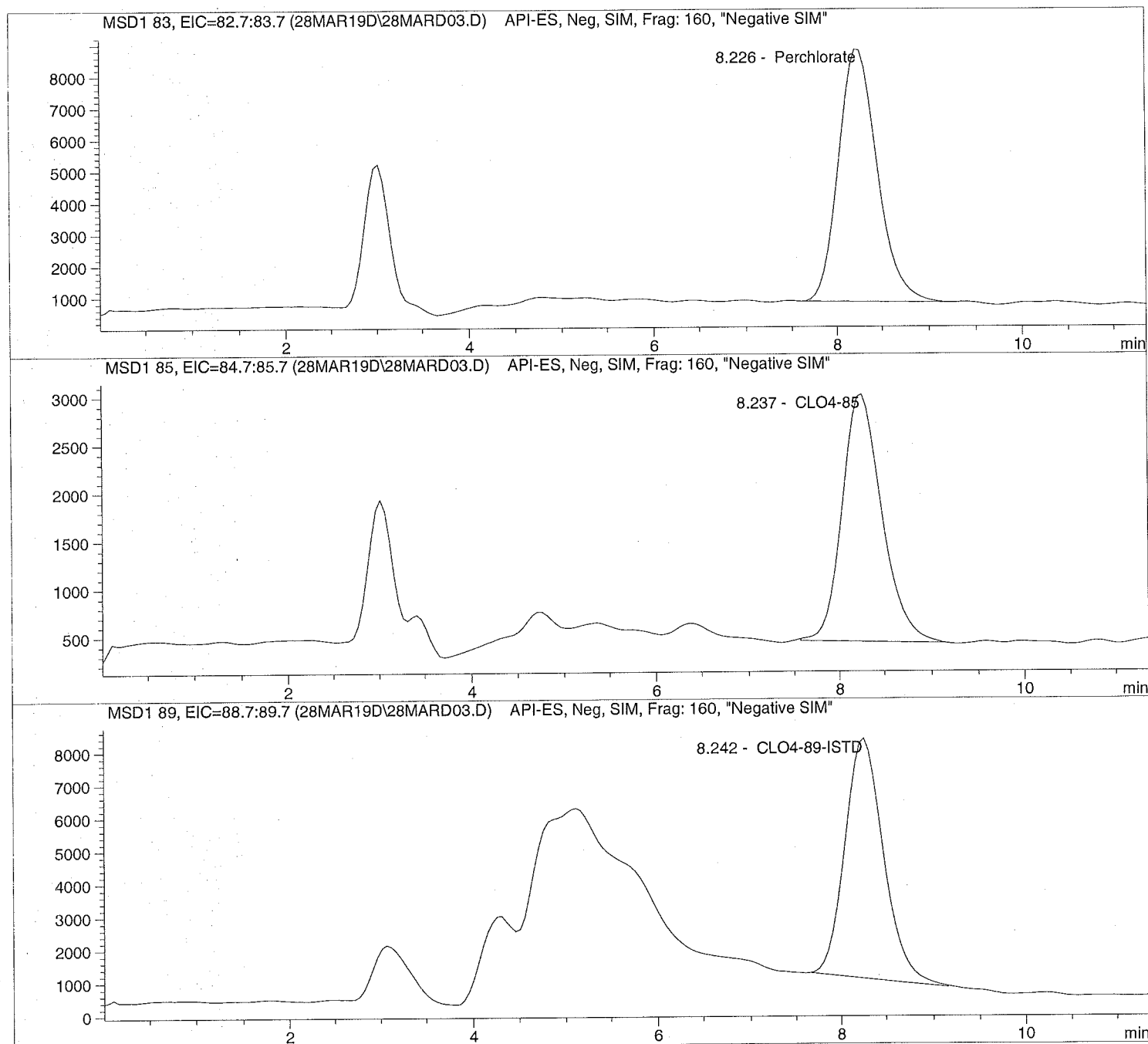
=====
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD03.D Sample Name: 645532 ICS@4.0

```
=====
Injection Date: 3/28/2019 09:00:54      Seq Line: 3
Sample Name:    645532 ICS@4.0          Location:  Vial 73
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD03.D Sample Name: 645532 ICS@4.0

```

=====
Injection Date:  3/28/2019  09:00:54      Seq Line:      3
Sample Name:    645532   ICS@4.0          Location:      Vial 73
Acq Operator:   TNB                        Inj. No.:     1
                                             Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:     1.000000
Sample Amount: 4.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.226	BBA	238654.8	3.6562	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.237	BBA	78833.9	3.9087	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.242	PBA	216079.7	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D

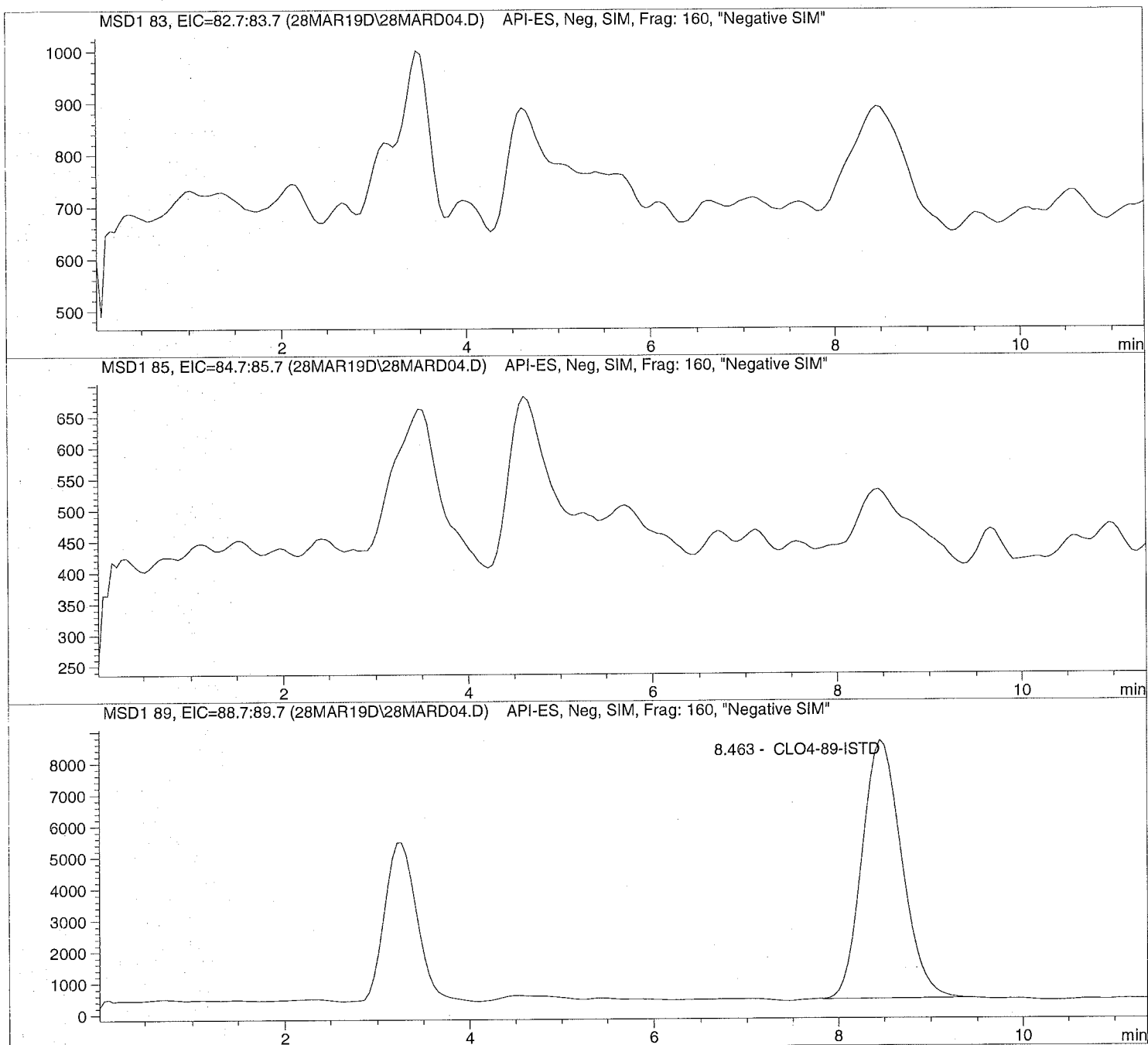
Sample Name: 645533 LMB

Injection Date: 3/28/2019 09:14:08
Sample Name: 645533 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D Sample Name: 645533 LMB

```

=====
Injection Date: 3/28/2019 09:14:08      Seq Line: 4
Sample Name: 645533 LMB                  Location: Vial 74
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.463	BBA	242742.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

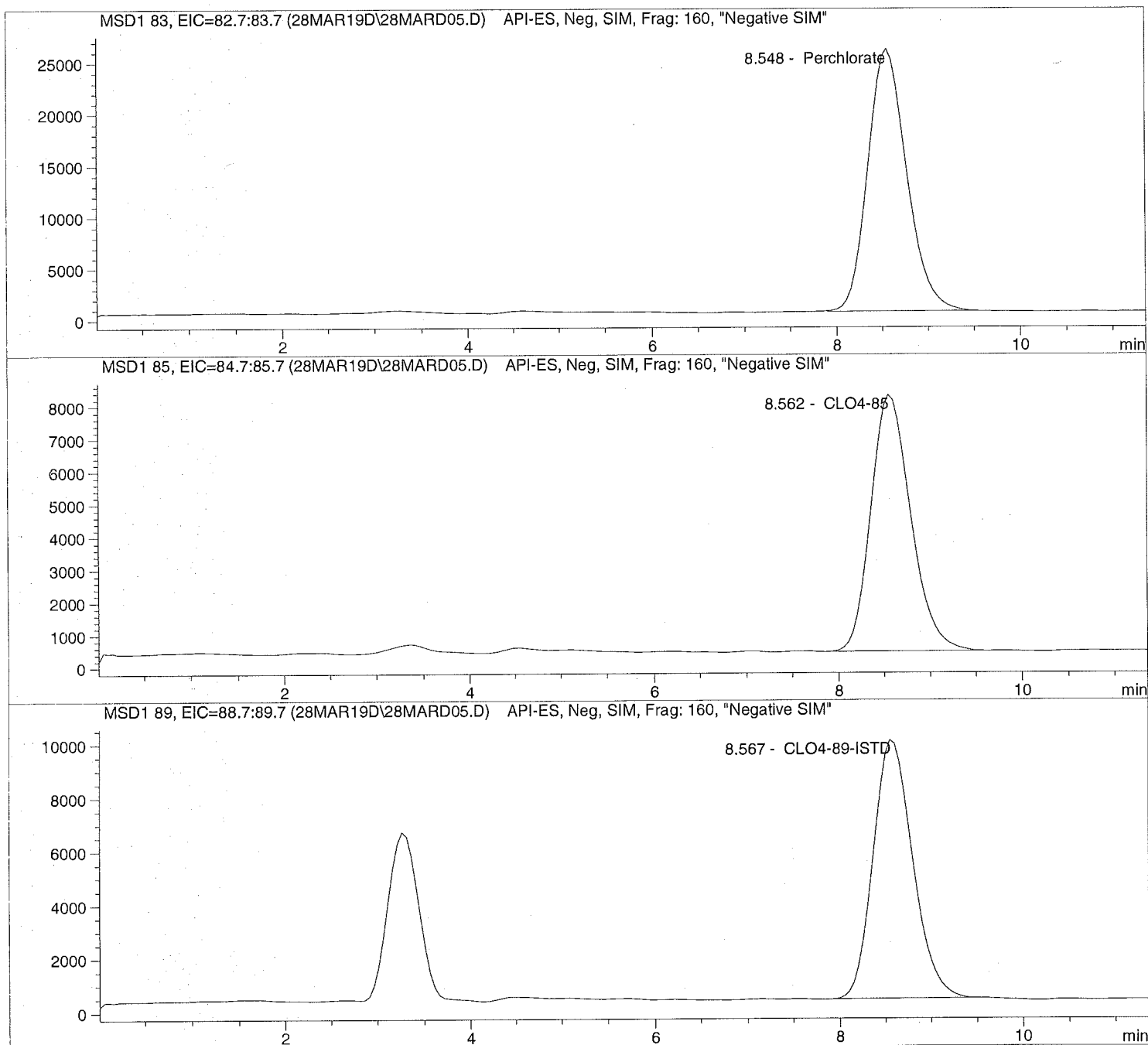
```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```
=====
Injection Date: 3/28/2019 09:28:25      Seq Line:      5
Sample Name:    1907867001 1K           Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```

=====
Injection Date: 3/28/2019 09:28:25      Seq Line:      5
Sample Name:    1907867001 1K           Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1000.000000
Sample Amount:  0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.548	BBA	755115.1	8344.7796	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.562	PBA	238826.8	8762.3570	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.567	PBA	289135.6	5000.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D

Sample Name: 1907869001

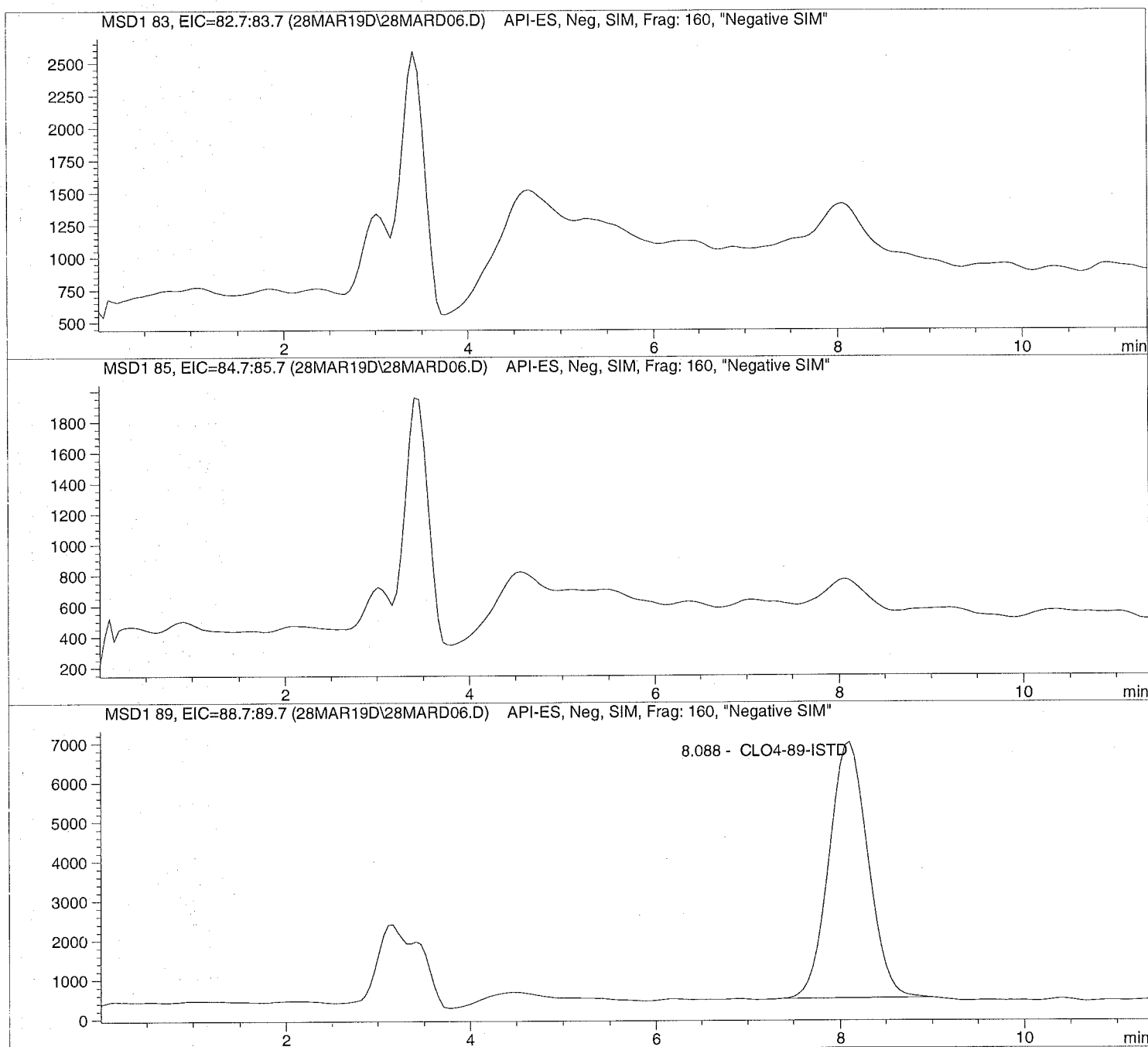
=====
Injection Date: 3/28/2019 09:41:37
Sample Name: 1907869001
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D Sample Name: 1907869001

```

=====
Injection Date: 3/28/2019 09:41:37      Seq Line: 6
Sample Name: 1907869001                Location: Vial 76
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.088	BBA	189925.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D

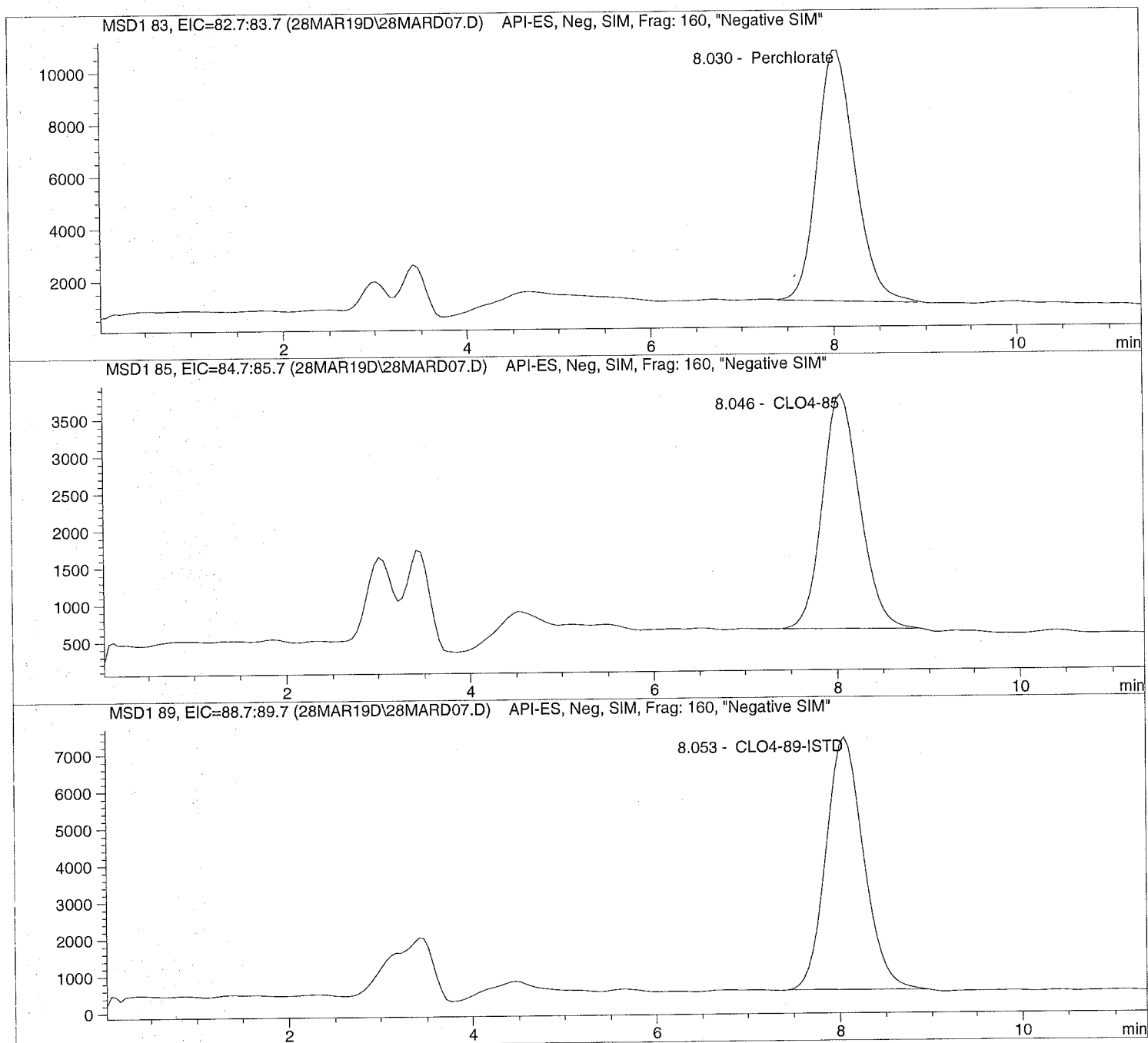
Sample Name: 645535 78691MS

=====
Injection Date: 3/28/2019 09:54:47
Sample Name: 645535 78691MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 25 µl

=====
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D Sample Name: 645535 78691MS

```

=====
Injection Date: 3/28/2019 09:54:47      Seq Line: 7
Sample Name:    645535 78691MS          Location:  Vial 77
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.030	BBA	277046.3	4.6616	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.046	BBA	89339.9	4.9137	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	194412.9	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D

Sample Name: 645536 78691SD

Injection Date: 3/28/2019 10:08:05
Sample Name: 645536 78691SD
Acq Operator: TNB

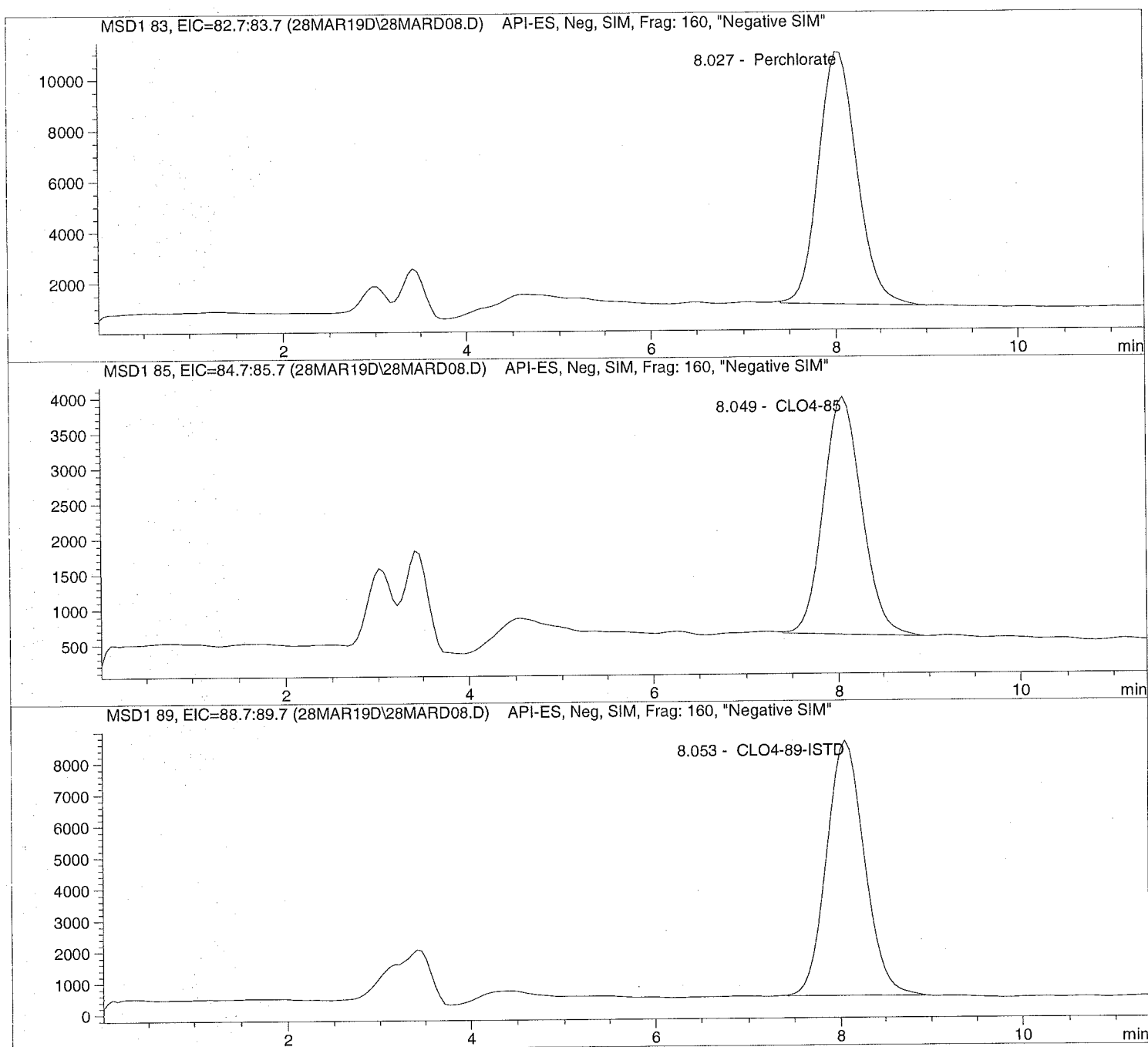
Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D Sample Name: 645536 78691SD

```

=====
Injection Date: 3/28/2019 10:08:05 Seq Line: 8
Sample Name: 645536 78691SD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.027	BBA	289383.7	4.0513	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.049	BBA	98569.0	4.4846	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	235219.8	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D

Sample Name: 1907871001

=====
Injection Date: 3/28/2019 10:21:14
Sample Name: 1907871001
Acq Operator: TNB

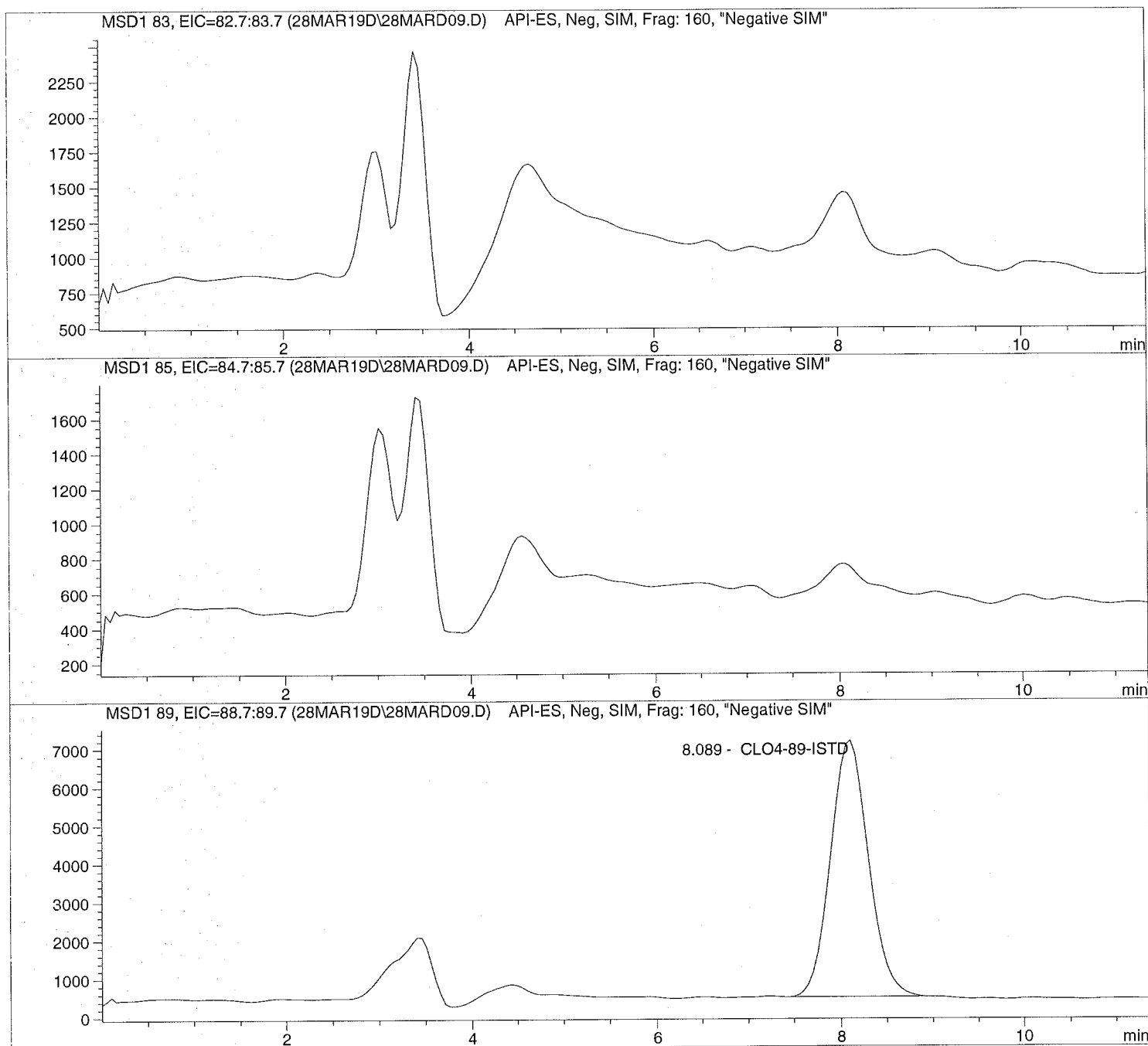
Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D

Sample Name: 1907871001

```

=====
Injection Date: 3/28/2019 10:21:14      Seq Line:          9
Sample Name:    1907871001              Location:         Vial 79
Acq Operator:   TNB                      Inj. No.:        1
                                           Inj. Vol.:       25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.089	PBA	189608.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD10.D

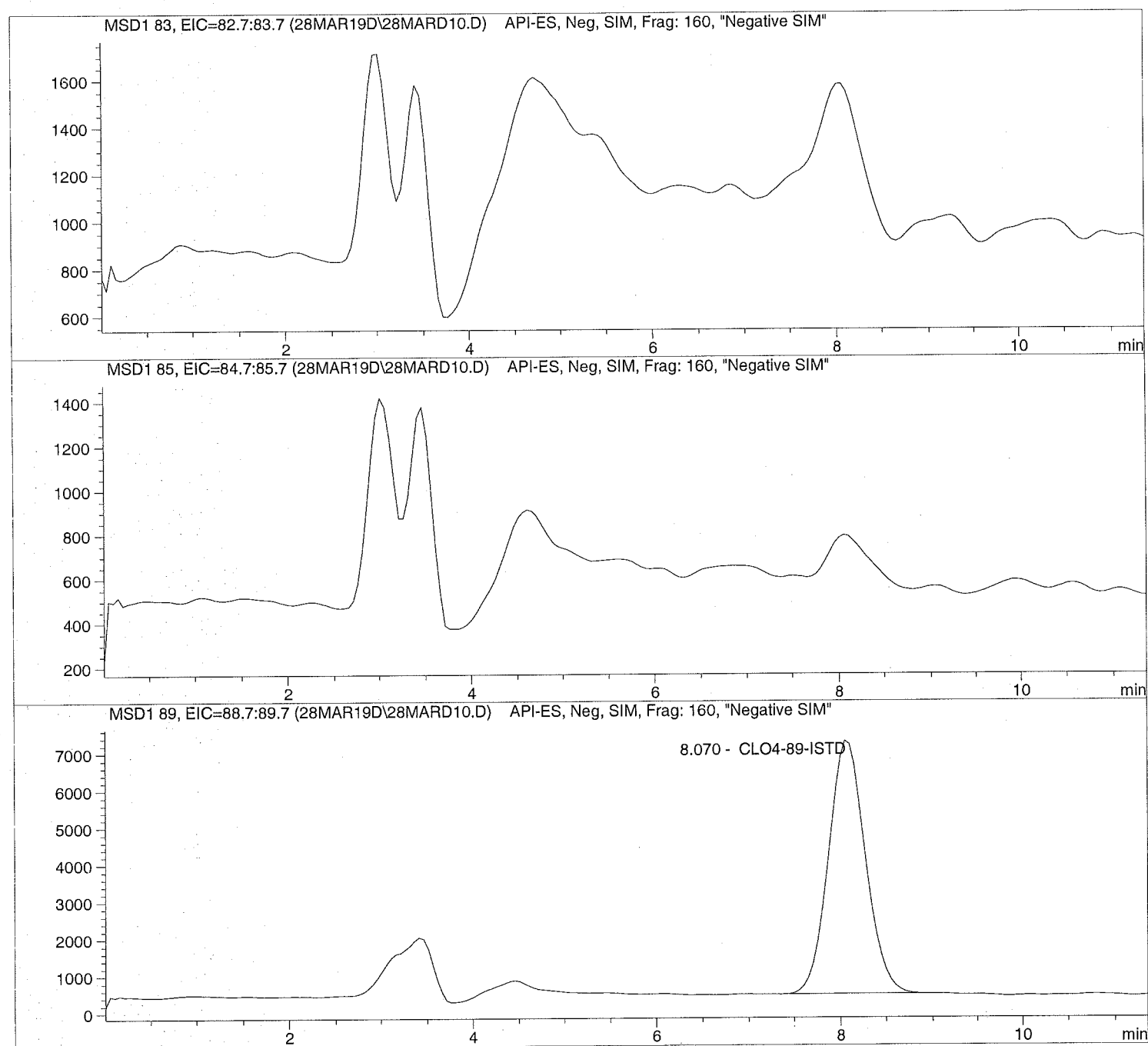
Sample Name: 1908794001

=====
Injection Date: 3/28/2019 10:34:27
Sample Name: 1908794001
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 25 µl

=====
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19\28MARD10.D Sample Name: 1908794001

```

=====
Injection Date: 3/28/2019 10:34:27      Seq Line: 10
Sample Name: 1908794001                 Location: Vial 80
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.070	PBA	192453.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D

Sample Name: 645537 CCV@25

Injection Date: 3/28/2019 10:49:41

Seq Line: 11

Sample Name: 645537 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

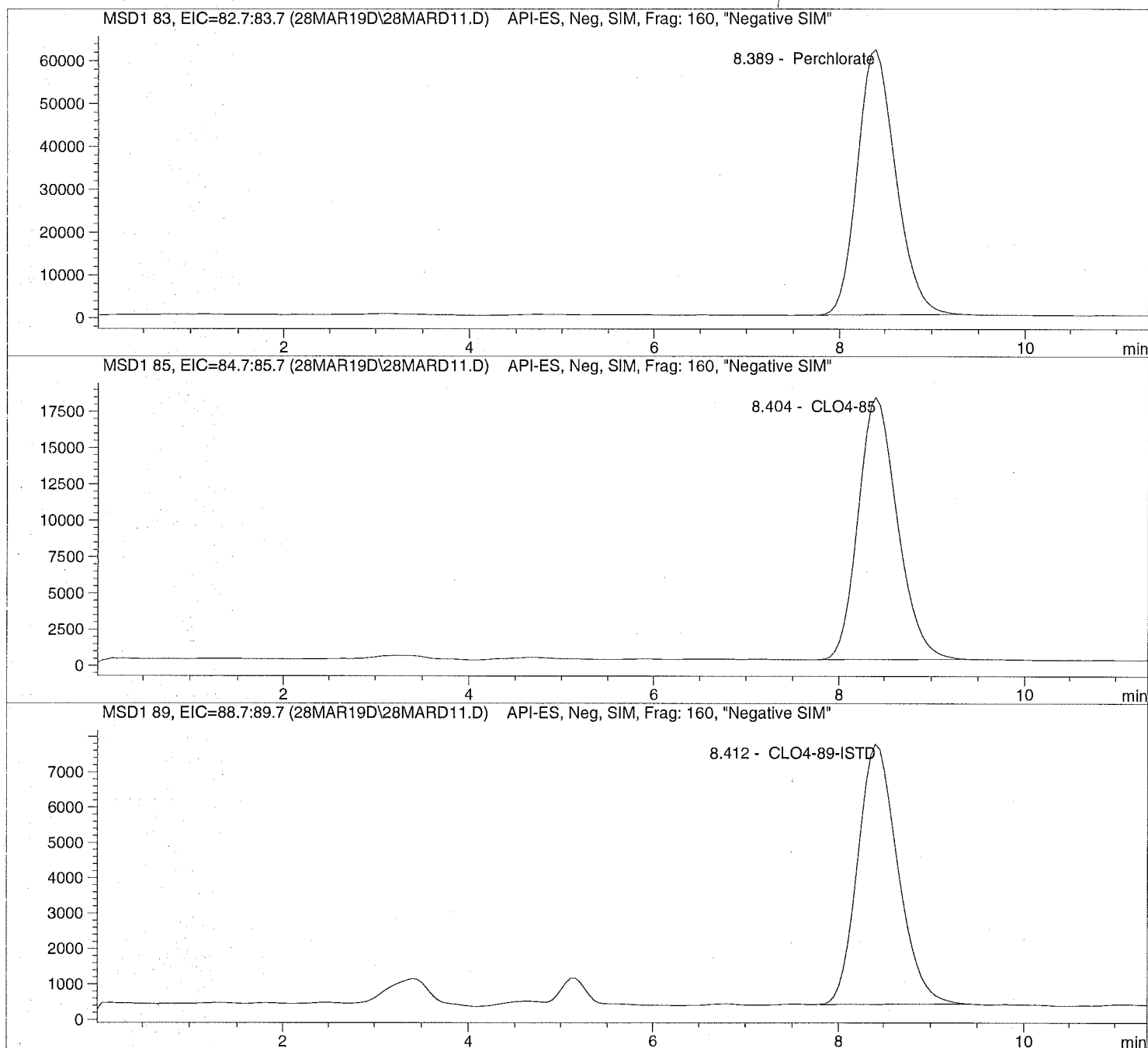
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D Sample Name: 645537 CCV@25

```

=====
Injection Date: 3/28/2019 10:49:41      Seq Line:      11
Sample Name:    645537  CCV@25          Location:      Vial 71
Acq Operator:   TNB                    Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.389	PBA	1797085.1	24.4912	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.404	PBA	535801.9	24.5911	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.412	BBA	223514.2	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
 Calibration Table
 =====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
 Based on : Peak Area

Rel. Reference Window : 20.000 %
 Abs. Reference Window : 0.000 min
 Rel. Non-ref. Window : 20.000 %
 Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
 Uncalibrated Peaks : not reported
 Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
 Origin : Ignored (some peaks differ, see below)
 Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
 Average Response : Average all calibrations
 Average Retention Time: Floating Average New 75%

Calibration Report Options :
 Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
 If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
 Signal 2: MSD1 85, EIC=84.7:85.7
 Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl	Amount	Area	Amt/Area	Ref	Grp	Name
8.744	1	1.00000	7.76074e4	1.28854e-5	1		Perchlorate
	2	2.00000	1.35273e5	1.47849e-5			
	3	5.00000	3.37764e5	1.48033e-5			
	4	10.00000	6.83454e5	1.46316e-5			
	5	25.00000	2.08433e6	1.19943e-5			
	6	50.00000	4.13334e6	1.20968e-5			
	7	75.00000	5.99313e6	1.25143e-5			
8.755	2	1.00000	2.36780e4	4.22333e-5	1		CLO4-85
	2	2.00000	4.69486e4	4.25998e-5			
	3	5.00000	1.06124e5	4.71147e-5			
	4	10.00000	2.13523e5	4.68335e-5			
	5	25.00000	6.14295e5	4.06971e-5			
	6	50.00000	1.19814e6	4.17315e-5			
	7	75.00000	1.78355e6	4.20509e-5			
8.766	3	5.00000	2.73208e5	1.83011e-5	+I1		CLO4-89-ISTD
	2	5.00000	2.24886e5	2.22335e-5			
	3	5.00000	2.33196e5	2.14412e-5			
	4	5.00000	2.34454e5	2.13262e-5			
	5	5.00000	2.50568e5	1.99547e-5			
	6	5.00000	2.30977e5	2.16472e-5			

RetTime [min]	Lvl Sig	Amount	Area	Amt/Area	Ref Grp Name
	7	5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-85

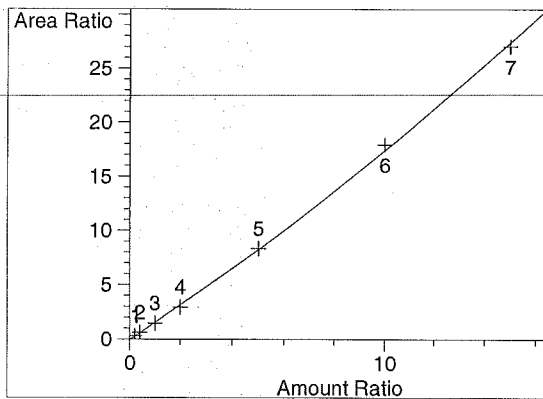
Time Window : From 6.650 min To 12.505 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-89-ISTD

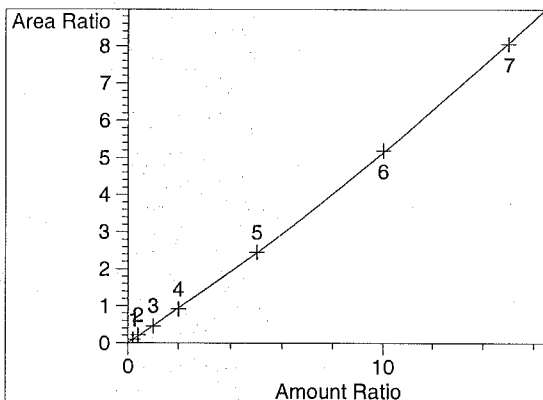
Time Window : From 6.659 min To 12.466 min
Curve Type : Linear
Origin : Included
Calibration Level Weights:/
Level 1 : 1
Level 2 : 1
Level 3 : 1
Level 4 : 1
Level 5 : 1
Level 6 : 1
Level 7 : 1

=====
Peak Sum Table
=====***No Entries in table***
=====

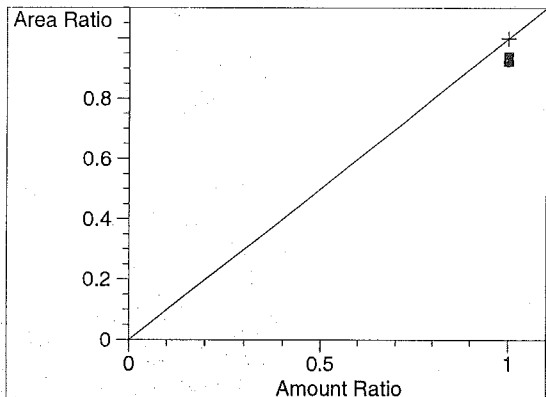
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

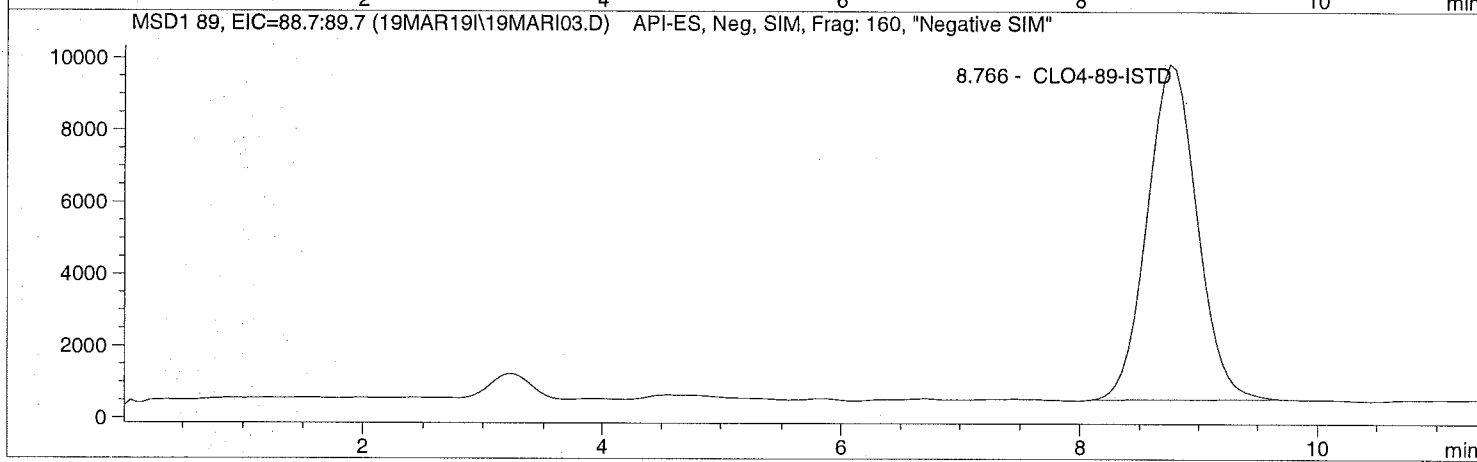
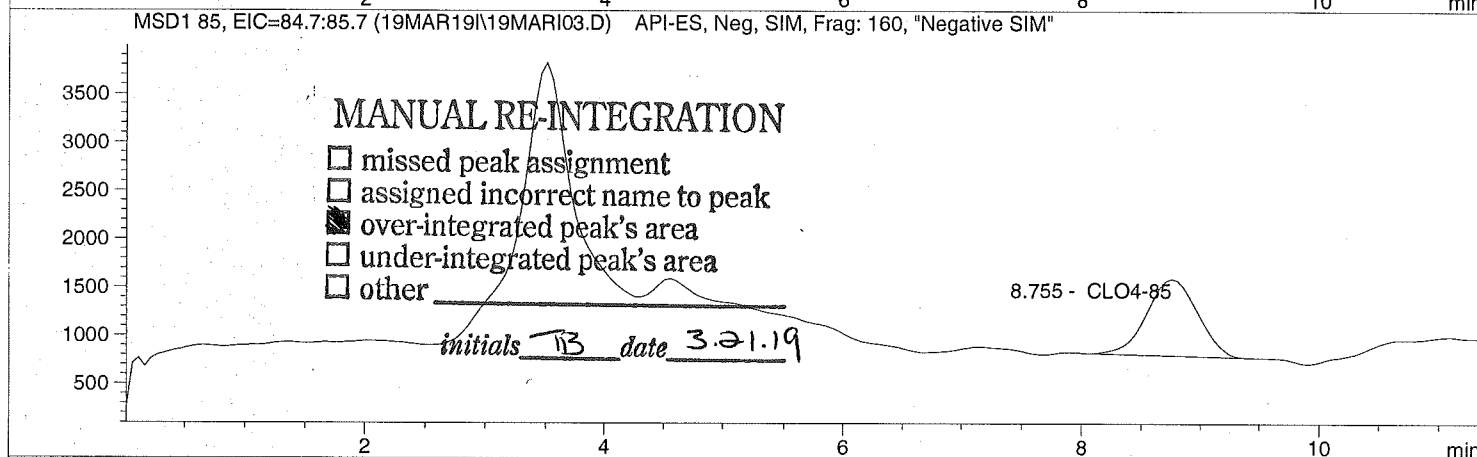
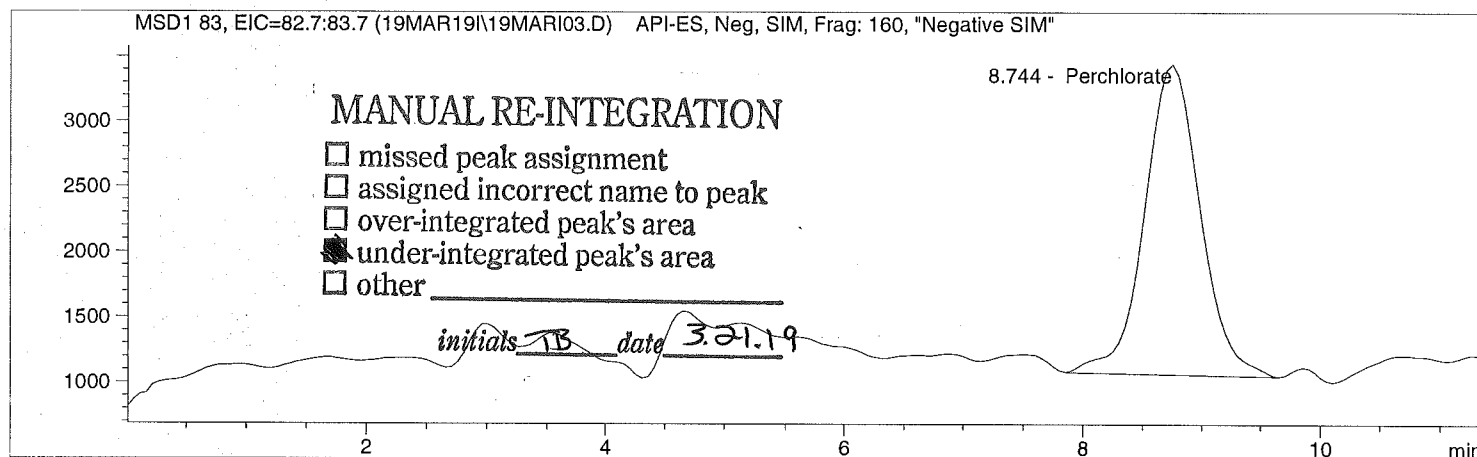
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:          3
Sample Name:    CLO4@ 1.0ug/L           Location:          Vial 73
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D

Sample Name: CLO4@ 2.0ug/L

Injection Date: 3/19/2019 09:53:00

Seq Line: 4

Sample Name: CLO4@ 2.0ug/L

Location: Vial 74

Acq Operator: TNB

Inj. No.: 1

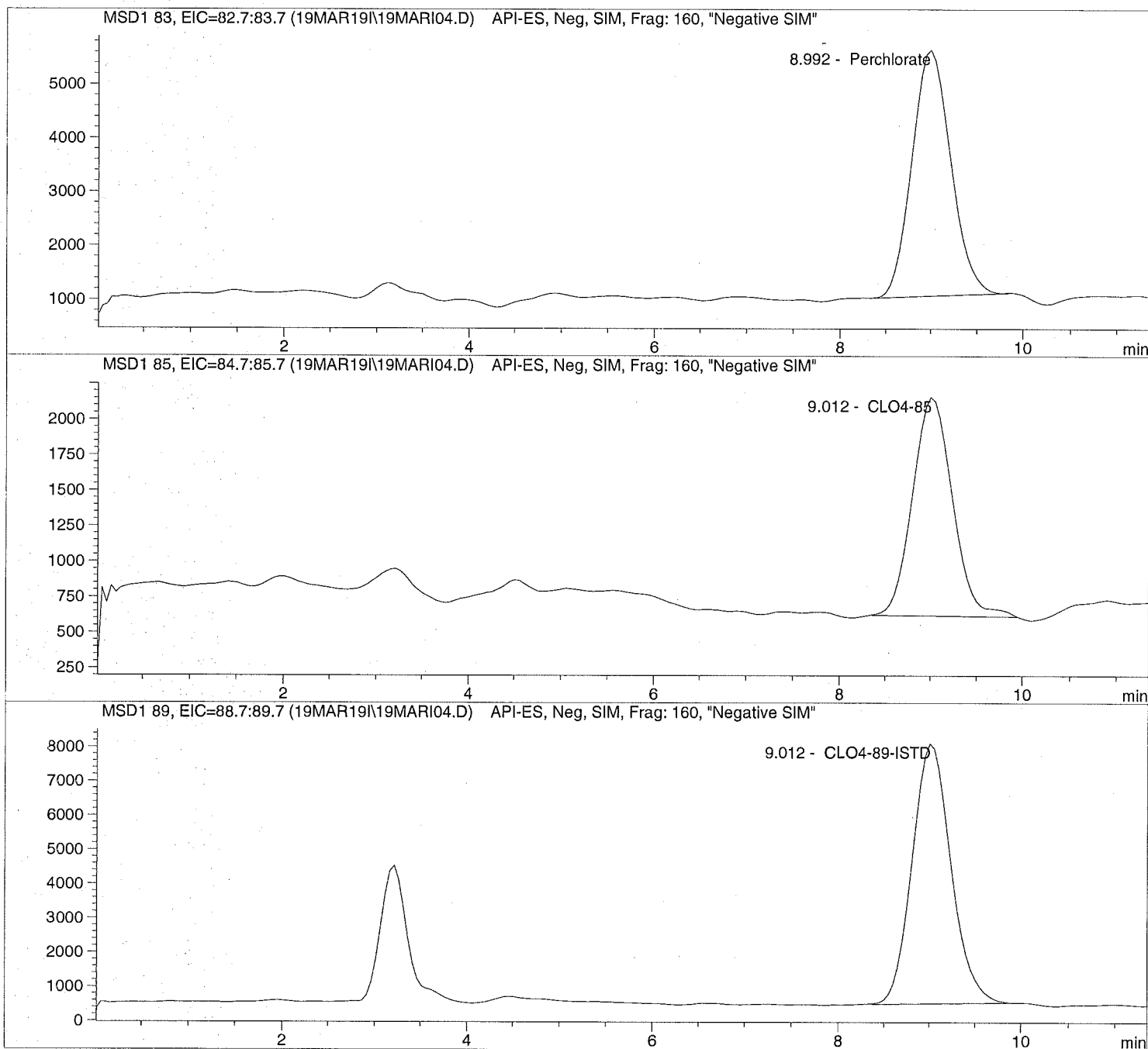
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line:      4
Sample Name:    CLO4@ 2.0ug/L           Location:      Vial 74
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D

Sample Name: CLO4@ 5.0ug/L

Injection Date: 3/19/2019 10:06:16

Seq Line: 5

Sample Name: CLO4@ 5.0ug/L

Location: Vial 75

Acq Operator: TNB

Inj. No.: 1

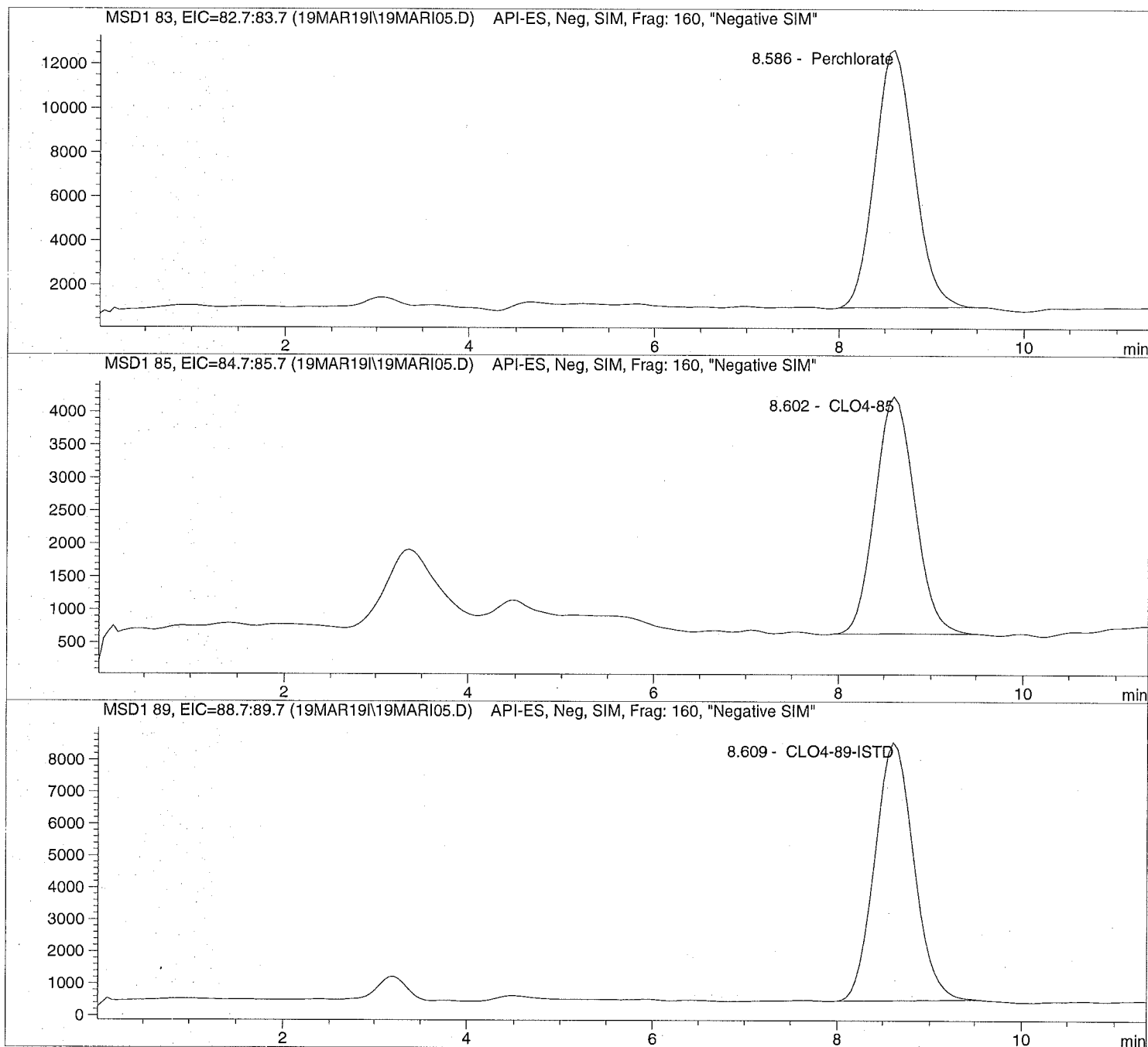
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line:      5
Sample Name:    CLO4@ 5.0ug/L           Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

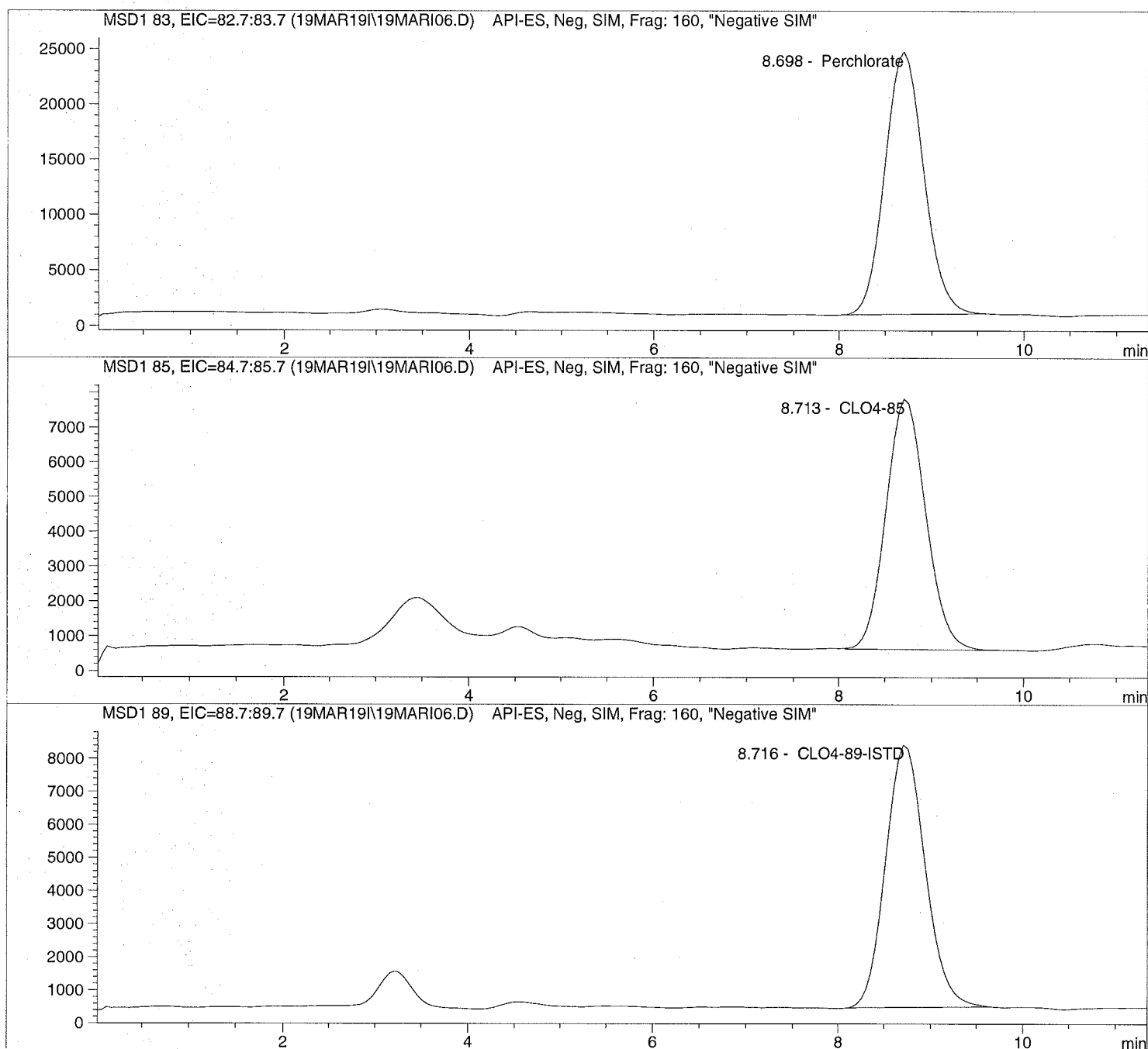
```


Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```
=====
Injection Date: 3/19/2019 10:19:32      Seq Line:      6
Sample Name:    CLO4@ 10.ug/L           Location:      Vial 76
Acq Operator:   TNB                     Inj. No.:    1
                                           Inj. Vol.:   30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line:          6
Sample Name:    CLO4@ 10.ug/L           Location:          Vial 76
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D

Sample Name: CLO4@ 25.ug/L

Injection Date: 3/19/2019 10:32:49

Seq Line: 7

Sample Name: CLO4@ 25.ug/L

Location: Vial 77

Acq Operator: TNB

Inj. No.: 1

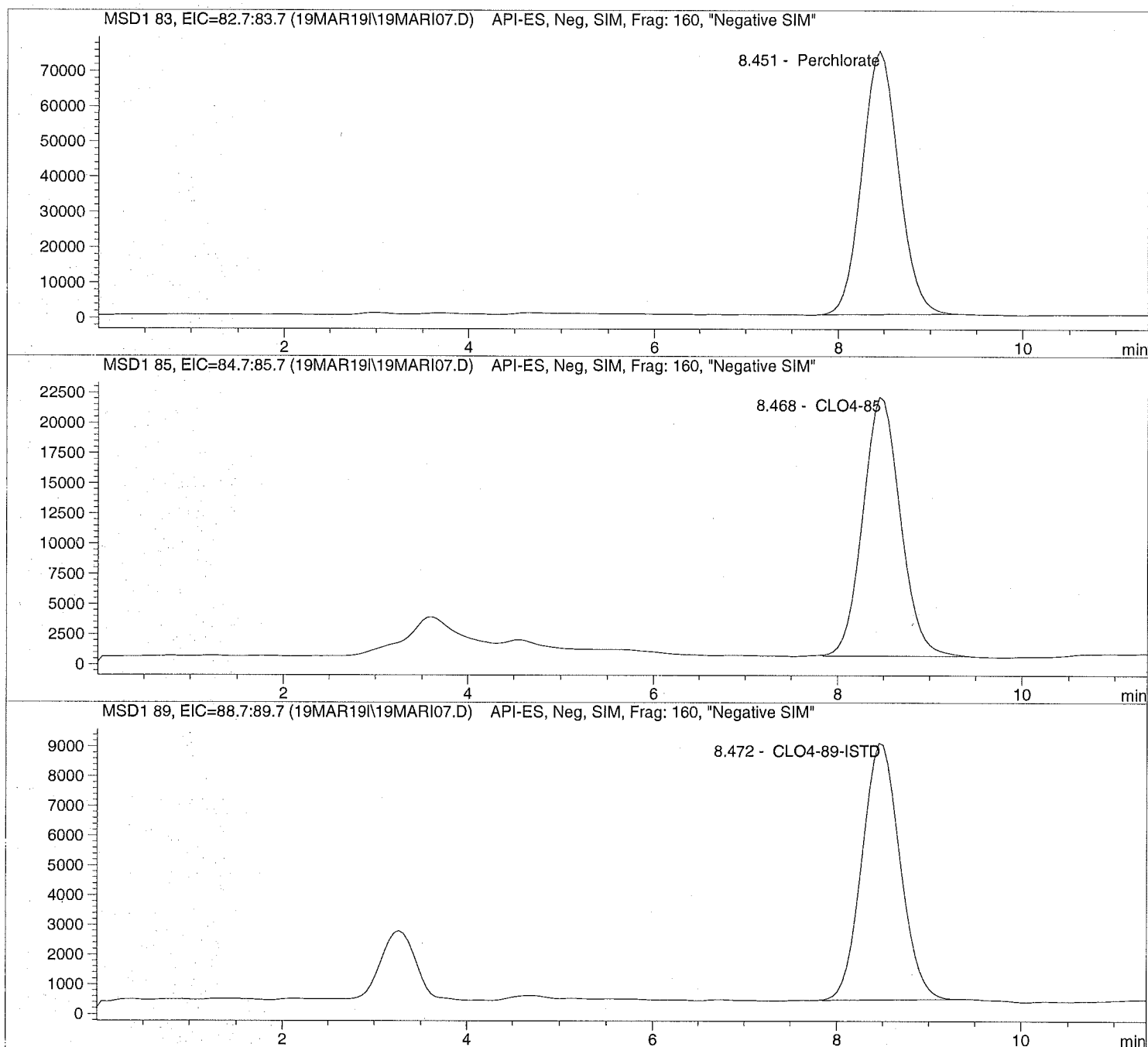
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 3/19/2019 10:32:49      Seq Line:            7
Sample Name:    CLO4@ 25.ug/L            Location:            Vial 77
Acq Operator:   TNB                      Inj. No.:            1
                                         Inj. Vol.:            30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:            Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:           1.000000
Dilution:             1.000000
Sample Amount:        25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

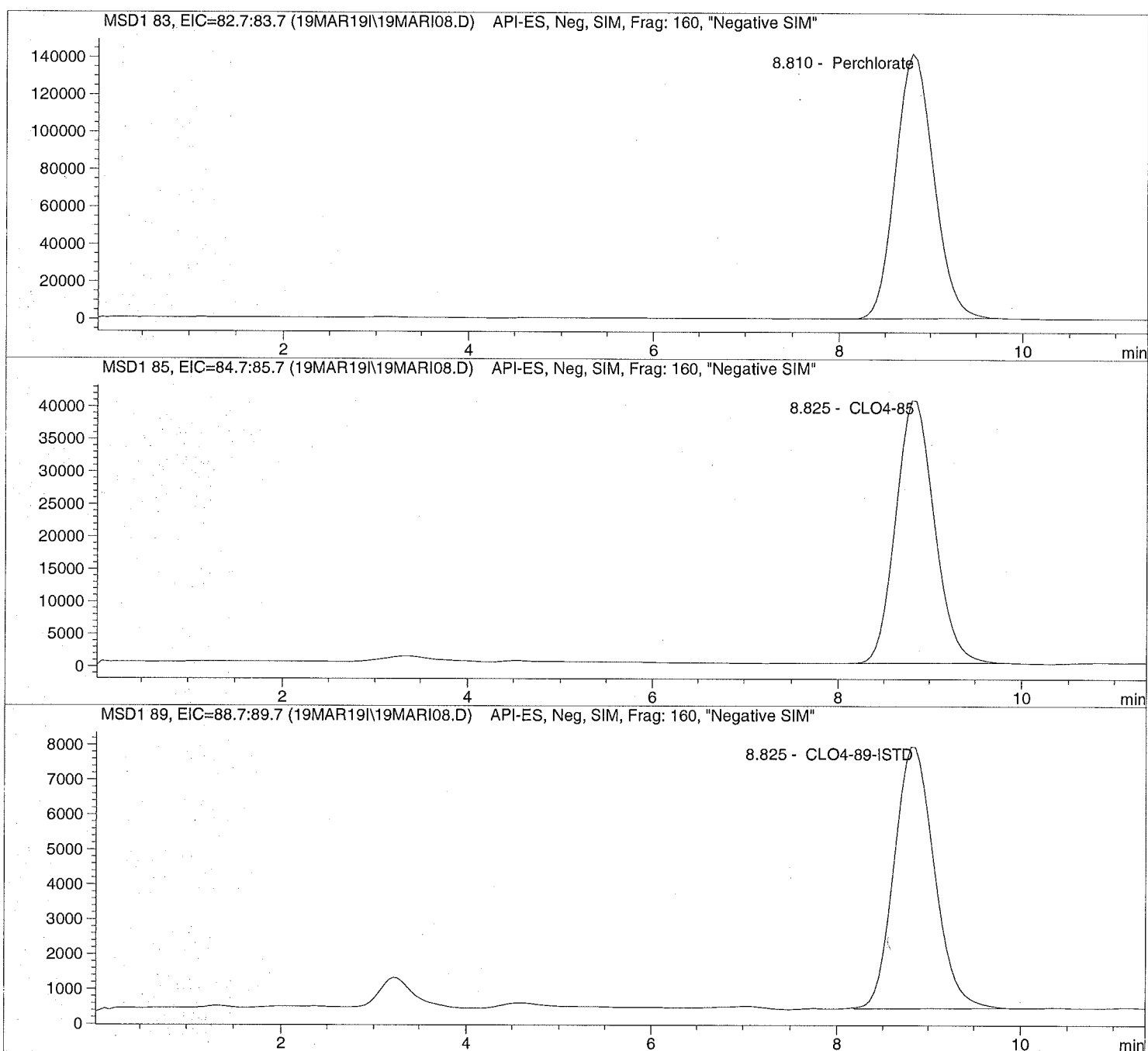
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI08.D Sample Name: CLO4@ 50.ug/L

```
=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 50.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 3/19/2019 10:59:22

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

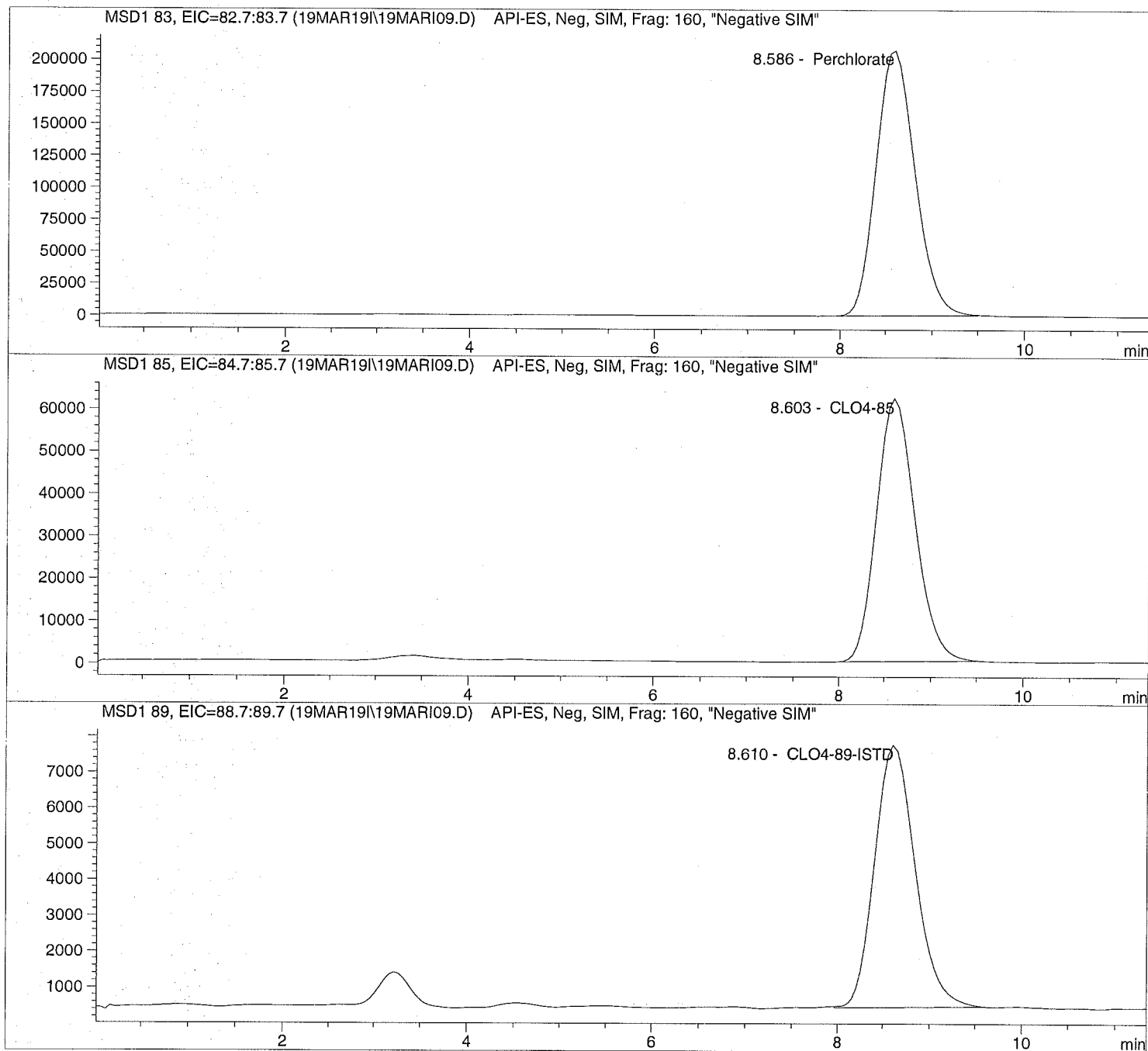
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date: 3/19/2019 10:59:22      Seq Line:          9
Sample Name:    CLO4@ 75.ug/L           Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  75.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D

Sample Name: ICAL Verf@10ug/L

Injection Date: 3/19/2019 11:12:42

Seq Line: 10

Sample Name: ICAL Verf@10ug/L

Location: Vial 80

Acq Operator: TNB

Inj. No.: 1

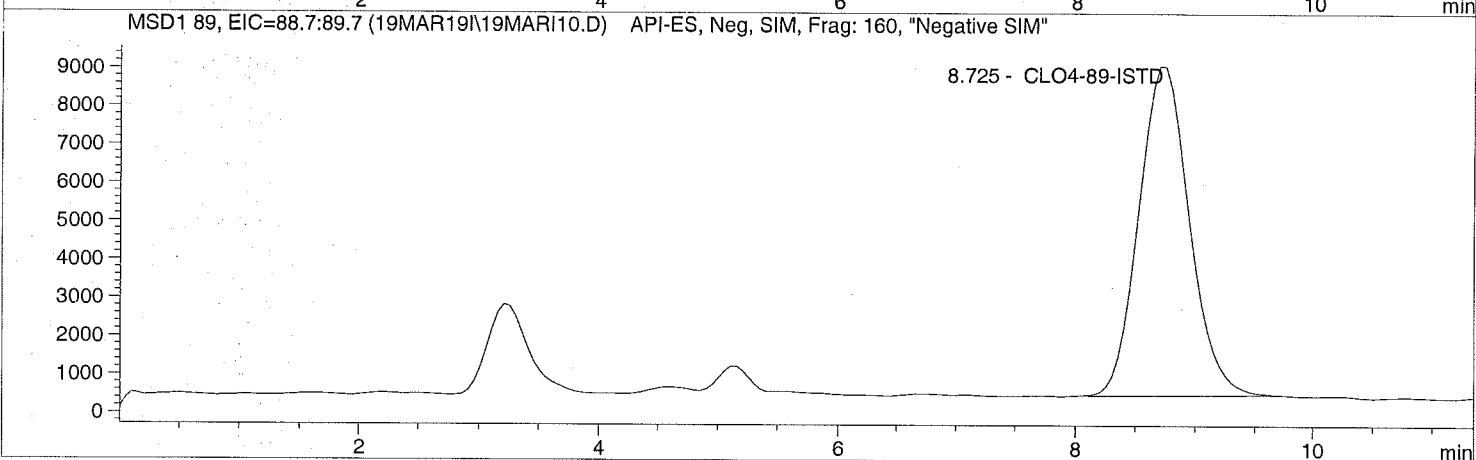
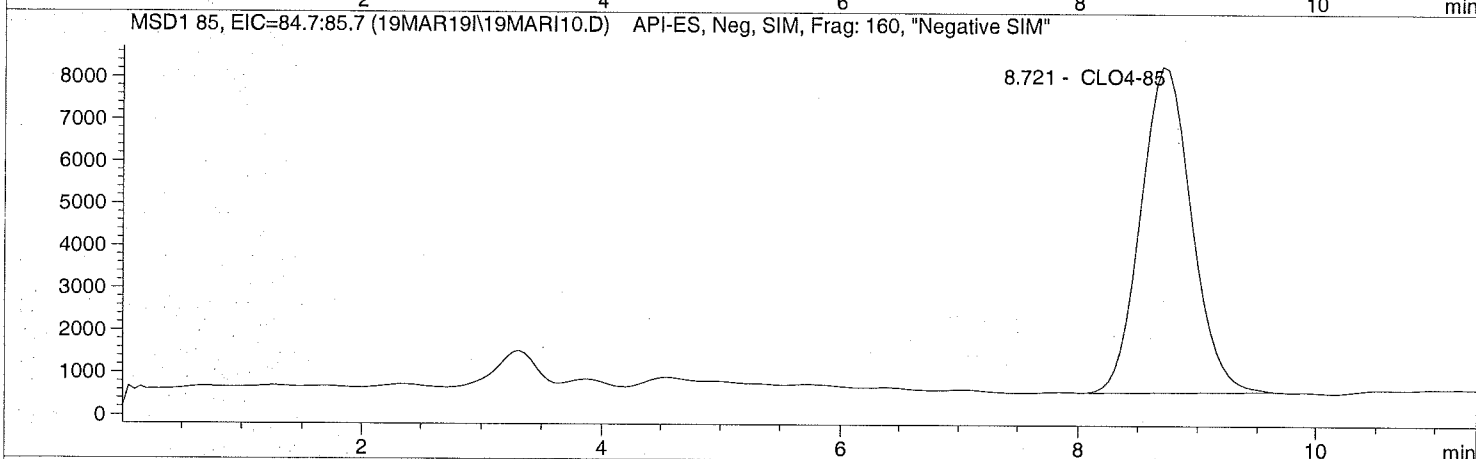
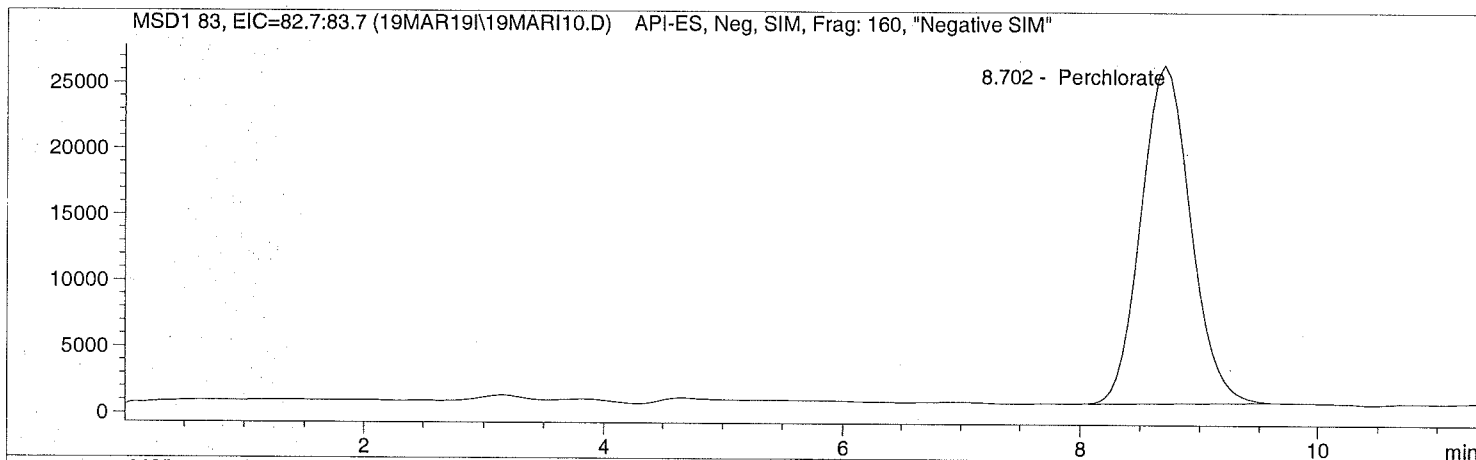
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 3/19/2019 11:12:42      Seq Line:      10
Sample Name:    ICAL Verf@10ug/L        Location:      Vial 80
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

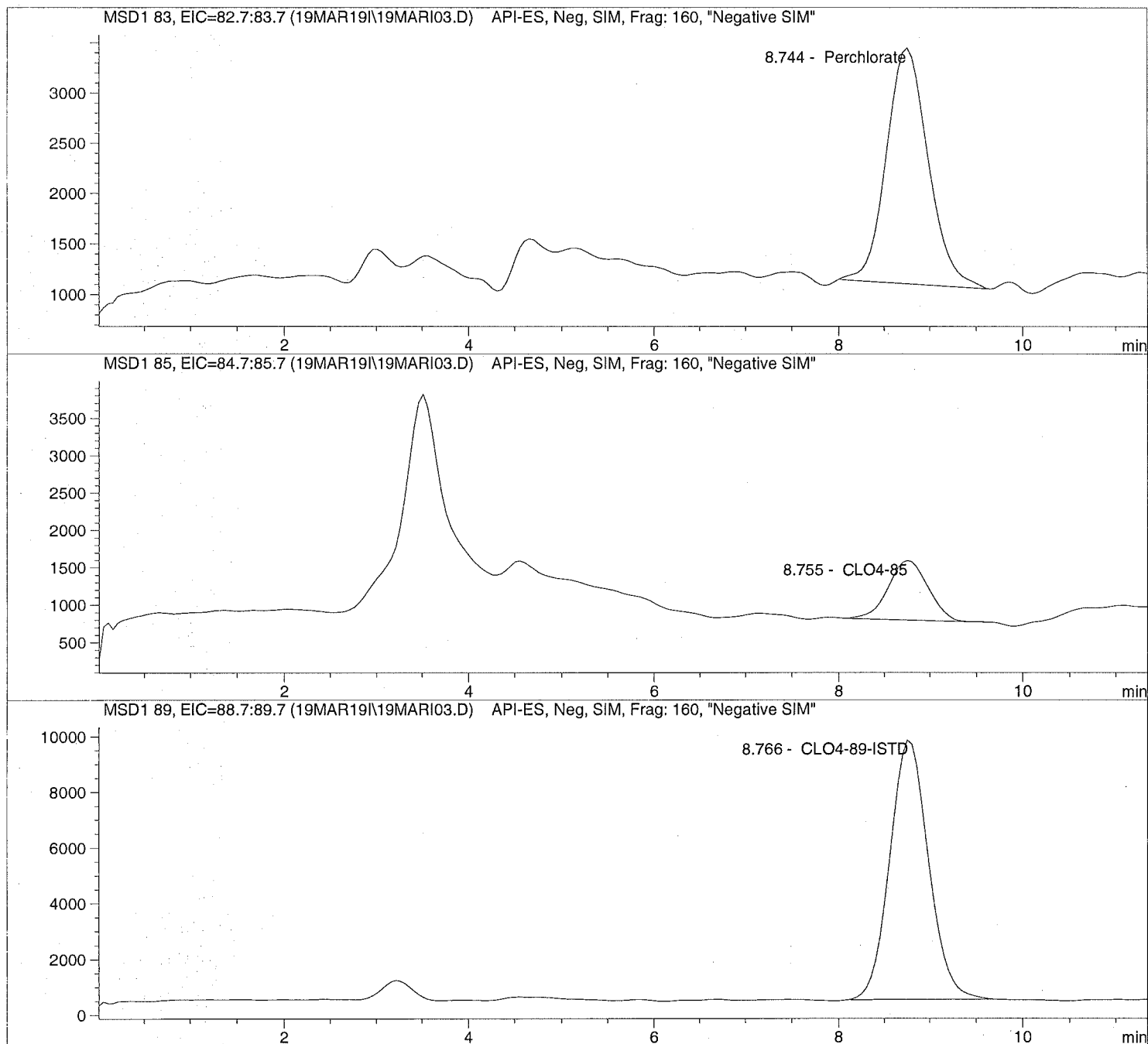
Inj. Vol.: 30 µl

Acq Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```
=====
Injection Date: 3/19/2019 09:39:40      Seq Line:                    3
Sample Name:    CLO4@ 1.0ug/L            Location:                Vial 73
Acq Operator:    TNB                      Inj. No.:                1
                                          Inj. Vol.:               30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:    3/19/2019 14:38:25
```

Perchlorate analysis

Sample Information

```
Sorted By:                    Signal
Calib. Data Modified:        Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                  1.000000
Dilution:                    1.000000
Sample Amount:                1.000
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

*** End of Report ***



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 31, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19030763**

Laboratory Results for: **LH18/24 GW Treatment Plant Monthly Influent Samples**

Dear Marcia,

ALS Environmental received 1 sample(s) on Mar 15, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
Work Order: HS19030763

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19030763-01	LH18/24-SP140_031419	Water		14-Mar-2019 14:00	15-Mar-2019 08:38	<input type="checkbox"/>

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
-

Metals by Method SW6020**Batch ID: 138939**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

WetChemistry by Method SW7196**Batch ID: R334751**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 GW Treatment Plant Monthly Influent Samples
 Sample ID: LH18/24-SP140_031419
 Collection Date: 14-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19030763
 Lab ID:HS19030763-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
ICP-MS METALS BY SW6020A		Method:SW6020				Prep:SW3010A / 21-Mar-2019		Analyst: JCJ
Selenium	0.00250	U	0.00110	0.00250	0.00200	mg/L	1	23-Mar-2019 01:07
Silver	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	23-Mar-2019 01:07
HEXAVALENT CHROMIUM BY SW7196A		Method:SW7196						Analyst: MZD
Chromium, Hexavalent	0.0100	U	0.00600	0.0100	0.0100	mg/L	1	15-Mar-2019 12:14
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	31-Mar-2019 12:13

Note: See Qualifiers Page for a list of qualifiers and their explanation.

WEIGHT LOG

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030763

Batch ID: 138939 **Method:** ICP-MS METALS BY SW6020A **Prep:** 3010A

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19030763-01	1	10	10 (mL)	1

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030763

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID 138939	Test Name : ICP-MS METALS BY SW6020A		Matrix: Water			
HS19030763-01	LH18/24-SP140_031419	14 Mar 2019 14:00		21 Mar 2019 10:00	23 Mar 2019 01:07	1
Batch ID R334751	Test Name : HEXAVALENT CHROMIUM BY SW7196A		Matrix: Water			
HS19030763-01	LH18/24-SP140_031419	14 Mar 2019 14:00			15 Mar 2019 12:14	1
Batch ID R335660	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19030763-01	LH18/24-SP140_031419	14 Mar 2019 14:00			31 Mar 2019 12:13	1

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030763

QC BATCH REPORT NEW

Batch ID: 138939 (0)		Instrument: ICPMS04		Method: ICP-MS METALS BY SW6020A						
MBLK	Sample ID: MBLK-138939	Units: mg/L		Analysis Date: 23-Mar-2019 00:40						
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003309		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.00250	0.00200							U	
Silver	0.000500	0.00200							U	
LCS	Sample ID: LCS-138939	Units: mg/L		Analysis Date: 23-Mar-2019 00:42						
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003310		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.05143	0.00200	0.05	0	103	80 - 120				
Silver	0.05011	0.00200	0.05	0	100	80 - 120				
MS	Sample ID: HS19030918-01MS	Units: mg/L		Analysis Date: 23-Mar-2019 00:49						
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003313		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.05006	0.00200	0.05	-0.000569	101	80 - 120				
Silver	0.04751	0.00200	0.05	-0.000012	95.0	80 - 120				
MSD	Sample ID: HS19030918-01MSD	Units: mg/L		Analysis Date: 23-Mar-2019 00:51						
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003314		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.0556	0.00200	0.05	-0.000569	112	80 - 120	0.05006	10.5	20	
Silver	0.04885	0.00200	0.05	-0.000012	97.7	80 - 120	0.04751	2.79	20	
PDS	Sample ID: HS19030918-01PDS	Units: mg/L		Analysis Date: 23-Mar-2019 00:53						
Client ID:	Run ID: ICPMS04_335069	SeqNo: 5003315		PrepDate: 21-Mar-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Selenium	0.1029	0.00200	0.1	-0.000569	103	75 - 125				
Silver	0.08679	0.00200	0.1	-0.000012	86.8	75 - 125				

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030763

QC BATCH REPORT NEW

Batch ID: 138939 (0) **Instrument:** ICPMS04 **Method:** ICP-MS METALS BY SW6020A

SD	Sample ID:	HS19030918-01SD			Units:	mg/L					Analysis Date:	23-Mar-2019 00:47	
Client ID:	Run ID:	ICPMS04_335069			SeqNo:	5003312		PrepDate:	21-Mar-2019		DF:	5	
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	Limit	Qual			
Selenium	0.0125	0.0100					-0.000569	0	10	U			
Silver	0.00250	0.0100					-0.000012	0	10	U			

The following samples were analyzed in this batch:

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
WorkOrder: HS19030763

QC BATCH REPORT NEW

Batch ID: R334751 (0)		Instrument: UV-2450		Method: HEXAVALENT CHROMIUM BY SW7196A						
MBLK	Sample ID: MBLK-334751	Units: mg/L		Analysis Date: 15-Mar-2019 12:14						
Client ID:	Run ID: UV-2450_334751	SeqNo: 4995026		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chromium, Hexavalent	0.0100	0.0100							U	
LCS	Sample ID: LCS-334751	Units: mg/L		Analysis Date: 15-Mar-2019 12:14						
Client ID:	Run ID: UV-2450_334751	SeqNo: 4995027		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chromium, Hexavalent	0.244	0.0100	0.25	0	97.6	90 - 111				
LCSD	Sample ID: LCSD-334751	Units: mg/L		Analysis Date: 15-Mar-2019 12:14						
Client ID:	Run ID: UV-2450_334751	SeqNo: 4995028		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chromium, Hexavalent	0.244	0.0100	0.25	0	97.6	90 - 111	0.244	0	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 31-Mar-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 GW Treatment Plant Monthly Influent Samples	
WorkOrder:	HS19030763	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

ALS Houston, US

Date: 31-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 GW Treatment Plant Monthly Influent Samples
Work Order: HS19030763

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19030763-01	LH18/24-SP140_031419	Login	3/15/2019 11:03:24 AM	JRM	WET176
HS19030763-01	LH18/24-SP140_031419	Login	3/15/2019 11:03:24 AM	JRM	WET176
HS19030763-01	LH18/24-SP140_031419	Login	3/15/2019 11:03:24 AM	JRM	MET068

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19030763

Date/Time Received: **15-Mar-2019 08:38**
 Received by: **JRM**

Checklist completed by: Jared R. Makan 15-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 15-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s):

1.3c/1.3c UC/C	IR25
----------------	------

Cooler(s)/Kit(s):

25780

Date/Time sample(s) sent to storage:

03/15/2019 11:10

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

--

Login Notes:

Client Contacted: _____ Date Contacted: _____ Person Contacted: _____
 Contacted By: _____ Regarding: _____

Comments:

--

Corrective Action:

--

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd., Suite 210 Houston, TX, 77099 (281) 530-5656 ATTN: RJ Modshia

Page 1 of 1

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001			Analyses										Remarks (Preservatives, etc.)	Lab I.D.#					
Job: GROUNDWATER TREATMENT PLANT MONTHLY INFLUENT SAMPLES			MS / MSD	No. OF CONTAINERS	SILVER & SELENIUM	HEXAVALENT CHROMIUM	PERCHLORATE															
Prepared By: Scott Beesinger		P.O. Number																				
Field Sample I.D.	Sample Matrix	Date / Time																				
LH18/24-SP140_031419	Water	03/14/19 / 14:00		1	X																	
LH18/24-SP140_031419	Water	03/14/19 / 14:00		2		X	X															

Additional Remarks: STANDARD TURN AROUND TIME

Relinquished By: <i>Scott Beesinger</i>	Date 03/14/19	Time 14:30	Received By:	Date	Time	Relinquished By:	Date	Time	Received By:	Date	Time
---	-------------------------	----------------------	---------------------	-------------	-------------	-------------------------	-------------	-------------	---------------------	-------------	-------------

For Lab Use Only									
Received At Lab By: <i>J. Wynn</i>	Date 3/15/19	Time 08:38	Airbill No.	Opened By:	Date	Time	Temp of Container	Seal No.	Condition
Remarks: <i>Cooler 25780 1K25</i> <i>Temp 1.3 CF0.0</i>									

HS19030763

Bhate Environmental Associates, Inc.
 18/24 GW Treatment Plant Monthly Influent Samples



	ALS 10450 Stanciff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5687	CU Date: 3/14/14 Name: Scott Company: B...
---	---	--

STODY SEAL Time: 1430 B...	Seal Broken By: SM Date: 3/15/14
---	---

FedEx
 TRK#
 0221 4380 9530 9490

FRI - 15 MAR 10:30A
 PRIORITY OVERNIGHT

AB SGRA

77099
 TX-US
 IAH



FID 3586291 14MAR19 GDA 553C174603/0C8A



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1907867; 1907869; 1907871;
1908794

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2230 (235529)

General Set Information: There were four field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Method QC data: The method blank (LMB 645533) was less than 1/2 the CRDL. The recovery for the LCS (645534) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0 μ l of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4. μ g/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in μ g/L. Results were calculated in μ g/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (μ g/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level of 4.0 μ g/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch March 29, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 29, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1907867**

Project ID: HS19030763

Purchase Order: HS19030763

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP140_031419	1907867001	03/14/19	03/19/19	



ANALYTICAL REPORT

Workorder: 34-1907867

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP140_031419	Sampling Site: NA	Collected: 03/14/2019				
Lab ID: 1907867001	Media: 125 mL Nalgene	Received: 03/19/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2230 (HBN: 235529) Analyzed: 03/28/2019 09:28	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	8300	1000	2000	4000	1000	

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 235529)

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/28/2019 14:08	/S/ Stephen Brose 03/29/2019 12:58

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1907867

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935413

Analysis Information

Workorder: 1907867

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2230 (HBN: 235529)
Analyzed By: Thomas Bosch

Blank

LMB: 645533 Analyzed: 03/28/2019 09:14 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 645534 Analyzed: 03/28/2019 08:47 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.00	4.00	99.9	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1907869001 Analyzed: 03/28/2019 09:41 Dilution: 1 Units: ug/L			MS: 645535 Analyzed: 03/28/2019 09:54 Dilution: 1 Units: ug/L			MSD: 645536 Analyzed: 03/28/2019 10:08 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	ND	4.66	4	117	78.8 123.8	4.05	101	14	0.0 20.0

Comments

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/28/2019 14:23	/S/ Stephen Brose 03/29/2019 12:58

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



18698/#2

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10918

1907867

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19030763
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19030763-01	LH18/24-SP140_031419	Water	14 Mar 2019 14:00
SUB_Perch-6850			29 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. [Signature]
Received By: Jamir [Signature]
Cooler ID(s): _____

Date/Time: 3/18/19 18:00
Date/Time: 03/19/19 9:50
Temperature(s): _____

ALS Environmental
CHAIN-OF-CUSTODY



Project / Job / Task: HS19030763		Workorder ID: 1907867		Level: ENV_LVL4		Requested Analysis	
Client: ALS Environmental (Houston)		Split:		Type: 125Poly			
		Account: 8101		Preservatives			
Comments:				Cool			
				Containers			
Item	Collect Date/Time	Sample ID	Lab ID	QC	Matrix	ID(s)	Count
1	03/14/2019 14:00	LH18/24-SP140_031419	1907867001		Water	A	1
2							
3							
4							
5							
6							
7							
8							
9							
10							

EP A 6850, D, D, OSM

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY				SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY			
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location	Prepared / Analyzed by:	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location
<i>Wraith</i>	03/19/2019 09:50	ALS Sample Receiving	Sample Login	Lab Notebook No.:			
<i>Wraith</i>	03/19/2019 11:45	<i>KSC</i>	<i>Storage</i>	Date / Time:			
<i>R-33.1</i>		<i>T. Brook</i>	<i>6850</i>				

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: ALS Houston Project/Task/Site: 1907867
 Date/Time of Receipt: 03-19-19 9:50 Number of Coolers Received: 1

Condition of Coolers: Acceptable/Unacceptable Temperature Control: Present/Not Included
 Cooler Custody Seals: Present/Absent/NA
 Intact/Broken/NA Location Temp Taken: Control/Between Samples
 Container Custody Seals: Present/Absent/NA
 Intact/Broken/NA Are all temperatures within project specific guidelines? Yes/No/NA
 Ice Present: Yes/No/NA VOA Headspace Present? Yes/No/NA
Frozen/Melted/NA

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9750</u>	<u>1</u> °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: Tamir Vassell Signature Tamir Vassell Printed Name 03-19-19 Date

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES No

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Part # 159469-434 RITZ EXP 11/19

ORIGIN ID: SGRA (281) 530-5656
SHIPPING DEPT
ALS LABORATORY GROUP
10450 STANCLIFF RD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

SHIP DATE: 18MAR19
ACTWT: 9.35 LB
CAD#: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

3001/EGEP/4/1C1/1155

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(01) 288-7700

REF: HS19030749/761/763 RJ



FedEx
Express



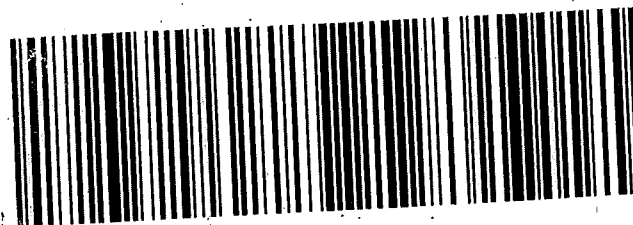
AN 10505080606011181J

TRK# 4809 7831 7725
0201

TUE - 19 MAR 3:00P
STANDARD OVERNIGHT

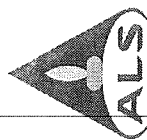
AX BTFA

84123
UT-US SLC



ALS
10450 Stancliff Rd., Suite 210
Houston, Texas 77099
Tel: +1 281 530 5656
Fax: +1 281 530 5887

Date
Name
Com



Batch Worklist

Batch: ELMS/ 2230 **Created:** 3/28/2019 07:45 **Instrument:** HBN: 235529
Rule: EPA 6850, DoD QSM Water **Analyst:** T. Bosch **Status:** WP



Workorder: 1907867 [ENV_LVL4]
Workorder: 1907869 [ENV_LVL4]
Workorder: 1907871 [ENV_LVL4]
Workorder: 1908794 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	64530	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	
2	64531	RLV'S for HBN 235529 [ELMS/2230]				RLV'S	3		E685041C3Q	5311		4/1/2019	
3	64532	ICS for HBN 235529 [ELMS/2230]				ICS	3		E6850.D3Q	5311		4/1/2019	
4	64533	LMB for HBN 235529 [ELMS/2230]				LMB	3		E6850Q413Q	5311		4/1/2019	
5	64534	LCS for HBN 235529 [ELMS/2230]				LCS	3		E6850Q413Q	5311		4/1/2019	
6	1907867001	LH18/24-SP140_031419				SAMPLE	3	1907867001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
7	1907869001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907869001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
8	64535	LH18/24-SP650...(1907869001MS)				MS	3		E6850Q413Q	5311		4/1/2019	
9	64536	LH18/24-SP65...(1907869001MSD)				MSD	3		E6850Q413Q	5311		4/1/2019	
10	1907871001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907871001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
11	1908794001	LH18/24-SP650_032119_BIX				SAMPLE	3	1908794001-A	E6850Q41.3	5480	4/18/2019	4/8/2019	
12	64537	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #'s: 1907867 (001); 1907869 (001); 1907871 (001); 1908794 (001)

ELMS Batch/HBN ID: 2230 (235529)

Prep Date: 03/27/2019 Analysis Date: 03/28/2019 Analyst: T. Bosch

Analyte: **Perchlorate** Matrix: **Water** Method: **6850**

Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\28MAR19D.s

Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by **TNB**. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 25µL
Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 645534; Target = 4.0µg/L. ASTM type II water was used for LMB 645533.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\235529-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
<u>Batch(es)/SDG: ELMS: 2230 HBN: 235529</u>		
<u>Sample Set IDs if Applicable: 1907867/1907869/1907871/1908794</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK		Description - 6850 WKG Std 100.ug/L			
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK		Description - 6850 Stock AccStd 1,000ug/mL	
Standard: 43659	Created By: Thomas Bosch	Amount: 100 mL	
MFG: AccuStandard	Create Date: 09/17/2018 09:09AM	Expires: 07/25/2020	
MFG Lot: 218065075		Usable: No	
Part ID: IC-PER-10X-1		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT		Description - 6850 Intermdt AccStd 10.ug/mL			
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK		Description - 6850 QC WKG STD 100ug/L			
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK		Description - 6850 QC Stock STD 1,000ug/mL	
Standard: 36748		Created By: Thomas Bosch	Amount: 100 mL
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	Expires: 03/31/2020
MFG Lot: CP-0860			Usable: Yes
Part ID: ICC-013			Lab Lot: CLO4 QC STOCK
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:

This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis



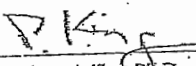
ISO Guide 34 Reference Material

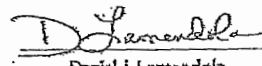
Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:

The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QA/RA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:
 ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
 (Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$
 Labeled CAS Number: NA
 Unlabeled CAS Number: 7601-89-0
 MW*: 130.4
 Chemical Formula: NaCl^+O_4
 Storage: Store at room temperature away from light and moisture.
 Stability: See storage and expiration data.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 $\mu\text{g/mL}$
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 \pm 2.8 $\mu\text{g/mL}$ ($k=2$)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
*	645530	CCV@25	Vial 71	1	Control	1	1.81499e6	24.13647
*	645534	QC@4.0	Vial 72	1	Control	2	2.89937e5	3.99582
*	645532	ICS@4.0	Vial 73	1	Control	3	2.38655e5	3.65624
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	7.55115e5	8344.77963
*	1907869001		Vial 76	1	Sample	6	0.00000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	2.77046e5	4.66163
*	645536	78691SD	Vial 78	1	Sample	8	2.89384e5	4.05130
*	1907871001		Vial 79	1	Sample	9	0.00000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	1.79709e6	24.49119

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
*	645530	CCV@25	Vial 71	1	Control	1	5.33395e5	23.90219
*	645534	QC@4.0	Vial 72	1	Control	2	9.52963e4	4.26699
*	645532	ICS@4.0	Vial 73	1	Control	3	7.88339e4	3.90865
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.38827e5	8762.35701
*	1907869001		Vial 76	1	Sample	6	0.00000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	8.93399e4	4.91371
*	645536	78691SD	Vial 78	1	Sample	8	9.85690e4	4.48457
*	1907871001		Vial 79	1	Sample	9	0.00000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	5.35802e5	24.59111

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
*	645530	CCV@25	Vial 71	1	Control	1	2.29255e5	5.00000
*	645534	QC@4.0	Vial 72	1	Control	2	2.39106e5	5.00000
*	645532	ICS@4.0	Vial 73	1	Control	3	2.16080e5	5.00000
*	645533	LMB	Vial 74	1	Control	4	2.42742e5	5.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.89136e5	5000.00000
*	1907869001		Vial 76	1	Sample	6	1.89925e5	5.00000
*	645535	78691MS	Vial 77	1	Sample	7	1.94413e5	5.00000
*	645536	78691SD	Vial 78	1	Sample	8	2.35220e5	5.00000
*	1907871001		Vial 79	1	Sample	9	1.89609e5	5.00000
*	1908794001		Vial 80	1	Sample	10	1.92453e5	5.00000
*	645537	CCV@25	Vial 71	1	Control	11	2.23514e5	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

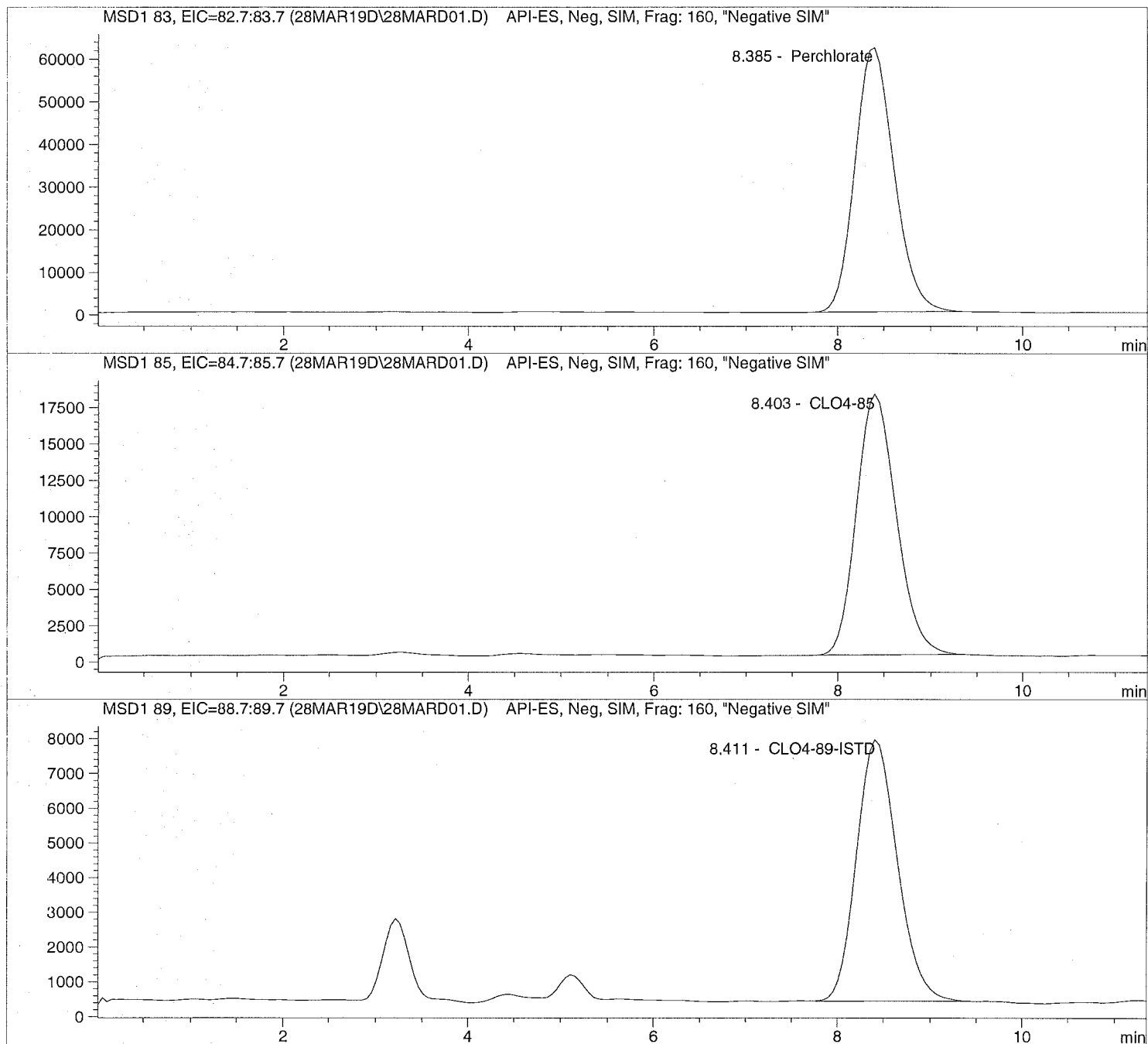
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	645530	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	645534	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	645532	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	645533	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1907867001	1K	CLO4-AQN	1	Sample	
6	Vial 76	1907869001		CLO4-AQN	1	Sample	
7	Vial 77	645535	78691MS	CLO4-AQN	1	Sample	
8	Vial 78	645536	78691SD	CLO4-AQN	1	Sample	
9	Vial 79	1907871001		CLO4-AQN	1	Sample	
10	Vial 80	1908794001		CLO4-AQN	1	Sample	
11	Vial 71	645537	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D Sample Name: 645530 CCV@25

```
=====
Injection Date: 3/28/2019 08:30:53      Seq Line: 1
Sample Name: 645530 CCV@25              Location: Vial 71
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D Sample Name: 645530 CCV025

```

=====
Injection Date: 3/28/2019 08:30:53      Seq Line: 1
Sample Name: 645530 CCV025              Location: Vial 71
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.385	PBA	1814992.7	24.1365	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.403	PBA	533395.5	23.9022	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.411	BBA	229255.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D

Sample Name: 645534 QC@4.0

Injection Date: 3/28/2019 08:47:42

Seq Line: 2

Sample Name: 645534 QC@4.0

Location: Vial 72

Acq Operator: TNB

Inj. No.: 1

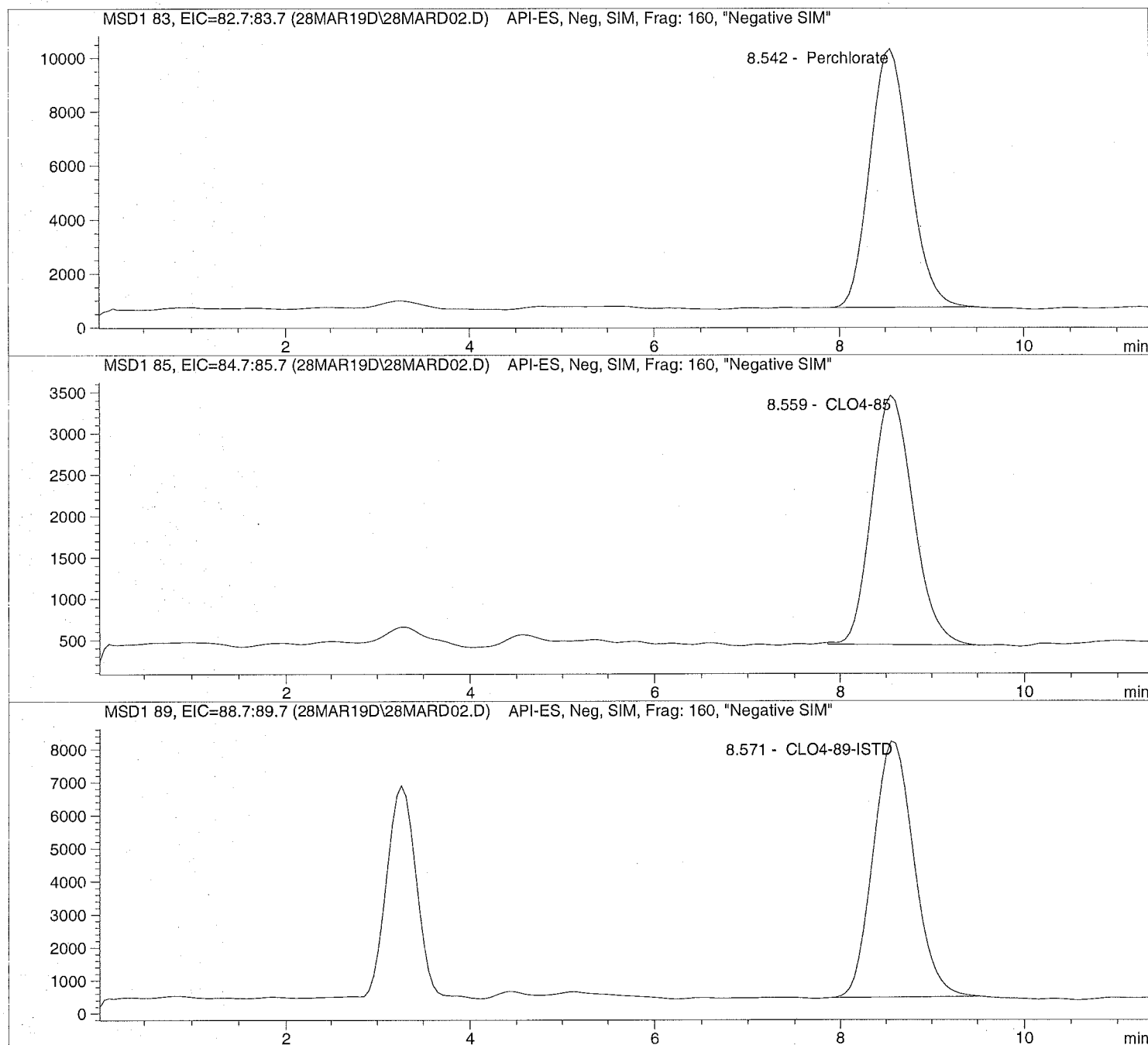
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D Sample Name: 645534 QC@4.0

```

=====
Injection Date: 3/28/2019 08:47:42      Seq Line: 2
Sample Name: 645534 QC@4.0              Location: Vial 72
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.542	PBA	289937.1	3.9958	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.559	BBA	95296.3	4.2670	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.571	PBA	239105.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

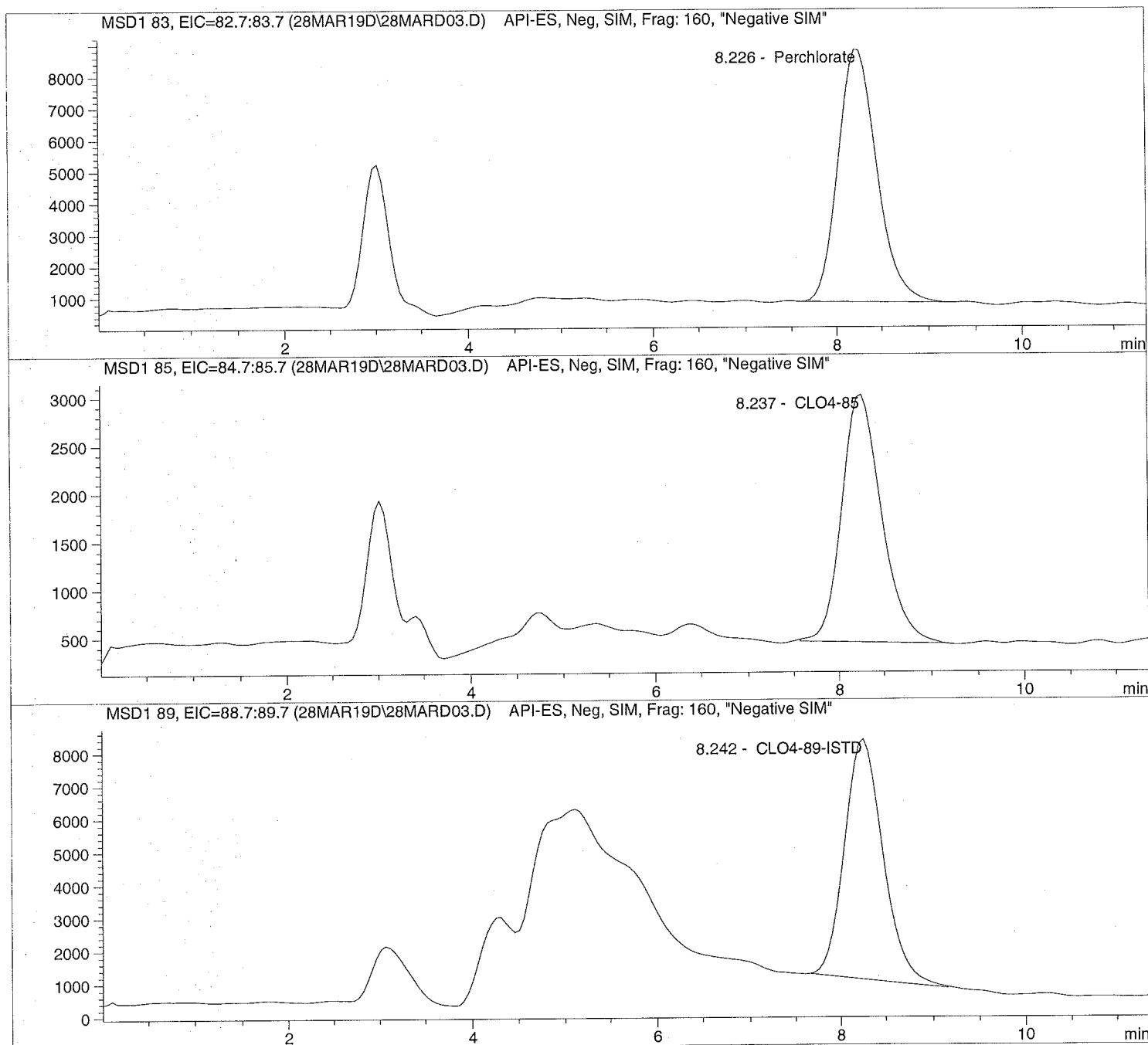
```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD03.D Sample Name: 645532 ICS@4.0

```
=====
Injection Date: 3/28/2019 09:00:54 Seq Line: 3
Sample Name: 645532 ICS@4.0 Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D

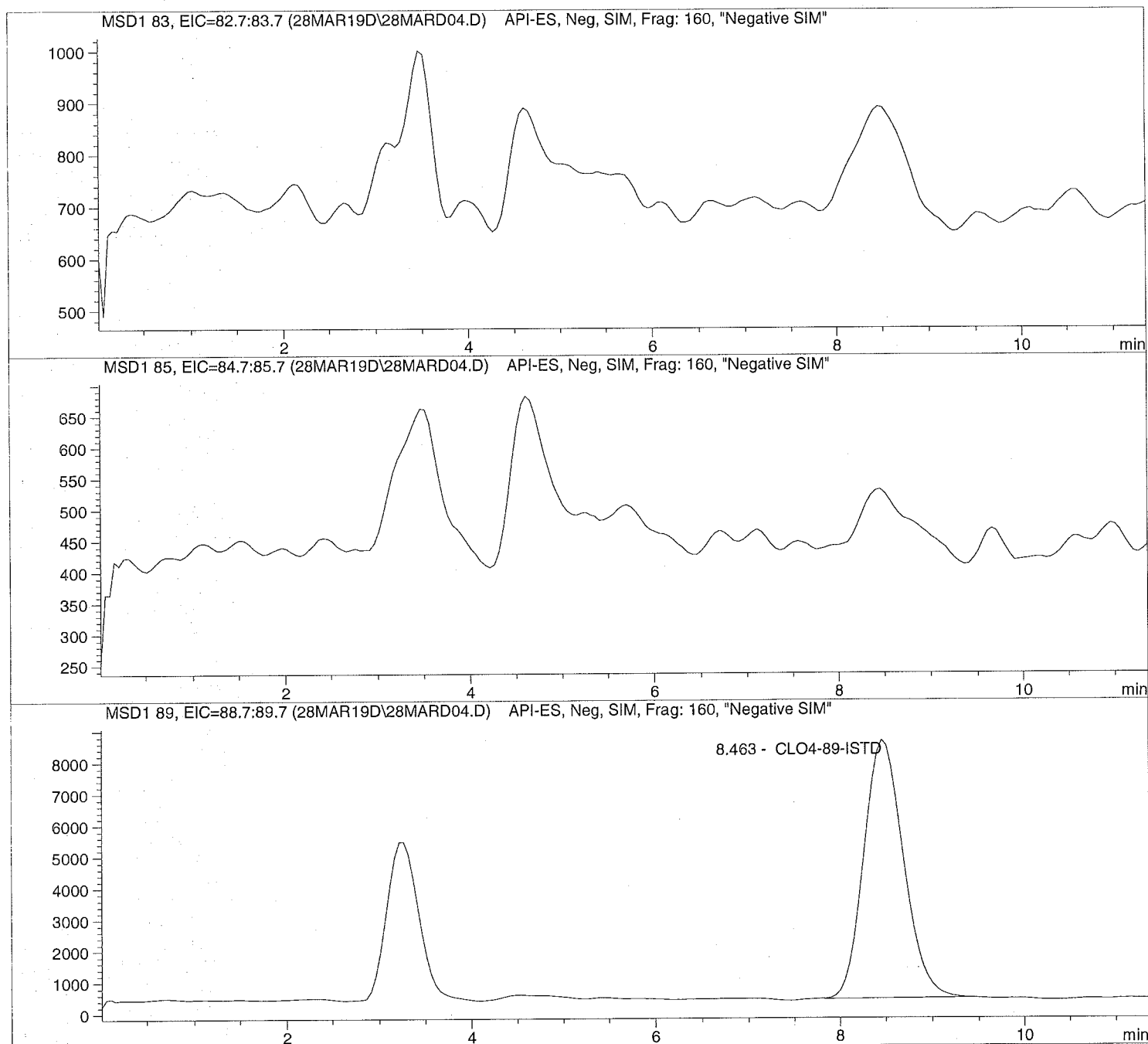
Sample Name: 645533 LMB

Injection Date: 3/28/2019 09:14:08
Sample Name: 645533 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D Sample Name: 645533 LMB

```

=====
Injection Date: 3/28/2019 09:14:08      Seq Line: 4
Sample Name: 645533 LMB                 Location: Vial 74
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.463	BBA	242742.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

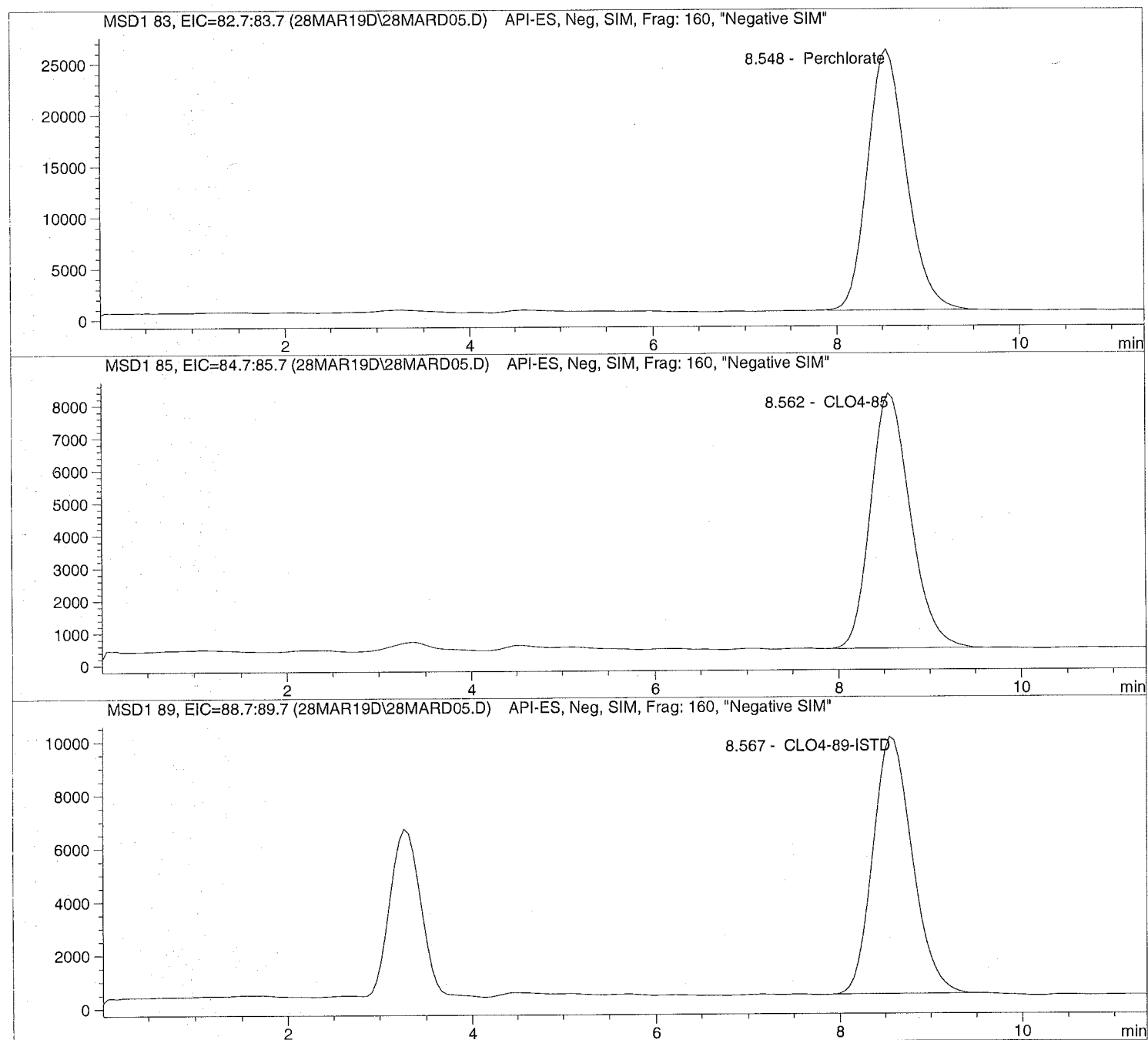
```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```
=====
Injection Date: 3/28/2019 09:28:25      Seq Line:      5
Sample Name:    1907867001 1K           Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```

=====
Injection Date: 3/28/2019 09:28:25      Seq Line:          5
Sample Name:    1907867001 1K           Location:          Vial 75
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1000.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.548	BBA	755115.1	8344.7796	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.562	PBA	238826.8	8762.3570	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.567	PBA	289135.6	5000.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D

Sample Name: 1907869001

Injection Date: 3/28/2019 09:41:37
Sample Name: 1907869001
Acq Operator: TNB

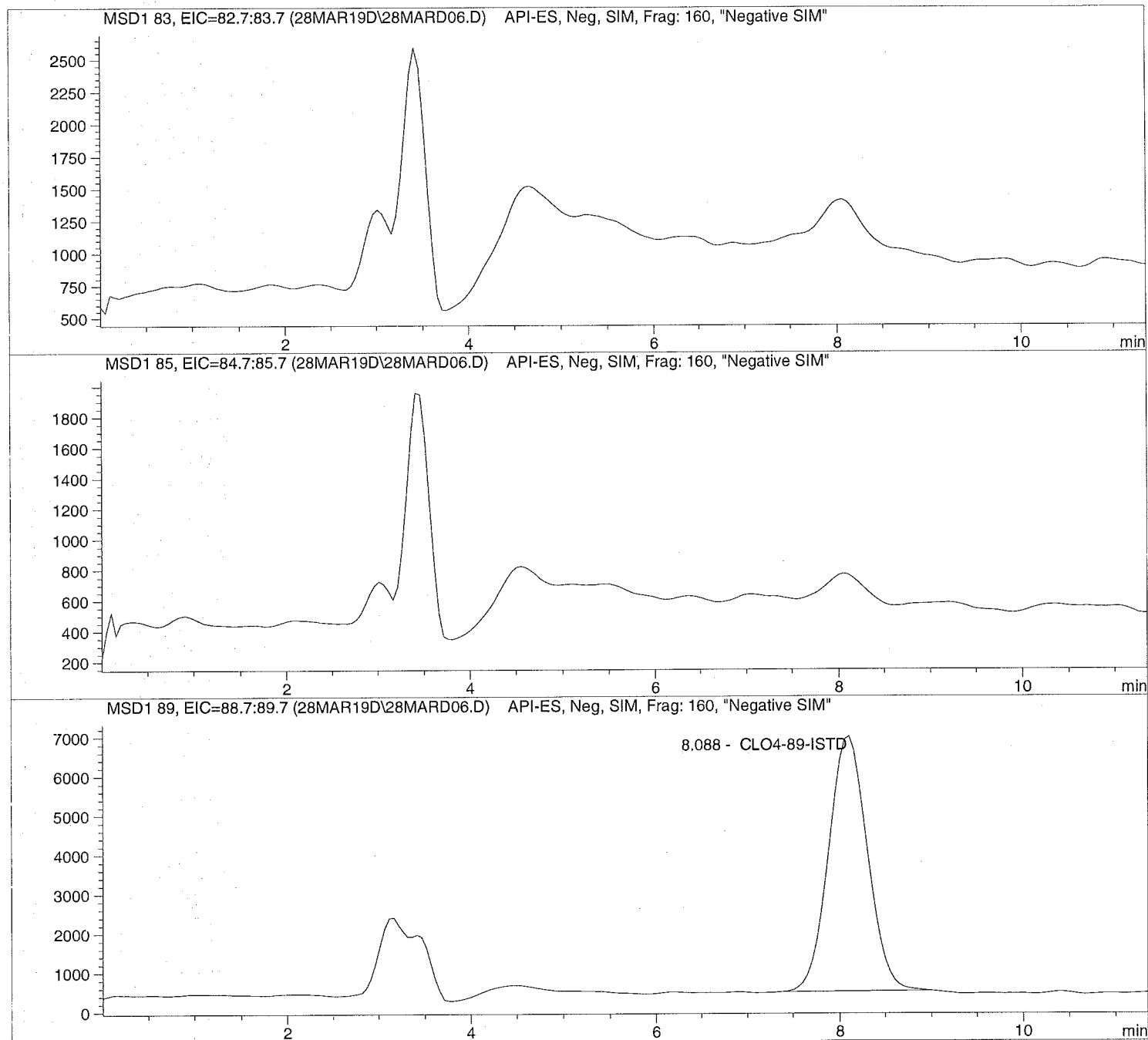
Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D Sample Name: 1907869001

```

=====
Injection Date: 3/28/2019 09:41:37      Seq Line: 6
Sample Name: 1907869001                Location: Vial 76
Acq Operator: TNB                      Inj. No.: 1
                                         Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.088	BBA	189925.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

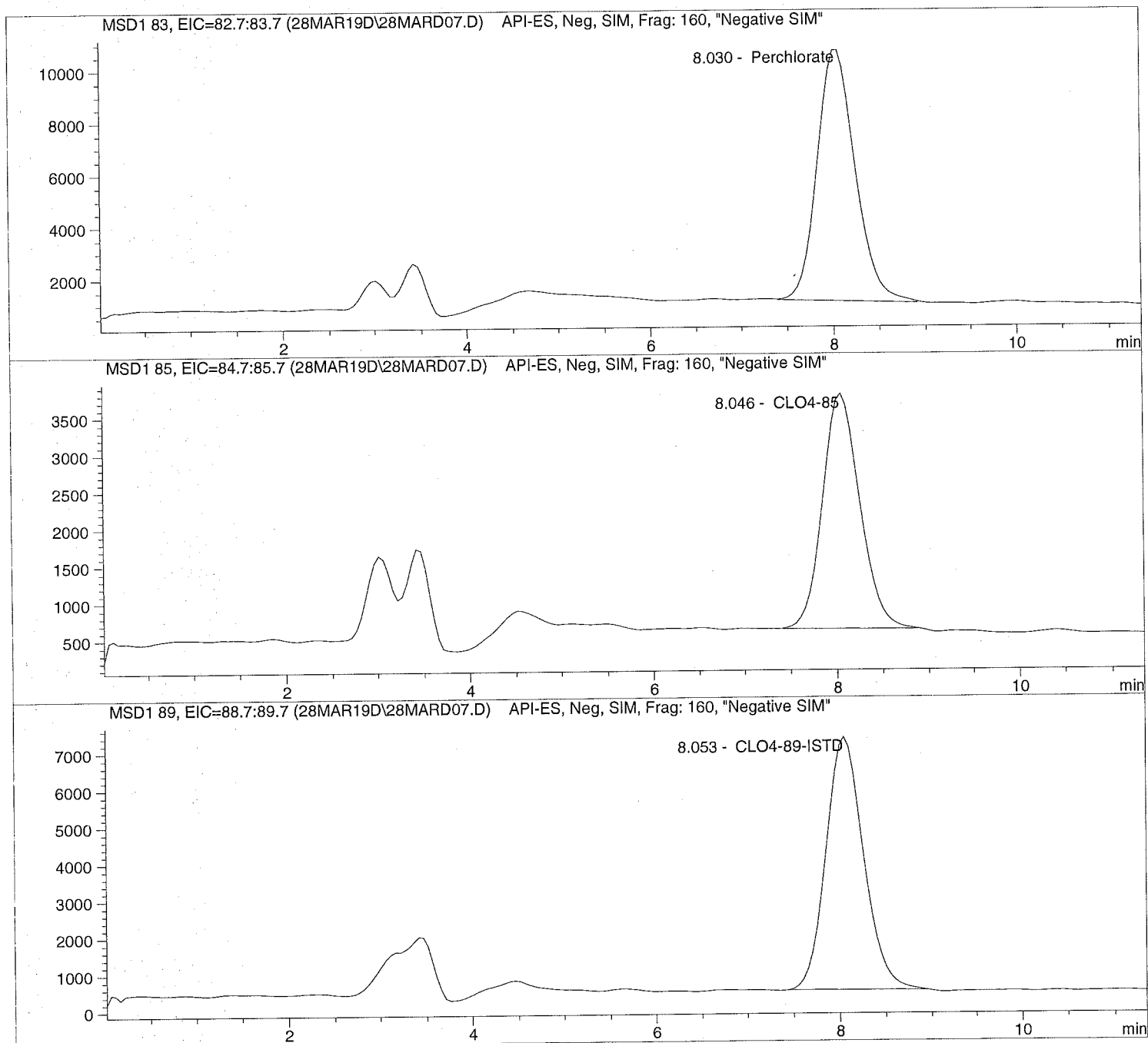
```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D Sample Name: 645535 78691MS

=====
Injection Date: 3/28/2019 09:54:47 Seq Line: 7
Sample Name: 645535 78691MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

=====
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis
=====



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D Sample Name: 645535 78691MS

```

=====
Injection Date: 3/28/2019 09:54:47      Seq Line: 7
Sample Name: 645535 78691MS      Location: Vial 77
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.030	BBA	277046.3	4.6616	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.046	BBA	89339.9	4.9137	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	194412.9	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D

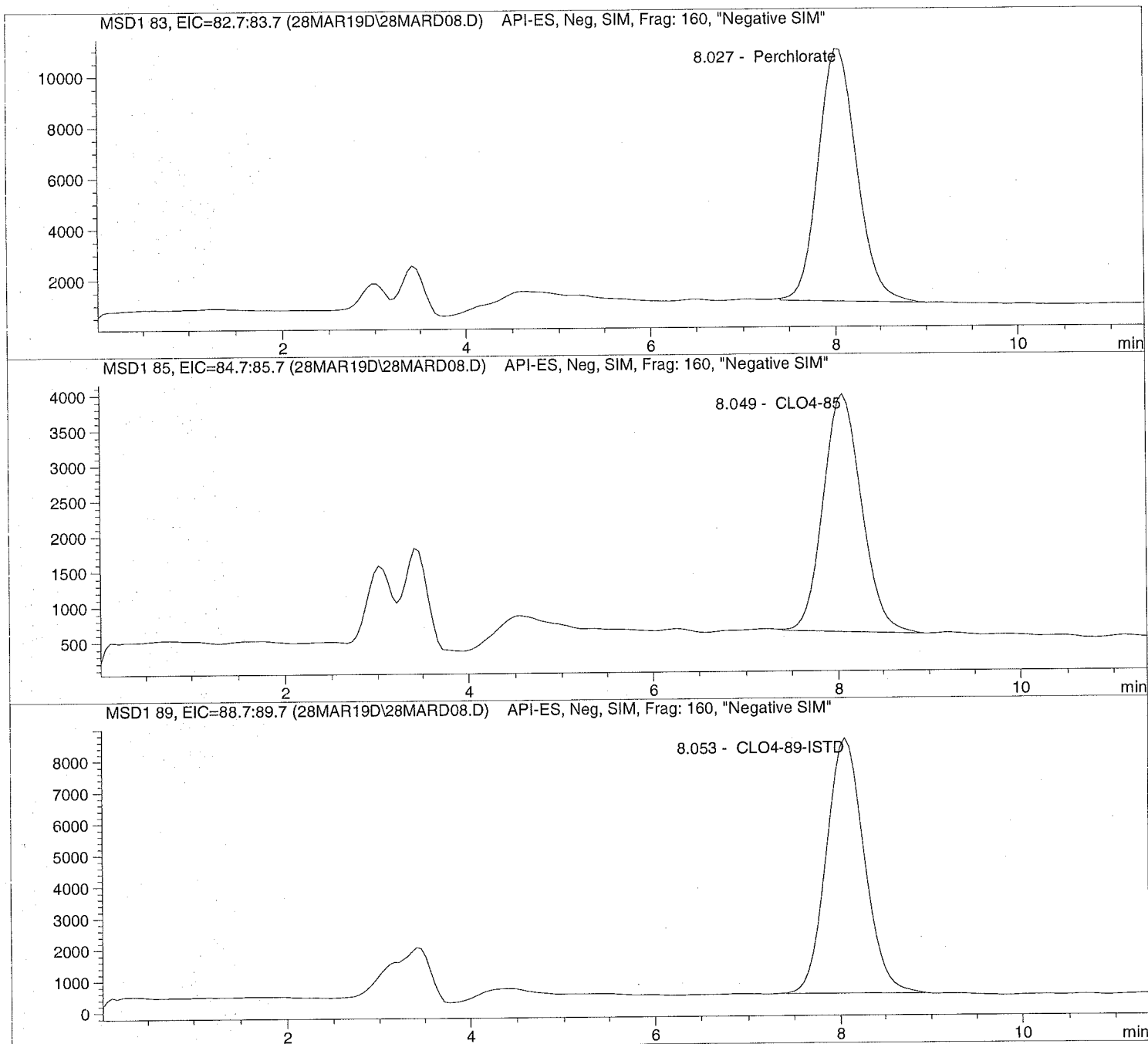
Sample Name: 645536 78691SD

Injection Date: 3/28/2019 10:08:05
Sample Name: 645536 78691SD
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D Sample Name: 645536 78691SD

```

=====
Injection Date: 3/28/2019 10:08:05 Seq Line: 8
Sample Name: 645536 78691SD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.027	BBA	289383.7	4.0513	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.049	BBA	98569.0	4.4846	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	235219.8	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D

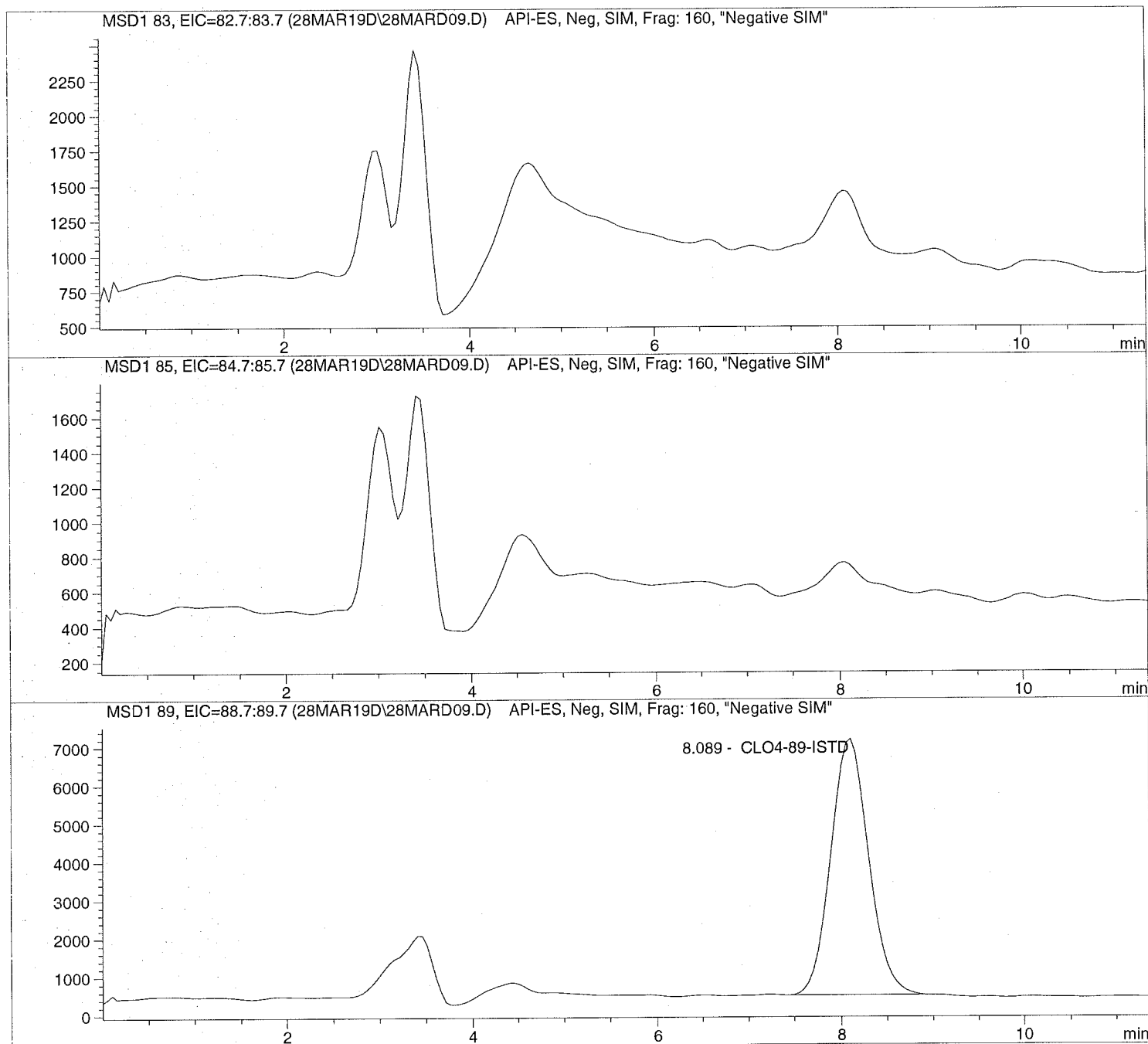
Sample Name: 1907871001

Injection Date: 3/28/2019 10:21:14
Sample Name: 1907871001
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D

Sample Name: 1907871001

```

=====
Injection Date: 3/28/2019 10:21:14      Seq Line: 9
Sample Name: 1907871001                Location: Vial 79
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.089	PBA	189608.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD10.D

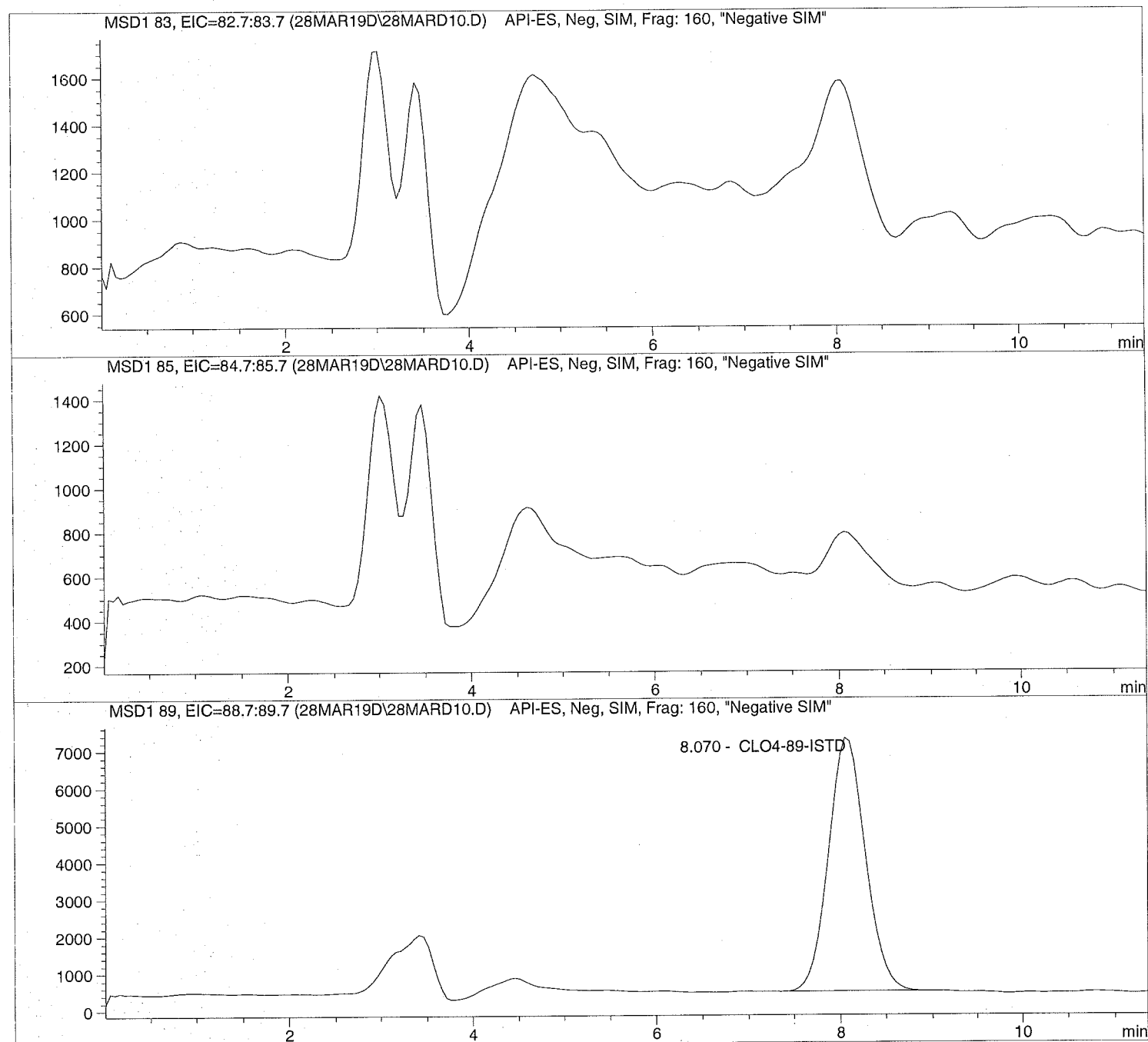
Sample Name: 1908794001

=====
Injection Date: 3/28/2019 10:34:27
Sample Name: 1908794001
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 25 µl

=====
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19\28MARD10.D Sample Name: 1908794001

```

=====
Injection Date: 3/28/2019 10:34:27      Seq Line: 10
Sample Name: 1908794001                Location: Vial 80
Acq Operator: TNB                      Inj. No.: 1
                                         Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.070	PBA	192453.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

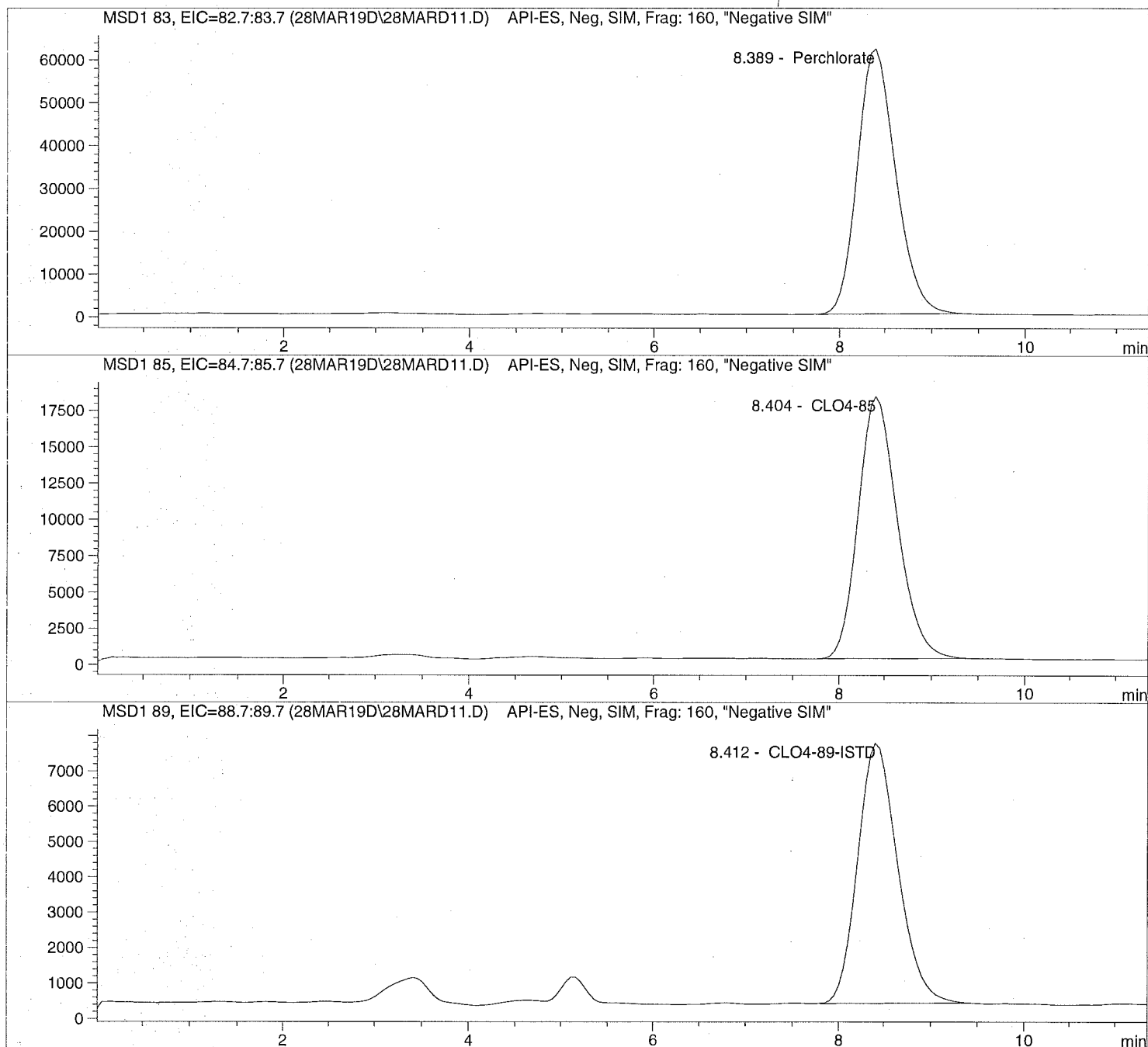
```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D Sample Name: 645537 CCV@25

=====
Injection Date: 3/28/2019 10:49:41 Seq Line: 11
Sample Name: 645537 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl

=====
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D Sample Name: 645537 CCV@25

```

=====
Injection Date: 3/28/2019 10:49:41 Seq Line: 11
Sample Name: 645537 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.389	PBA	1797085.1	24.4912	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.404	PBA	535801.9	24.5911	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.412	BBA	223514.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
 Calibration Table
 =====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
 Based on : Peak Area

Rel. Reference Window : 20.000 %
 Abs. Reference Window : 0.000 min
 Rel. Non-ref. Window : 20.000 %
 Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
 Uncalibrated Peaks : not reported
 Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
 Origin : Ignored (some peaks differ, see below)
 Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
 Average Response : Average all calibrations
 Average Retention Time: Floating Average New 75%

Calibration Report Options :
 Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
 If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
 Signal 2: MSD1 85, EIC=84.7:85.7
 Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl	Amount	Area	Amt/Area	Ref Grp	Name
8.744	1	1.00000	7.76074e4	1.28854e-5	1	Perchlorate
	2	2.00000	1.35273e5	1.47849e-5		
	3	5.00000	3.37764e5	1.48033e-5		
	4	10.00000	6.83454e5	1.46316e-5		
	5	25.00000	2.08433e6	1.19943e-5		
	6	50.00000	4.13334e6	1.20968e-5		
	7	75.00000	5.99313e6	1.25143e-5		
8.755	2	1.00000	2.36780e4	4.22333e-5	1	CLO4-85
	2	2.00000	4.69486e4	4.25998e-5		
	3	5.00000	1.06124e5	4.71147e-5		
	4	10.00000	2.13523e5	4.68335e-5		
	5	25.00000	6.14295e5	4.06971e-5		
	6	50.00000	1.19814e6	4.17315e-5		
	7	75.00000	1.78355e6	4.20509e-5		
8.766	3	5.00000	2.73208e5	1.83011e-5	+I1	CLO4-89-ISTD
	2	5.00000	2.24886e5	2.22335e-5		
	3	5.00000	2.33196e5	2.14412e-5		
	4	5.00000	2.34454e5	2.13262e-5		
	5	5.00000	2.50568e5	1.99547e-5		
	6	5.00000	2.30977e5	2.16472e-5		

RetTime	Lvl	Amount	Area	Amt/Area	Ref Grp Name
[min]	Sig				
7		5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-85

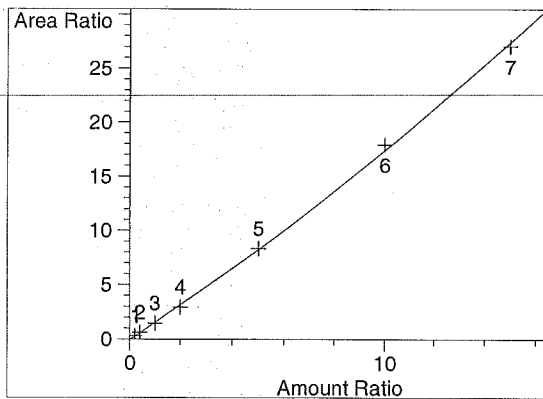
Time Window : From 6.650 min To 12.505 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-89-ISTD

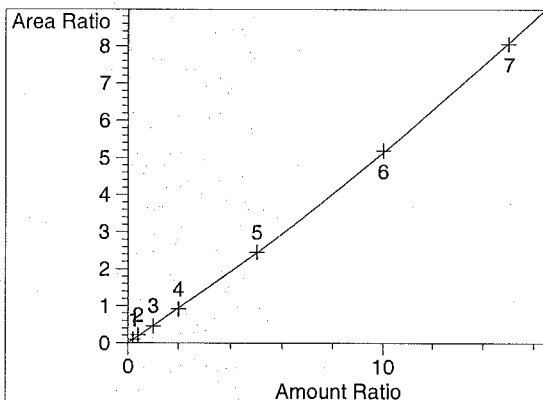
Time Window : From 6.659 min To 12.466 min
Curve Type : Linear
Origin : Included
Calibration Level Weights:/
Level 1 : 1
Level 2 : 1
Level 3 : 1
Level 4 : 1
Level 5 : 1
Level 6 : 1
Level 7 : 1

=====
Peak Sum Table
=====***No Entries in table***
=====

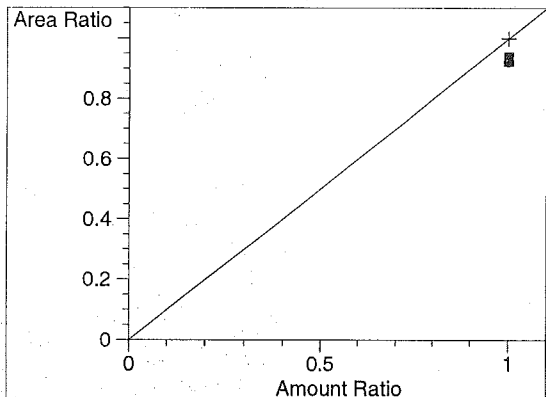
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

##	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
##	-----	-----	---	-----	---	-----	-----	-----
##	CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
##	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
##	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
##	CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
##	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
##	CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
##	CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
##	ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

##	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
##	-----	-----	---	-----	---	-----	-----	-----
##	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
##	CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
##	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
##	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
##	CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
##	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
##	CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
##	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

##	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
##	-----	-----	---	-----	---	-----	-----	-----
##	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
##	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
##	CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
##	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
##	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
##	CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
##	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
##	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

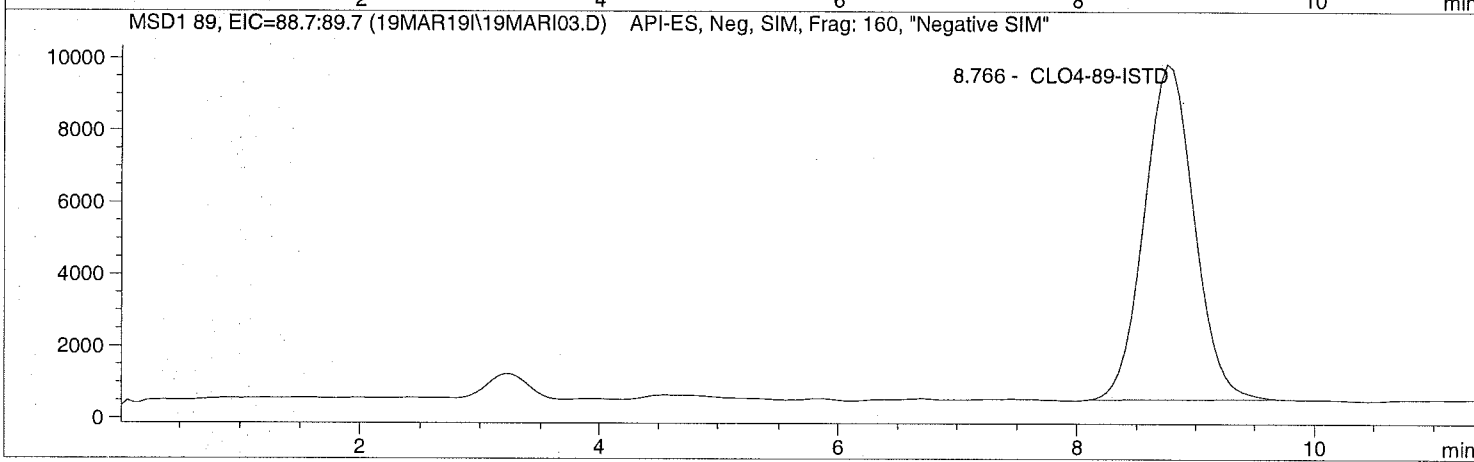
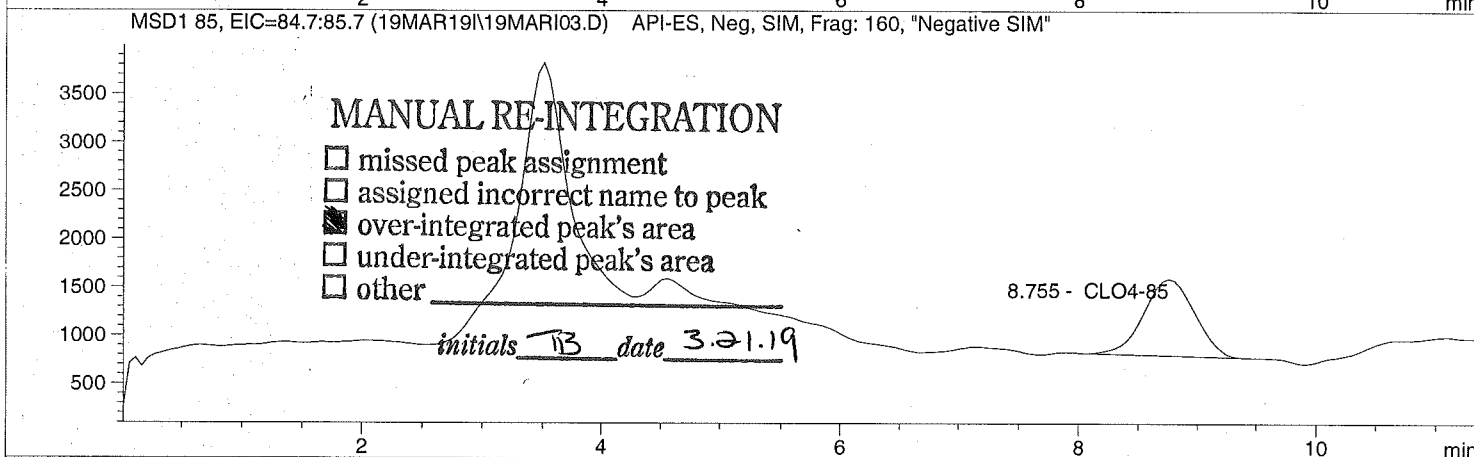
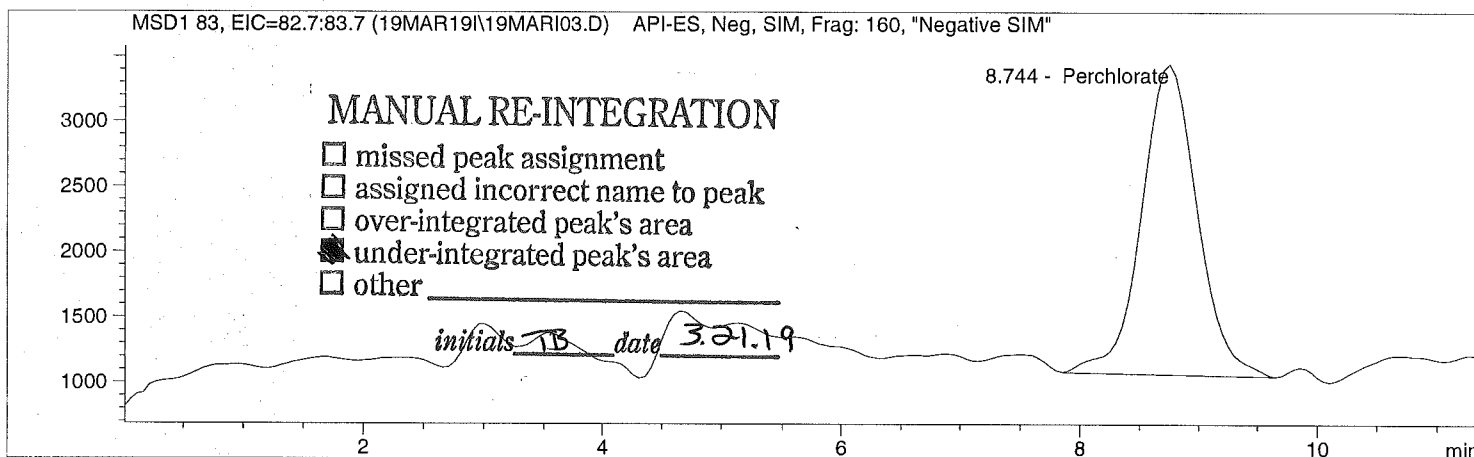
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:          3
Sample Name:    CLO4@ 1.0ug/L           Location:          Vial 73
Acq Operator:  TNB                      Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 1.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D

Sample Name: CLO4@ 2.0ug/L

Injection Date: 3/19/2019 09:53:00

Seq Line: 4

Sample Name: CLO4@ 2.0ug/L

Location: Vial 74

Acq Operator: TNB

Inj. No.: 1

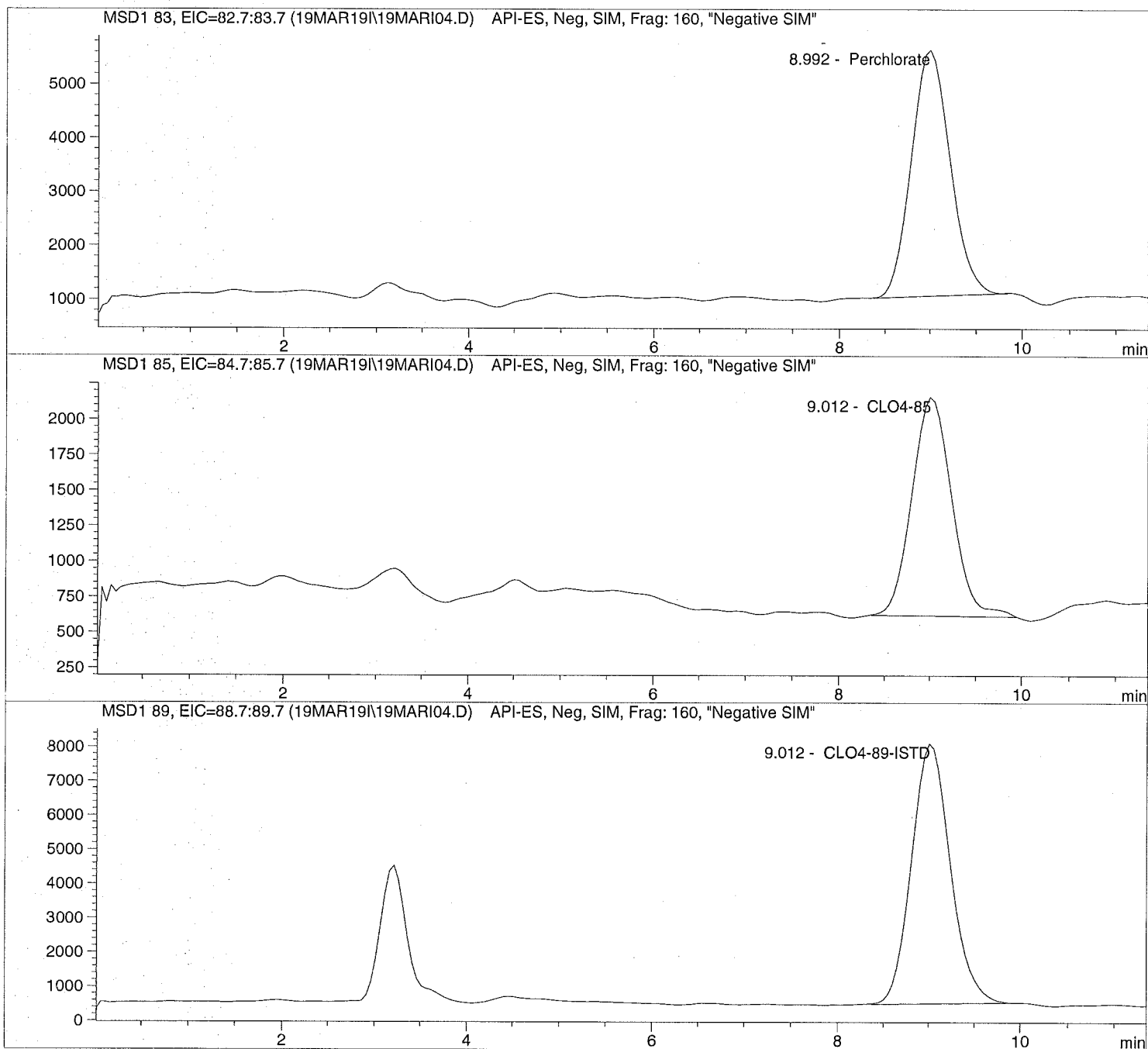
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line:      4
Sample Name:    CLO4@ 2.0ug/L          Location:      Vial 74
Acq Operator:   TNB                   Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI05.D

Sample Name: CLO4@ 5.0ug/L

Injection Date: 3/19/2019 10:06:16

Seq Line: 5

Sample Name: CLO4@ 5.0ug/L

Location: Vial 75

Acq Operator: TNB

Inj. No.: 1

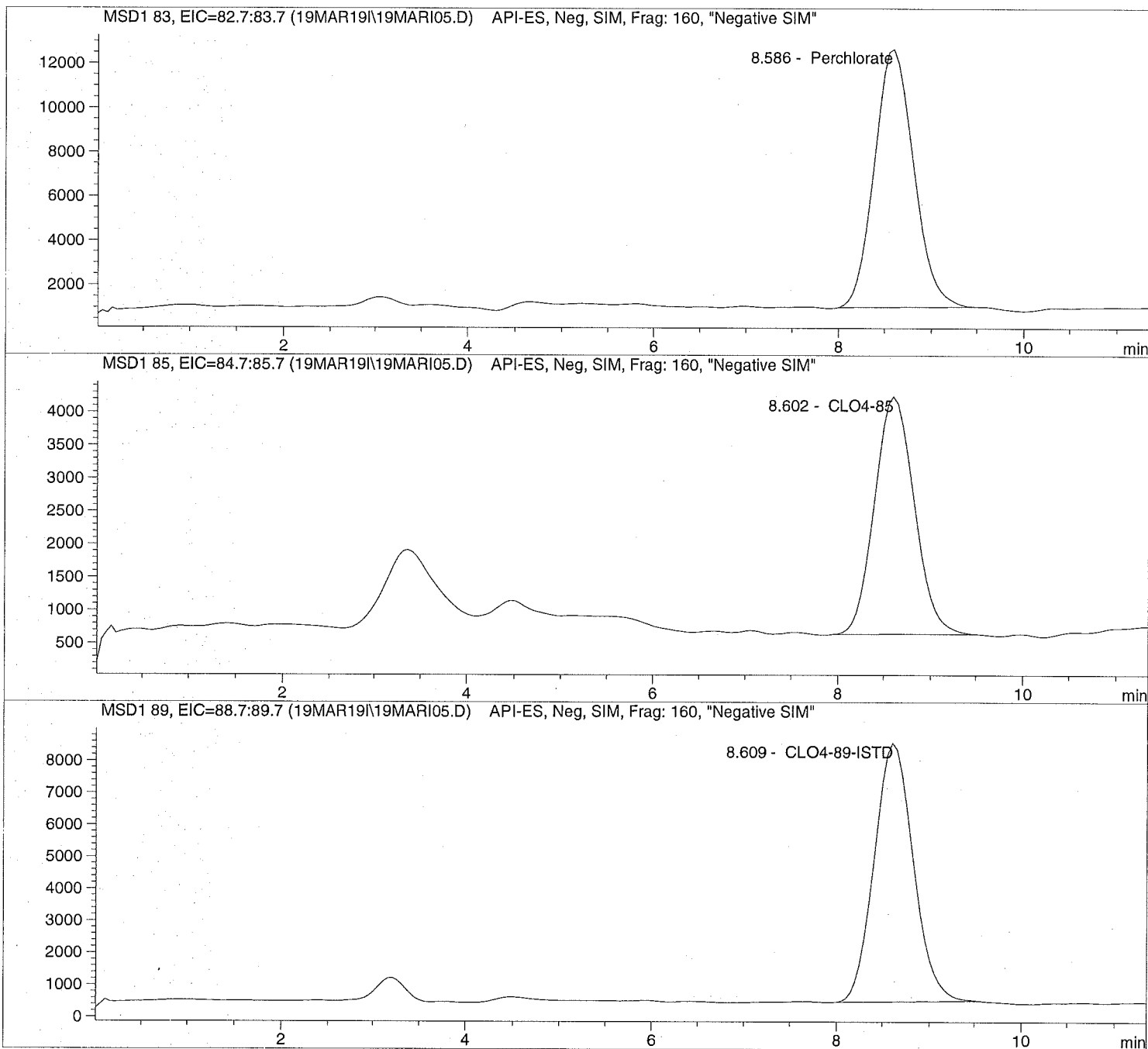
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line:      5
Sample Name:    CLO4@ 5.0ug/L           Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D

Sample Name: CLO4@ 10.ug/L

Injection Date: 3/19/2019 10:19:32

Seq Line: 6

Sample Name: CLO4@ 10.ug/L

Location: Vial 76

Acq Operator: TNB

Inj. No.: 1

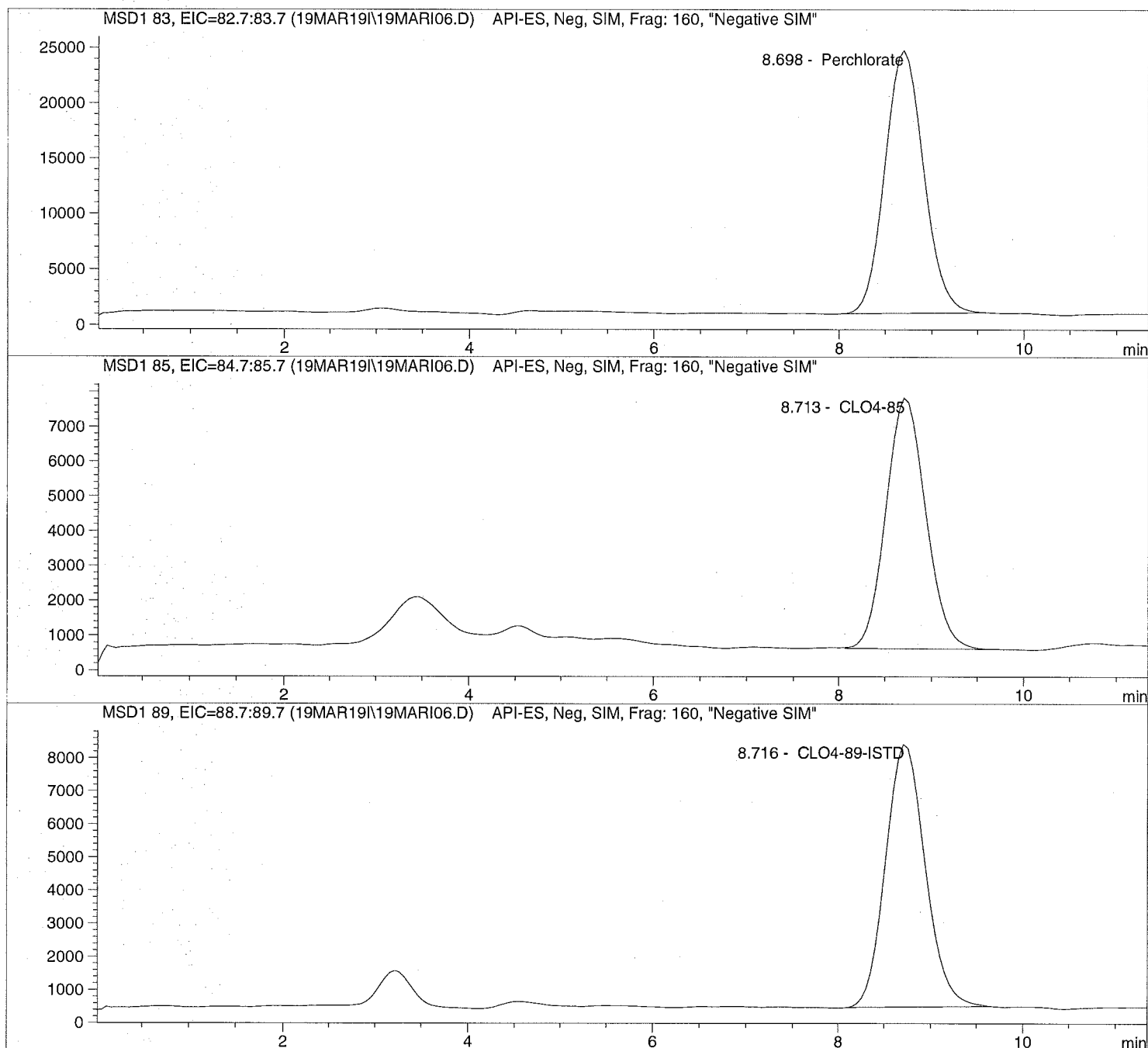
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line:          6
Sample Name:    CLO4@ 10.ug/L           Location:          Vial 76
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D

Sample Name: CLO4@ 25.ug/L

Injection Date: 3/19/2019 10:32:49

Seq Line: 7

Sample Name: CLO4@ 25.ug/L

Location: Vial 77

Acq Operator: TNB

Inj. No.: 1

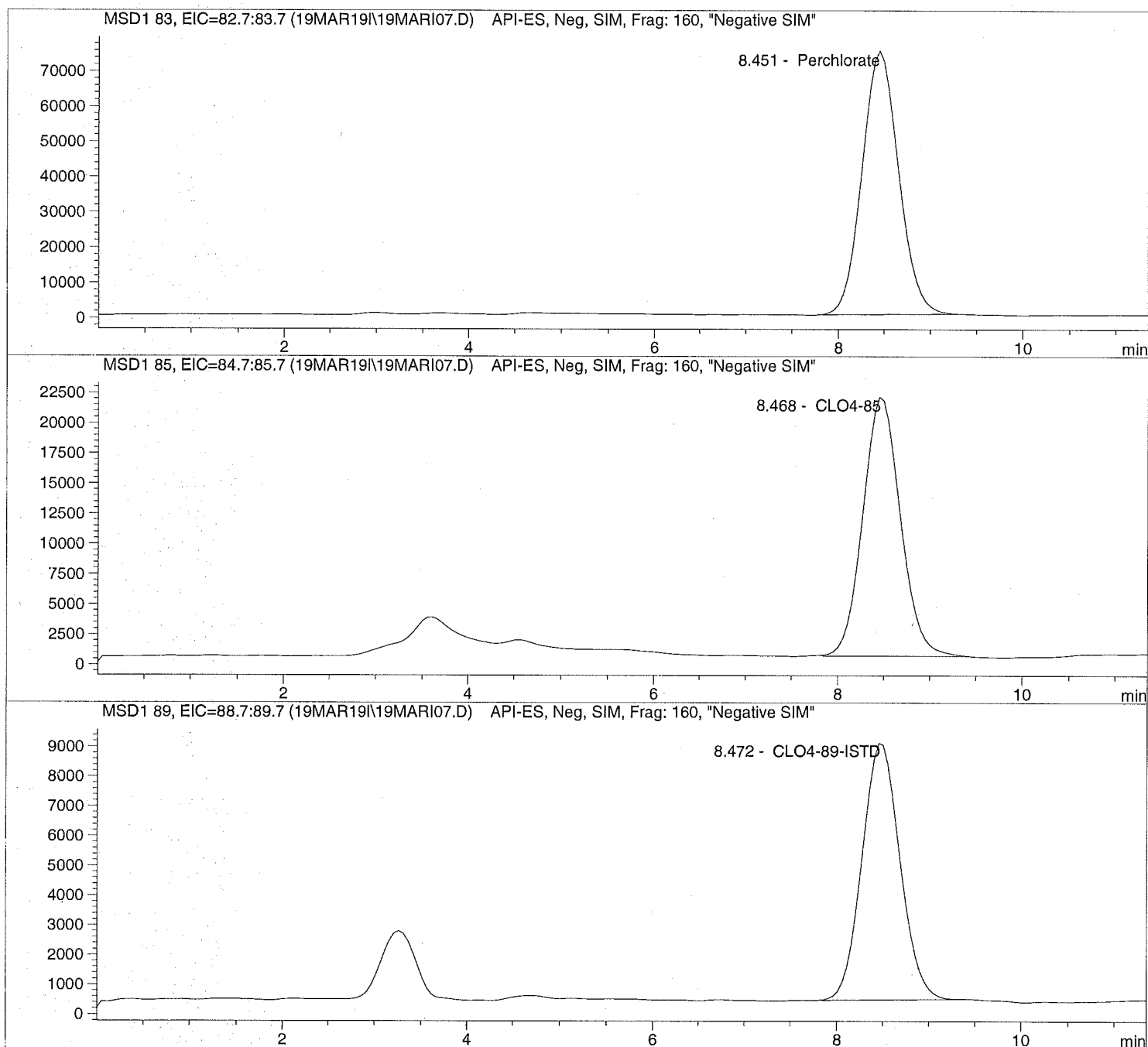
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 3/19/2019 10:32:49      Seq Line:      7
Sample Name:    CLO4@ 25.ug/L           Location:      Vial 77
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI08.D

Sample Name: CLO4@ 50.ug/L

Injection Date: 3/19/2019 10:46:05

Seq Line: 8

Sample Name: CLO4@ 50.ug/L

Location: Vial 78

Acq Operator: TNB

Inj. No.: 1

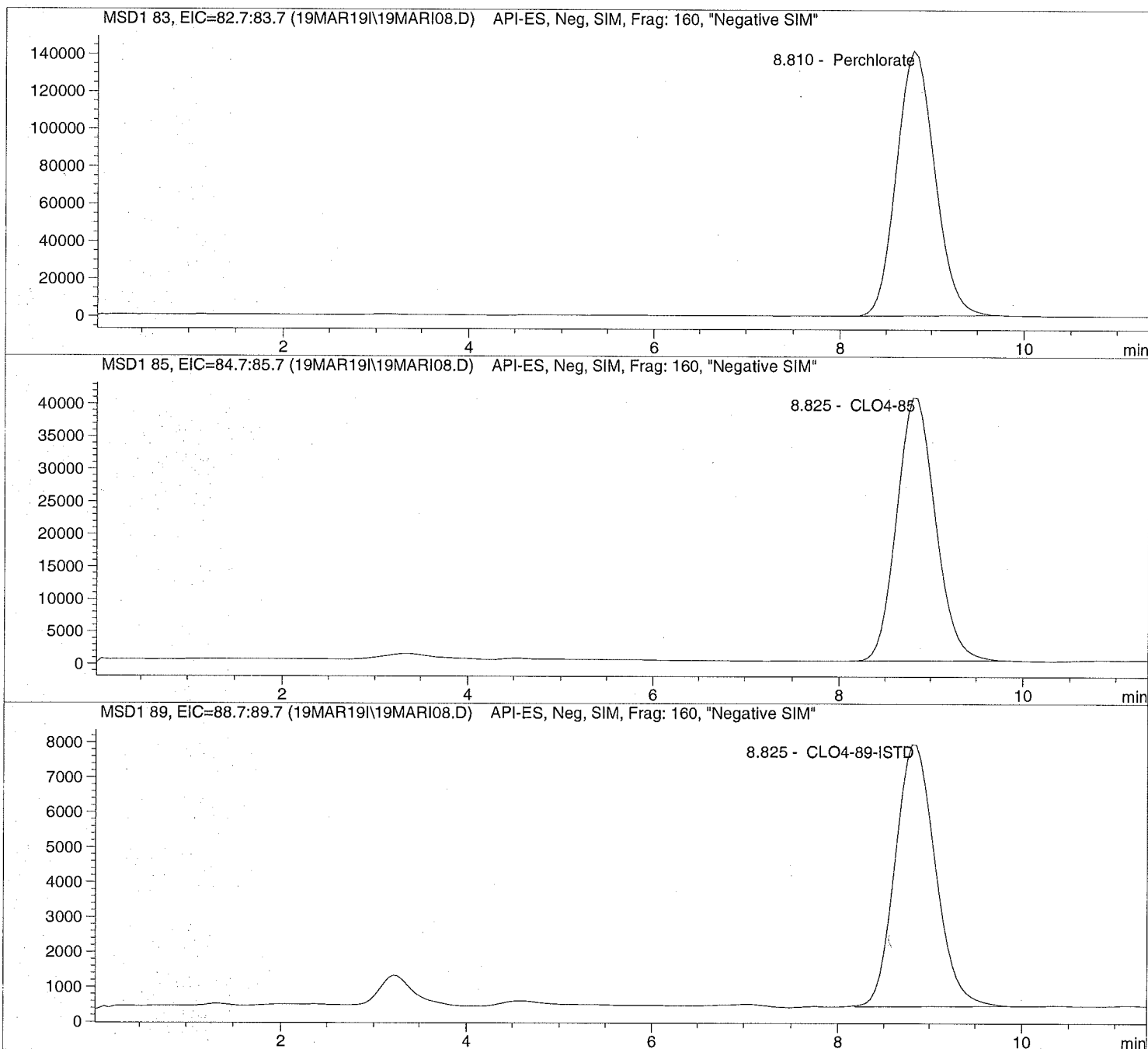
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```
=====
Injection Date: 3/19/2019 10:46:05      Seq Line:            8
Sample Name:    CLO4@ 50.ug/L            Location:            Vial 78
Acq Operator:   TNB                      Inj. No.:            1
                                         Inj. Vol.:            30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis

=====
Sample Information
=====

```
Sorted By:            Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:           1.000000
Dilution:             1.000000
Sample Amount:         50.000
```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

=====
*** End of Report ***
=====

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 3/19/2019 10:59:22

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

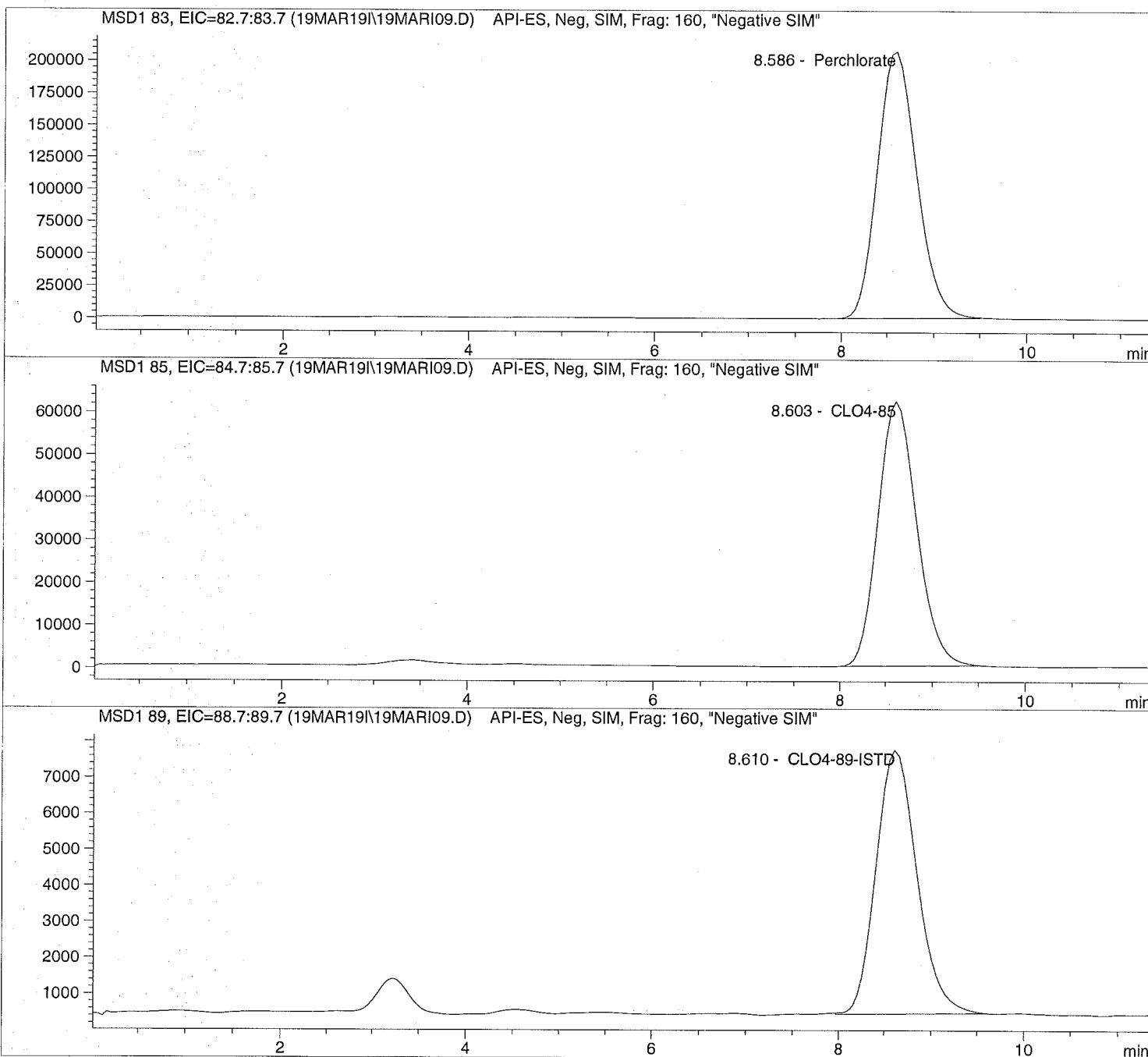
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```
=====
Injection Date:  3/19/2019  10:59:22      Seq Line:           9
Sample Name:    CLO4@ 75.ug/L           Location:           Vial 79
Acq Operator:   TNB                      Inj. No.:          1
                                           Inj. Vol.:         30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:35:22
```

Perchlorate analysis

Sample Information

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:      1.000000
Sample Amount:  75.000
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

*** End of Report ***

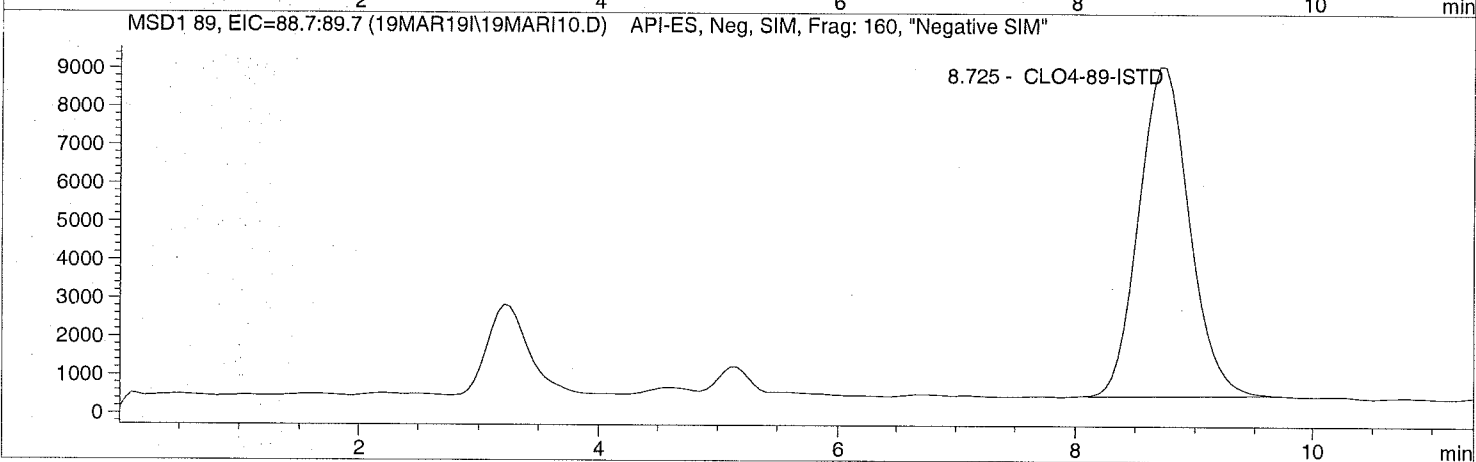
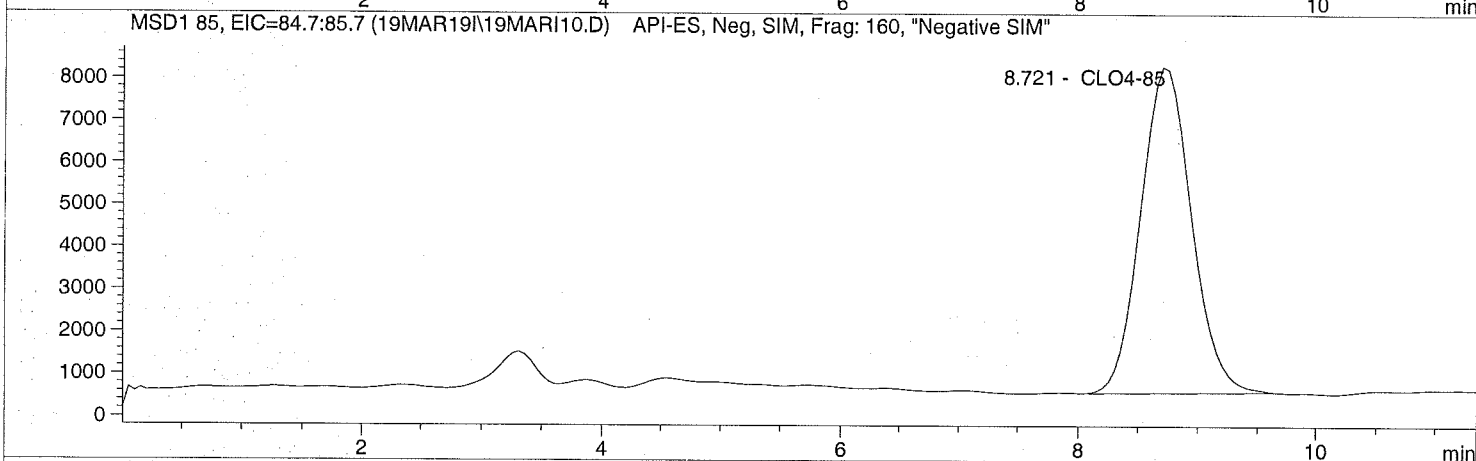
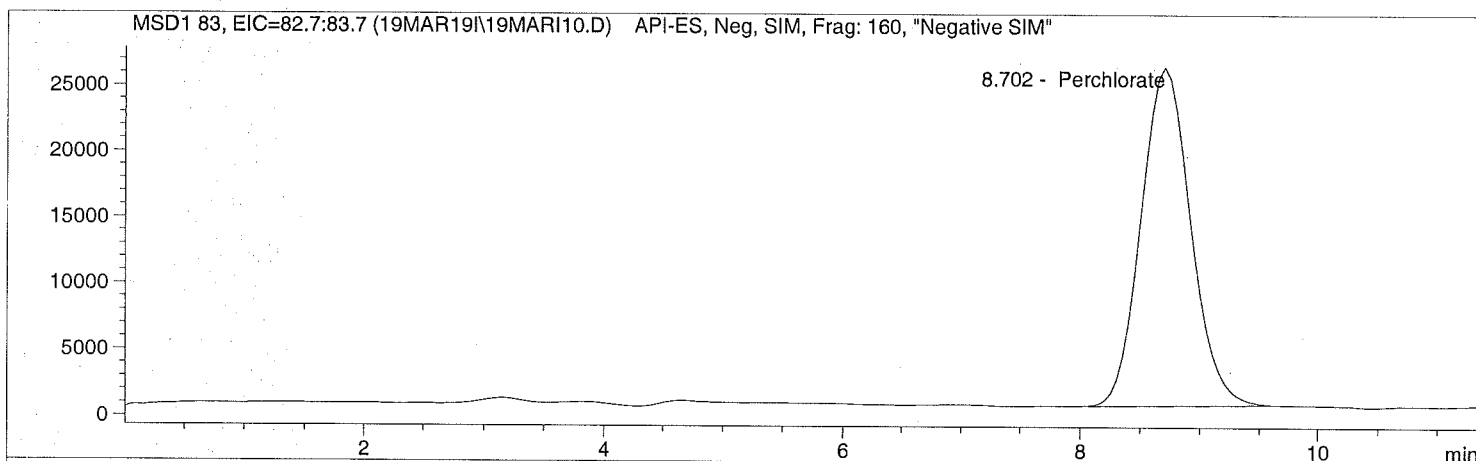
Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

=====
Injection Date: 3/19/2019 11:12:42 Seq Line: 10
Sample Name: ICAL Verf@10ug/L Location: Vial 80
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 3/19/2019 11:12:42      Seq Line:      10
Sample Name:    ICAL Verf@10ug/L        Location:      Vial 80
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

=====
Injection Date: 3/19/2019 09:39:40
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

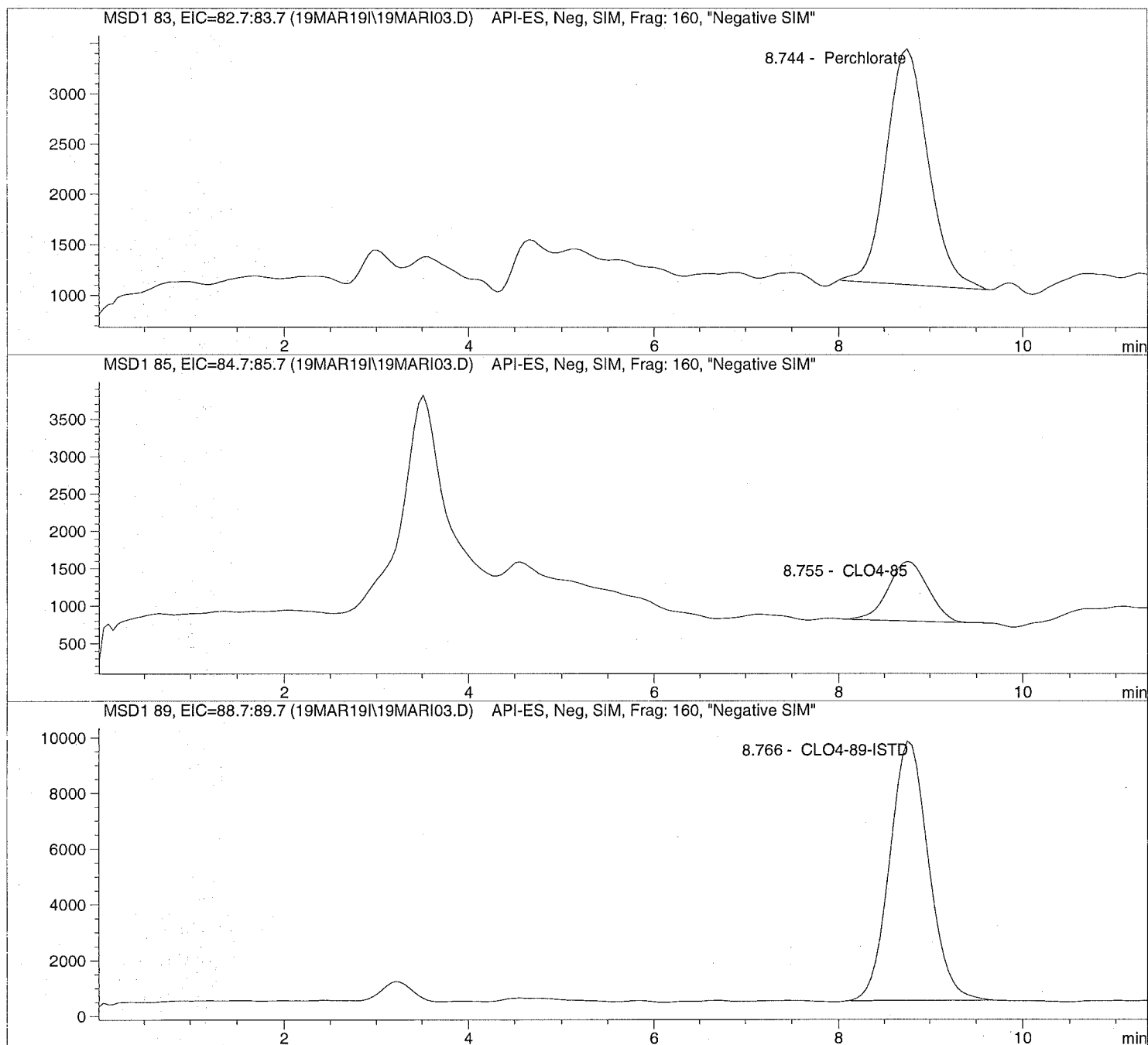
Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 30 µl

Acq Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:          3
Sample Name:    CLO4@ 1.0ug/L          Location:         Vial 73
Acq Operator:   TNB                   Inj. No.:        1
                                           Inj. Vol.:       30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

April 01, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19031160**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 22, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031160

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19031160-01	LH18/24-SP650_032119	Water		21-Mar-2019 14:00	22-Mar-2019 08:52	<input type="checkbox"/>
HS19031160-02	LH18/24-SP650_032119_BIX	Water		21-Mar-2019 14:00	22-Mar-2019 08:52	<input type="checkbox"/>

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE**Work Order Comments**

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
 - The analysis for TOC was subcontracted to ALS Kelso, WA. Final report attached.
-

WetChemistry by Method E350.3**Batch ID: R335466**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

WetChemistry by Method E365.3**Batch ID: R335143**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032119
 Collection Date: 21-Mar-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19031160
 Lab ID:HS19031160-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
AMMONIA AS N BY E350.3(ISE)								Analyst: MZD
	Method:E350.3							
Nitrogen, Ammonia (As N)	16		0.20	0.20	0.20	mg/L	1	28-Mar-2019 10:45
ORTHO PHOSPHATE (PO4) AS P BY E365.3								Analyst: MZD
	Method:E365.3							
Phosphorus, Total Orthophosphate (As P)	4.15		0.100	0.250	0.250	mg/L	10	22-Mar-2019 17:28
SUBCONTRACT ANALYSIS - TOC ANALYSIS								Analyst: SUBK
	Method:NA							
Subcontract Analysis	See Attached		0	0		NA	1	01-Apr-2019 18:17

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032119_BIX
 Collection Date: 21-Mar-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19031160
 Lab ID:HS19031160-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA		Analyst: SUB				
Subcontract Analysis	See Attached		0	0		NA	1	31-Mar-2019 12:13

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031160

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R335143	Test Name : ORTHO PHOSPHATE (PO4) AS P BY E365.3		Matrix: Water			
HS19031160-01	LH18/24-SP650_032119	21 Mar 2019 14:00			22 Mar 2019 17:28	10
Batch ID R335466	Test Name : AMMONIA AS N BY E350.3(ISE)		Matrix: Water			
HS19031160-01	LH18/24-SP650_032119	21 Mar 2019 14:00			28 Mar 2019 10:45	1
Batch ID R335660	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19031160-02	LH18/24-SP650_032119_BIX	21 Mar 2019 14:00			31 Mar 2019 12:13	1
Batch ID R335735	Test Name : SUBCONTRACT ANALYSIS - TOC ANALYSIS		Matrix: Water			
HS19031160-01	LH18/24-SP650_032119	21 Mar 2019 14:00			01 Apr 2019 18:17	1

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031160

QC BATCH REPORT NEW

Batch ID:	R335143 (0)	Instrument:	UV-2450	Method:	ORTHO PHOSPHATE (PO4) AS P BY E365.3					
MBLK	Sample ID: MBLK-335143	Units:	mg/L	Analysis Date:	22-Mar-2019 17:28					
Client ID:	Run ID: UV-2450_335143	SeqNo:	5003495	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	0.0250	0.0250								U
LCS	Sample ID: LCS-335143	Units:	mg/L	Analysis Date:	22-Mar-2019 17:28					
Client ID:	Run ID: UV-2450_335143	SeqNo:	5003496	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	0.233	0.0250	0.25	0	93.2	85 - 115				
MS	Sample ID: HS19031160-01MS	Units:	mg/L	Analysis Date:	22-Mar-2019 17:28					
Client ID: LH18/24-SP650_032119	Run ID: UV-2450_335143	SeqNo:	5003498	PrepDate:	DF: 10					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	6.4	0.250	2.5	4.15	90.0	80 - 120				
MSD	Sample ID: HS19031160-01MSD	Units:	mg/L	Analysis Date:	22-Mar-2019 17:28					
Client ID: LH18/24-SP650_032119	Run ID: UV-2450_335143	SeqNo:	5003499	PrepDate:	DF: 10					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	6.51	0.250	2.5	4.15	94.4	80 - 120	6.4	1.7	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031160

QC BATCH REPORT NEW

Batch ID:	R335466 (0)	Instrument:	WetChem_HS	Method:	AMMONIA AS N BY E350.3(ISE)					
MBLK	Sample ID: MBLK-335466	Units:	mg/L	Analysis Date:	28-Mar-2019 10:45					
Client ID:	Run ID: WetChem_HS_335466	SeqNo:	5010825	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	0.20	0.20								U
LCS	Sample ID: LCS-335466	Units:	mg/L	Analysis Date:	28-Mar-2019 10:45					
Client ID:	Run ID: WetChem_HS_335466	SeqNo:	5010826	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	8.922	0.20	10	0	89.2	80 - 120				
MS	Sample ID: HS19031270-01MS	Units:	mg/L	Analysis Date:	28-Mar-2019 10:45					
Client ID:	Run ID: WetChem_HS_335466	SeqNo:	5010828	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	10.01	0.20	10	0.5221	94.9	80 - 120				
MSD	Sample ID: HS19031270-01MSD	Units:	mg/L	Analysis Date:	28-Mar-2019 10:45					
Client ID:	Run ID: WetChem_HS_335466	SeqNo:	5010829	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Nitrogen, Ammonia (As N)	9.997	0.20	10	0.5221	94.7	80 - 120	10.01	0.13	20	

The following samples were analyzed in this batch: HS19031160-01

ALS Houston, US

Date: 01-Apr-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19031160	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

ALS Houston, US

Date: 01-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031160

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19031160-01	LH18/24-SP650_032119	Login	3/22/2019 4:19:21 PM	JRM	WET069
HS19031160-01	LH18/24-SP650_032119	Login	3/22/2019 4:19:21 PM	JRM	WET069
HS19031160-01	LH18/24-SP650_032119	Login	3/22/2019 4:19:21 PM	JRM	Sub
HS19031160-02	LH18/24-SP650_032119_BIX	Login	3/22/2019 4:19:21 PM	JRM	Sub

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19031160

Date/Time Received: **22-Mar-2019 08:52**
 Received by: **NDR**

Checklist completed by: Jared R. Makan 22-Mar-2019
 eSignature Date

Reviewed by: Corey Grandits 26-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 0.7c/0.7c UC/C IR11
 Cooler(s)/Kit(s): 43950
 Date/Time sample(s) sent to storage: 03/22/2019 16:25

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

CHAIN OF CUSTODY


Name Of Lab Shipping To: ALS 10450 Stancliff Rd. Suite 210 Houston, TX, 77099 (281) 530-5656 ATTN: R.J Modshia

Page 1 of 1

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001			Analyses										Remarks (Preservatives, etc.)	Lab I.D.#
Job: GROUNDWATER TREATMENT PLANT WEEKLY SAMPLES																	
Prepared By: Scott Beesinger						P.O. Number											
Field Sample I.D.	Sample Matrix	Date / Time	MS / MSD	No. OF CONTAINERS	AMMONIA-N	TOTAL ORGANIC CARBON	ORTHO-PHOSPHATE	PERCHLORATE									
LH18/24-SP650_032119	Water	03/21/19 / 14:00		2	X	X											H2SO4
LH18/24-SP650_032119	Water	03/21/19 / 14:00		1			X										NONE
LH18/24-SP650_032119_BIX	Water	03/21/19 / 14:00		1				X									NONE
Additional Remarks: Standard TAT on all parameters																	
Relinquished By: <i>Scott Beesinger</i>		Date 03/21/19	Time 14:30	Received By:		Date	Time	Relinquished By:		Date	Time	Received By: N/A		Date 3/21/19	Time 08:52		
For Lab Use Only																	
Received At Lab By:				Date	Time	Airbill No.	Opened By:			Date	Time	Temp of Container	Seal No.	Condition			
Remarks: HS19031160 Bhate Environmental Associates, Inc. 18/24 Longhorn GW Treatment Plant Weekly Sampl 43950 Temp 41c 0.7 12/11																	

(Word) S:\1-ces\Forms\Chain of Custody - Bi\Weekl



 ALS Environmental 10450 Standliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CUSTODY SEAL		Seal Broken By:
	Date: 3/22/19	Time: 1430	RM
	Name: Scott Beesinger	Company: BH&T	Date: 03/22/19

43950 MAR 22 2019

Must Deliver Next Business Day
Time and Temperature Sensitive!



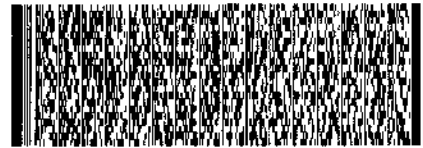
ORIGIN ID: SGRA (303) 597-2450
 SCOTT BEESINGER
 BH&T ENVIRONMENTAL ASSOCIATES
 1203-B EAST GRAND AVE. PMB202
 MARSHALL, TX 75670
 UNITED STATES US

917
 B03
 10:30 A
 3100
 03/22

TO CLIENT SERVICES
 ALS LABORATORY GROUP
 10450 STANCLIFF ROAD
 SUITE 210
 HOUSTON TX 77099

(281) 630-6668
 REF: LHAAP-58-BD 64113-RJ

RMA: ||| ||| |||



RETURNS MON - SAT

FedEx
 TRK# 4809 7831 3400
 10721

FRI - 22 MAR 10:30A
PRIORITY OVERNIGHT

AB SGRA

77099
 TX-US
 IAH



FTD 162785 21MAR19 66GA 55311/4603/NCBA



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1907867; 1907869; 1907871;
1908794

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2230 (235529)

General Set Information: There were four field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Method QC data: The method blank (LMB 645533) was less than 1/2 the CRDL. The recovery for the LCS (645534) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.µg/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in µg/L. Results were calculated in µg/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (µg/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level of 4.0µg/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch March 29, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 29, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1908794**

Project ID: HS19031160

Purchase Order: HS19031160

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP650_032119_BIX	1908794001	03/21/19	03/26/19	



ANALYTICAL REPORT

Workorder: 34-1908794

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP650_032119_BIX	Sampling Site: NA	Collected: 03/21/2019				
Lab ID: 1908794001	Media: 125 mL Nalgene	Received: 03/26/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2230 (HBN: 235529) Analyzed: 03/28/2019 10:34	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 235529)

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/28/2019 14:08	/S/ Stephen Brose 03/29/2019 12:58

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1908794

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935505

Analysis Information

Workorder: 1908794

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2230 (HBN: 235529)
Analyzed By: Thomas Bosch

Blank

LMB: 645533 Analyzed: 03/28/2019 09:14 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 645534 Analyzed: 03/28/2019 08:47 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.00	4.00	99.9	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1907869001 Analyzed: 03/28/2019 09:41 Dilution: 1 Units: ug/L		MS: 645535 Analyzed: 03/28/2019 09:54 Dilution: 1 Units: ug/L				MSD: 645536 Analyzed: 03/28/2019 10:08 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	ND	4.66	4	117	78.8 123.8	4.05	101	14	0.0 20.0

Comments

Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/28/2019 14:23	/S/ Stephen Brose 03/29/2019 12:58

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



1908794

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10960

18698/#2
1908794

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19031160
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19031160-02	LH18/24-SP650_032119_BIX	Water	21 Mar 2019 14:00
SUB_Perch-6850			01 Apr 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: _____
 Received By: _____
 Cooler ID(s): _____

Date/Time: 3/25/19 1800
 Date/Time: 3-26-19 10:05
 Temperature(s): _____

RIGHT SOLUTIONS | RIGHT PARTNER

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

W32719
1908794

Client Name: ALS HOUSTON Project/Task/Site: HS19031160
 Date/Time of Receipt: 3-26-19 10:05 Number of Coolers Received: 1

Condition of Coolers: Acceptable/Unacceptable Temperature Control: Present/Not Included
 Cooler Custody Seals: Present/Absent/NA Location Temp Taken: Control/Between Samples
 Container Custody Seals: Present/Absent/NA Intact/Broken/NA
 Intact/Broken/NA
 Ice Present: Yes/No/NA Are all temperatures within project specific guidelines? Yes/No/NA
Frozen/Melted/NA VOA Headspace Present? Yes/No/NA

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9266</u>	1 °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: *Gayleen Coates* Signature GAYLEEN COATES Printed Name 3-26-19 Date

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

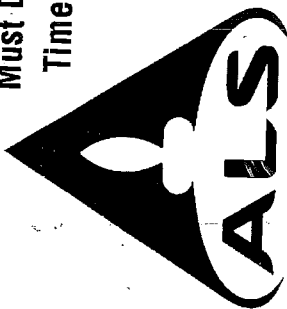
PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature

Part # 159460-434 RIT2 EXP 11/19

SS1CL/45D3/1043

**Must Deliver Next Business Day
Time and Temperature Sensitive!**



ORIGIN ID: S6RA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

SHIP DATE: 25MAR19
ACT WT: 6.90 LB
CAD: 300130/CAFE3211
DIRS: 14X11X10 IN
BILL THIRD PARTY

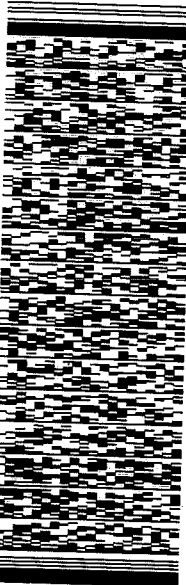
**TO SAMPLE RECEIVING
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE**

SALT LAKE CITY UT 84123

(801) 286-7700

REF: HS19031160 - RJ

FedEx
Express



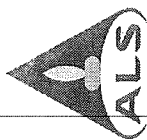
**TUE - 26 MAR 3:00P
STANDARD OVERNIGHT**

TRK# 4809 7832 0971

AX BTFA

84123
UT-US SLC





Batch Worklist

Batch: ELMS/ 2230 **Created:** 3/28/2019 07:45 **Instrument:** HBN: 235529
Rule: EPA 6850, DoD QSM Water **Analyst:** T. Bosch **Status:** WP



Workorder: 1907867 [ENV_LVL4]
Workorder: 1907869 [ENV_LVL4]
Workorder: 1907871 [ENV_LVL4]
Workorder: 1908794 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	64530	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	
2	64531	RLV'S for HBN 235529 [ELMS/2230]				RLV'S	3		E685041C3Q	5311		4/1/2019	
3	64532	ICS for HBN 235529 [ELMS/2230]				ICS	3		E6850.D3Q	5311		4/1/2019	
4	64533	LMB for HBN 235529 [ELMS/2230]				LMB	3		E6850Q413Q	5311		4/1/2019	
5	64534	LCS for HBN 235529 [ELMS/2230]				LCS	3		E6850Q413Q	5311		4/1/2019	
6	1907867001	LH18/24-SP140_031419				SAMPLE	3	1907867001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
7	1907869001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907869001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
8	64535	LH18/24-SP650...(1907869001MS)				MS	3		E6850Q413Q	5311		4/1/2019	
9	64536	LH18/24-SP65...(1907869001MSD)				MSD	3		E6850Q413Q	5311		4/1/2019	
10	1907871001	LH18/24-SP650_031419_BIX				SAMPLE	3	1907871001-A	E6850Q41.3	5480	4/11/2019	4/1/2019	
11	1908794001	LH18/24-SP650_032119_BIX				SAMPLE	3	1908794001-A	E6850Q41.3	5480	4/18/2019	4/8/2019	
12	64537	CCV for HBN 235529 [ELMS/2230]				CCV	3		E685041C3Q	5311		4/1/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1907867 (001); 1907869 (001); 1907871 (001); 1908794 (001)

ELMS Batch/HBN ID: 2230 (235529)

Prep Date: 03/27/2019 Analysis Date: 03/28/2019 Analyst: T. Bosch

Analyte: **Perchlorate** Matrix: **Water** Method: **6850**

Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\28MAR19D.s

Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by **TNB**. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 25µL
Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 645534; Target = 4.0µg/L. ASTM type II water was used for LMB 645533.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1907869001 (Client ID's: LH18/24-SP650_031419_BIX). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field sample 1907867001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\235529-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 645531) is reported from the analysis of the Laboratory Control Sample (LCS – 645534) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
<u>Batch(es)/SDG: E LMS: 2230 HBN: 235529</u>		
<u>Sample Set IDs if Applicable: 1907867/1907869/1907871/1908794</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK		Description - 6850 WKG Std 100.ug/L			
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK		Description - 6850 Stock AccStd 1,000ug/mL	
Standard: 43659	Created By: Thomas Bosch	Amount: 100 mL	
MFG: AccuStandard	Create Date: 09/17/2018 09:09AM	Expires: 07/25/2020	
MFG Lot: 218065075		Usable: No	
Part ID: IC-PER-10X-1		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT		Description - 6850 Intermdt AccStd 10.ug/mL			
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK		Description - 6850 QC WKG STD 100ug/L			
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte:	Name:	Concentration:
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK		Description: 6850 QC Stock STD 1,000ug/mL	
Standard: 36748		Created By: Thomas Bosch	
MFG: Ultra Scientific		Amount: 100 mL	
MFG Lot: CP-0860		Create Date: 05/11/2017 01:05PM	
Part ID: ICC-013		Expires: 03/31/2020	
		Usable: Yes	
		Lab Lot: CLO4 QC STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

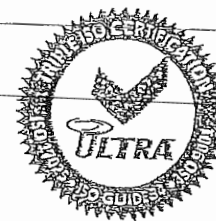
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:

This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, <50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis



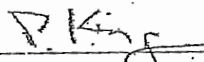
ISO Guide 34 Reference Material

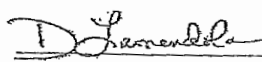
Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:

The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QA/RA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:
ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$

Labeled CAS Number: NA

Unlabeled CAS Number: 7601-89-0

MW*: 130.4

Chemical Formula: $\text{NaCl}^+\text{O}_4^-$

Storage: Store at room temperature away from light and moisture.

Stability: See storage and expiration data.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 $\mu\text{g/mL}$
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 \pm 2.8 $\mu\text{g/mL}$ (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount	
*	645530	CCV@25	Vial 71	1	Control	1	1.81499e6	8.385	24.13647
*	645534	QC@4.0	Vial 72	1	Control	2	2.89937e5	8.542	3.99582
*	645532	ICS@4.0	Vial 73	1	Control	3	2.38655e5	8.226	3.65624
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	7.55115e5	8.548	8344.77963
*	1907869001		Vial 76	1	Sample	6	0.00000	0.000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	2.77046e5	8.030	4.66163
*	645536	78691SD	Vial 78	1	Sample	8	2.89384e5	8.027	4.05130
*	1907871001		Vial 79	1	Sample	9	0.00000	0.000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	1.79709e6	8.389	24.49119

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount	
*	645530	CCV@25	Vial 71	1	Control	1	5.33395e5	8.403	23.90219
*	645534	QC@4.0	Vial 72	1	Control	2	9.52963e4	8.559	4.26699
*	645532	ICS@4.0	Vial 73	1	Control	3	7.88339e4	8.237	3.90865
*	645533	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.38827e5	8.562	8762.35701
*	1907869001		Vial 76	1	Sample	6	0.00000	0.000	0.00000
*	645535	78691MS	Vial 77	1	Sample	7	8.93399e4	8.046	4.91371
*	645536	78691SD	Vial 78	1	Sample	8	9.85690e4	8.049	4.48457
*	1907871001		Vial 79	1	Sample	9	0.00000	0.000	0.00000
*	1908794001		Vial 80	1	Sample	10	0.00000	0.000	0.00000
*	645537	CCV@25	Vial 71	1	Control	11	5.35802e5	8.404	24.59111

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount	
*	645530	CCV@25	Vial 71	1	Control	1	2.29255e5	8.411	5.00000
*	645534	QC@4.0	Vial 72	1	Control	2	2.39106e5	8.571	5.00000
*	645532	ICS@4.0	Vial 73	1	Control	3	2.16080e5	8.242	5.00000
*	645533	LMB	Vial 74	1	Control	4	2.42742e5	8.463	5.00000
*	1907867001	1K	Vial 75	1	Sample	5	2.89136e5	8.567	5000.00000
*	1907869001		Vial 76	1	Sample	6	1.89925e5	8.088	5.00000
*	645535	78691MS	Vial 77	1	Sample	7	1.94413e5	8.053	5.00000
*	645536	78691SD	Vial 78	1	Sample	8	2.35220e5	8.053	5.00000
*	1907871001		Vial 79	1	Sample	9	1.89609e5	8.089	5.00000
*	1908794001		Vial 80	1	Sample	10	1.92453e5	8.070	5.00000
*	645537	CCV@25	Vial 71	1	Control	11	2.23514e5	8.412	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
1	Vial 71	645530	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	645534	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	645532	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	645533	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1907867001	1K	CLO4-AQN	1	Sample	
6	Vial 76	1907869001		CLO4-AQN	1	Sample	
7	Vial 77	645535	78691MS	CLO4-AQN	1	Sample	
8	Vial 78	645536	78691SD	CLO4-AQN	1	Sample	
9	Vial 79	1907871001		CLO4-AQN	1	Sample	
10	Vial 80	1908794001		CLO4-AQN	1	Sample	
11	Vial 71	645537	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D

Sample Name: 645530 CCV@25

Injection Date: 3/28/2019 08:30:53

Seq Line: 1

Sample Name: 645530 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

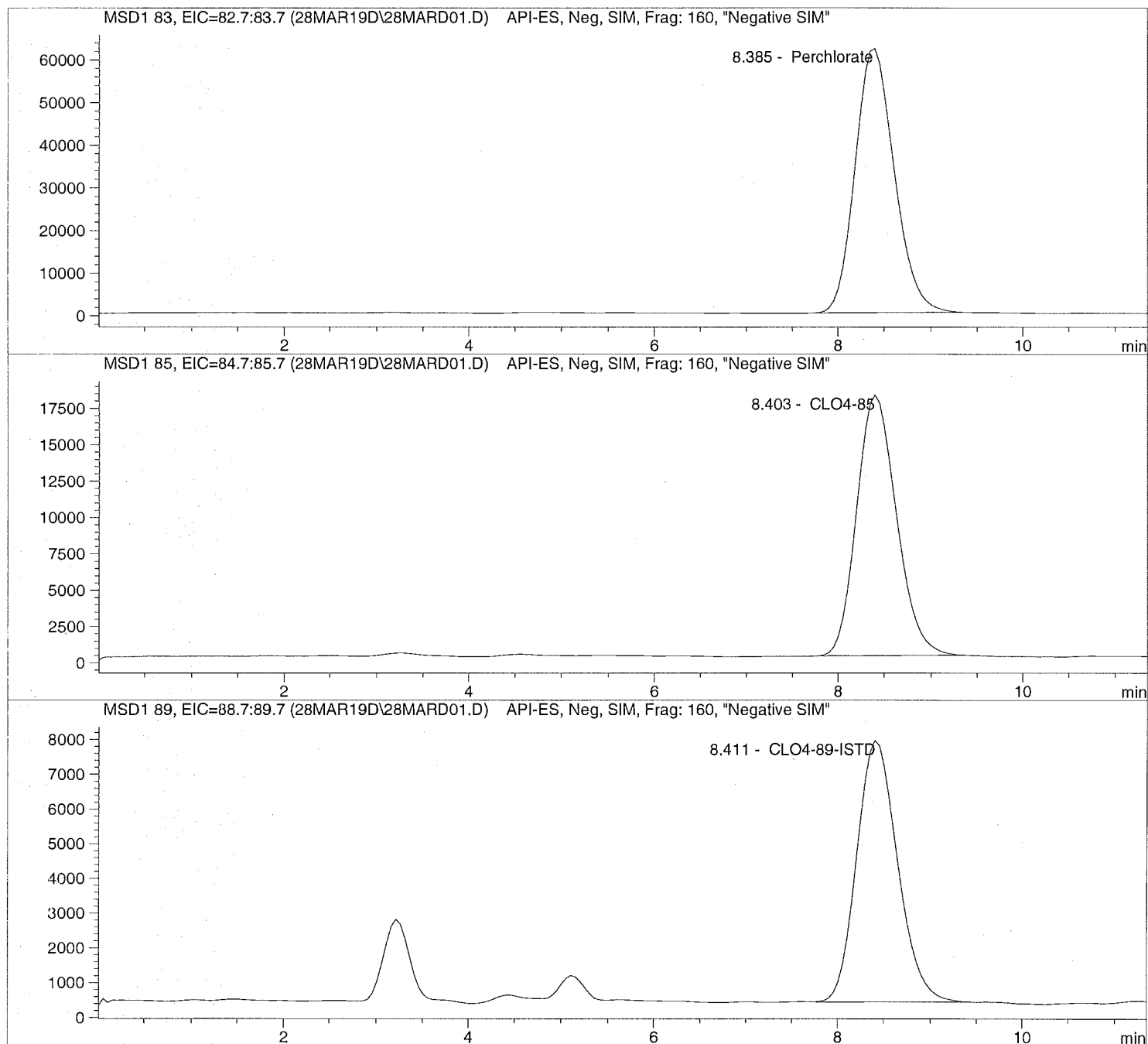
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD01.D Sample Name: 645530 CCV025

```

=====
Injection Date: 3/28/2019 08:30:53      Seq Line: 1
Sample Name: 645530 CCV025             Location: Vial 71
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.385	PBA	1814992.7	24.1365	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.403	PBA	533395.5	23.9022	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.411	BBA	229255.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D

Sample Name: 645534 QC@4.0

Injection Date: 3/28/2019 08:47:42

Seq Line: 2

Sample Name: 645534 QC@4.0

Location: Vial 72

Acq Operator: TNB

Inj. No.: 1

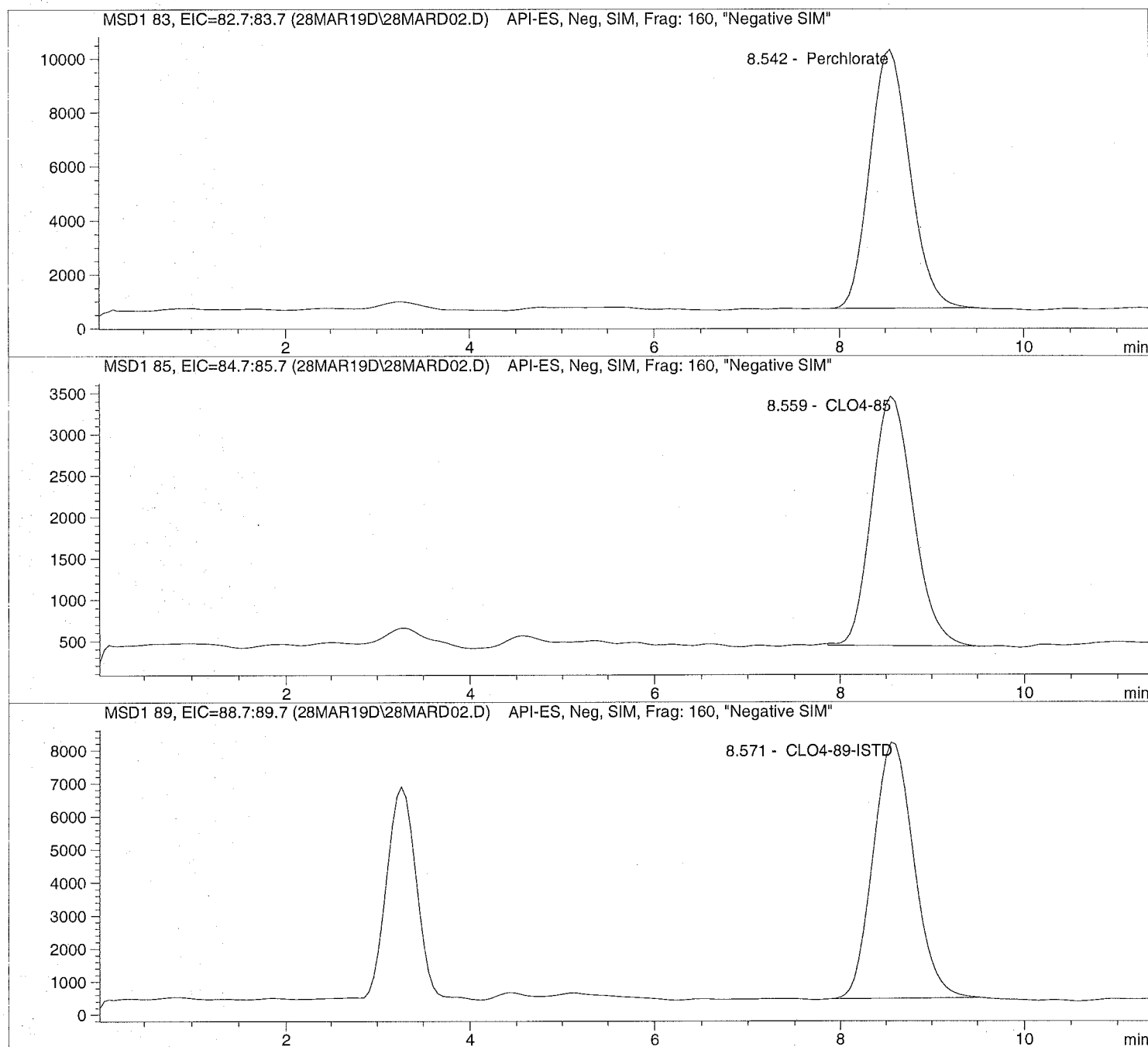
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD02.D Sample Name: 645534 QC@4.0

```

=====
Injection Date: 3/28/2019 08:47:42      Seq Line: 2
Sample Name: 645534 QC@4.0             Location: Vial 72
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.542	PBA	289937.1	3.9958	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.559	BBA	95296.3	4.2670	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.571	PBA	239105.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD03.D

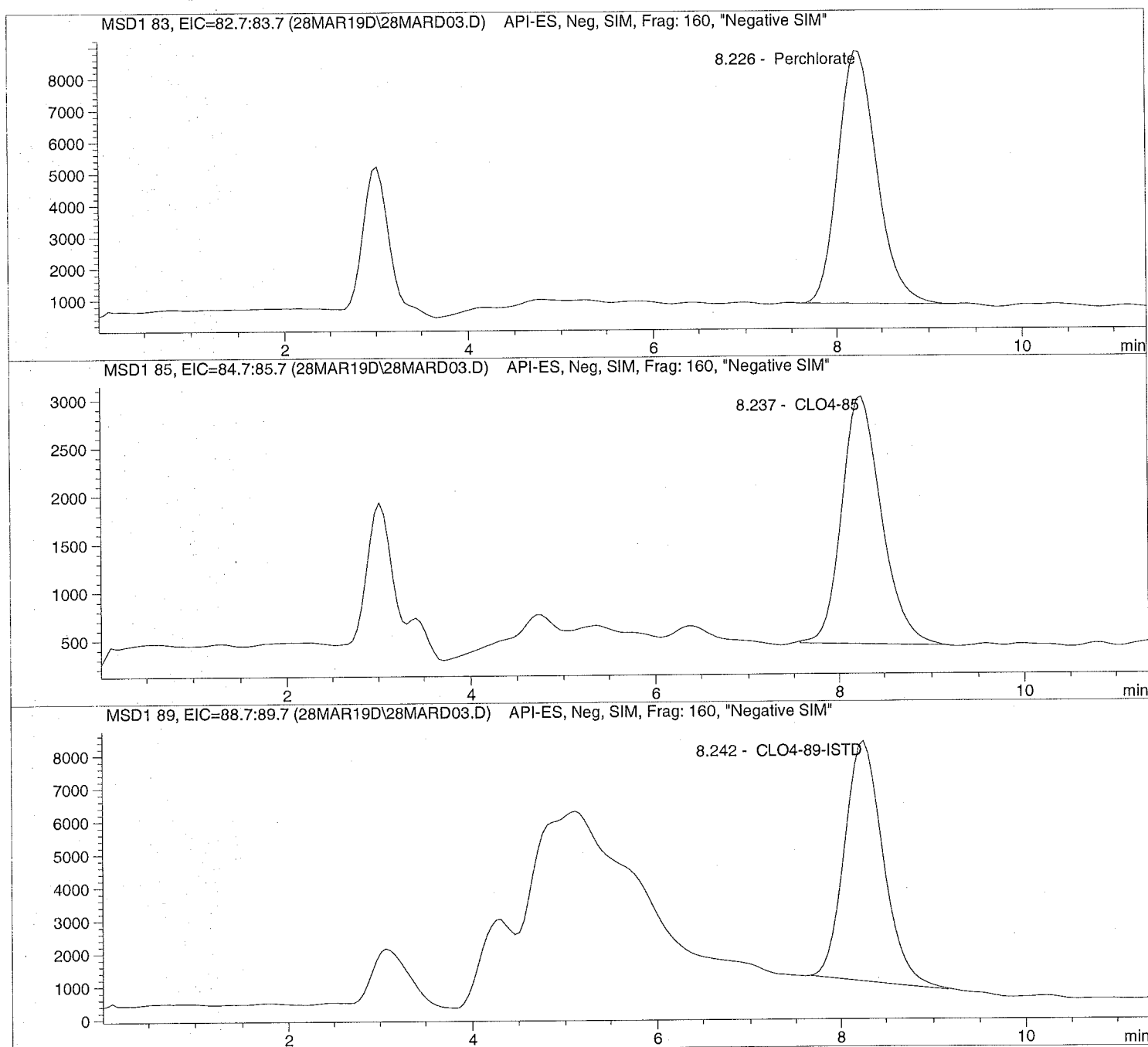
Sample Name: 645532 ICS@4.0

Injection Date: 3/28/2019 09:00:54
Sample Name: 645532 ICS@4.0
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD03.D Sample Name: 645532 ICS@4.0

```

=====
Injection Date:  3/28/2019  09:00:54      Seq Line:      3
Sample Name:    645532   ICS@4.0      Location:      Vial 73
Acq Operator:   TNB                      Inj. No.:     1
                                         Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 4.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.226	BBA	238654.8	3.6562	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.237	BBA	78833.9	3.9087	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.242	PBA	216079.7	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D

Sample Name: 645533 LMB

Injection Date: 3/28/2019 09:14:08

Seq Line: 4

Sample Name: 645533 LMB

Location: Vial 74

Acq Operator: TNB

Inj. No.: 1

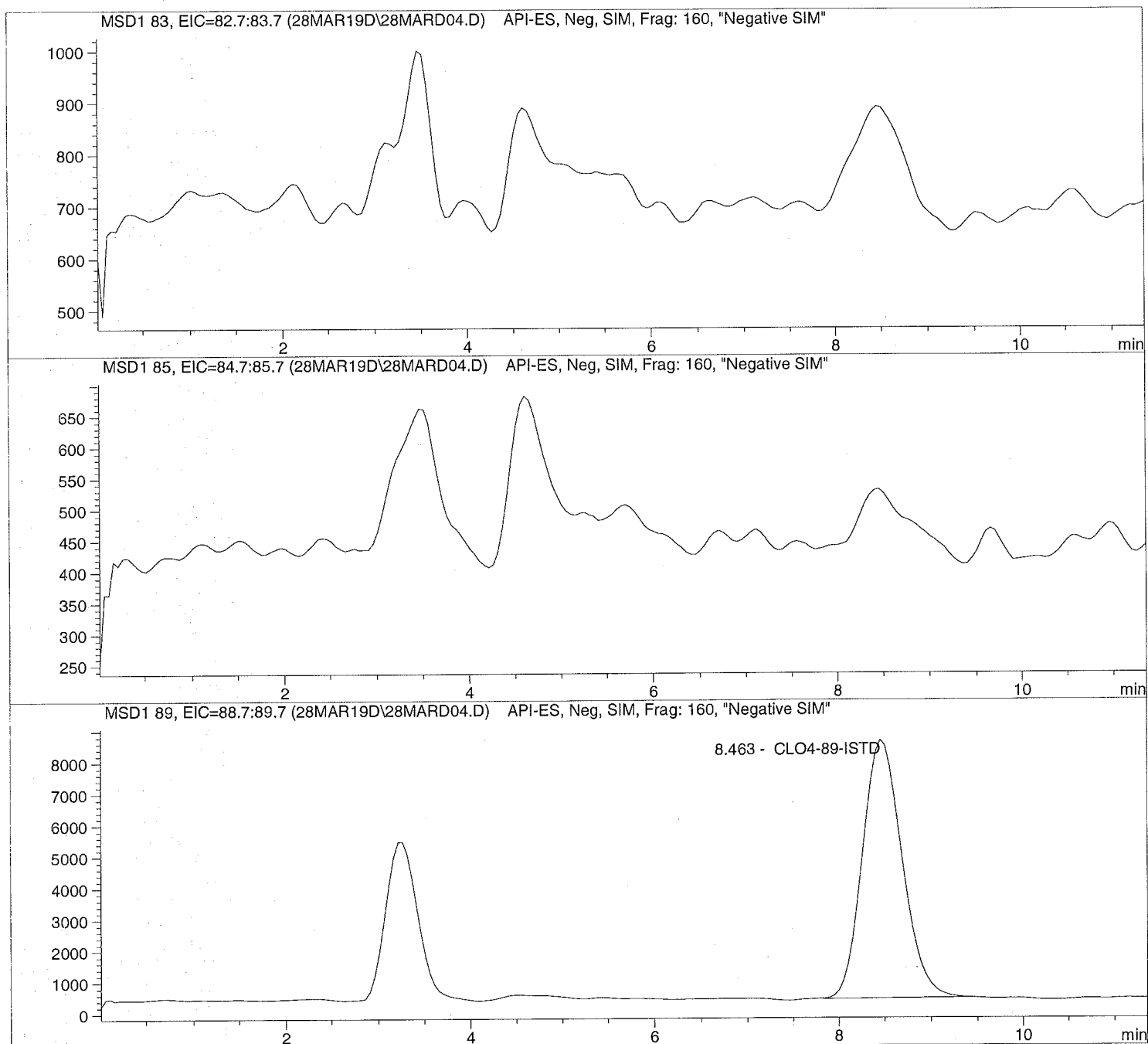
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD04.D Sample Name: 645533 LMB

```

=====
Injection Date: 3/28/2019 09:14:08      Seq Line: 4
Sample Name: 645533 LMB                  Location: Vial 74
Acq Operator: TNB                        Inj. No.: 1
                                         Inj. Vol.: 25 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.463	BBA	242742.3	5.0000	CLO4-89-ISTD

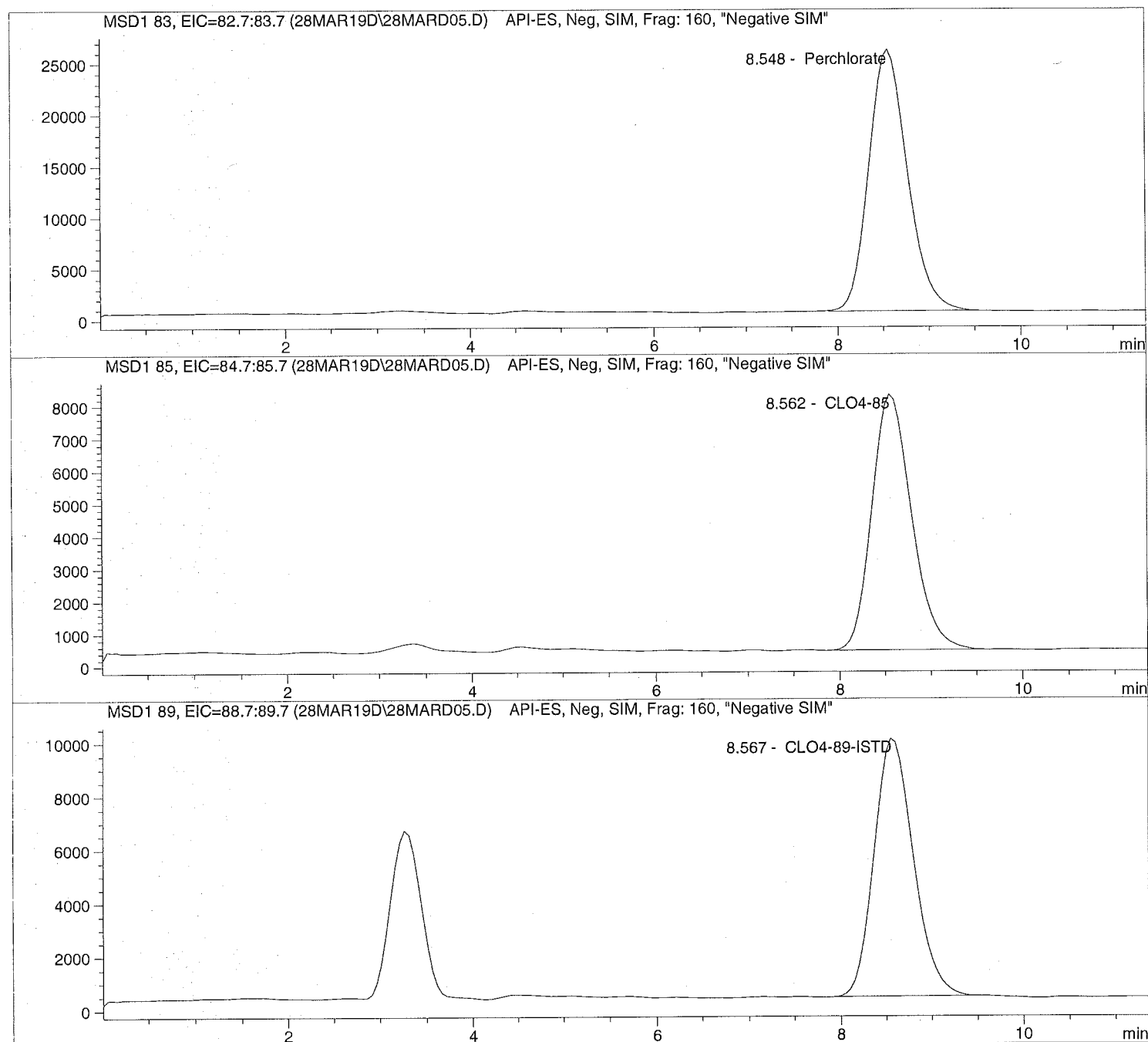
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```
=====
Injection Date: 3/28/2019 09:28:25      Seq Line:      5
Sample Name:    1907867001 1K           Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD05.D Sample Name: 1907867001 1K

```

=====
Injection Date:  3/28/2019  09:28:25      Seq Line:      5
Sample Name:    1907867001  1K           Location:      Vial 75
Acq Operator:   TNB                Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1000.000000
Sample Amount:  0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.548	BBA	755115.1	8344.7796	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.562	PBA	238826.8	8762.3570	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.567	PBA	289135.6	5000.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D

Sample Name: 1907869001

Injection Date: 3/28/2019 09:41:37
Sample Name: 1907869001
Acq Operator: TNB

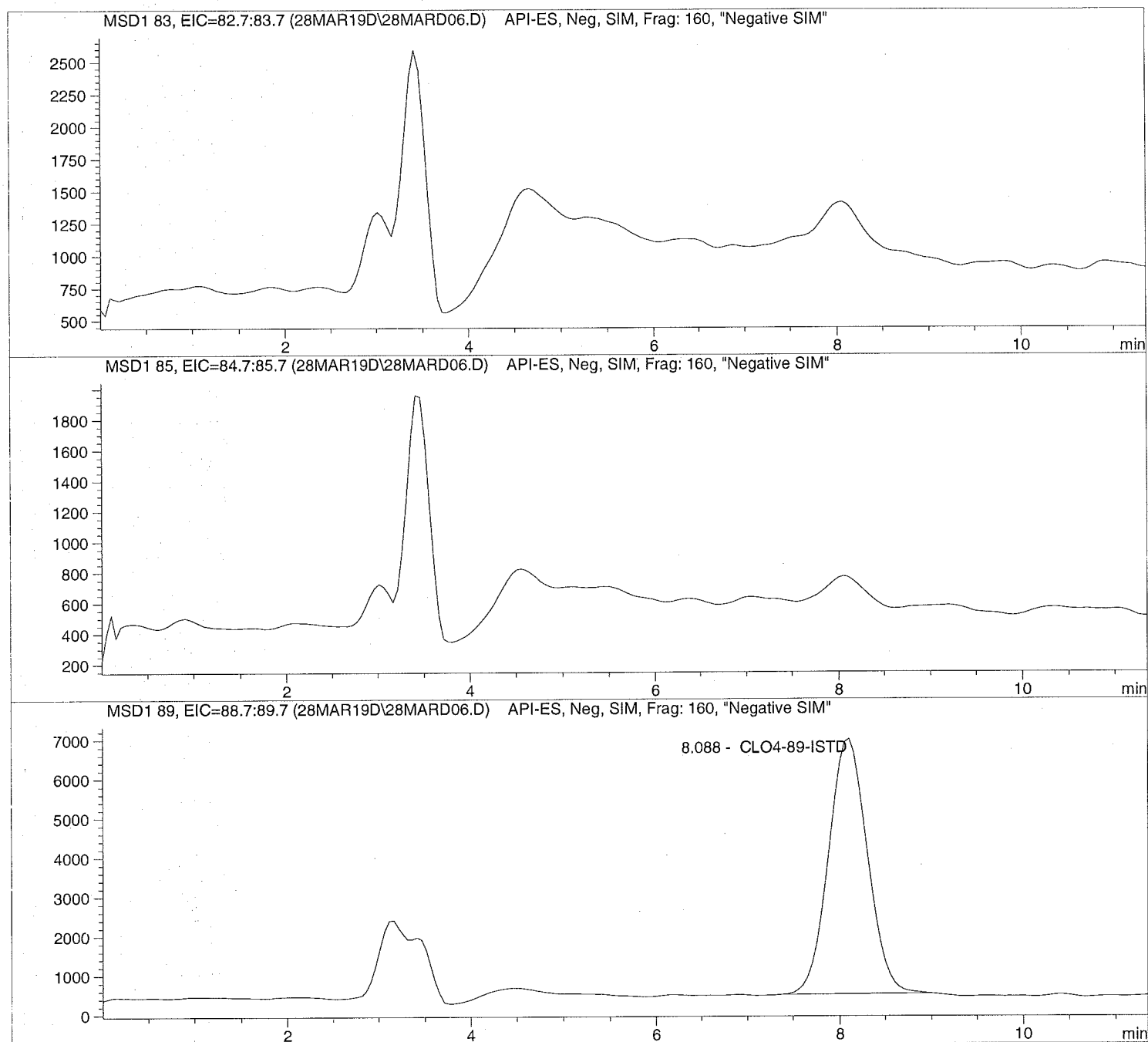
Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD06.D Sample Name: 1907869001

```

=====
Injection Date: 3/28/2019 09:41:37      Seq Line: 6
Sample Name: 1907869001                Location: Vial 76
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.088	BBA	189925.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D

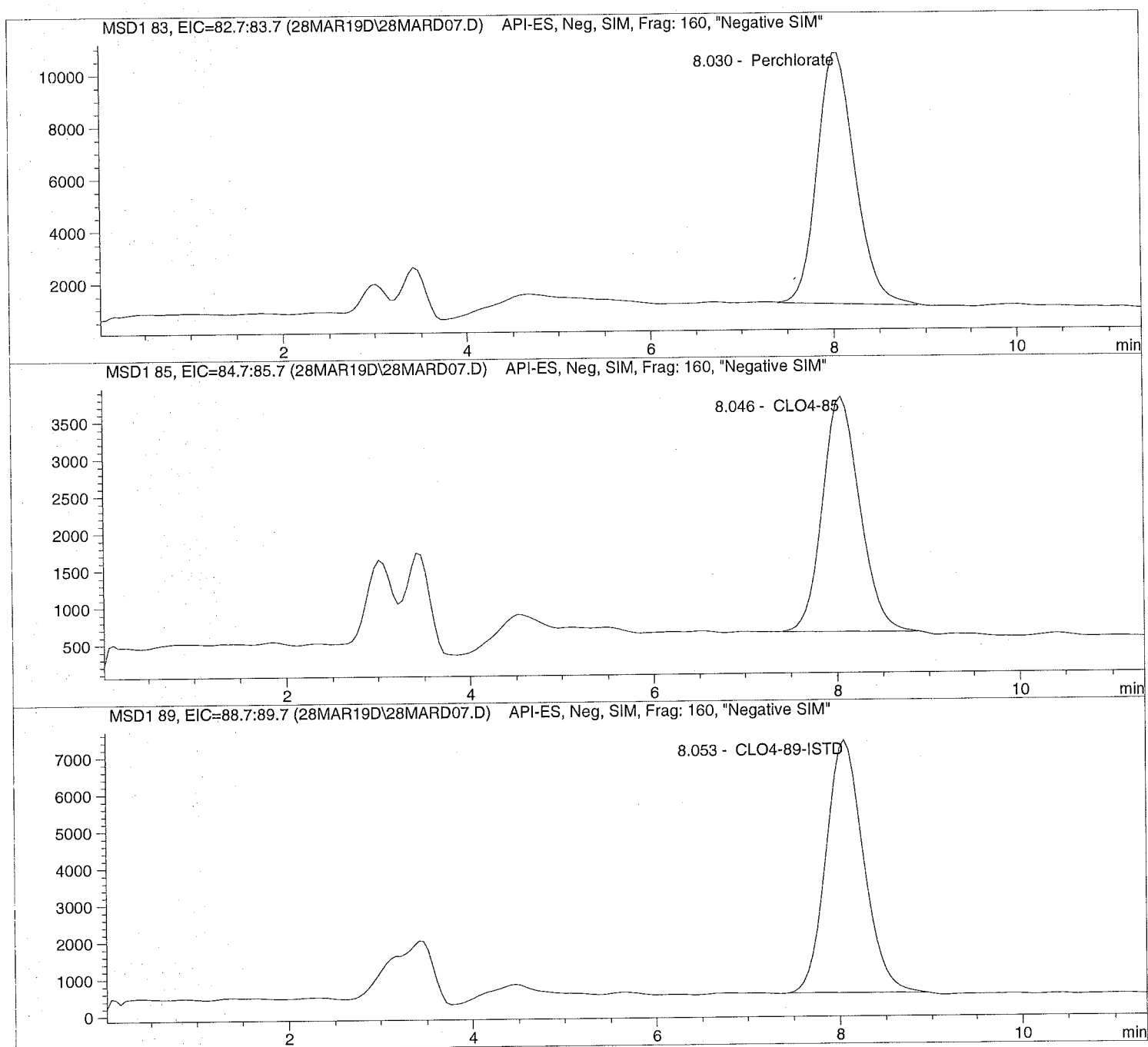
Sample Name: 645535 78691MS

Injection Date: 3/28/2019 09:54:47
Sample Name: 645535 78691MS
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD07.D Sample Name: 645535 78691MS

```

=====
Injection Date: 3/28/2019 09:54:47 Seq Line: 7
Sample Name: 645535 78691MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.030	BBA	277046.3	4.6616	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.046	BBA	89339.9	4.9137	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	194412.9	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D

Sample Name: 645536 78691SD

Injection Date: 3/28/2019 10:08:05
Sample Name: 645536 78691SD
Acq Operator: TNB

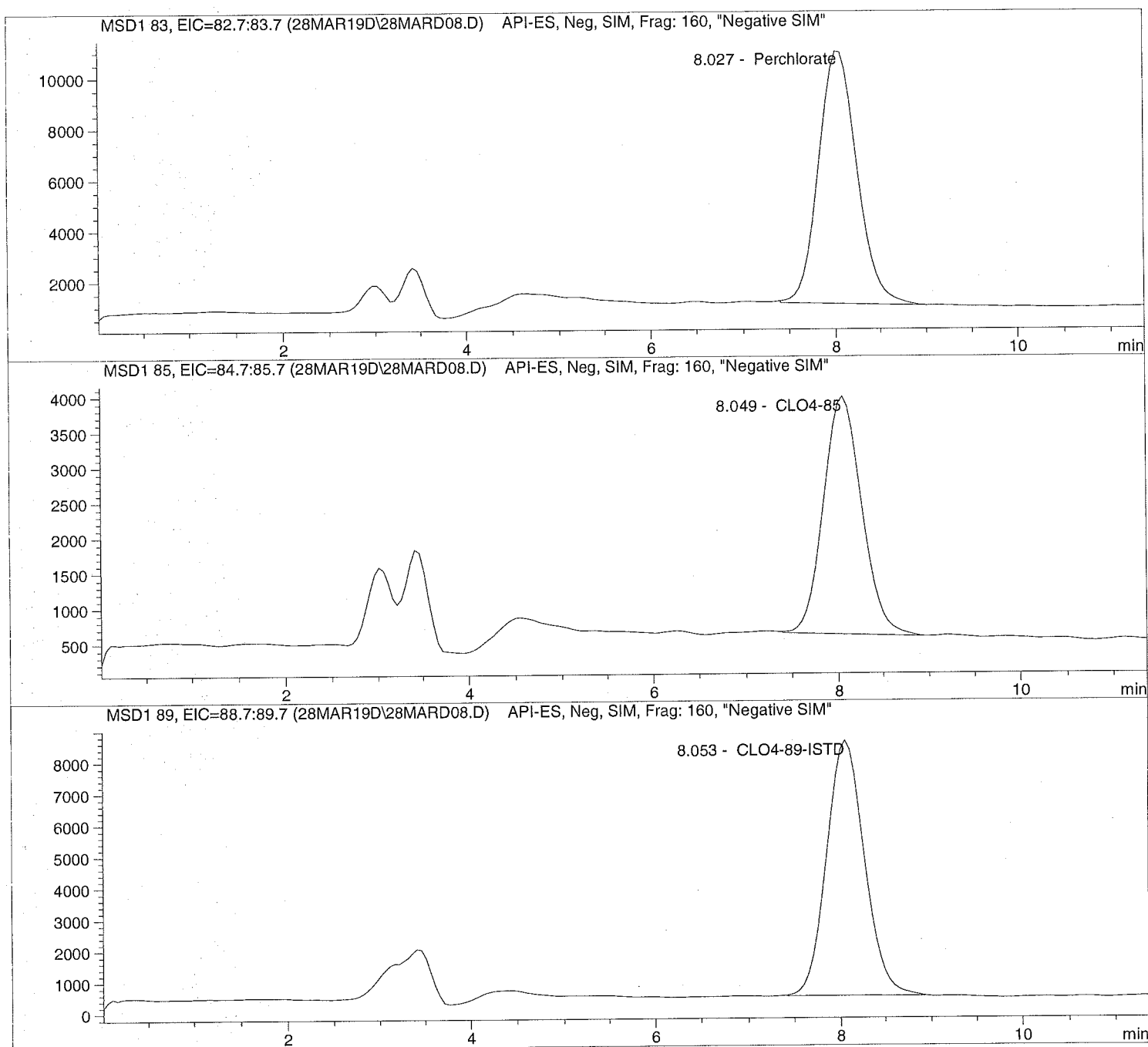
Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD08.D Sample Name: 645536 78691SD

```

=====
Injection Date: 3/28/2019 10:08:05 Seq Line: 8
Sample Name: 645536 78691SD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.027	BBA	289383.7	4.0513	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.049	BBA	98569.0	4.4846	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.053	BBA	235219.8	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D

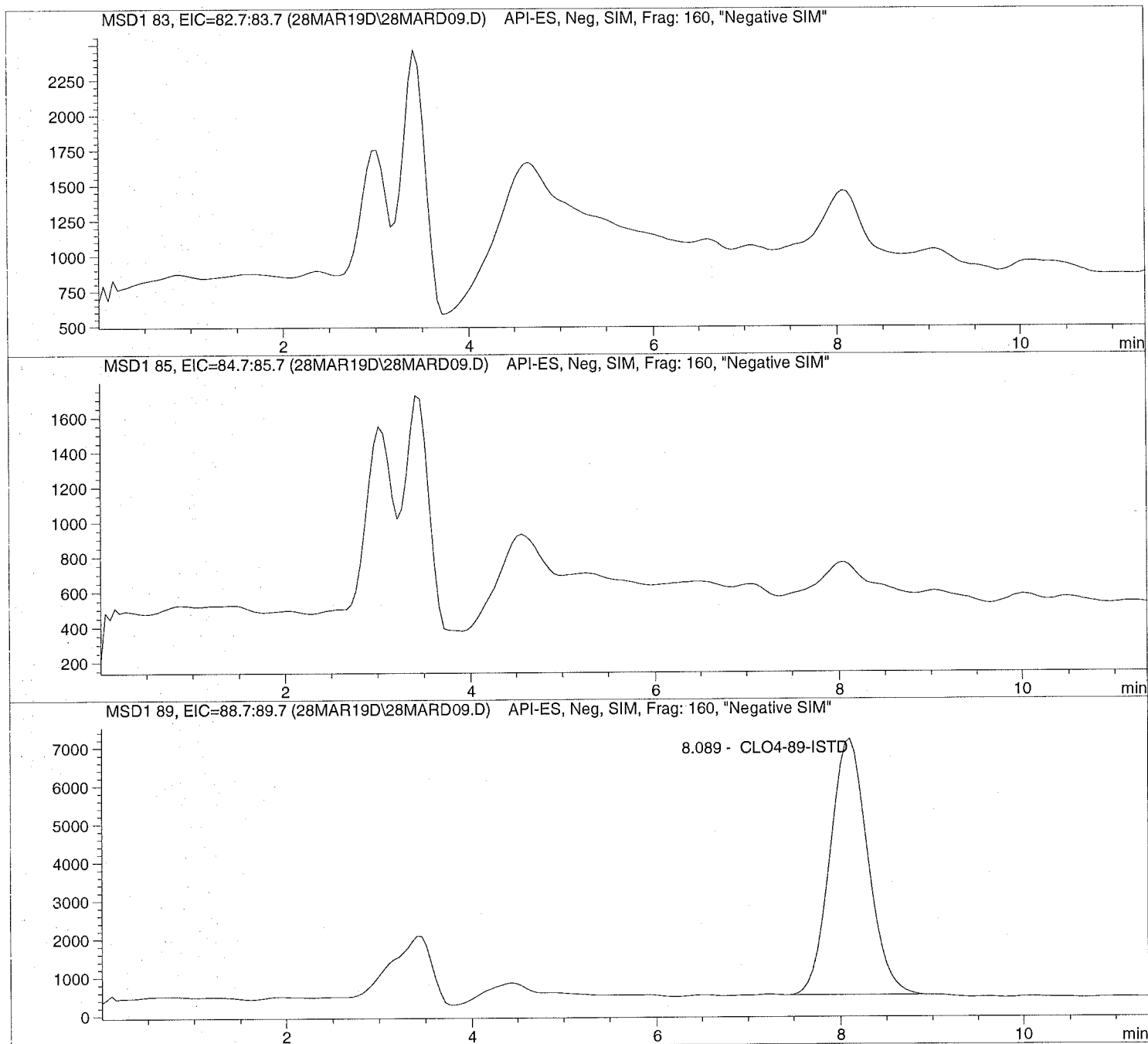
Sample Name: 1907871001

Injection Date: 3/28/2019 10:21:14
Sample Name: 1907871001
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 25 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD09.D Sample Name: 1907871001

```

=====
Injection Date: 3/28/2019 10:21:14      Seq Line:          9
Sample Name:    1907871001              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:   0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.089	PBA	189608.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD10.D

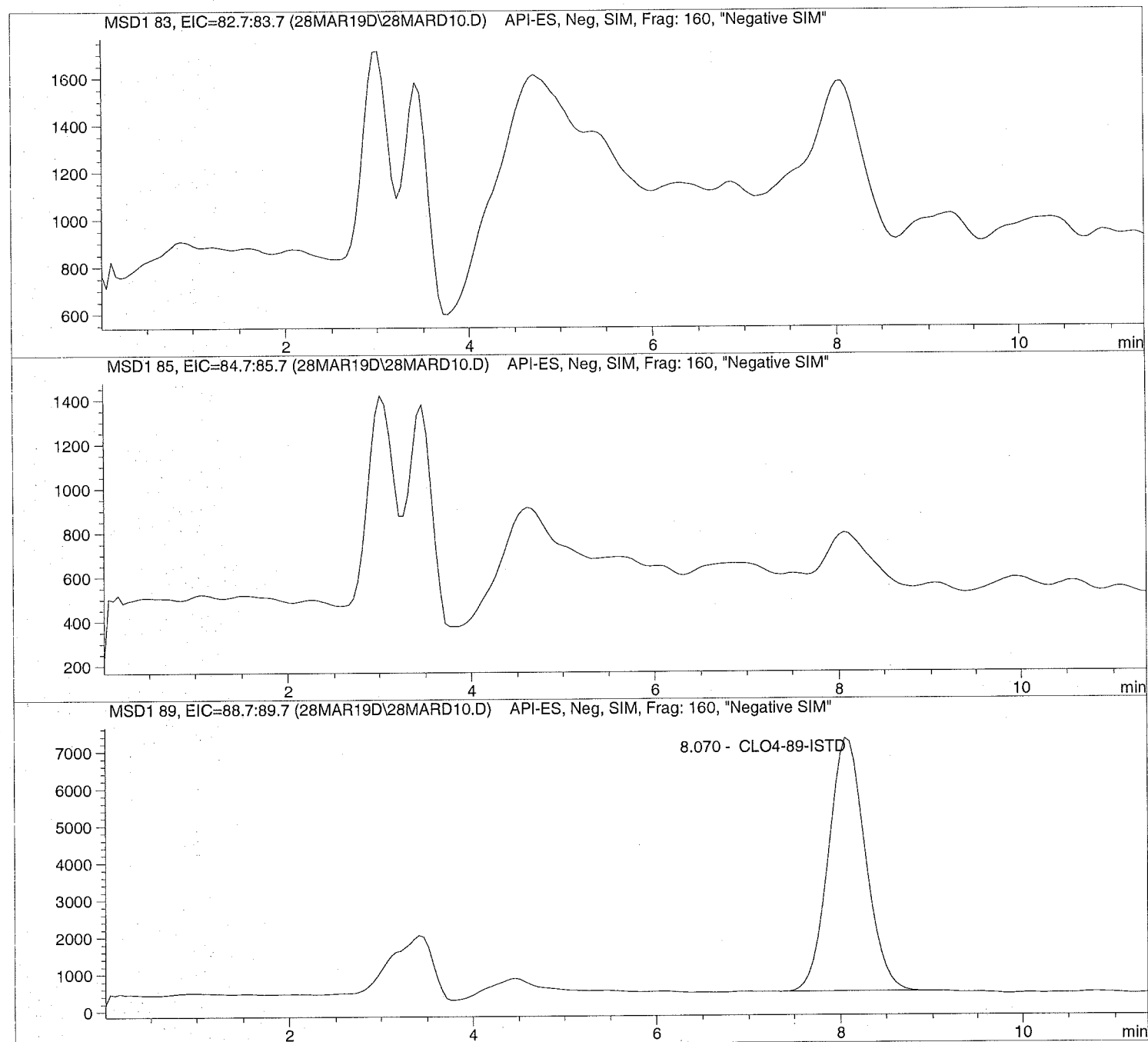
Sample Name: 1908794001

=====
Injection Date: 3/28/2019 10:34:27
Sample Name: 1908794001
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 25 µl

=====
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD10.D Sample Name: 1908794001

```

=====
Injection Date: 3/28/2019 10:34:27      Seq Line:      10
Sample Name:   1908794001              Location:      Vial 80
Acq Operator:  TNB                     Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====

```

```

Acq. Method:  CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.070	PBA	192453.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D

Sample Name: 645537 CCV@25

Injection Date: 3/28/2019 10:49:41

Seq Line: 11

Sample Name: 645537 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

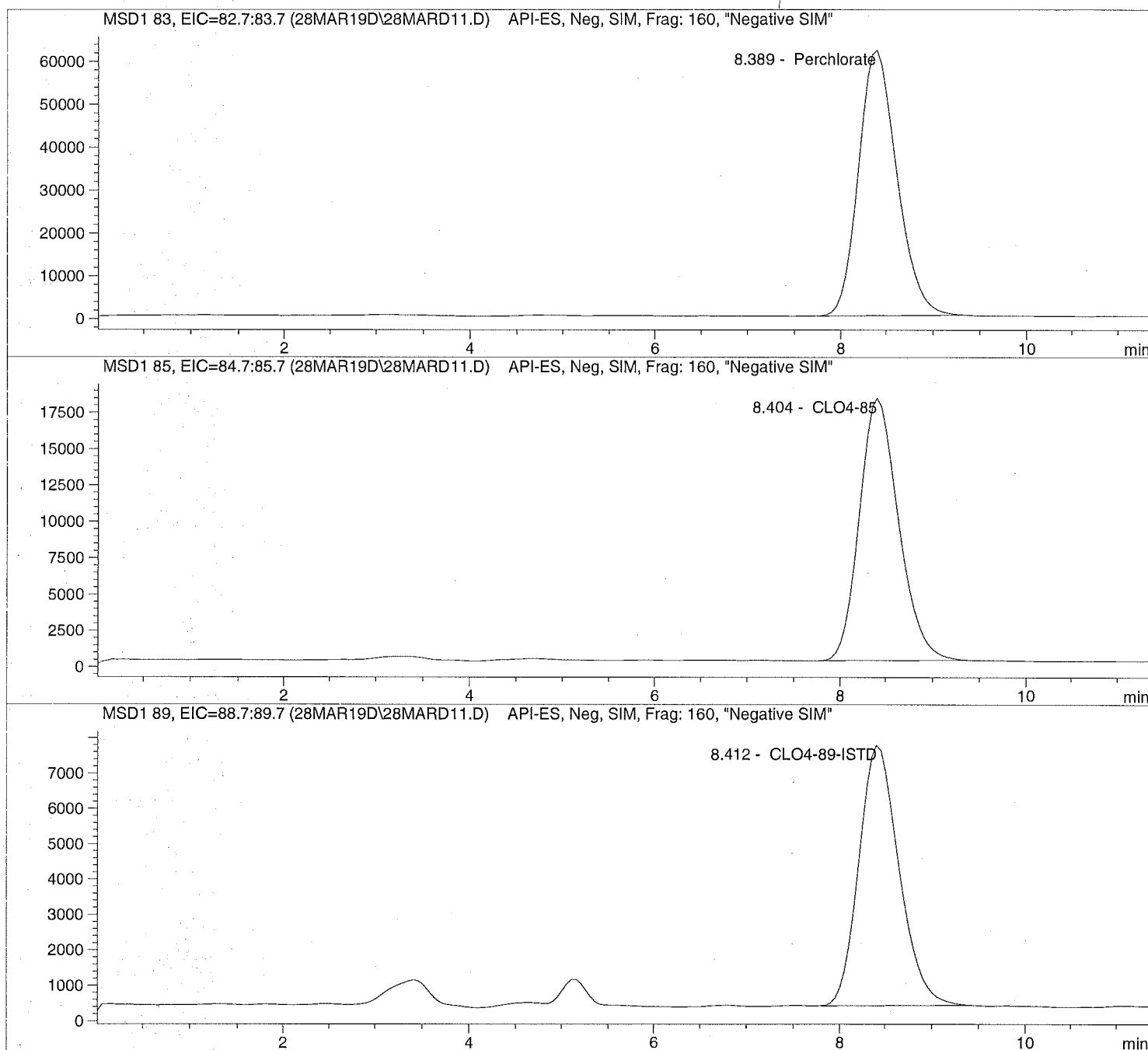
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\28MAR19D\28MARD11.D Sample Name: 645537 CCV@25

```

=====
Injection Date: 3/28/2019 10:49:41      Seq Line:      11
Sample Name:   645537  CCV@25           Location:      Vial 71
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.389	PBA	1797085.1	24.4912	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.404	PBA	535801.9	24.5911	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.412	BBA	223514.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
 Calibration Table
 =====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
 Based on : Peak Area

Rel. Reference Window : 20.000 %
 Abs. Reference Window : 0.000 min
 Rel. Non-ref. Window : 20.000 %
 Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
 Uncalibrated Peaks : not reported
 Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
 Origin : Ignored (some peaks differ, see below)
 Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
 Average Response : Average all calibrations
 Average Retention Time: Floating Average New 75%

Calibration Report Options :
 Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
 If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
 Signal 2: MSD1 85, EIC=84.7:85.7
 Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl	Amount	Area	Amt/Area	Ref Grp	Name
8.744	1	1.00000	7.76074e4	1.28854e-5	1	Perchlorate
	2	2.00000	1.35273e5	1.47849e-5		
	3	5.00000	3.37764e5	1.48033e-5		
	4	10.00000	6.83454e5	1.46316e-5		
	5	25.00000	2.08433e6	1.19943e-5		
	6	50.00000	4.13334e6	1.20968e-5		
	7	75.00000	5.99313e6	1.25143e-5		
8.755	2	1.00000	2.36780e4	4.22333e-5	1	CLO4-85
	2	2.00000	4.69486e4	4.25998e-5		
	3	5.00000	1.06124e5	4.71147e-5		
	4	10.00000	2.13523e5	4.68335e-5		
	5	25.00000	6.14295e5	4.06971e-5		
	6	50.00000	1.19814e6	4.17315e-5		
	7	75.00000	1.78355e6	4.20509e-5		
8.766	3	5.00000	2.73208e5	1.83011e-5	+I1	CLO4-89-ISTD
	2	5.00000	2.24886e5	2.22335e-5		
	3	5.00000	2.33196e5	2.14412e-5		
	4	5.00000	2.34454e5	2.13262e-5		
	5	5.00000	2.50568e5	1.99547e-5		
	6	5.00000	2.30977e5	2.16472e-5		

RetTime [min]	Lvl Sig	Amount	Area	Amt/Area	Ref Grp Name
7		5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-85

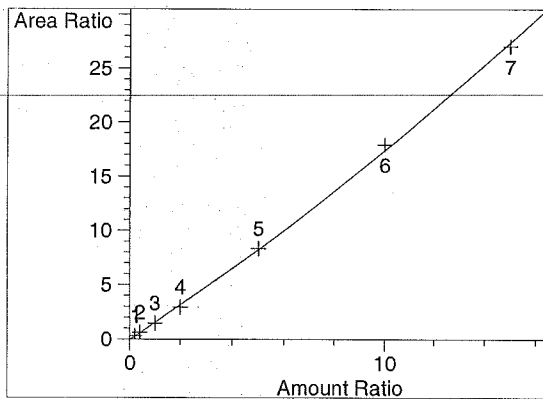
Time Window : From 6.650 min To 12.505 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-89-ISTD

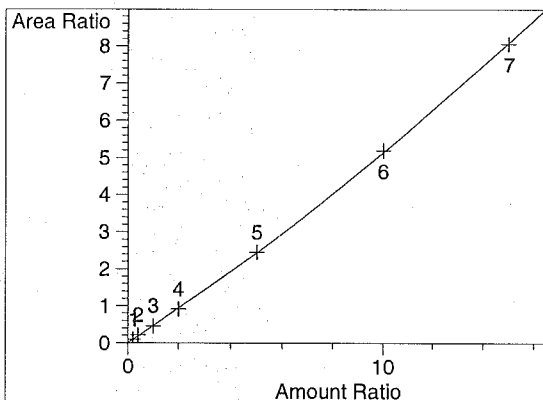
Time Window : From 6.659 min To 12.466 min
Curve Type : Linear
Origin : Included
Calibration Level Weights:/
Level 1 : 1
Level 2 : 1
Level 3 : 1
Level 4 : 1
Level 5 : 1
Level 6 : 1
Level 7 : 1

=====
Peak Sum Table
=====***No Entries in table***
=====

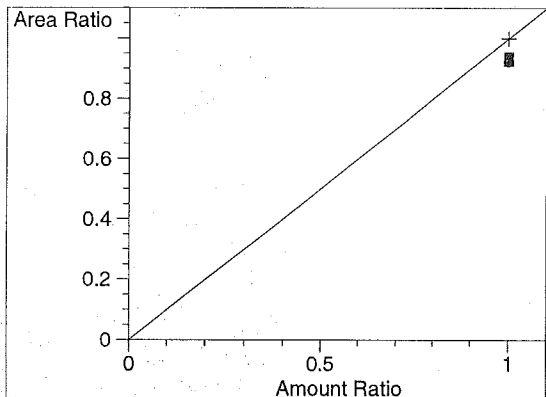
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

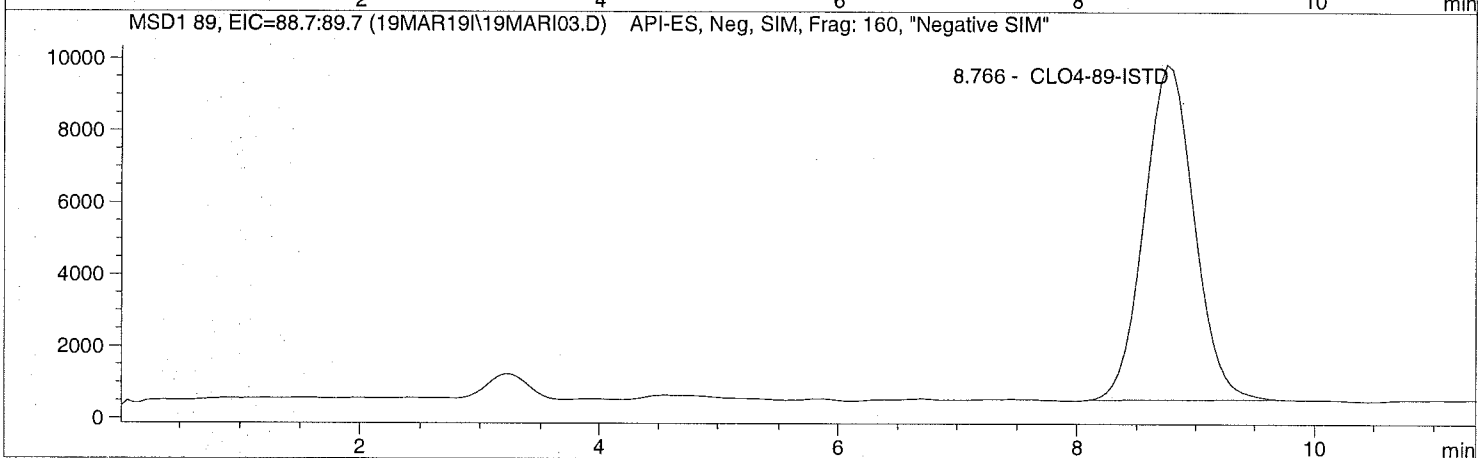
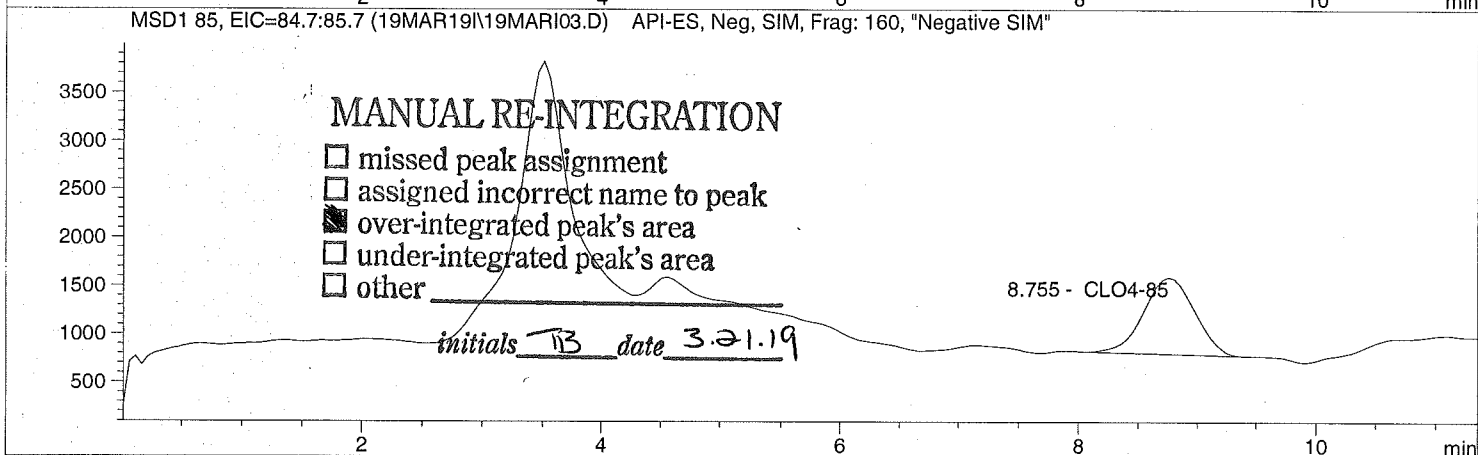
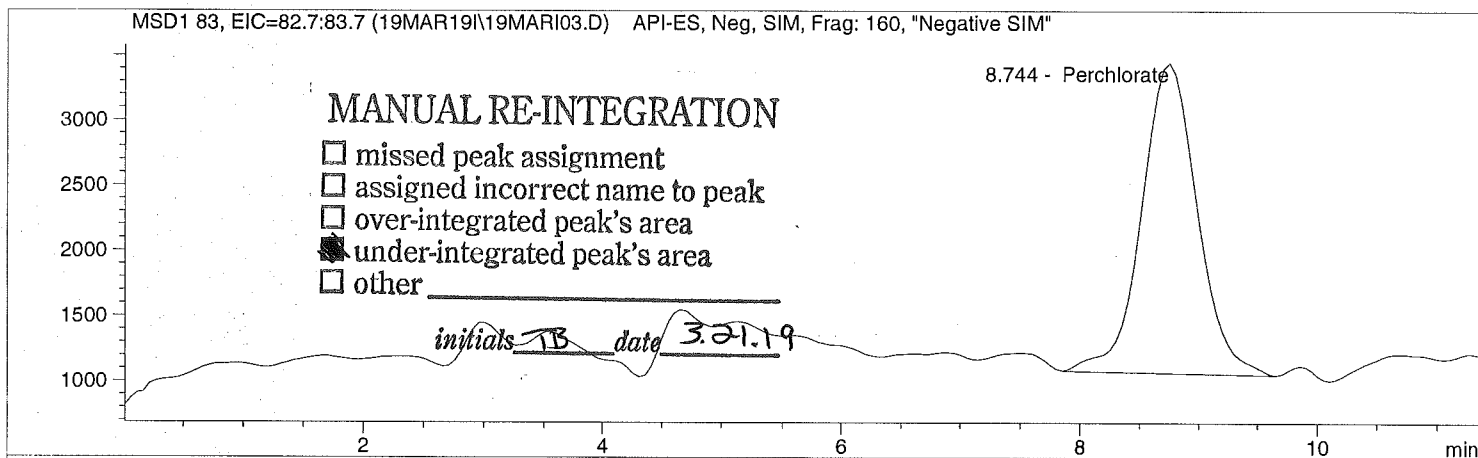
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:      3
Sample Name:    CLO4@ 1.0ug/L           Location:      Vial 73
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D

Sample Name: CLO4@ 2.0ug/L

Injection Date: 3/19/2019 09:53:00

Seq Line: 4

Sample Name: CLO4@ 2.0ug/L

Location: Vial 74

Acq Operator: TNB

Inj. No.: 1

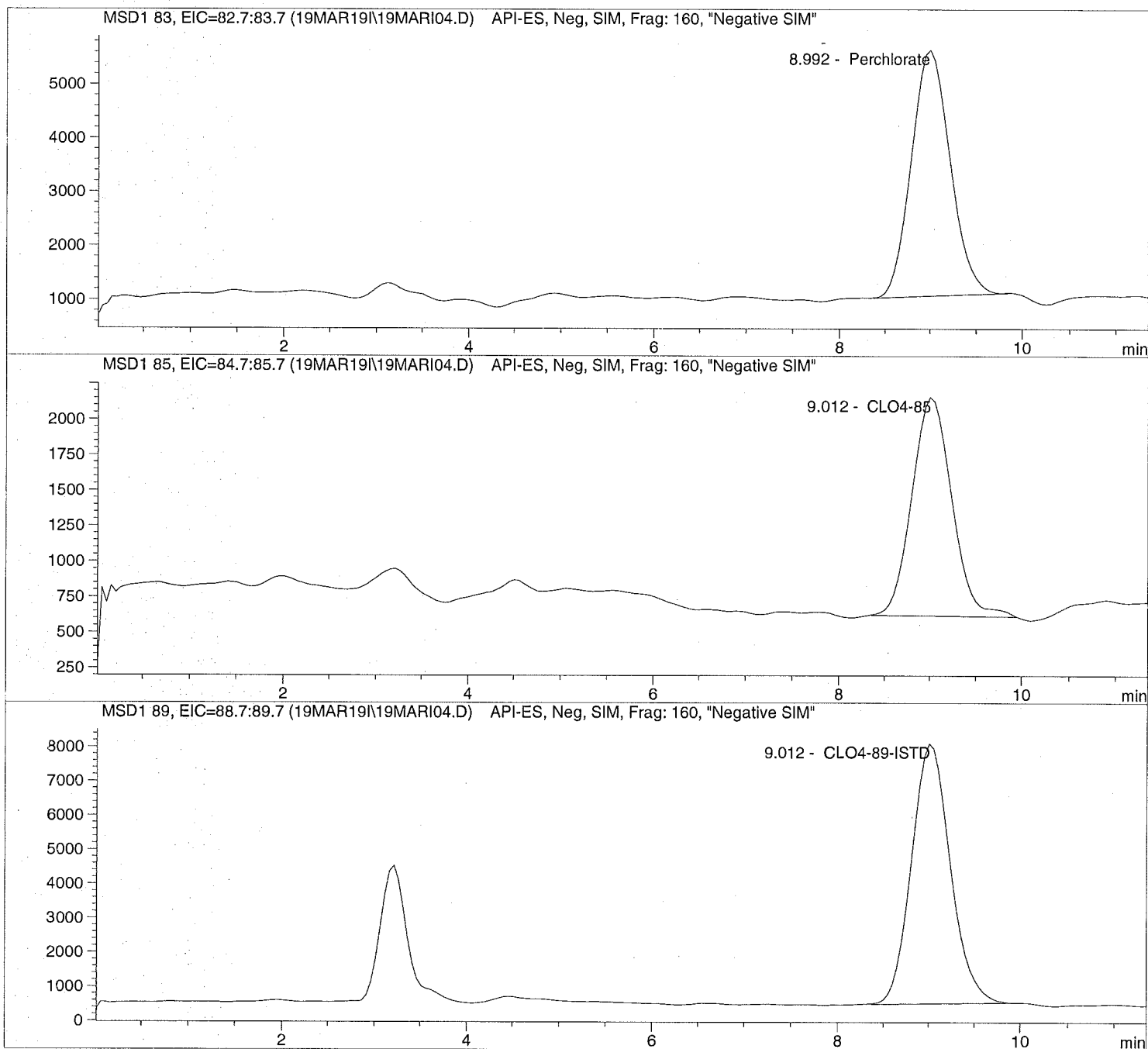
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line:      4
Sample Name:    CLO4@ 2.0ug/L          Location:      Vial 74
Acq Operator:   TNB                   Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D

Sample Name: CLO4@ 5.0ug/L

Injection Date: 3/19/2019 10:06:16

Seq Line: 5

Sample Name: CLO4@ 5.0ug/L

Location: Vial 75

Acq Operator: TNB

Inj. No.: 1

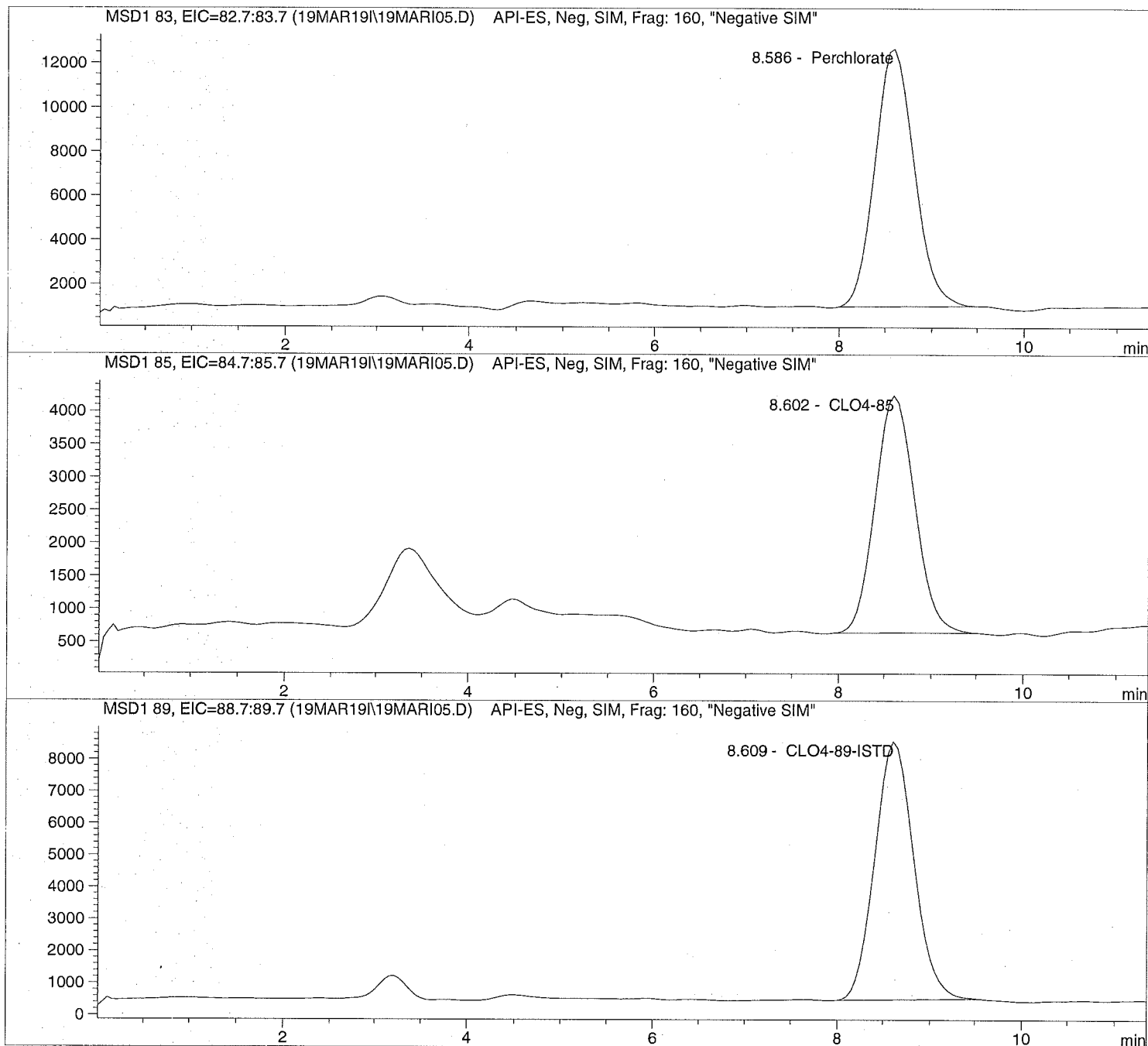
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line:          5
Sample Name:    CLO4@ 5.0ug/L           Location:          Vial 75
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:       30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D

Sample Name: CLO4@ 10.ug/L

Injection Date: 3/19/2019 10:19:32

Seq Line: 6

Sample Name: CLO4@ 10.ug/L

Location: Vial 76

Acq Operator: TNB

Inj. No.: 1

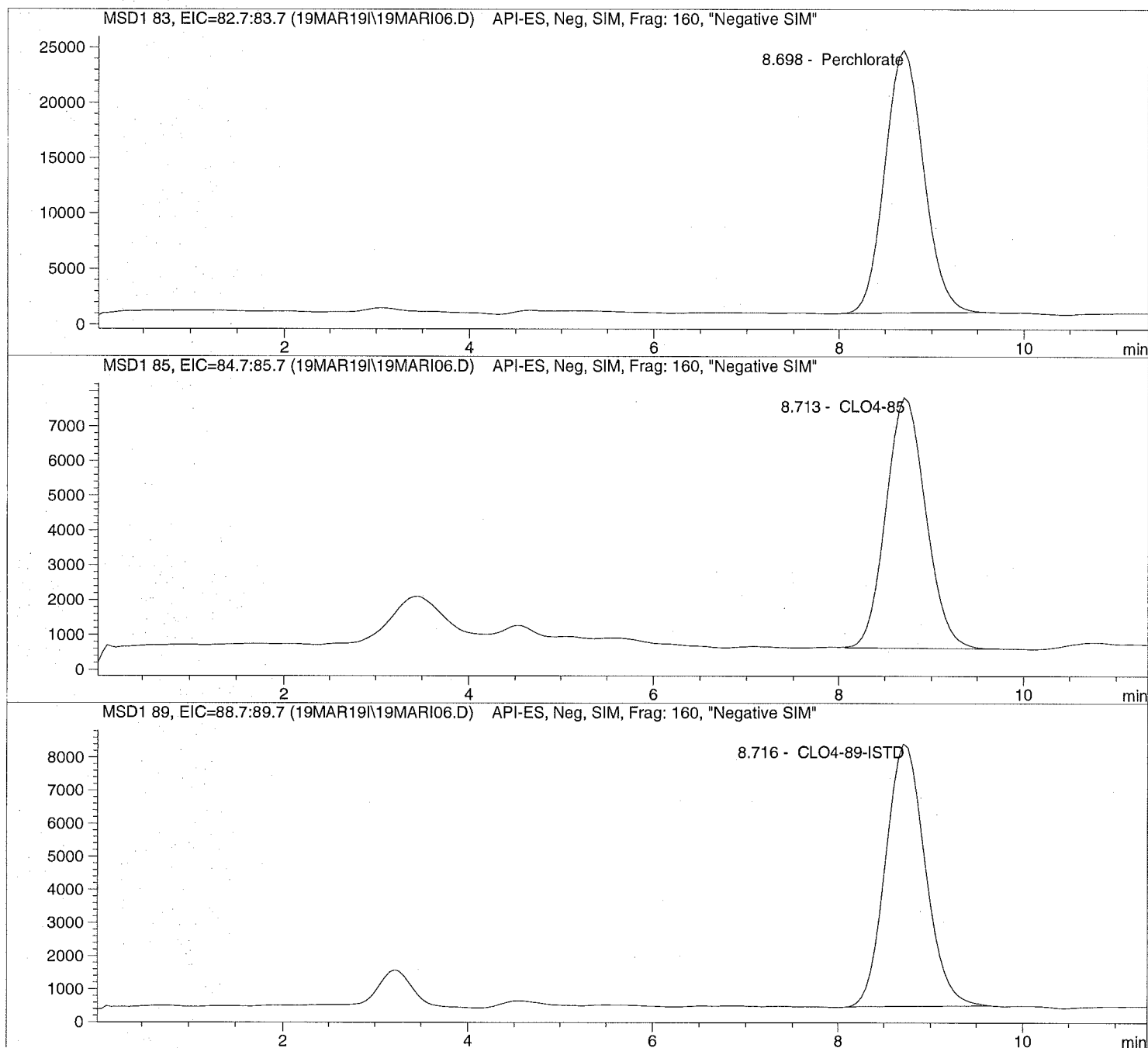
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line:                 6
Sample Name:    CLO4@ 10.ug/L            Location:                Vial 76
Acq Operator:   TNB                      Inj. No.:                1
                                         Inj. Vol.:                30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:                    Signal
Calib. Data Modified:      Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                 1.000000
Dilution:                    1.000000
Sample Amount:              10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D

Sample Name: CLO4@ 25.ug/L

Injection Date: 3/19/2019 10:32:49

Seq Line: 7

Sample Name: CLO4@ 25.ug/L

Location: Vial 77

Acq Operator: TNB

Inj. No.: 1

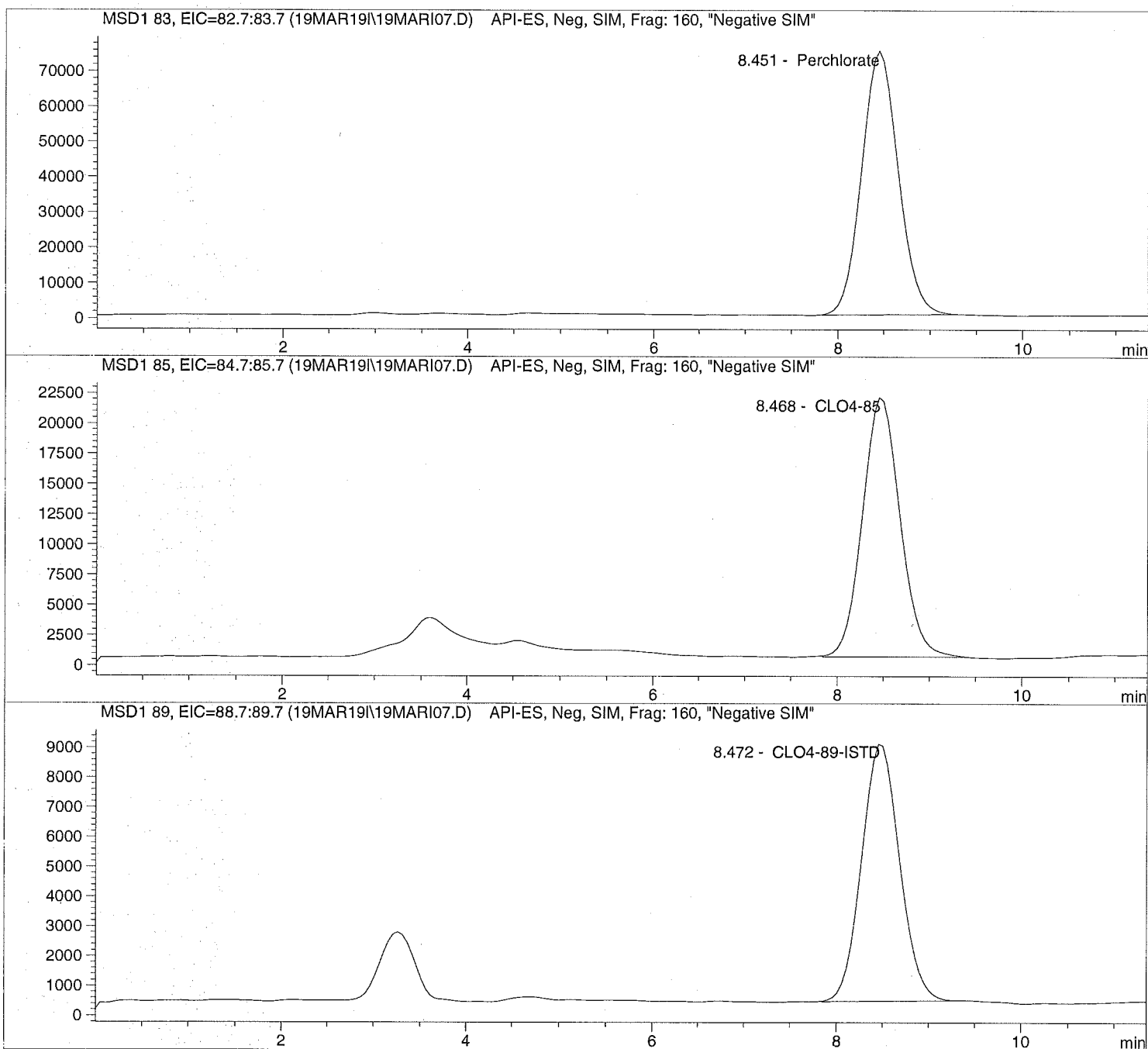
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 3/19/2019 10:32:49      Seq Line:          7
Sample Name:    CLO4@ 25.ug/L           Location:          Vial 77
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI08.D

Sample Name: CLO4@ 50.ug/L

Injection Date: 3/19/2019 10:46:05

Seq Line: 8

Sample Name: CLO4@ 50.ug/L

Location: Vial 78

Acq Operator: TNB

Inj. No.: 1

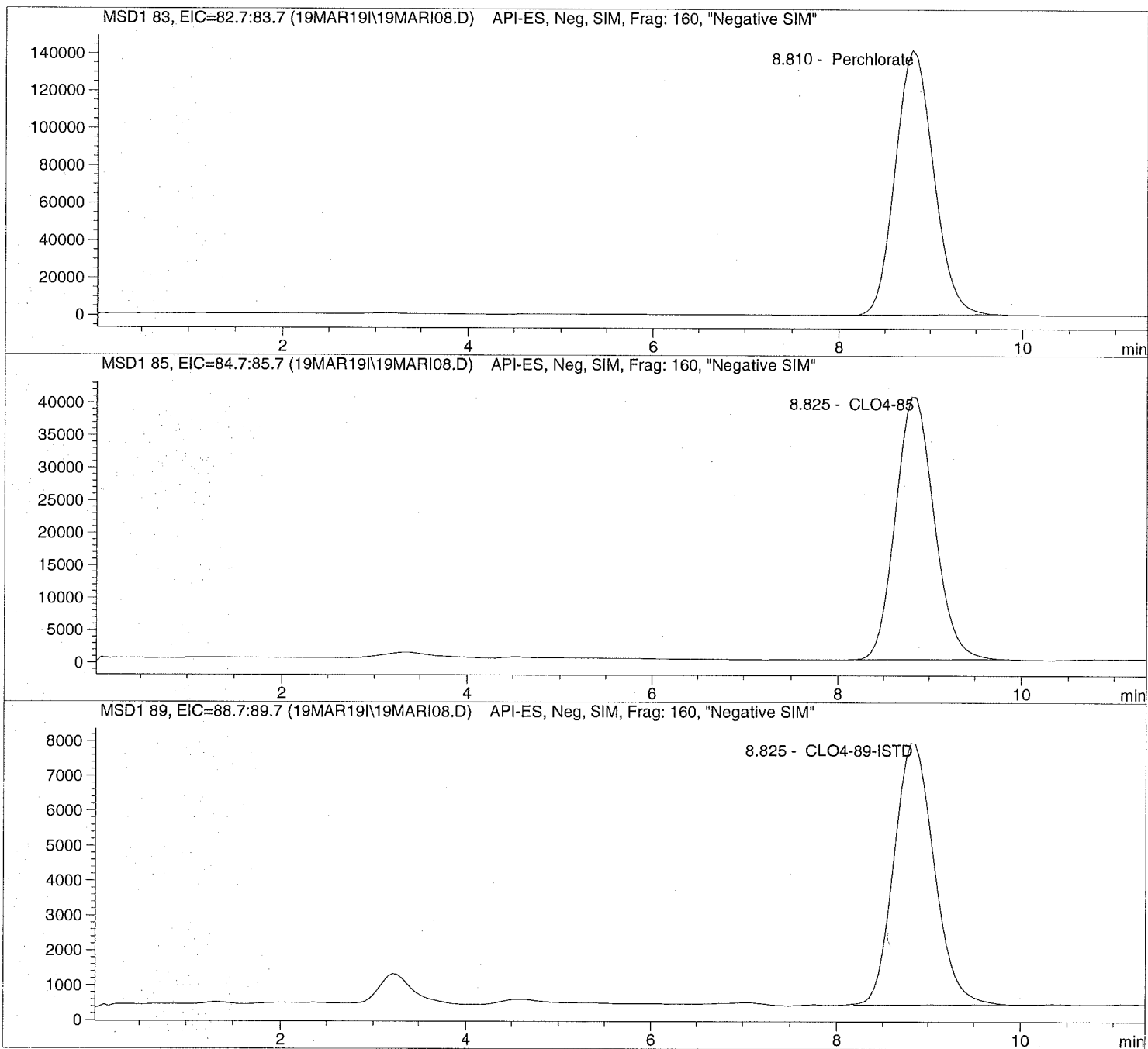
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:   CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:  TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 50.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 3/19/2019 10:59:22

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

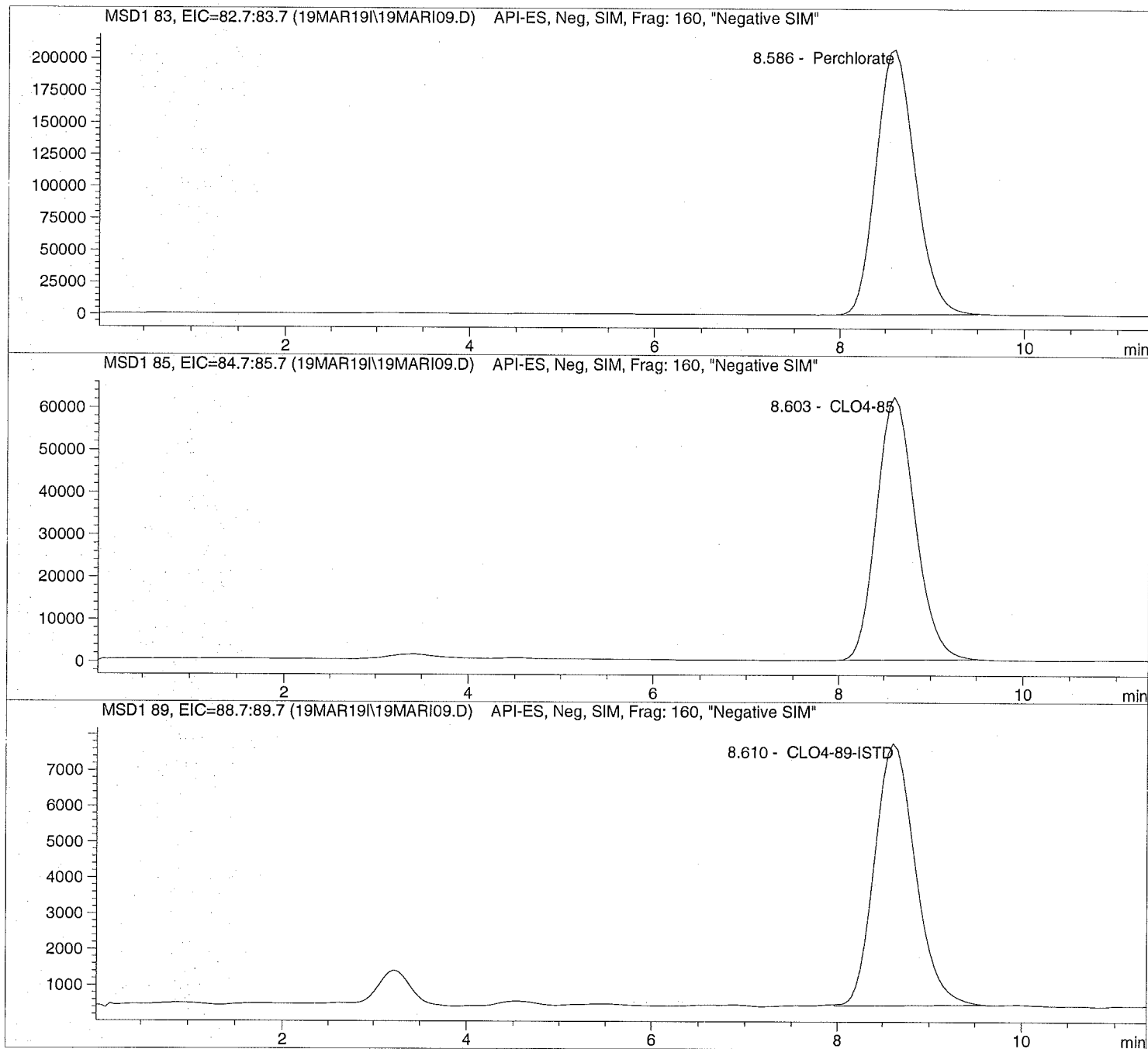
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date:  3/19/2019  10:59:22           Seq Line:           9
Sample Name:    CLO4@ 75.ug/L                 Location:           Vial 79
Acq Operator:   TNB                           Inj. No.:          1
                                           Inj. Vol.:         30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 75.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D

Sample Name: ICAL Verf@10ug/L

Injection Date: 3/19/2019 11:12:42

Seq Line: 10

Sample Name: ICAL Verf@10ug/L

Location: Vial 80

Acq Operator: TNB

Inj. No.: 1

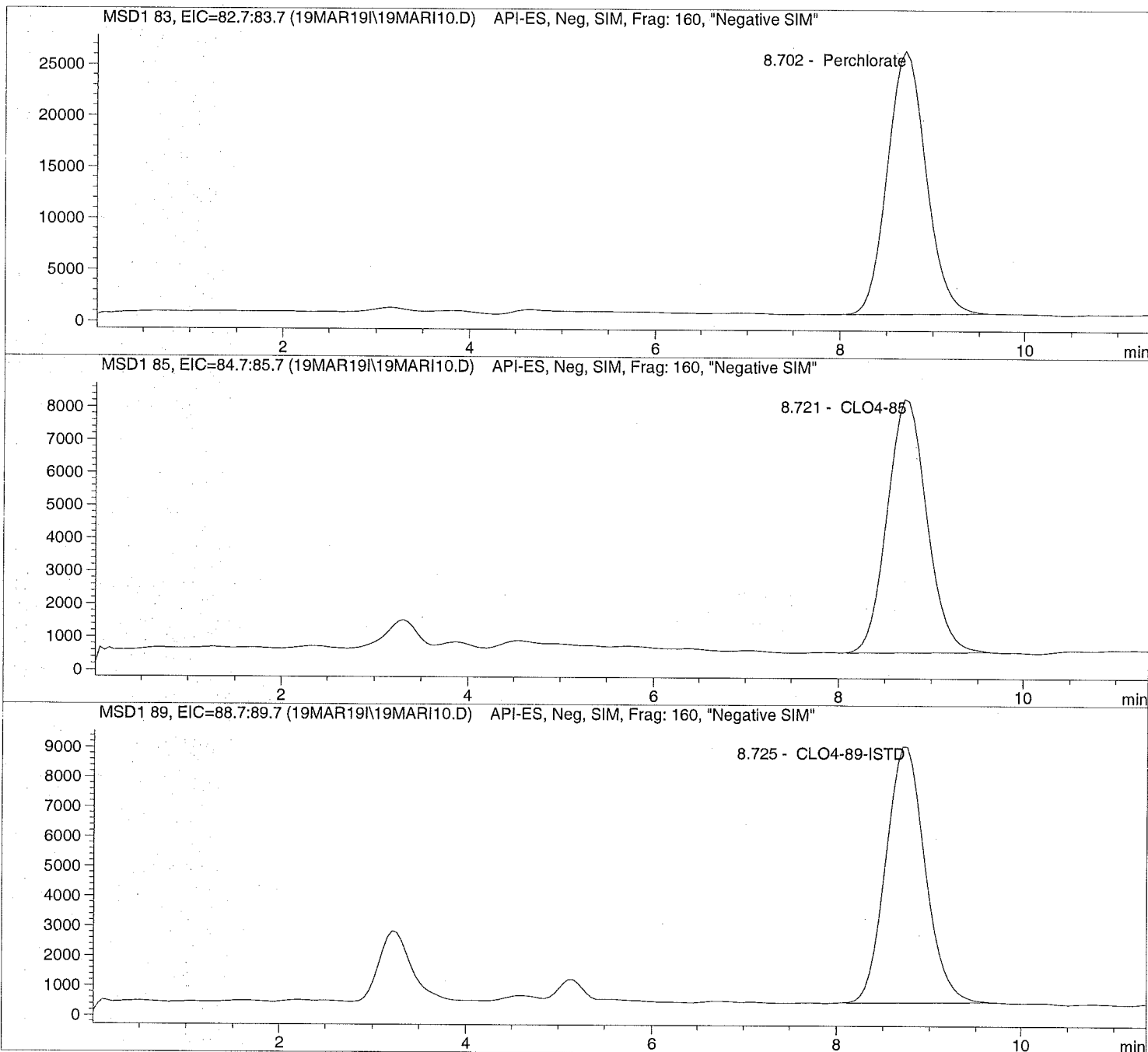
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```
=====
Injection Date: 3/19/2019 11:12:42      Seq Line: 10
Sample Name: ICAL Verf@10ug/L      Location: Vial 80
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22
```

Perchlorate analysis

Sample Information

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 10.000
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

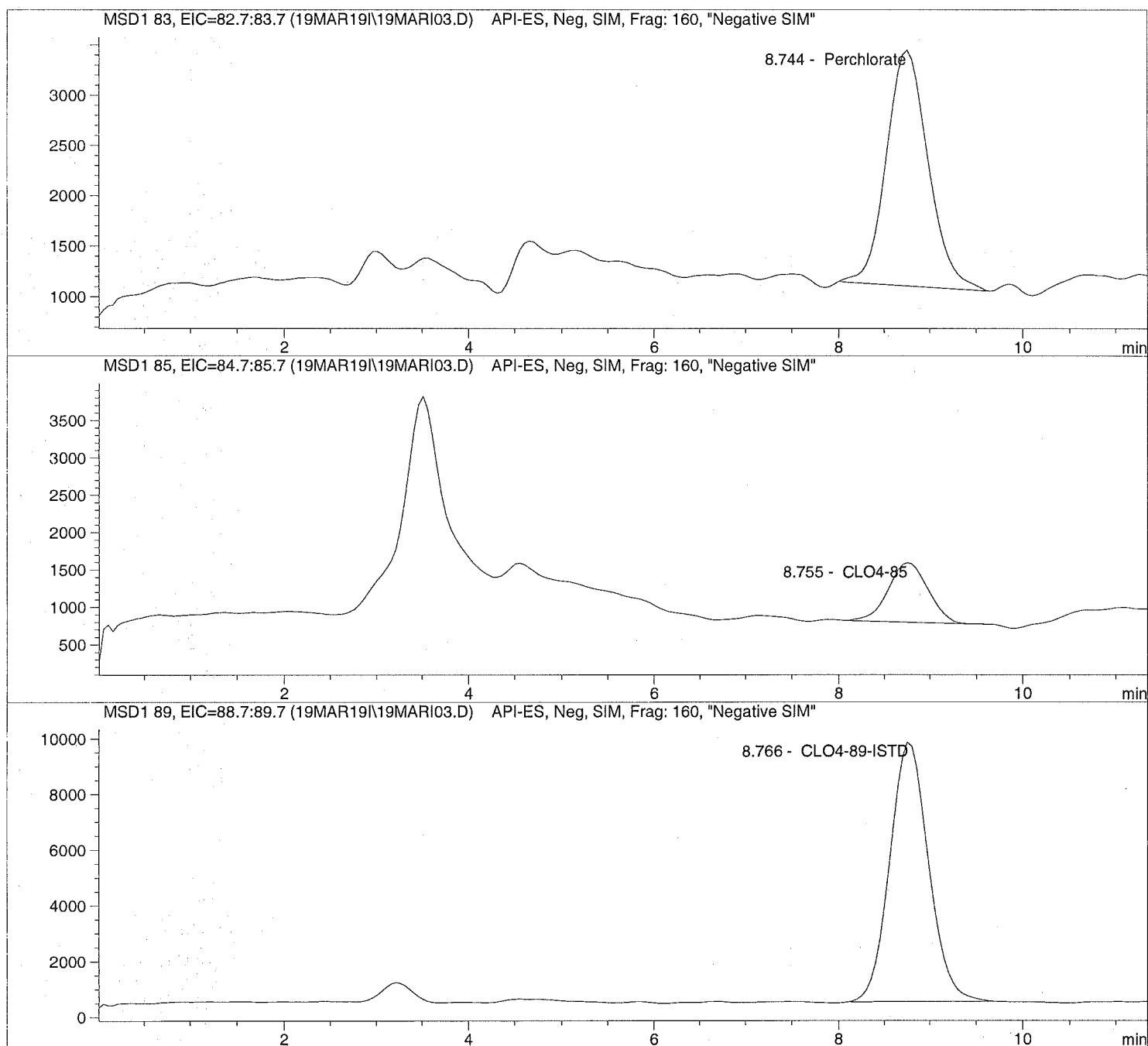
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line:          3
Sample Name:    CLO4@ 1.0ug/L           Location:          Vial 73
Acq Operator:  TNB                      Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:38:25
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 1.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



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www.alsglobal.com

April 01, 2019

Analytical Report for Service Request No: K1902558

RJ Modashia
ALS Laboratory Group
10450 Stancliff Road
Suite 210
Houston, TX 77099-4338

RE: ALS Houston DOD TOC

Dear RJ,

Enclosed are the results of the sample(s) submitted to our laboratory March 26, 2019
For your reference, these analyses have been assigned our service request number **K1902558**.

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. The test results meet requirements of the current NELAP standards, where applicable, and except as noted in the laboratory case narrative provided. For a specific list of NELAP-accredited analytes, refer to the certifications section at www.alsglobal.com. All results are intended to be considered in their entirety, and ALS Group USA Corp. dba ALS Environmental (ALS) is not responsible for use of less than the complete report. Results apply only to the items submitted to the laboratory for analysis and individual items (samples) analyzed, as listed in the report.

Please contact me if you have any questions. My extension is 3350. You may also contact me via email at Kelley.Lovejoy@alsglobal.com.

Respectfully submitted,

ALS Group USA, Corp. dba ALS Environmental

Kelley Lovejoy
Project Manager



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Table of Contents

Acronyms

Qualifiers

State Certifications, Accreditations, And Licenses

Case Narrative

Chain of Custody

General Chemistry

Raw Data

 General Chemistry

Acronyms

ASTM	American Society for Testing and Materials
A2LA	American Association for Laboratory Accreditation
CARB	California Air Resources Board
CAS Number	Chemical Abstract Service registry Number
CFC	Chlorofluorocarbon
CFU	Colony-Forming Unit
DEC	Department of Environmental Conservation
DEQ	Department of Environmental Quality
DHS	Department of Health Services
DOE	Department of Ecology
DOH	Department of Health
EPA	U. S. Environmental Protection Agency
ELAP	Environmental Laboratory Accreditation Program
GC	Gas Chromatography
GC/MS	Gas Chromatography/Mass Spectrometry
LOD	Limit of Detection
LOQ	Limit of Quantitation
LUFT	Leaking Underground Fuel Tank
M	Modified
MCL	Maximum Contaminant Level is the highest permissible concentration of a substance allowed in drinking water as established by the USEPA.
MDL	Method Detection Limit
MPN	Most Probable Number
MRL	Method Reporting Limit
NA	Not Applicable
NC	Not Calculated
NCASI	National Council of the Paper Industry for Air and Stream Improvement
ND	Not Detected
NIOSH	National Institute for Occupational Safety and Health
PQL	Practical Quantitation Limit
RCRA	Resource Conservation and Recovery Act
SIM	Selected Ion Monitoring
TPH	Total Petroleum Hydrocarbons
tr	Trace level is the concentration of an analyte that is less than the PQL but greater than or equal to the MDL.

Inorganic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- E The result is an estimate amount because the value exceeded the instrument calibration range.
- J The result is an estimated value.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.
- H The holding time for this test is immediately following sample collection. The samples were analyzed as soon as possible after receipt by the laboratory.

Metals Data Qualifiers

- # The control limit criteria is not applicable. See case narrative.
- J The result is an estimated value.
- E The percent difference for the serial dilution was greater than 10%, indicating a possible matrix interference in the sample.
- M The duplicate injection precision was not met.
- N The Matrix Spike sample recovery is not within control limits. See case narrative.
- S The reported value was determined by the Method of Standard Additions (MSA).
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- W The post-digestion spike for furnace AA analysis is out of control limits, while sample absorbance is less than 50% of spike absorbance.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- + The correlation coefficient for the MSA is less than 0.995.
- Q See case narrative. One or more quality control criteria was outside the limits.

Organic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- A A tentatively identified compound, a suspected aldol-condensation product.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- C The analyte was qualitatively confirmed using GC/MS techniques, pattern recognition, or by comparing to historical data.
- D The reported result is from a dilution.
- E The result is an estimated value.
- J The result is an estimated value.
- N The result is presumptive. The analyte was tentatively identified, but a confirmation analysis was not performed.
- P The GC or HPLC confirmation criteria was exceeded. The relative percent difference is greater than 40% between the two analytical results.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a chromatographic interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.

Additional Petroleum Hydrocarbon Specific Qualifiers

- F The chromatographic fingerprint of the sample matches the elution pattern of the calibration standard.
- L The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of lighter molecular weight constituents than the calibration standard.
- H The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of heavier molecular weight constituents than the calibration standard.
- O The chromatographic fingerprint of the sample resembles an oil, but does not match the calibration standard.
- Y The chromatographic fingerprint of the sample resembles a petroleum product eluting in approximately the correct carbon range, but the elution pattern does not match the calibration standard.
- Z The chromatographic fingerprint does not resemble a petroleum product.

**ALS Group USA Corp. dba ALS Environmental (ALS) - Kelso
State Certifications, Accreditations, and Licenses**

Agency	Web Site	Number
Alaska DEH	http://dec.alaska.gov/eh/lab/cs/csapproval.htm	UST-040
Arizona DHS	http://www.azdhs.gov/lab/license/env.htm	AZ0339
Arkansas - DEQ	http://www.adeq.state.ar.us/techsvs/labcert.htm	88-0637
California DHS (ELAP)	http://www.cdph.ca.gov/certlic/labs/Pages/ELAP.aspx	2795
DOD ELAP	http://www.denix.osd.mil/edqw/Accreditation/AccreditedLabs.cfm	L16-58-R4
Florida DOH	http://www.doh.state.fl.us/lab/EnvLabCert/WaterCert.htm	E87412
Hawaii DOH	http://health.hawaii.gov/	-
ISO 17025	http://www.pjllabs.com/	L16-57
Louisiana DEQ	http://www.deq.louisiana.gov/page/la-lab-accreditation	03016
Maine DHS	http://www.maine.gov/dhhs/	WA01276
Minnesota DOH	http://www.health.state.mn.us/accreditation	053-999-457
Nevada DEP	http://ndep.nv.gov/bsdw/labservice.htm	WA01276
New Jersey DEP	http://www.nj.gov/dep/enforcement/oqa.html	WA005
New York - DOH	https://www.wadsworth.org/regulatory/elap	12060
North Carolina DEQ	https://deq.nc.gov/about/divisions/water-resources/water-resources-data/water-sciences-home-page/laboratory-certification-branch/non-field-lab-certification	605
Oklahoma DEQ	http://www.deq.state.ok.us/CSDnew/labcert.htm	9801
Oregon – DEQ (NELAP)	http://public.health.oregon.gov/LaboratoryServices/EnvironmentalLaboratoryAccreditation/Pages/index.aspx	WA100010
South Carolina DHEC	http://www.scdhec.gov/environment/EnvironmentalLabCertification/	61002
Texas CEQ	http://www.tceq.texas.gov/field/qa/env_lab_accreditation.html	T104704427
Washington DOE	http://www.ecy.wa.gov/programs/eap/labs/lab-accreditation.html	C544
Wyoming (EPA Region 8)	https://www.epa.gov/region8-waterops/epa-region-8-certified-drinking-water	-
Kelso Laboratory Website	www.alsglobal.com	NA

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. A complete listing of specific NELAP-certified analytes, can be found in the certification section at www.ALSGlobal.com or at the accreditation bodies web site.

Please refer to the certification and/or accreditation body's web site if samples are submitted for compliance purposes. The states highlighted above, require the analysis be listed on the state certification if used for compliance purposes and if the method/analyte is offered by that state.



Case Narrative

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com



Client: ALS Environmental - US
Project: ALS Houston DOD TOC
Sample Matrix: Water

Service Request: K1902558
Date Received: 03/26/2019

CASE NARRATIVE

All analyses were performed consistent with the quality assurance program of ALS Group. This report contains analytical results and quality control parameters for samples designated for Tier IV validation deliverables, which includes summary forms and all of the associated raw data for each of the analyses. When appropriate to the method, method blank results have been reported with each analytical test.

Sample Receipt:

One water sample was received for analysis at ALS laboratory on 03/26/2019. The sample was received in good condition and consistent with the accompanying chain of custody form. Unless otherwise noted, samples were stored in a refrigerator at 4°C upon receipt at the laboratory.

General Chemistry:

No significant anomalies were noted with this analysis.

Approved by Kelley Lovejoy

Date 04/01/2019



Chain of Custody

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
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10450 Stancliff Rd, Ste 210
 Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

K1902558

Subcontract Chain of Custody

COC ID: 10959

SUBCONTRACT TO:

ALS Environmental Kelso
 1317 S. 13th Avenue
 Kelso, WA 98626

Phone: +1 360 501 3312

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19031160
TSR: Danielle Winnings

	LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
	ANALYSIS REQUESTED			DUE DATE
1.	HS19031160-01	LH18/24-SP650_032119	Water	21 Mar 2019 14:00
	TOC Analysis for DOD Level IV			01 Apr 2019

Comments: Please analyze for the analysis listed above.
 Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: *R. C. Age* Date/Time: *3/25/19*
 Received By: *H. Morrow ALS-Kelso* Date/Time: *3/26/19 0940*
 Cooler ID(s): _____ Temperature(s): _____

RIGHT SOLUTIONS | RIGHT PARTNER



PC KL

Cooler Receipt and Preservation Form

Client ALS Houston Service Request K1902558

Received: 3/26/19 Opened: 3/26/19 By: Km Unloaded: 3/26/19 By: Km

- 1. Samples were received via? USPS Fed Ex UPS DHL PDX Courier Hand Delivered
- 2. Samples were received in: (circle) Cooler Box Envelope Other NA
- 3. Were custody seals on coolers? NA Y N If yes, how many and where? 1 Front
- If present, were custody seals intact? Y N If present, were they signed and dated? Y N

Raw Cooler Temp	Corrected Cooler Temp	Raw Temp Blank	Corrected Temp Blank	Corr. Factor	Thermometer ID	Cooler/COC ID	Tracking Number	NA	Filed
<u>0.6</u>	<u>0.3</u>	<u>N/A</u>	<u>N/A</u>	<u>+0.3</u>	<u>384</u>	<u>10959</u>	<u>4809 7832 DFB</u>		

- 4. Packing material: Inserts Baggies Bubble Wrap Gel Packs Wet Ice Dry Ice Sleeves
- 5. Were custody papers properly filled out (ink, signed, etc.)? NA Y N
- 6. Were samples received in good condition (temperature, unbroken)? Indicate in the table below. NA Y N
If applicable, tissue samples were received: Frozen Partially Thawed Thawed
- 7. Were all sample labels complete (i.e analysis, preservation, etc.)? NA Y N
- 8. Did all sample labels and tags agree with custody papers? Indicate major discrepancies in the table on page 2. NA Y N
- 9. Were appropriate bottles/containers and volumes received for the tests indicated? NA Y N
- 10. Were the pH-preserved bottles (see SMO GEN SOP) received at the appropriate pH? Indicate in the table below NA Y N
- 11. Were VOA vials received without headspace? Indicate in the table below. NA Y N
- 12. Was C12/Res negative? NA Y N

Sample ID on Bottle	Sample ID on COC	Identified by:

Sample ID	Bottle Count	Bottle Type	Out of Temp	Head-space	Broke	pH	Reagent	Volume added	Reagent Lot Number	Initials	Time

Notes, Discrepancies, & Resolutions: _____

RUSH



General Chemistry

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Analytical Report

Client: ALS Environmental - US
Project: ALS Houston DOD TOC
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1902558
Date Collected: 03/21/19
Date Received: 03/26/19
Units: mg/L
Basis: NA

Carbon, Total Organic

Sample Name	Lab Code	Result	LOQ	LOD	MDL	Dil.	Date Analyzed	Q
LH18/24-SP650_032119	K1902558-001	2.50	0.50	0.20	0.07	1	03/28/19 22:24	
Method Blank	K1902558-MB	ND U	0.50	0.20	0.07	1	03/29/19 00:17	

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: ALS Houston DOD TOC
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1902558
Date Collected: 03/21/19
Date Received: 03/26/19

Units: mg/L
Basis: NA

Replicate Sample Summary
Carbon, Total Organic

Sample Name:	Lab Code:	MRL	LOQ	MDL	Sample Result	Duplicate Result	Average	RPD	RPD Limit	Date Analyzed
Batch QC	K1902526-001DUP	50	20	7	353	348	350	2	10	03/28/19
Batch QC	K1902526-002DUP	50	20	7	378	377	378	<1	10	03/28/19
LH18/24-SP650_032119	K1902558-001DUP	0.50	0.20	0.07	2.50	2.49	2.49	<1	10	03/28/19
Batch QC	K1902567-001DUP	0.50	0.20	0.07	11.3	11.2	11.3	<1	10	03/29/19
Batch QC	K1902567-002DUP	0.50	0.20	0.07	3.20	3.10	3.15	3	10	03/29/19
Batch QC	K1902567-003DUP	0.50	0.20	0.07	3.30	3.25	3.27	1	10	03/29/19
Batch QC	K1902567-004DUP	0.50	0.20	0.07	1.33	1.32	1.32	<1	10	03/29/19
Batch QC	K1902567-005DUP	0.50	0.20	0.07	2.44	2.37	2.41	3	10	03/29/19
Batch QC	K1902603-001DUP	0.50	0.20	0.07	3.10	3.04	3.07	2	10	03/29/19
Batch QC	K1902603-002DUP	0.50	0.20	0.07	1.46	1.46	1.46	<1	10	03/29/19
Batch QC	K1902603-003DUP	0.50	0.20	0.07	1.61	1.65	1.63	2	10	03/29/19
Batch QC	K1902603-004DUP	0.50	0.20	0.07	1.69	1.65	1.67	3	10	03/29/19

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: ALS Houston DOD TOC
Sample Matrix: Water

Service Request: K1902558
Date Collected: N/A
Date Received: N/A
Date Analyzed: 03/29/19
Date Extracted: NA

Matrix Spike Summary
Carbon, Total Organic

Sample Name: Batch QC
Lab Code: K1902603-002
Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA

Matrix Spike
K1902603-002MS

Analyte Name	Sample Result	Result	Spike Amount	% Rec	% Rec Limits
Carbon, Total Organic	1.46	29.3	25.0	112	83-117

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: ALS Houston DOD TOC
Sample Matrix: Water

Service Request: K1902558
Date Analyzed: 03/29/19
Date Extracted: NA

Lab Control Sample Summary
Carbon, Total Organic

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA
Analysis Lot: 629967

Sample Name	Lab Code	Result	Spike Amount	% Rec	% Rec Limits
Lab Control Sample	K1902558-LCS	26.6	25.0	106	83-117

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: ALS Houston DOD TOC

Service Request: K1902558

Continuing Calibration Verification (CCV) Summary

Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis		Date	True	Measured	Percent	Acceptance Limits
	Lot	Lab Code	Analyzed	Value	Value	Recovery	
CCV1	629967	KQ1904109-05	03/28/19 18:38	25.0	26.9	107	90-110
CCV2	629967	KQ1904109-06	03/28/19 23:48	25.0	26.0	104	90-110
CCV3	629967	KQ1904109-07	03/29/19 04:59	25.0	26.0	104	90-110
CCV4	629967	KQ1904109-08	03/29/19 09:43	25.0	26.3	105	90-110

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: ALS Houston DOD TOC

Service Request: K1902558

Continuing Calibration Blank (CCB) Summary
Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis Lot	Lab Code	Date Analyzed	LOQ	LOD	MDL	Result	Q
CCB1	629967	KQ1904109-01	03/28/19 18:52	0.50	0.20	0.07	0.09	J
CCB2	629967	KQ1904109-02	03/29/19 00:03	0.50	0.20	0.07	ND	U
CCB3	629967	KQ1904109-03	03/29/19 05:14	0.50	0.20	0.07	ND	U
CCB4	629967	KQ1904109-04	03/29/19 09:58	0.50	0.20	0.07	ND	U



Raw Data

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General Chemistry

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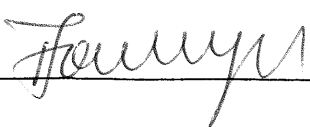
Work Request # ^{Original} () K1902469, 2514, 2526, 2558, 2567, 2603, 2607, 2623
 Tier: II IV II IV II II I II
 Date Analyzed: 3/28/19 TOC: 629966,
629967
 Analyst: BCD Run # DOC: 629968
 Analysis: TOC/DOC

DATA QUALITY REPORT INORGANICS

Explain any "no" responses to questions below, and any corrective actions in the comments section below.

1. Is the method name and number correct and appropriate? yes/no/NA
2. Holding times met for all analyses and for all samples? yes/no/NA
3. Are calculations correct? yes/no/NA
4. Is the reporting basis correct? (Dry Weight) yes/no/NA
5. All quality control criteria met? yes/no
6. Is the calibration curve correlation coefficient ≥ 0.995 ? yes/no/NA
7. MBs, CCVs, CCBs, LCSs, Dups, and Spikes, analyzed at proper frequency? yes/no/NA
8. Are ICVs, CCVs, and CCBs all within acceptance limits? yes/no/NA
9. Are results for methods blanks all ND? yes/no/NA
10. Are all QC samples within acceptance criteria? (LCS % rec, MS/DMS % rec, DUP or MS/DMS RPDs, etc.) yes/no/NA
11. Are all exceptions explained? yes/no/NA
12. Have all applicable service requests been reviewed? yes/no/NA
13. Are all samples labeled correctly? yes/no/NA
14. Have all instructions on the service request been followed? (e.g. Special MRLs, QC on a specific sample, Form V) yes/no/NA
15. Are detection limits and units reported correctly? yes/no/NA
16. Is the unused space on the benchsheet crossed out? yes/no/NA
17. Was analysis turned in by the due date? (n-2) (If not record SR#) yes/no/NA

COMMENTS: K1902607-1/14 report a high %RSD. However, these samples are less than 5x the MRL.

Final Approved by: 

Date: 03/29/19

DQREPORT

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629966 Method/Testcode: 415.1/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902469-001	Carbon, Total Organic	N/A		Water	3.83 mg/L	10 ml	3.83 mg/L	1	0.07	0.50			3/28/19 19:07	N	II
K1902469-002	Carbon, Total Organic	N/A		Water	7.70 mg/L	10 ml	7.70 mg/L	1	0.07	0.50			3/28/19 19:35	N	II
K1902469-003	Carbon, Total Organic	N/A		Water	6.83 mg/L	10 ml	6.83 mg/L	1	0.07	0.50			3/28/19 20:03	N	II
K1902469-004	Carbon, Total Organic	N/A		Water	3.65 mg/L	10 ml	3.65 mg/L	1	0.07	0.50			3/28/19 20:31	N	II
K1902469-005	Carbon, Total Organic	N/A		Water	7.13 mg/L	10 ml	7.13 mg/L	1	0.07	0.50			3/28/19 20:59	N	II
K1902469-006	Carbon, Total Organic	N/A		Water	6.53 mg/L	10 ml	6.53 mg/L	1	0.07	0.50			3/28/19 21:27	N	II
K1902514-001	Carbon, Total Organic	N/A		Water	0.33 mg/L	10 ml	0.33 mg/L	J 1	0.07	0.50			3/28/19 16:16	Y	IV
K1902514-002	Carbon, Total Organic	N/A		Water	0.56 mg/L	10 ml	0.56 mg/L	1	0.07	0.50			3/28/19 17:13	N	IV
K1902514-003	Carbon, Total Organic	N/A		Water	0.21 mg/L	10 ml	0.21 mg/L	J 1	0.07	0.50			3/28/19 17:41	N	IV
K1902514-004	Carbon, Total Organic	N/A		Water	0.36 mg/L	10 ml	0.36 mg/L	J 1	0.07	0.50			3/28/19 18:09	N	IV
KQ1904107-01	Carbon, Total Organic	CCB		Water	0.10 mg/L	10 ml	0.10 mg/L	J 1	0.07	0.50			3/28/19 14:49	N	IV
KQ1904107-02	Carbon, Total Organic	CCB		Water	0.09 mg/L	10 ml	0.09 mg/L	J 1	0.07	0.50			3/28/19 18:52	N	IV
KQ1904107-03	Carbon, Total Organic	CCB		Water	0.00 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 00:03	N	IV
KQ1904107-04	Carbon, Total Organic	CCV		Water	26.99 mg/L	10 ml	27.0 mg/L	1			108		3/28/19 14:34	N	IV
KQ1904107-05	Carbon, Total Organic	CCV		Water	26.85 mg/L	10 ml	26.9 mg/L	1			108		3/28/19 18:38	N	IV
KQ1904107-06	Carbon, Total Organic	CCV		Water	25.98 mg/L	10 ml	26.0 mg/L	1			104		3/28/19 23:48	N	IV
KQ1904107-07	Carbon, Total Organic	MB		Water	0.10 mg/L	10 ml	0.10 mg/L	J 1	0.07	0.50			3/28/19 15:04	N	IV
KQ1904107-08	Carbon, Total Organic	LCS		Water	27.22 mg/L	10 ml	27.2 mg/L	1	0.07	0.50	109		3/28/19 15:18	N	IV
KQ1904107-09	Carbon, Total Organic	MS	K1902514-001	Water	28.37 mg/L	10 ml	28.4 mg/L	1	0.07	0.50	112		3/28/19 16:44	N	IV
KQ1904107-10	Carbon, Total Organic	DUP	K1902514-001	Water	0.35 mg/L	10 ml	0.35 mg/L	J 1	0.07	0.50		5	3/28/19 16:16	N	IV
KQ1904107-11	Carbon, Total Organic	DUP	K1902514-002	Water	0.51 mg/L	10 ml	0.51 mg/L	1	0.07	0.50		8	3/28/19 17:13	N	IV
KQ1904107-12	Carbon, Total Organic	DUP	K1902514-003	Water	0.21 mg/L	10 ml	0.21 mg/L	J 1	0.07	0.50		1	3/28/19 17:41	N	IV
KQ1904107-13	Carbon, Total Organic	DUP	K1902514-004	Water	0.35 mg/L	10 ml	0.35 mg/L	J 1	0.07	0.50		<1	3/28/19 18:09	N	IV
KQ1904107-14	Carbon, Total Organic	DUP	K1902469-001	Water	3.71 mg/L	10 ml	3.71 mg/L	1	0.07	0.50		3	3/28/19 19:07	N	II
KQ1904107-15	Carbon, Total Organic	DUP	K1902469-002	Water	7.63 mg/L	10 ml	7.63 mg/L	1	0.07	0.50		<1	3/28/19 19:35	N	II
KQ1904107-16	Carbon, Total Organic	DUP	K1902469-003	Water	6.77 mg/L	10 ml	6.77 mg/L	1	0.07	0.50		<1	3/28/19 20:03	N	II
KQ1904107-17	Carbon, Total Organic	DUP	K1902469-004	Water	3.53 mg/L	10 ml	3.53 mg/L	1	0.07	0.50		3	3/28/19 20:31	N	II
KQ1904107-18	Carbon, Total Organic	DUP	K1902469-005	Water	7.18 mg/L	10 ml	7.18 mg/L	1	0.07	0.50		<1	3/28/19 20:59	N	II
KQ1904107-19	Carbon, Total Organic	DUP	K1902469-006	Water	6.45 mg/L	10 ml	6.45 mg/L	1	0.07	0.50		1	3/28/19 21:27	N	II

03/29/19
Freezell

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629967 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902526-001	Carbon, Total Organic	N/A		Water	3.53 mg/L	10 ml	353 mg/L	100	7	50			3/28/19 22:52	N	II
K1902526-002	Carbon, Total Organic	N/A		Water	3.78 mg/L	10 ml	378 mg/L	100	7	50			3/28/19 23:20	N	II
K1902558-001	Carbon, Total Organic	N/A		Water	2.50 mg/L	10 ml	2.50 mg/L	1	0.07	0.50			3/28/19 22:24	N	IV
K1902567-001	Carbon, Total Organic	N/A		Ground Water	11.32 mg/L	10 ml	11.3 mg/L	1	0.07	0.50			3/29/19 00:47	N	II
K1902567-002	Carbon, Total Organic	N/A		Ground Water	3.20 mg/L	10 ml	3.20 mg/L	1	0.07	0.50			3/29/19 01:15	N	II
K1902567-003	Carbon, Total Organic	N/A		Ground Water	3.30 mg/L	10 ml	3.30 mg/L	1	0.07	0.50			3/29/19 01:43	N	II
K1902567-004	Carbon, Total Organic	N/A		Ground Water	1.33 mg/L	10 ml	1.33 mg/L	1	0.07	0.50			3/29/19 02:11	N	II
K1902567-005	Carbon, Total Organic	N/A		Ground Water	2.45 mg/L	10 ml	2.44 mg/L	1	0.07	0.50			3/29/19 03:07	N	II
K1902603-001	Carbon, Total Organic	N/A		Ground Water	3.10 mg/L	10 ml	3.10 mg/L	1	0.07	0.50			3/29/19 04:03	N	II
K1902603-002	Carbon, Total Organic	N/A		Ground Water	1.46 mg/L	10 ml	1.46 mg/L	1	0.07	0.50			3/29/19 05:28	N	II
K1902603-003	Carbon, Total Organic	N/A		Ground Water	1.61 mg/L	10 ml	1.61 mg/L	1	0.07	0.50			3/29/19 06:26	N	II
K1902603-004	Carbon, Total Organic	N/A		Ground Water	1.69 mg/L	10 ml	1.69 mg/L	1	0.07	0.50			3/29/19 06:54	N	II
KQ1904109-01	Carbon, Total Organic	CCB		Water	0.09 mg/L	10 ml	0.09 mg/L	J 1	0.07	0.50			3/28/19 18:52	N	IV
KQ1904109-02	Carbon, Total Organic	CCB		Water	0.00 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 00:03	N	IV
KQ1904109-03	Carbon, Total Organic	CCB		Water	-0.07 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 05:14	N	IV
KQ1904109-04	Carbon, Total Organic	CCB		Water	-0.07 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 09:58	N	IV
KQ1904109-05	Carbon, Total Organic	CCV		Water	26.85 mg/L	10 ml	26.9 mg/L	1					3/28/19 18:38	N	IV
KQ1904109-06	Carbon, Total Organic	CCV		Water	25.98 mg/L	10 ml	26.0 mg/L	1			108		3/28/19 23:48	N	IV
KQ1904109-07	Carbon, Total Organic	CCV		Water	25.99 mg/L	10 ml	26.0 mg/L	1			104		3/29/19 04:59	N	IV
KQ1904109-08	Carbon, Total Organic	CCV		Water	26.27 mg/L	10 ml	26.3 mg/L	1			105		3/29/19 09:43	N	IV
KQ1904109-09	Carbon, Total Organic	LCS		Water	26.62 mg/L	10 ml	26.6 mg/L	1	0.07	0.50	106		3/29/19 00:32	N	IV
KQ1904109-10	Carbon, Total Organic	MB		Water	0.00 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 00:17	N	IV
KQ1904109-11	Carbon, Total Organic	MS	K1902603-002	Ground Water	29.33 mg/L	10 ml	29.3 mg/L	1	0.07	0.50	112		3/29/19 05:57	N	II
KQ1904109-12	Carbon, Total Organic	DUP	K1902558-001	Water	2.49 mg/L	10 ml	2.49 mg/L	1	0.07	0.50		<1	3/28/19 22:24	N	IV
KQ1904109-13	Carbon, Total Organic	DUP	K1902526-001	Water	3.48 mg/L	10 ml	348 mg/L	100	7	50		2	3/28/19 22:52	N	II
KQ1904109-14	Carbon, Total Organic	DUP	K1902526-002	Water	3.77 mg/L	10 ml	377 mg/L	100	7	50		<1	3/28/19 23:20	N	II
KQ1904109-15	Carbon, Total Organic	DUP	K1902567-001	Ground Water	11.24 mg/L	10 ml	11.2 mg/L	1	0.07	0.50		<1	3/29/19 00:47	N	II
KQ1904109-16	Carbon, Total Organic	DUP	K1902567-002	Ground Water	3.10 mg/L	10 ml	3.10 mg/L	1	0.07	0.50		3	3/29/19 01:15	N	II
KQ1904109-17	Carbon, Total Organic	DUP	K1902567-003	Ground Water	3.25 mg/L	10 ml	3.25 mg/L	1	0.07	0.50		1	3/29/19 01:43	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

03/29/19
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Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629967 Method/Testcode: SM 5310 C/TOC T

<u>Lab Code</u>	<u>Target Analytes</u>	<u>QC</u>	<u>Parent Sample</u>	<u>Matrix</u>	<u>Raw Result</u>	<u>Sample Amt.</u>	<u>Final Result</u>	<u>Dil</u>	<u>MDL</u>	<u>PQL</u>	<u>% Rec</u>	<u>% RSD</u>	<u>Date Analyzed</u>	<u>QC?</u>	<u>Tier</u>
KQ1904109-18	Carbon, Total Organic	DUP	K1902567-004	Ground Water	1.32 mg/L	10 ml	1.32 mg/L	1	0.07	0.50		<1	3/29/19 02:11	N	II
KQ1904109-19	Carbon, Total Organic	DUP	K1902567-005	Ground Water	2.37 mg/L	10 ml	2.37 mg/L	1	0.07	0.50		3	3/29/19 03:07	N	II
KQ1904109-20	Carbon, Total Organic	DUP	K1902603-001	Ground Water	3.04 mg/L	10 ml	3.04 mg/L	1	0.07	0.50		2	3/29/19 04:03	N	II
KQ1904109-21	Carbon, Total Organic	DUP	K1902603-002	Ground Water	1.46 mg/L	10 ml	1.46 mg/L	1	0.07	0.50		<1	3/29/19 05:28	N	II
KQ1904109-22	Carbon, Total Organic	DUP	K1902603-003	Ground Water	1.65 mg/L	10 ml	1.65 mg/L	1	0.07	0.50		2	3/29/19 06:26	N	II
KQ1904109-23	Carbon, Total Organic	DUP	K1902603-004	Ground Water	1.65 mg/L	10 ml	1.65 mg/L	1	0.07	0.50		3	3/29/19 06:54	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 629968 Method/Testcode: SM 5310 C/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902607-001	Carbon, Dissolved Organic (DOC)	N/A		Water	1.09 mg/L	10 ml	1.09 mg/L	1	0.07	0.50			3/29/19 08:46	N	I
K1902607-002	Carbon, Dissolved Organic (DOC)	N/A		Water	16.60 mg/L	10 ml	33.2 mg/L	2	0.2	1.0			3/29/19 10:42	N	I
K1902623-001	Carbon, Dissolved Organic (DOC)	N/A		Water	0.74 mg/L	10 ml	74 mg/L	100	7	50			3/29/19 11:10	N	II
K1902623-002	Carbon, Dissolved Organic (DOC)	N/A		Water	1.93 mg/L	10 ml	1.93 mg/L	1	0.07	0.50			3/29/19 12:06	N	II
KQ1904111-01	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.07 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 05:14	N	I
KQ1904111-02	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.07 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 09:58	N	I
KQ1904111-03	Carbon, Dissolved Organic (DOC)	CCB		Water	-0.04 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			3/29/19 13:17	N	I
KQ1904111-04	Carbon, Dissolved Organic (DOC)	CCV		Water	25.99 mg/L	10 ml	26.0 mg/L	1			104		3/29/19 04:59	N	I
KQ1904111-05	Carbon, Dissolved Organic (DOC)	CCV		Water	26.27 mg/L	10 ml	26.3 mg/L	1			105		3/29/19 09:43	N	I
KQ1904111-06	Carbon, Dissolved Organic (DOC)	CCV		Water	26.43 mg/L	10 ml	26.4 mg/L	1			106		3/29/19 13:02	N	I
KQ1904111-07	Carbon, Dissolved Organic (DOC)	MB		Water	-0.07 mg/L	10 mL	0.50 mg/L	U 1	0.07	0.50			3/29/19 10:12:00	N	I
KQ1904111-08	Carbon, Dissolved Organic (DOC)	LCS		Water	26.73 mg/L	10 ml	26.7 mg/L	1	0.07	0.50	107		3/29/19 10:27	N	I
KQ1904111-09	Carbon, Dissolved Organic (DOC)	MS	K1902607-001	Water	29.41 mg/L	10 ml	29.4 mg/L	1	0.07	0.50	113		3/29/19 09:14	N	I
KQ1904111-10	Carbon, Dissolved Organic (DOC)	DUP	K1902607-001	Water	1.27 mg/L	10 ml	1.27 mg/L	1	0.07	0.50		15*	3/29/19 08:46	N	I
KQ1904111-11	Carbon, Dissolved Organic (DOC)	DUP	K1902607-002	Water	16.05 mg/L	10 ml	32.1 mg/L	2	0.2	1.0		3	3/29/19 10:42	N	I
KQ1904111-12	Carbon, Dissolved Organic (DOC)	DUP	K1902623-001	Water	0.76 mg/L	10 ml	76 mg/L	100	7	50		2	3/29/19 11:10	N	II
KQ1904111-13	Carbon, Dissolved Organic (DOC)	DUP	K1902623-002	Water	1.92 mg/L	10 ml	1.92 mg/L	1	0.07	0.50		<1	3/29/19 12:06	N	II

03/29/19
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indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

TOC: 629966,
629967
DOC: 629968

Schedule: 03282019

Version: 6

Instrument: Fusion1

Last Saved by: Fusion1 (Fusion1)

Last Saved on: 2019/03/29 10:38 - Friday

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Blank)	Blank	Reagent/Acid Blank		1	True	Ready
D	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
1	Sample	MB1	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
2	Sample	ICS	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
3	Sample	H2SO4 W142572	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
4	Sample	K1902514-001.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
5	Sample	K1902514-001.05 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
6	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
7	Sample	K1902514-002.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
8	Sample	K1902514-003.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
9	Sample	K1902514-004.05	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
10	Sample	K1902469-001.16	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
11	Sample	K1902469-002.15	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
12	Sample	K1902469-003.15	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
13	Sample	K1902469-004.15	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
14	Sample	K1902469-005.16	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
15	Sample	K1902469-006.16	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
16	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
17	Sample	K1902558-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
18	Sample	K1902526-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
19	Sample	K1902526-002.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
20	Sample	MB2	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
21	Sample	K1902567-001.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
22	Sample	K1902567-002.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
23	Sample	K1902567-003.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
24	Sample	K1902567-004.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
25	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
26	Sample	K1902567-005.03	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
27	Sample	K1902603-001.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
28	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
29	Sample	K1902603-002.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
30	Sample	K1902603-002.03 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
31	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
32	Sample	K1902603-003.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
33	Sample	K1902603-004.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
34	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
35	Sample	FB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
36	Sample	K1902607-001.03 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
37	Sample	K1902607-001.03 ms doc	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
38	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready

Printed on: March 29, 2019 13:37:28

03/29/19
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Schedule: 03282019

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
39	Sample	MB3	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
40	Sample	K1902607-002.03 doc 2x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
41	Sample	K1902623-001.07 doc 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
42	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
43	Sample	K1902623-002.07 doc	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
44	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
					False	

Fusion Report - 03282019

Thursday, March 28, 2019 12:22 PM

(View - Reps, Unused Reps, Meta-Data, Signature, History)
 Printed on 2019/03/29 13:37 -
 Friday

Report Summary Information

Company Location: Gen Chem Lab
 Schedule Name: 03282019
 Instrument Name: Fusion1
 Report Version: 1 of 1
 Report Creation by Operators (schedule version): Fusion1 (Fusion1) (v2)
 Fusion1 (Fusion1) (v3)
 Fusion1 (Fusion1) (v4)
 Fusion1 (Fusion1) (v5)
 Fusion1 (Fusion1) (v6)
 Comment:

Engine 1.1.5.1
 Version:
 Firmware 1.2.0696
 Version:
 Connection: RS232 COM1

Report Results

03/29/19
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Sample Type: Clean From Schedule Version 2

Pos	Analysis Type	Sample ID	Start Time
♦ (clean)		Clean	2019/03/28 12:22

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	12.77	15.74	2.97	49.67	05:24
2	TC Clean	5.47	8.34	2.86	49.86	07:16
3	TC Clean	2.02	5.15	3.13	50.03	03:57
4	TC Clean	1.43	4.57	3.14	50.04	03:56

Sample Type: Clean From Schedule Version 3

Pos	Analysis Type	Sample ID	Start Time
♦ (clean)		Clean	2019/03/28 12:48

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	10.76	13.67	2.90	49.66	05:24
2	TC Clean	3.58	6.55	2.98	50.02	04:05
3	TC Clean	1.57	4.51	2.94	50.03	03:48

4	TC Clean	1.27	4.29	3.02	50.01	03:56
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Sample Type: Clean From Schedule Version 4

Pos	Analysis Type	Sample ID	Start Time
♦ (clean)		Clean	2019/03/28 13:20

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	10.78	13.56	2.78	49.70	05:25
2	TC Clean	4.15	7.11	2.95	50.00	04:05
3	TC Clean	1.96	4.96	3.00	50.00	03:48
4	TC Clean	1.27	4.27	3.00	50.03	03:56

Sample Type: Blank (Creating v1240) From Schedule Version 4

Pos	Analysis Type	Sample ID	Start Time
♦ (blank)		Reagent/Acid Blank	2019/03/28 13:42

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	0.65	3.57	2.91	49.63	05:24
2	TC Clean	3.66	6.78	3.12	50.02	04:06
3	TC Clean	2.11	5.18	3.07	50.01	03:58
4	TC Clean	1.45	4.47	3.01	50.04	03:55
5	Reagent Blank	7.18	9.96	2.77	49.83	08:13
6	Acid Blank	1.31	4.25	2.94	49.74	05:33

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ D	TOC	RB	0.5296 ppm	0.0000 ppm	0.0000%	2019/03/28 14:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5296	5.2960	12.59	15.63	3.04	50.04	10:30

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 8.9951 (IC) (v1240)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	27.0597 ppm (PASS)	0.0000 ppm	0%	2019/03/28 14:34

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	27.0597	270.5968	193.14	196.16	3.02	50.04	10:30

Completion State **Success Action** **Method** **Calibration** **STD Conc - Pos B**
 Success - Criteria met. Do Nothing CAS_salt_010711 (v4) CAS_salt_010711 (v30) 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.1693 ppm (PASS)	0.0000 ppm	0%	2019/03/28 14:49

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.1693	1.6935	10.61	13.62	3.00	50.02	10:32

Completion State **Success Action** **Method** **Calibration** **STD Conc - Pos D**
 Success - Criteria met. Do Nothing CAS_salt_010711 (v4) CAS_salt_010711 (v30) 0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 1	TOC	MB1	0.1693 ppm	0.0000 ppm	0.0000%	2019/03/28 15:04

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1693	1.6926	10.14	12.87	2.72	50.00	10:32

Dilution **Blank Contribution** **Method** **Calibration**
 1:10 (TC) 8.9951 (IC) (v1240) CAS_salt_010711 (v4) CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	27.2866 ppm (PASS)	0.0000 ppm	0%	2019/03/28 15:18

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	27.2866	272.8655	194.68	197.53	2.85	49.98	10:29
Completion State		Success Action		Method		Calibration		STD Conc - Pos C		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		25 ppmC		

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
* 2	TOC	ICS	0.5331 ppm	0.0000 ppm	0.0000%	2019/03/28 15:33

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5331	5.3314	12.61	15.59	2.97	49.96	10:31

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
* 3	TOC	H2SO4 W142572	0.3003 ppm	0.0120 ppm	3.9900%	2019/03/28 15:48

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2918	2.9183	10.98	13.78	2.80	49.93	10:26
2	TOC	0.3088	3.0877	11.09	13.91	2.82	49.96	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
* 4	TOC	K1902514-001.05	0.4081 ppm	0.0110 ppm	2.7100%	2019/03/28 16:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4003	4.0025	11.71	14.55	2.83	49.90	10:25
2	TOC	0.4159	4.1587	11.82	14.78	2.96	49.88	10:30

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
* 5	TOC	K1902514-001.05 ms	28.4356 ppm	0.0000 ppm	0.0000%	2019/03/28 16:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	28.4356	284.3556	202.01	205.02	3.01	49.86	10:32

Dilution **Blank Contribution** **Method** **Calibration**

1:10 (TC) 8.9951 (IC) CAS_salt_010711 CAS_salt_010711
(v1240) (v4) (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
6	TOC	RB	0.3207 ppm	0.0000 ppm	0.0000%	2019/03/28 16:58

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3207	3.2070	11.17	14.08	2.91	49.90	10:32

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Sample

From Schedule Version 5

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
7	TOC	K1902514-002.05	0.6042 ppm	0.0301 ppm	4.9800%	2019/03/28 17:13

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.6255	6.2551	13.24	16.17	2.93	49.81	10:27
2	TOC	0.5829	5.8293	12.95	15.87	2.92	49.82	10:26

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
8	TOC	K1902514-003.05	0.2805 ppm	0.0019 ppm	0.6700%	2019/03/28 17:41

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2792	2.7916	10.89	13.82	2.93	49.85	10:29
2	TOC	0.2818	2.8181	10.91	13.94	3.04	49.71	10:25

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
9	TOC	K1902514-004.05	0.4246 ppm	0.0015 ppm	0.3400%	2019/03/28 18:09

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4256	4.2559	11.88	14.84	2.96	49.79	10:29
2	TOC	0.4235	4.2353	11.87	14.76	2.89	49.79	10:32

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.9205 ppm (PASS)	0.0000 ppm	0%	2019/03/28 18:38

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.9205	269.2046	192.20	195.11	2.92	49.79	10:32

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.1587 ppm (PASS)	0.0000 ppm	0%	2019/03/28 18:52

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.1587	1.5874	10.54	13.59	3.05	49.79	10:31

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample

From Schedule Version 5

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 10	TOC	K1902469-001.16	3.8357 ppm	0.0843 ppm	2.2000%	2019/03/28 19:07

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.8953	38.9527	35.44	38.16	2.73	49.80	10:31
2	TOC	3.7761	37.7609	34.63	37.52	2.90	49.82	10:31

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 11	TOC	K1902469-002.15	7.7316 ppm	0.0454 ppm	0.5900%	2019/03/28 19:35

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.7638	77.6375	61.70	64.74	3.05	49.85	10:26
2	TOC	7.6995	76.9952	61.26	64.12	2.86	49.87	10:27

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
12	TOC	K1902469-003.15	6.8677 ppm	0.0418 ppm	0.6100%	2019/03/28 20:03

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.8972	68.9722	55.81	58.63	2.82	49.88	10:26
2	TOC	6.8381	68.3814	55.41	58.28	2.86	49.92	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
13	TOC	K1902469-004.15	3.6574 ppm	0.0881 ppm	2.4100%	2019/03/28 20:31

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.7197	37.1967	34.24	37.13	2.88	49.94	10:28
2	TOC	3.5950	35.9504	33.40	36.42	3.02	49.96	10:29

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
14	TOC	K1902469-005.16	7.2204 ppm	0.0342 ppm	0.4700%	2019/03/28 20:59

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	7.1963	71.9628	57.84	60.75	2.90	49.99	10:26
2	TOC	7.2446	72.4460	58.17	61.12	2.95	50.02	10:25

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
15	TOC	K1902469-006.16	6.5588 ppm	0.0523 ppm	0.8000%	2019/03/28 21:27

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	6.5958	65.9580	53.77	56.71	2.95	50.03	10:28
2	TOC	6.5218	65.2184	53.26	56.21	2.95	50.08	10:28

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
16	TOC	RB	0.1506 ppm	0.0436 ppm	28.9800%	2019/03/28 21:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1198	1.1976	9.81	12.79	2.98	50.10	10:29
2	TOC	0.1815	1.8148	10.23	13.03	2.80	50.12	10:31

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
17	TOC	K1902558-001.01	2.5634 ppm	0.0089 ppm	0.3500%	2019/03/28 22:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.5697	25.6969	26.44	29.30	2.86	50.15	10:27
2	TOC	2.5572	25.5717	26.35	29.31	2.95	50.17	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
18	TOC	K1902526-001.01 100x	3.5724 ppm	0.0395 ppm	1.1100%	2019/03/28 22:52

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.6003	36.0034	33.43	36.40	2.97	50.17	10:27
2	TOC	3.5445	35.4451	33.06	36.09	3.03	50.20	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
19	TOC	K1902526-002.01 100x	3.8439 ppm	0.0019 ppm	0.0500%	2019/03/28 23:20

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.8452	38.4519	35.10	38.10	3.00	50.23	10:26
2	TOC	3.8425	38.4253	35.08	38.06	2.99	50.20	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.0455 ppm (PASS)	0.0000 ppm	0%	2019/03/28 23:48

Pos	Base Analysis	ID	Rep	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run
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	Type		#							Time
B	TOC	25 ppm	1	26.0455	260.4553	186.26	189.19	2.93	50.24	10:32
Completion State		Success Action		Method		Calibration		STD Conc - Pos B		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		50 ppmC		

Sample Type: Check Standard --> CCB From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0671 ppm (PASS)	0.0000 ppm	0%	2019/03/29 00:03	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0671	0.6711	9.92	12.89	2.97	50.26	10:32
Completion State		Success Action		Method		Calibration		STD Conc - Pos D		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		0 ppmC		

Sample Type: Sample From Schedule Version 5

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
♦ 20	TOC	MB2	0.0704 ppm	0.0000 ppm	0.0000%	2019/03/29 00:17		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0704	0.7040	9.47	12.51	3.04	50.23	10:32
Dilution		Blank Contribution		Method		Calibration		
1:10		(TC) 8.9951 (IC) (v1240)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Sample Type: Check Standard --> LCS From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
♦ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.6903 ppm (PASS)	0.0000 ppm	0%	2019/03/29 00:32	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.6903	266.9035	190.64	193.77	3.14	50.23	10:33
Completion State		Success Action		Method		Calibration		STD Conc - Pos C		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		25 ppmC		

Sample Type: Sample

From Schedule Version 5

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
21	TOC	K1902567-001.03	11.3478 ppm	0.0566 ppm	0.5000%	2019/03/29 00:47

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	11.3878	113.8783	86.29	89.36	3.06	50.27	10:26
2	TOC	11.3078	113.0783	85.75	88.77	3.02	50.24	10:23

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
22	TOC	K1902567-002.03	3.2184 ppm	0.0726 ppm	2.2600%	2019/03/29 01:15

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.2698	32.6975	31.19	34.11	2.92	50.26	10:29
2	TOC	3.1671	31.6707	30.49	33.44	2.95	50.28	10:24

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
23	TOC	K1902567-003.03	3.3409 ppm	0.0346 ppm	1.0400%	2019/03/29 01:43

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.3654	33.6536	31.84	34.76	2.92	50.23	10:31
2	TOC	3.3165	33.1645	31.51	34.40	2.89	50.22	10:29

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
24	TOC	K1902567-004.03	1.3922 ppm	0.0033 ppm	0.2400%	2019/03/29 02:11

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.3945	13.9452	18.46	21.31	2.85	50.27	10:30
2	TOC	1.3898	13.8980	18.43	21.34	2.92	50.25	10:27

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
25	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/29 02:39

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time

1	TOC	0.0000	0.0000	8.82	11.77	2.95	50.26	10:27
2	TOC	0.0000	0.0000	8.84	11.82	2.98	50.30	10:27

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
26	TOC	K1902567-005.03	2.4650 ppm	0.0327 ppm	1.3300%	2019/03/29 03:07

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.5137	25.1371	26.06	28.90	2.84	50.28	10:27
2	TOC	2.4434	24.4344	25.58	28.53	2.95	50.30	10:24
3	TOC	2.4492	24.4918	25.62	28.78	3.16	50.31	10:30
4	TOC	2.4536	24.5360	25.65	28.58	2.93	50.29	10:30

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
27	TOC	K1902603-001.03	3.1388 ppm	0.0402 ppm	1.2800%	2019/03/29 04:03

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.1672	31.6722	30.49	33.41	2.92	50.31	10:30
2	TOC	3.1104	31.1035	30.11	33.07	2.96	50.30	10:27

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
28	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/29 04:31

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.62	11.73	3.10	50.30	10:28
2	TOC	0.0000	0.0000	8.78	11.76	2.98	50.32	10:24

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.0566 ppm (PASS)	0.0000 ppm	0%	2019/03/29 04:59

Base	Rep	Run

Pos	Analysis Type	ID	#	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Time
B	TOC	25 ppm	1	26.0566	260.5658	186.33	189.40	3.06	50.33	10:32

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/29 05:14

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	9.26	12.30	3.04	50.31	10:33

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample From Schedule Version 5

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 29	TOC	K1902603-002.03	1.5256 ppm	0.0021 ppm	0.1400%	2019/03/29 05:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.5271	15.2710	19.36	22.47	3.11	50.32	10:29
2	TOC	1.5242	15.2416	19.34	22.44	3.10	50.34	10:28

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 30	TOC	K1902603-002.03 ms	29.4024 ppm	0.0000 ppm	0.0000%	2019/03/29 05:57

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	29.4024	294.0242	208.58	211.47	2.89	50.36	10:29

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 31	TOC	RB	0.1231 ppm	0.0000 ppm	0.0000%	2019/03/29 06:11

Rep	Base	ppm	µg	Adjusted	NDIR (Abs)	Baseline	Pressure	Run
-----	------	-----	----	----------	------------	----------	----------	-----

#	Analysis Type			(Abs)		(Abs)	(psig)	Time
1	TOC	0.1231	1.2314	9.83	12.73	2.90	50.39	10:29

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
32	TOC	K1902603-003.03	1.7012 ppm	0.0280 ppm	1.6500%	2019/03/29 06:26

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.6813	16.8135	20.41	23.51	3.10	50.30	10:26
2	TOC	1.7210	17.2098	20.68	23.66	2.98	50.23	10:24

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
33	TOC	K1902603-004.03	1.7390 ppm	0.0300 ppm	1.7300%	2019/03/29 06:54

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.7602	17.6016	20.94	23.65	2.71	50.22	10:26
2	TOC	1.7177	17.1774	20.66	23.64	2.99	50.18	10:28

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
34	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/29 07:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.75	11.63	2.88	50.14	10:30
2	TOC	0.0000	0.0000	8.46	11.37	2.91	50.13	10:27
3	TOC	0.0000	0.0000	8.56	11.44	2.89	50.13	10:29
4	TOC	0.0000	0.0000	8.53	11.42	2.89	50.08	10:30

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
35	TOC	FB	0.0297 ppm	0.0175 ppm	58.8400%	2019/03/29 08:17

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0421	0.4212	9.28	12.01	2.73	49.99	10:31
2	TOC	0.0174	0.1737	9.11	11.94	2.83	50.08	10:24

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

1:10 (TC) 8.9951 (IC) CAS_salt_010711 CAS_salt_010711
(v1240) (v4) (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
36	TOC	K1902607-001.03 doc	1.2481 ppm	0.1250 ppm	10.0200%	2019/03/29 08:46

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.1597	11.5969	16.87	19.76	2.90	50.15	10:27
2	TOC	1.3365	13.3647	18.07	21.09	3.02	50.23	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
37	TOC	K1902607-001.03 ms doc	29.4740 ppm	0.0000 ppm	0.0000%	2019/03/29 09:14

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	29.4740	294.7402	209.06	212.00	2.94	50.26	10:33

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
38	TOC	RB	0.1616 ppm	0.0000 ppm	0.0000%	2019/03/29 09:29

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1616	1.6159	10.09	13.05	2.96	50.28	10:31

Dilution 1:10 **Blank Contribution** (TC) 8.9951 (IC) (v1240) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.3431 ppm (PASS)	0.0000 ppm	0%	2019/03/29 09:43

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.3431	263.4312	188.28	191.09	2.81	50.26	10:32

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30) **STD Conc - Pos B** 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/03/29 09:58

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	9.36	12.36	3.00	50.26	10:30

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample

From Schedule Version 5

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 39	TOC	MB3	0.0000 ppm	0.0000 ppm	0.0000%	2019/03/29 10:12

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.76	11.71	2.95	50.24	10:31

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS

From Schedule Version 5

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
♦ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.7989 ppm (PASS)	0.0000 ppm	0%	2019/03/29 10:27

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.7989	267.9892	191.37	194.35	2.98	50.25	10:29

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos C 25 ppmC

Sample Type: Sample

From Schedule Version 6

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
♦ 40	TOC	K1902607-002.03 doc 2x	16.3952 ppm	0.3933 ppm	2.4000%	2019/03/29 10:42

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time

1	TOC	16.6734	166.7338	122.17	125.20	3.02	50.23	10:29
2	TOC	16.1171	161.1710	118.40	121.48	3.08	50.19	10:25

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
41	TOC	K1902623-001.07 doc 100x	0.8175 ppm	0.0091 ppm	1.1100%	2019/03/29 11:10

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.8111	8.1113	14.50	17.48	2.98	50.19	10:26
2	TOC	0.8239	8.2395	14.59	17.54	2.95	50.15	10:26

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
42	TOC	RB	0.1606 ppm	0.0653 ppm	40.6600%	2019/03/29 11:38

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1145	1.1445	9.77	12.62	2.85	50.15	10:26
2	TOC	0.2068	2.0682	10.40	13.47	3.07	50.15	10:25

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
43	TOC	K1902623-002.07 doc	1.9932 ppm	0.0042 ppm	0.2100%	2019/03/29 12:06

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.9962	19.9617	22.54	25.71	3.17	50.14	10:27
2	TOC	1.9903	19.9028	22.50	25.48	2.98	50.16	10:26

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
44	TOC	RB	0.0225 ppm	0.0318 ppm	141.4200%	2019/03/29 12:34

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.86	11.78	2.92	50.17	10:28
2	TOC	0.0449	0.4492	9.30	12.23	2.93	50.16	10:24

Dilution 1:10
Blank Contribution (TC) 8.9951 (IC) (v1240)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 6

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	26.4972 ppm (PASS)	0.0000 ppm	0%	2019/03/29 13:02

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	26.4972	264.9721	189.32	192.19	2.87	50.13	10:30

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 6

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0300 ppm (PASS)	0.0000 ppm	0%	2019/03/29 13:17

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0300	0.2998	9.67	12.46	2.79	50.12	10:30

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Meta Data Used in this Report

Blanks

Version	Reagent (Abs)	Acid (Abs)	DI IC (Abs)	DI TC (Abs)	DI TOC (Abs)	Save Time	Operator
v1239	1.1963	1.0700	0.0000	0.0000	0.0000	2019/03/26 13:09	Fusion1 (Fusion1)
v1240	2.3943	1.3110	0.0000	0.0000	0.0000	2019/03/28 14:20	Fusion1 (Fusion1)

Calibrations

Name: CAS_salt_010711 (TOC)

Version: v30

Calibration curve formula: TOC: $y = 6.788x + 9.463$

Ver Creation: 2019/03/05 17:42 r² value: TOC: r² = 0.99963
 Comment:
 Operator: Fusion1 (Fusion1)
 Basic Analysis Type TOC

Basic Analysis Type: TOC

Sample ID	Y Raw Value	X Expected	Message	End Time
DI Water	7.8970	0.0000		2019/03/05 16:15
0.500 ppm	11.5280	0.5000		2019/03/05 16:29
1.0 ppm	14.9760	1.0000		2019/03/05 16:44
5.0 ppm	43.6500	5.0000		2019/03/05 16:58
10 ppm	79.6020	10.0000		2019/03/05 17:12
25 ppm	183.3580	25.0000		2019/03/05 17:26
50 ppm	346.3230	50.0000		2019/03/05 17:40

Methods**Name: CAS_salt_010711 (TOC)**

Version: v4 Operator: Fusion1 (Fusion1)
 Ver Creation: 2019/02/21 17:57
 Comment:

Parameter	Value	Advanced Parameter	Value
SampleVolume	10.0 mL	NeedleRinseVolume	5.0 ml
Dilution	1:10	VialPrimeVolume	2.0 ml
AcidVolume	0.5 ml	ICSamplePrimeVolume	2.0 ml
ReagentVolume	2.0 ml	ICSpurgeRinseVolume	12.0 ml
UVReactorPrerinse	Off	BaselineStabilizeTime	0.70 min
UVReactorPrerinseVolume	5.0	DetectorPressureFlow	150 ml/min
NumberOfUVReactorPrerinses	1	SyringeSpeedWaste	10
ICSpurgeTime	1.00 mins	SyringeSpeedAcid	7
DetectorSweepFlow	500 ml/min	SyringeSpeedReagent	7
PreSpurgeTime	2.00 mins	SyringeSpeedDIWater	7
SystemFlow	500 ml/min	NDIRPressurization	60 psig
		SyringeSpeedSampleDispense	5
		SyringeSpeedSampleAspirate	4
		SyringeSpeedUVDispense	5
		SyringeSpeedUVAspirate	5
		SyringeSpeedICDispense	5
		SyringeSpeedICAspirate	5
		NDIRPressureStabilize	1.75 min
		SampleMixing	Off
		SampleMixingCycles	1
		SampleMixingVolume	10.0
		LowLevelFilterNDIR	Off

Acceptance / Approval

Electronic Signatures

Report Version	User Name	Acceptance	Reason	Date
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Report History

Report History

Report Version	User Name	System Reason	User Reason	Date
1	Fusion1 (Fusion1)	Schedule completed	Schedule completed	2019/03/29 13:33

StarLIMS Run: 629966, 629967, 629968
 Analysis: TOC/DOC
 Method: 9060, 415.1, SM 5310 C, 9060A

CCV: 11-GEN-05-77C 50 ppm LCS: 11-GEN-05-77D 25.0 ppm

ICAL Date: 3/6/19

ICAL ID: 11-GEN-05-76H

ICS ID: 11-GEN-05-74A

ICS TV: 25.0 ppm ICS % R = 2

Spike ID: 11-GEN-05-700 0.05 ml of 5000 ppm stock ---> 10.0 ml = 25.0 ppm x dilution factor

Sodium Persulfate: 11-GEN-05-77E

21 % H3PO4: 11-GEN-05-77A

Equipment ID: K-TOC-03

PIPETTE ID: 124276B, 129001F, N11314F, Marge

FILTER ID: NA

CBA: 0.06875

Analyzed By: <u>BCD</u>	Date Analyzed: <u>3/28/19</u>
Reviewed By: <u>Freeper</u>	Date Reviewed: <u>03/29/19</u>



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April 03, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19031189**

Laboratory Results for: **Longhorn GW Treatment Plant**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 22, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: DAYNA.FISHER
RJ Modashia
Project Manager

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
Work Order: HS19031189

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19031189-01	LH18/24-SP650_032119	Water		21-Mar-2019 14:00	22-Mar-2019 08:52	<input type="checkbox"/>
HS19031189-02	Trip Blank	Water	ALS-022719-71	21-Mar-2019 00:00	22-Mar-2019 08:52	<input type="checkbox"/>

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
Work Order: HS19031189

CASE NARRATIVE

GCMS Volatiles by Method SW8260**Batch ID: R335378****Sample ID: CCV_END**

- Benzene and Toluene exceeded %D recoveries for closing CCV due to carryover from previous sample .

Sample ID: HS19031013-07MS

- MS and MSD are for an unrelated sample

Sample ID: VLCSW-190326

- 1,2,4_Trimethyl benzene exceeded QC limits for LCS. CCV is OK.
-

WetChemistry by Method SW9056**Batch ID: R335875**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: LH18/24-SP650_032119
 Collection Date: 21-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031189
 Lab ID:HS19031189-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
VOLATILES ORGANICS BY METHOD		Method:SW8260						
8260C								Analyst: AKP
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	26-Mar-2019 20:42
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	26-Mar-2019 20:42
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	26-Mar-2019 20:42
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	26-Mar-2019 20:42
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	26-Mar-2019 20:42
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: LH18/24-SP650_032119
 Collection Date: 21-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031189
 Lab ID:HS19031189-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: AKP	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
cis-1,2-Dichloroethene	1.8		0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	26-Mar-2019 20:42	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	26-Mar-2019 20:42	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	26-Mar-2019 20:42	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Trichloroethene	0.71	J	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 20:42	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>90.5</i>			0	<i>81-118</i>	%REC	1	26-Mar-2019 20:42	
<i>Surr: 4-Bromofluorobenzene</i>	<i>97.6</i>			0	<i>85-114</i>	%REC	1	26-Mar-2019 20:42	
<i>Surr: Dibromofluoromethane</i>	<i>92.4</i>			0	<i>80-119</i>	%REC	1	26-Mar-2019 20:42	
<i>Surr: Toluene-d8</i>	<i>108</i>			0	<i>89-112</i>	%REC	1	26-Mar-2019 20:42	
ANIONS BY SW9056A		Method:SW9056						Analyst: KMU	
Chloride	314		2.00	5.00	5.00	mg/L	10	03-Apr-2019 03:29	
Sulfate	26.8		2.00	5.00	5.00	mg/L	10	03-Apr-2019 03:29	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: Trip Blank
 Collection Date: 21-Mar-2019 00:00

ANALYTICAL REPORT

WorkOrder:HS19031189
 Lab ID:HS19031189-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
VOLATILES ORGANICS BY METHOD		Method:SW8260						
8260C								Analyst: AKP
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	26-Mar-2019 15:51
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	26-Mar-2019 15:51
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	26-Mar-2019 15:51
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	26-Mar-2019 15:51
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	26-Mar-2019 15:51
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: Longhorn GW Treatment Plant
 Sample ID: Trip Blank
 Collection Date: 21-Mar-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19031189
 Lab ID:HS19031189-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: AKP
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	26-Mar-2019 15:51	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	26-Mar-2019 15:51	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	26-Mar-2019 15:51	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Trichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	26-Mar-2019 15:51	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>90.9</i>			<i>0</i>	<i>81-118</i>	<i>%REC</i>	<i>1</i>	<i>26-Mar-2019 15:51</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>97.8</i>			<i>0</i>	<i>85-114</i>	<i>%REC</i>	<i>1</i>	<i>26-Mar-2019 15:51</i>	
<i>Surr: Dibromofluoromethane</i>	<i>91.1</i>			<i>0</i>	<i>80-119</i>	<i>%REC</i>	<i>1</i>	<i>26-Mar-2019 15:51</i>	
<i>Surr: Toluene-d8</i>	<i>107</i>			<i>0</i>	<i>89-112</i>	<i>%REC</i>	<i>1</i>	<i>26-Mar-2019 15:51</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R335378					Test Name : VOLATILES ORGANICS BY METHOD 8260C	Matrix: Water
HS19031189-01	LH18/24-SP650_032119	21 Mar 2019 14:00				26 Mar 2019 20:42 1
HS19031189-02	Trip Blank	21 Mar 2019 00:00				26 Mar 2019 15:51 1
Batch ID R335875					Test Name : ANIONS BY SW9056A	Matrix: Water
HS19031189-01	LH18/24-SP650_032119	21 Mar 2019 14:00				03 Apr 2019 03:29 10

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335378 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190325	Units: UG/L			Analysis Date: 26-Mar-2019 15:03					
Client ID:	Run ID: VOA6_335378	SeqNo: 5008738	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	45.09	1.0	50	0	90.2	81 - 118				
Surr: 4-Bromofluorobenzene	48.89	1.0	50	0	97.8	85 - 114				
Surr: Dibromofluoromethane	45.31	1.0	50	0	90.6	80 - 119				
Surr: Toluene-d8	52.99	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335378 (0)		Instrument: VOA6			Method: VOLATILES ORGANICS BY METHOD 8260C					
LCS	Sample ID: VLCSW-190326	Units: UG/L			Analysis Date: 26-Mar-2019 13:26					
Client ID:	Run ID: VOA6_335378	SeqNo: 5008736		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	59.18	1.0	50	0	118	78 - 124				
1,1,1-Trichloroethane	49.91	1.0	50	0	99.8	74 - 131				
1,1,2,2-Tetrachloroethane	56.51	1.0	50	0	113	71 - 121				
1,1,2-Trichloroethane	56.76	1.0	50	0	114	80 - 119				
1,1-Dichloroethane	48.82	1.0	50	0	97.6	77 - 125				
1,1-Dichloroethene	48.97	1.0	50	0	97.9	71 - 131				
1,1-Dichloropropene	50.25	1.0	50	0	100	78 - 125				
1,2,3-Trichlorobenzene	62.38	1.0	50	0	125	69 - 129				
1,2,3-Trichloropropane	60.16	1.0	50	0	120	73 - 122				
1,2,4-Trichlorobenzene	62.08	1.0	50	0	124	69 - 130				
1,2,4-Trimethylbenzene	62.05	1.0	50	0	124	76 - 124				S
1,2-Dibromo-3-chloropropane	60.85	1.0	50	0	122	62 - 128				
1,2-Dibromoethane	59.41	1.0	50	0	119	77 - 121				
1,2-Dichlorobenzene	57.13	1.0	50	0	114	80 - 119				
1,2-Dichloroethane	53.57	1.0	50	0	107	73 - 128				
1,2-Dichloropropane	52.36	1.0	50	0	105	78 - 122				
1,3,5-Trimethylbenzene	60.55	1.0	50	0	121	75 - 124				
1,3-Dichlorobenzene	57.8	1.0	50	0	116	80 - 119				
1,3-Dichloropropane	57.1	1.0	50	0	114	80 - 119				
1,4-Dichlorobenzene	56.85	1.0	50	0	114	79 - 118				
2,2-Dichloropropane	51.22	1.0	50	0	102	60 - 139				
2-Butanone	109.7	2.0	100	0	110	56 - 143				
2-Chlorotoluene	56	1.0	50	0	112	79 - 122				
2-Hexanone	117.1	2.0	100	0	117	57 - 139				
4-Chlorotoluene	57.36	1.0	50	0	115	78 - 122				
4-Isopropyltoluene	60.43	1.0	50	0	121	77 - 127				
4-Methyl-2-pentanone	115.8	2.0	100	0	116	67 - 130				
Acetone	104.5	2.0	100	0	104	39 - 160				
Benzene	54.3	1.0	50	0	109	79 - 120				
Bromobenzene	59.62	1.0	50	0	119	80 - 120				
Bromochloromethane	51.72	1.0	50	0	103	78 - 123				
Bromodichloromethane	56.5	1.0	50	0	113	79 - 125				
Bromoform	60.49	1.0	50	0	121	66 - 130				
Bromomethane	52.77	1.0	50	0	106	53 - 141				

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335378 (0)		Instrument: VOA6			Method: VOLATILES ORGANICS BY METHOD 8260C					
LCS	Sample ID: VLCSW-190326	Units: UG/L			Analysis Date: 26-Mar-2019 13:26					
Client ID:	Run ID: VOA6_335378	SeqNo: 5008736			PrepDate:			DF: 1		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	94.6	2.0	100	0	94.6	64 - 133				
Carbon tetrachloride	55.46	1.0	50	0	111	72 - 136				
Chlorobenzene	54.64	1.0	50	0	109	82 - 118				
Chloroethane	47.94	1.0	50	0	95.9	60 - 138				
Chloroform	50.48	1.0	50	0	101	79 - 124				
Chloromethane	51.79	1.0	50	0	104	50 - 139				
cis-1,2-Dichloroethene	50.02	1.0	50	0	100	78 - 123				
cis-1,3-Dichloropropene	53.85	1.0	50	0	108	75 - 124				
Dibromochloromethane	59.63	1.0	50	0	119	74 - 126				
Dibromomethane	55.72	1.0	50	0	111	79 - 123				
Dichlorodifluoromethane	59.44	1.0	50	0	119	32 - 152				
Ethylbenzene	58.6	1.0	50	0	117	79 - 121				
Hexachlorobutadiene	62.03	1.0	50	0	124	66 - 134				
Isopropylbenzene	58.17	1.0	50	0	116	72 - 131				
m,p-Xylene	114.5	2.0	100	0	115	80 - 121				
Methylene chloride	50.68	2.0	50	0	101	74 - 124				
Naphthalene	63.48	1.0	50	0	127	61 - 128				
n-Butylbenzene	62.6	1.0	50	0	125	75 - 128				
n-Propylbenzene	58.67	1.0	50	0	117	76 - 126				
o-Xylene	56.01	1.0	50	0	112	78 - 122				
sec-Butylbenzene	58.01	1.0	50	0	116	77 - 126				
Styrene	59.49	1.0	50	0	119	78 - 123				
tert-Butylbenzene	57.39	1.0	50	0	115	78 - 124				
Tetrachloroethene	57.94	1.0	50	0	116	74 - 129				
Toluene	57.21	1.0	50	0	114	80 - 121				
trans-1,2-Dichloroethene	49.24	1.0	50	0	98.5	75 - 124				
trans-1,3-Dichloropropene	58.56	1.0	50	0	117	73 - 127				
Trichloroethene	51.9	1.0	50	0	104	79 - 123				
Trichlorofluoromethane	49.99	1.0	50	0	100.0	65 - 141				
Vinyl chloride	48.66	1.0	50	0	97.3	58 - 137				
Surr: 1,2-Dichloroethane-d4	45.08	1.0	50	0	90.2	81 - 118				
Surr: 4-Bromofluorobenzene	49.32	1.0	50	0	98.6	85 - 114				
Surr: Dibromofluoromethane	46.14	1.0	50	0	92.3	80 - 119				
Surr: Toluene-d8	52.93	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335378 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19031013-07MS	Units: UG/L			Analysis Date: 26-Mar-2019 17:03					
Client ID:	Run ID: VOA6_335378	SeqNo: 5008743	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	11.16	1.0	20	0	55.8	78 - 124				S
1,1,1-Trichloroethane	11.15	1.0	20	0	55.7	74 - 131				S
1,1,2,2-Tetrachloroethane	15.76	1.0	20	0	78.8	71 - 121				
1,1,2-Trichloroethane	53.04	1.0	20	37.94	75.5	80 - 119				S
1,1-Dichloroethane	185.5	1.0	20	186	-2.54	77 - 125				SO
1,1-Dichloroethene	1050	1.0	20	1097	-235	71 - 131				SEO
1,1-Dichloropropene	11.14	1.0	20	0	55.7	78 - 125				S
1,2,3-Trichlorobenzene	10.79	1.0	20	0	53.9	69 - 129				S
1,2,3-Trichloropropane	16.08	1.0	20	0	80.4	73 - 122				
1,2,4-Trichlorobenzene	10.13	1.0	20	0	50.7	69 - 130				S
1,2,4-Trimethylbenzene	11.2	1.0	20	0	56.0	76 - 124				S
1,2-Dibromo-3-chloropropane	19.5	1.0	20	0	97.5	62 - 128				
1,2-Dibromoethane	12.4	1.0	20	0	62.0	77 - 121				S
1,2-Dichlorobenzene	20.35	1.0	20	7.577	63.9	80 - 119				S
1,2-Dichloroethane	23.92	1.0	20	13.75	50.8	73 - 128				S
1,2-Dichloropropane	9.635	1.0	20	0	48.2	78 - 122				S
1,3,5-Trimethylbenzene	10.98	1.0	20	0	54.9	75 - 124				S
1,3-Dichlorobenzene	10.86	1.0	20	0	54.3	80 - 119				S
1,3-Dichloropropane	11.74	1.0	20	0	58.7	80 - 119				S
1,4-Dichlorobenzene	11.93	1.0	20	0	59.7	79 - 118				S
2,2-Dichloropropane	10.85	1.0	20	0	54.2	60 - 139				S
2-Butanone	33.46	2.0	40	0	83.6	56 - 143				
2-Chlorotoluene	10.92	1.0	20	0	54.6	79 - 122				S
2-Hexanone	38.21	2.0	40	0	95.5	57 - 139				
4-Chlorotoluene	10.67	1.0	20	0	53.4	78 - 122				S
4-Isopropyltoluene	10.47	1.0	20	0	52.3	77 - 127				S
4-Methyl-2-pentanone	35.59	2.0	40	0	89.0	67 - 130				
Acetone	31.41	2.0	40	0	78.5	39 - 160				
Benzene	13.32	1.0	20	3.04	51.4	79 - 120				S
Bromobenzene	11.36	1.0	20	0	56.8	80 - 120				S
Bromochloromethane	9.213	1.0	20	0	46.1	78 - 123				S
Bromodichloromethane	9.838	1.0	20	0	49.2	79 - 125				S
Bromoform	13.52	1.0	20	0	67.6	66 - 130				
Bromomethane	10.32	1.0	20	0	51.6	53 - 141				S

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335378 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19031013-07MS	Units: UG/L			Analysis Date: 26-Mar-2019 17:03					
Client ID:	Run ID: VOA6_335378	SeqNo: 5008743	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	23.57	2.0	40	0	58.9	64 - 133				S
Carbon tetrachloride	13.73	1.0	20	0	68.6	72 - 136				S
Chlorobenzene	10.51	1.0	20	0	52.6	82 - 118				S
Chloroethane	175	1.0	20	0	875	60 - 138				S
Chloroform	9.489	1.0	20	0	47.4	79 - 124				S
Chloromethane	7.613	1.0	20	0	38.1	50 - 139				S
cis-1,2-Dichloroethene	55.42	1.0	20	49.43	30.0	78 - 123				S
cis-1,3-Dichloropropene	9.595	1.0	20	0	48.0	75 - 124				S
Dibromochloromethane	11.47	1.0	20	0	57.4	74 - 126				S
Dibromomethane	10.62	1.0	20	0	53.1	79 - 123				S
Dichlorodifluoromethane	13.94	1.0	20	0	69.7	32 - 152				S
Ethylbenzene	10.95	1.0	20	0	54.8	79 - 121				S
Hexachlorobutadiene	8.847	1.0	20	0	44.2	66 - 134				S
Isopropylbenzene	10.98	1.0	20	0	54.9	72 - 131				S
m,p-Xylene	21.6	2.0	40	0	54.0	80 - 121				S
Methylene chloride	9.478	2.0	20	0	47.4	74 - 124				S
Naphthalene	14.29	1.0	20	0	71.4	61 - 128				S
n-Butylbenzene	9.619	1.0	20	0	48.1	75 - 128				S
n-Propylbenzene	11.04	1.0	20	0	55.2	76 - 126				S
o-Xylene	11.05	1.0	20	0	55.2	78 - 122				S
sec-Butylbenzene	10.57	1.0	20	0	52.8	77 - 126				S
Styrene	10.37	1.0	20	0	51.8	78 - 123				S
tert-Butylbenzene	11.34	1.0	20	0	56.7	78 - 124				S
Tetrachloroethene	12.65	1.0	20	0	63.3	74 - 129				S
Toluene	11.44	1.0	20	0	57.2	80 - 121				S
trans-1,2-Dichloroethene	14.44	1.0	20	4.915	47.6	75 - 124				S
trans-1,3-Dichloropropene	10.16	1.0	20	0	50.8	73 - 127				S
Trichloroethene	178.6	1.0	20	177.3	6.25	79 - 123				SO
Trichlorofluoromethane	13.2	1.0	20	0	66.0	65 - 141				S
Vinyl chloride	61.81	1.0	20	53.81	40.0	58 - 137				S
Surr: 1,2-Dichloroethane-d4	47.29	1.0	50	0	94.6	81 - 118				
Surr: 4-Bromofluorobenzene	47.34	1.0	50	0	94.7	85 - 114				
Surr: Dibromofluoromethane	45.97	1.0	50	0	91.9	80 - 119				
Surr: Toluene-d8	54.67	1.0	50	0	109	89 - 112				

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335378 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C							
MSD	Sample ID: HS19031013-07MSD	Units: UG/L			Analysis Date: 26-Mar-2019 17:27						
Client ID:	Run ID: VOA6_335378	SeqNo: 5008744	PrepDate:	DF: 1							
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual	
1,1,1,2-Tetrachloroethane	14.29	1.0	20	0	71.5	78 - 124	11.16	24.6	20	SR	
1,1,1-Trichloroethane	13.73	1.0	20	0	68.7	74 - 131	11.15	20.7	20	SR	
1,1,2,2-Tetrachloroethane	16.44	1.0	20	0	82.2	71 - 121	15.76	4.24	20		
1,1,2-Trichloroethane	66.89	1.0	20	37.94	145	80 - 119	53.04	23.1	20	SR	
1,1-Dichloroethane	248.9	1.0	20	186	315	77 - 125	185.5	29.2	20	SREO	
1,1-Dichloroethene	1262	1.0	20	1097	824	71 - 131	1050	18.3	20	SEO	
1,1-Dichloropropene	13.99	1.0	20	0	70.0	78 - 125	11.14	22.7	20	SR	
1,2,3-Trichlorobenzene	13.99	1.0	20	0	69.9	69 - 129	10.79	25.8	20	R	
1,2,3-Trichloropropane	16.44	1.0	20	0	82.2	73 - 122	16.08	2.23	20		
1,2,4-Trichlorobenzene	13.67	1.0	20	0	68.3	69 - 130	10.13	29.7	20	SR	
1,2,4-Trimethylbenzene	13.83	1.0	20	0	69.1	76 - 124	11.2	21	20	SR	
1,2-Dibromo-3-chloropropane	18.23	1.0	20	0	91.1	62 - 128	19.5	6.76	20		
1,2-Dibromoethane	15.92	1.0	20	0	79.6	77 - 121	12.4	24.9	20	R	
1,2-Dichlorobenzene	25.18	1.0	20	7.577	88.0	80 - 119	20.35	21.2	20	R	
1,2-Dichloroethane	33.16	1.0	20	13.75	97.1	73 - 128	23.92	32.4	20	R	
1,2-Dichloropropane	13.43	1.0	20	0	67.1	78 - 122	9.635	32.9	20	SR	
1,3,5-Trimethylbenzene	13.61	1.0	20	0	68.1	75 - 124	10.98	21.4	20	SR	
1,3-Dichlorobenzene	13.83	1.0	20	0	69.1	80 - 119	10.86	24.1	20	SR	
1,3-Dichloropropane	14.82	1.0	20	0	74.1	80 - 119	11.74	23.2	20	SR	
1,4-Dichlorobenzene	14.9	1.0	20	0	74.5	79 - 118	11.93	22.1	20	SR	
2,2-Dichloropropane	13.44	1.0	20	0	67.2	60 - 139	10.85	21.4	20	R	
2-Butanone	35.67	2.0	40	0	89.2	56 - 143	33.46	6.39	20		
2-Chlorotoluene	13.5	1.0	20	0	67.5	79 - 122	10.92	21.1	20	SR	
2-Hexanone	38.9	2.0	40	0	97.2	57 - 139	38.21	1.77	20		
4-Chlorotoluene	13.3	1.0	20	0	66.5	78 - 122	10.67	22	20	SR	
4-Isopropyltoluene	13.64	1.0	20	0	68.2	77 - 127	10.47	26.3	20	SR	
4-Methyl-2-pentanone	37.71	2.0	40	0	94.3	67 - 130	35.59	5.77	20		
Acetone	31.97	2.0	40	0	79.9	39 - 160	31.41	1.77	20		
Benzene	17.94	1.0	20	3.04	74.5	79 - 120	13.32	29.6	20	SR	
Bromobenzene	13.7	1.0	20	0	68.5	80 - 120	11.36	18.7	20	S	
Bromochloromethane	13.51	1.0	20	0	67.5	78 - 123	9.213	37.8	20	SR	
Bromodichloromethane	14.1	1.0	20	0	70.5	79 - 125	9.838	35.6	20	SR	
Bromoform	16.39	1.0	20	0	82.0	66 - 130	13.52	19.2	20		
Bromomethane	13.02	1.0	20	0	65.1	53 - 141	10.32	23.1	20	R	

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335378 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C							
MSD	Sample ID: HS19031013-07MSD	Units: UG/L			Analysis Date: 26-Mar-2019 17:27						
Client ID:	Run ID: VOA6_335378	SeqNo: 5008744	PrepDate:	DF: 1							
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual	
Carbon disulfide	29.42	2.0	40	0	73.5	64 - 133	23.57	22.1	20	R	
Carbon tetrachloride	15.62	1.0	20	0	78.1	72 - 136	13.73	12.9	20		
Chlorobenzene	13.94	1.0	20	0	69.7	82 - 118	10.51	28	20	SR	
Chloroethane	213.7	1.0	20	0	1070	60 - 138	175	19.9	20	SE	
Chloroform	13.25	1.0	20	0	66.3	79 - 124	9.489	33.1	20	SR	
Chloromethane	11.04	1.0	20	0	55.2	50 - 139	7.613	36.7	20	R	
cis-1,2-Dichloroethene	77.1	1.0	20	49.43	138	78 - 123	55.42	32.7	20	SR	
cis-1,3-Dichloropropene	13.7	1.0	20	0	68.5	75 - 124	9.595	35.2	20	SR	
Dibromochloromethane	14.83	1.0	20	0	74.2	74 - 126	11.47	25.6	20	R	
Dibromomethane	14.69	1.0	20	0	73.4	79 - 123	10.62	32.1	20	SR	
Dichlorodifluoromethane	14.89	1.0	20	0	74.5	32 - 152	13.94	6.58	20		
Ethylbenzene	14.37	1.0	20	0	71.9	79 - 121	10.95	27	20	SR	
Hexachlorobutadiene	12.89	1.0	20	0	64.4	66 - 134	8.847	37.2	20	SR	
Isopropylbenzene	14.32	1.0	20	0	71.6	72 - 131	10.98	26.4	20	SR	
m,p-Xylene	28.13	2.0	40	0	70.3	80 - 121	21.6	26.2	20	SR	
Methylene chloride	13.2	2.0	20	0	66.0	74 - 124	9.478	32.8	20	SR	
Naphthalene	16.74	1.0	20	0	83.7	61 - 128	14.29	15.8	20		
n-Butylbenzene	13.53	1.0	20	0	67.7	75 - 128	9.619	33.8	20	SR	
n-Propylbenzene	13.92	1.0	20	0	69.6	76 - 126	11.04	23.1	20	SR	
o-Xylene	14.08	1.0	20	0	70.4	78 - 122	11.05	24.1	20	SR	
sec-Butylbenzene	13.79	1.0	20	0	68.9	77 - 126	10.57	26.5	20	SR	
Styrene	13.9	1.0	20	0	69.5	78 - 123	10.37	29.1	20	SR	
tert-Butylbenzene	13.72	1.0	20	0	68.6	78 - 124	11.34	19	20	S	
Tetrachloroethene	15.63	1.0	20	0	78.1	74 - 129	12.65	21.1	20	R	
Toluene	14.2	1.0	20	0	71.0	80 - 121	11.44	21.5	20	SR	
trans-1,2-Dichloroethene	19.45	1.0	20	4.915	72.7	75 - 124	14.44	29.6	20	SR	
trans-1,3-Dichloropropene	14.54	1.0	20	0	72.7	73 - 127	10.16	35.5	20	SR	
Trichloroethene	235.7	1.0	20	177.3	292	79 - 123	178.6	27.6	20	SREO	
Trichlorofluoromethane	14.65	1.0	20	0	73.2	65 - 141	13.2	10.4	20		
Vinyl chloride	74.06	1.0	20	53.81	101	58 - 137	61.81	18	20		
Surr: 1,2-Dichloroethane-d4	46.8	1.0	50	0	93.6	81 - 118	47.29	1.04	20		
Surr: 4-Bromofluorobenzene	50.69	1.0	50	0	101	85 - 114	47.34	6.83	20		
Surr: Dibromofluoromethane	46.32	1.0	50	0	92.6	80 - 119	45.97	0.744	20		
Surr: Toluene-d8	50.95	1.0	50	0	102	89 - 112	54.67	7.05	20		

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW**Batch ID:** R335378 (0)**Instrument:** VOA6**Method:** VOLATILES ORGANICS BY METHOD
8260C

The following samples were analyzed in this batch: HS19031189-01 HS19031189-02

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

QC BATCH REPORT NEW

Batch ID: R335875 (0)		Instrument: ICS2100		Method: ANIONS BY SW9056A						
MBLK	Sample ID: WBLKW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:16					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020414		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	0.500	0.500							U	
Sulfate	0.500	0.500							U	
LCS	Sample ID: WLCSW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:31					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020415		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	20.22	0.500	20	0	101	80 - 120				
Sulfate	19.98	0.500	20	0	99.9	80 - 120				
LCSD	Sample ID: WLCSDW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:46					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020416		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	18.95	0.500	20	0	94.8	80 - 120	20.22	6.51	20	
Sulfate	18.67	0.500	20	0	93.3	80 - 120	19.98	6.79	20	
MS	Sample ID: HS19031013-07MS	Units: mg/L			Analysis Date: 03-Apr-2019 04:28					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020427		PrepDate:			DF: 50			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	1535	25.0	500	1090	89.1	80 - 120				
Sulfate	1565	25.0	500	1119	89.2	80 - 120				
MSD	Sample ID: HS19031013-07MSD	Units: mg/L			Analysis Date: 03-Apr-2019 04:43					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020428		PrepDate:			DF: 50			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	1605	25.0	500	1090	103	80 - 120	1535	4.46	20	
Sulfate	1638	25.0	500	1119	104	80 - 120	1565	4.57	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 03-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: Longhorn GW Treatment Plant
WorkOrder: HS19031189

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

Unit Reported	Description
mg/L	Milligrams per Liter

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19031189

Date/Time Received: 22-Mar-2019 08:52
 Received by: NDR

Checklist completed by: Paresh M. Giga 22-Mar-2019
 eSignature Date

Reviewed by: Bernadette A. Fini 25-Mar-2019
 eSignature Date

Matrices: Water

Carrier name: FedEx

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:None
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 0.7c/0.7c U/c IR11
 Cooler(s)/Kit(s): 43950
 Date/Time sample(s) sent to storage: 3/22/19 19:30

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A


pH adjusted by:

Login Notes:

Client Contacted: _____ Date Contacted: _____ Person Contacted: _____
 Contacted By: _____ Regarding: _____

Comments:

Corrective Action:

 ALS Environmental 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CUSTODY SEAL		Seal Broken By: FM
	Date: 2/21/19	Time: 1430	Date: 03/22/19
	Name: Scott Beesiner	Company: BHATT	

439.50 MAR 22 2019



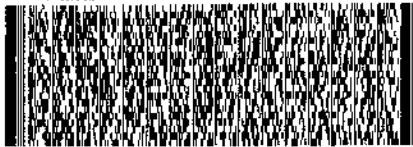
Must Deliver Next Business Day
Time and Temperature Sensitive!

ORIGIN ID: SGRA (303) 597-2450
 SCOTT BEESINER
 BHATE ENVIRONMENTAL ASSOCIATES
 1203-B EAST GRAND AVE. PH202
 MARSHALL, TX 75670
 UNITED STATES US

917
 B03
 1
 10:30
 A
 03/22

10 CLIENT SERVICES
 ALS LABORATORY GROUP
 10450 STANCLIFF ROAD
 SUITE 210
 HOUSTON TX 77099
 (281) 530-6056
 REF: LHAAP-58-80 64113-BJ

AMA: 011111



RETURNS MON - SAT

FedEx
 TRK#
 0221 4809 7831 3400

FRI - 22 MAR 10:30A
 PRIORITY OVERNIGHT

AB SGRA

77099
 TX-US
 IAH



FTD 162785 21MAR19 00GA 553C1/4603/800A



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

April 12, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19031492**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 28, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031492

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19031492-01	LH18/24-SP650_032719	Water		27-Mar-2019 14:00	28-Mar-2019 08:30	<input type="checkbox"/>
HS19031492-02	LH18/24-SP650_032719_BIX	Water		27-Mar-2019 14:00	28-Mar-2019 08:30	<input type="checkbox"/>

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.
-

Work Order Comments

- The analysis for TOC was subcontracted to ALS Environmental in Kelso, WA. Final report attached.
-

WetChemistry by Method E350.3**Batch ID: R336046**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

WetChemistry by Method E365.3**Batch ID: R335561**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032719
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19031492
 Lab ID:HS19031492-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
AMMONIA AS N BY E350.3(ISE)								
								Analyst: KVL
Nitrogen, Ammonia (As N)	11		0.20	0.20	0.20	mg/L	1	05-Apr-2019 11:45
ORTHO PHOSPHATE (PO4) AS P BY E365.3								
								Analyst: MZD
Phosphorus, Total Orthophosphate (As P)	2.42		0.100	0.250	0.250	mg/L	10	29-Mar-2019 12:03
SUBCONTRACT ANALYSIS - TOC ANALYSIS								
								Analyst: SUBK
Subcontract Analysis	See Attached		0	0		NA	1	05-Apr-2019 15:17

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032719_BIX
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19031492
 Lab ID:HS19031492-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA		Analyst: SUB				
Subcontract Analysis	See Attached		0	0		NA	1	12-Apr-2019 18:22

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031492

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R335561	Test Name : ORTHO PHOSPHATE (PO4) AS P BY E365.3		Matrix: Water			
HS19031492-01	LH18/24-SP650_032719	27 Mar 2019 14:00			29 Mar 2019 12:03	10
Batch ID R336046	Test Name : AMMONIA AS N BY E350.3(ISE)		Matrix: Water			
HS19031492-01	LH18/24-SP650_032719	27 Mar 2019 14:00			05 Apr 2019 11:45	1
Batch ID R336059	Test Name : SUBCONTRACT ANALYSIS - TOC ANALYSIS		Matrix: Water			
HS19031492-01	LH18/24-SP650_032719	27 Mar 2019 14:00			05 Apr 2019 15:17	1
Batch ID R336536	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Matrix: Water			
HS19031492-02	LH18/24-SP650_032719_BIX	27 Mar 2019 14:00			12 Apr 2019 18:22	1

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031492

QC BATCH REPORT NEW

Batch ID:	R335561 (0)	Instrument:	UV-2450	Method:	ORTHO PHOSPHATE (PO4) AS P BY E365.3					
MBLK	Sample ID: MBLK-335561	Units:	mg/L	Analysis Date:	29-Mar-2019 12:03					
Client ID:	Run ID: UV-2450_335561	SeqNo:	5012948	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	0.0250	0.0250								U
LCS	Sample ID: LCS-335561	Units:	mg/L	Analysis Date:	29-Mar-2019 12:03					
Client ID:	Run ID: UV-2450_335561	SeqNo:	5012949	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	0.252	0.0250	0.25	0	101	85 - 115				
MS	Sample ID: HS19031492-01MS	Units:	mg/L	Analysis Date:	29-Mar-2019 12:03					
Client ID: LH18/24-SP650_032719	Run ID: UV-2450_335561	SeqNo:	5012951	PrepDate:	DF: 10					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	5.01	0.250	2.5	2.42	104	80 - 120				
MSD	Sample ID: HS19031492-01MSD	Units:	mg/L	Analysis Date:	29-Mar-2019 12:03					
Client ID: LH18/24-SP650_032719	Run ID: UV-2450_335561	SeqNo:	5012952	PrepDate:	DF: 10					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Phosphorus, Total Orthophosphate (As P)	4.89	0.250	2.5	2.42	98.8	80 - 120	5.01	2.42	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031492

QC BATCH REPORT NEW

Batch ID:	R336046 (0)	Instrument:	WetChem_HS	Method:	AMMONIA AS N BY E350.3(ISE)					
MBLK	Sample ID: MBLK-R336046	Units:	mg/L	Analysis Date:	05-Apr-2019 11:45					
Client ID:	Run ID: WetChem_HS_336046	SeqNo:	5024111	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	0.20	0.20							U	
LCS	Sample ID: LCS-R336046	Units:	mg/L	Analysis Date:	05-Apr-2019 11:45					
Client ID:	Run ID: WetChem_HS_336046	SeqNo:	5024110	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	10.1	0.20	10	0	101	85 - 115				
MS	Sample ID: HS19031533-01MS	Units:	mg/L	Analysis Date:	05-Apr-2019 11:45					
Client ID:	Run ID: WetChem_HS_336046	SeqNo:	5024113	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	15.09	0.20	10	4.085	110	80 - 120				
MSD	Sample ID: HS19031533-01MSD	Units:	mg/L	Analysis Date:	05-Apr-2019 11:45					
Client ID:	Run ID: WetChem_HS_336046	SeqNo:	5024112	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Nitrogen, Ammonia (As N)	14.88	0.20	10	4.085	108	80 - 120	15.09	1.4	20	

The following samples were analyzed in this batch: HS19031492-01

ALS Houston, US

Date: 12-Apr-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19031492	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031492

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19031492-01	LH18/24-SP650_032719	Login	3/29/2019 7:48:43 AM	JRM	WET069
HS19031492-01	LH18/24-SP650_032719	Login	3/29/2019 7:48:43 AM	JRM	WET069
HS19031492-01	LH18/24-SP650_032719	Login	3/29/2019 7:48:43 AM	JRM	Sub
HS19031492-02	LH18/24-SP650_032719_BIX	Login	3/29/2019 7:48:43 AM	JRM	Sub

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19031492

Date/Time Received: **28-Mar-2019 08:30**
 Received by: **PMG**

Checklist completed by: Jared R. Makan 29-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 29-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.5c/1.5c UC/C IR25
 Cooler(s)/Kit(s): 44581
 Date/Time sample(s) sent to storage: 03/28/2019 19:50

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:


Login Notes: Sample refrigerated prior to login.

Client Contacted: _____ Date Contacted: _____ Person Contacted: _____

Contacted By: _____ Regarding: _____

Comments:

Corrective Action:

 ALS Environmental 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5856 Fax: +1 281 530 5887	CUSTODY SEAL		Seal Broken By: <i>SM</i>
	Date: <u>3/27/19</u>	Time: <u>1430</u>	Date: <u>03/28/19</u>
	Name: <u>Scott Bessinger</u>		
	Company: <u>DELTA</u>		

44581 MAR 28 2019

Must Deliver Next Business Day
Time and Temperature Sensitive!



44581

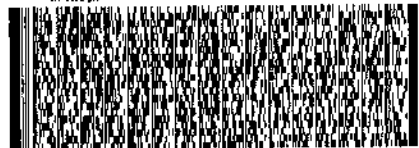
ORIGIN ID:SGRA (309) 587-2450
 SCOTT BESSINGER
 SHATE ENVIRONMENTAL ASSOCIATES
 1203-B EAST GRAND AVE. PMB202
 MARSHALL, TX 75670
 UNITED STATES US

SHIP DATE: DEMAR19
 ACTWT: 1.00 LB NWT
 CAD: 900130/CAF3211
 DIMS: 26x14x14 IN

TO **CLIENT SERVICES**
ALS LABORATORY GROUP
 10450 STANCLIFF ROAD
 SUITE 210
 HOUSTON TX 77099

(281) 530-6868
 REF: LHAAP-68-80 64113-RJ

RMA: 000000



FedEx Express



FedEx

TRK# 6221 4809 7831 3421

THU - 28 MAR 10:30A
PRIORITY OVERNIGHT

AB SGRA

77099
TX-US
IAH



FID 162785 27MAR19 GGA 553C1/4603/010A



ALS Environmental
ALS Group USA, Corp
1317 South 13th Avenue
Kelso, WA 98626
T : +1 360 577 7222
F : +1 360 636 1068
www.alsglobal.com

April 05, 2019

Analytical Report for Service Request No: K1902766

RJ Modashia
ALS Laboratory Group
10450 Stancliff Road
Suite 210
Houston, TX 77099-4338

RE: HS19031492

Dear RJ,

Enclosed are the results of the sample(s) submitted to our laboratory March 30, 2019
For your reference, these analyses have been assigned our service request number **K1902766**.

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. The test results meet requirements of the current NELAP standards, where applicable, and except as noted in the laboratory case narrative provided. For a specific list of NELAP-accredited analytes, refer to the certifications section at www.alsglobal.com. All results are intended to be considered in their entirety, and ALS Group USA Corp. dba ALS Environmental (ALS) is not responsible for use of less than the complete report. Results apply only to the items submitted to the laboratory for analysis and individual items (samples) analyzed, as listed in the report.

Please contact me if you have any questions. My extension is 3350. You may also contact me via email at Kelley.Lovejoy@alsglobal.com.

Respectfully submitted,

ALS Group USA, Corp. dba ALS Environmental

Kelley Lovejoy
Project Manager



ALS Environmental
ALS Group USA, Corp
1317 South 13th Avenue
Kelso, WA 98626
T : +1 360 577 7222
F : +1 360 636 1068
www.alsglobal.com

Table of Contents

Acronyms

Qualifiers

State Certifications, Accreditations, And Licenses

Case Narrative

Chain of Custody

General Chemistry

Raw Data

 General Chemistry

Acronyms

ASTM	American Society for Testing and Materials
A2LA	American Association for Laboratory Accreditation
CARB	California Air Resources Board
CAS Number	Chemical Abstract Service registry Number
CFC	Chlorofluorocarbon
CFU	Colony-Forming Unit
DEC	Department of Environmental Conservation
DEQ	Department of Environmental Quality
DHS	Department of Health Services
DOE	Department of Ecology
DOH	Department of Health
EPA	U. S. Environmental Protection Agency
ELAP	Environmental Laboratory Accreditation Program
GC	Gas Chromatography
GC/MS	Gas Chromatography/Mass Spectrometry
LOD	Limit of Detection
LOQ	Limit of Quantitation
LUFT	Leaking Underground Fuel Tank
M	Modified
MCL	Maximum Contaminant Level is the highest permissible concentration of a substance allowed in drinking water as established by the USEPA.
MDL	Method Detection Limit
MPN	Most Probable Number
MRL	Method Reporting Limit
NA	Not Applicable
NC	Not Calculated
NCASI	National Council of the Paper Industry for Air and Stream Improvement
ND	Not Detected
NIOSH	National Institute for Occupational Safety and Health
PQL	Practical Quantitation Limit
RCRA	Resource Conservation and Recovery Act
SIM	Selected Ion Monitoring
TPH	Total Petroleum Hydrocarbons
tr	Trace level is the concentration of an analyte that is less than the PQL but greater than or equal to the MDL.

Inorganic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- E The result is an estimate amount because the value exceeded the instrument calibration range.
- J The result is an estimated value.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.
- H The holding time for this test is immediately following sample collection. The samples were analyzed as soon as possible after receipt by the laboratory.

Metals Data Qualifiers

- # The control limit criteria is not applicable. See case narrative.
- J The result is an estimated value.
- E The percent difference for the serial dilution was greater than 10%, indicating a possible matrix interference in the sample.
- M The duplicate injection precision was not met.
- N The Matrix Spike sample recovery is not within control limits. See case narrative.
- S The reported value was determined by the Method of Standard Additions (MSA).
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- W The post-digestion spike for furnace AA analysis is out of control limits, while sample absorbance is less than 50% of spike absorbance.
- i The MRL/MDL or LOQ/LOD is elevated due to a matrix interference.
- X See case narrative.
- + The correlation coefficient for the MSA is less than 0.995.
- Q See case narrative. One or more quality control criteria was outside the limits.

Organic Data Qualifiers

- * The result is an outlier. See case narrative.
- # The control limit criteria is not applicable. See case narrative.
- A A tentatively identified compound, a suspected aldol-condensation product.
- B The analyte was found in the associated method blank at a level that is significant relative to the sample result as defined by the DOD or NELAC standards.
- C The analyte was qualitatively confirmed using GC/MS techniques, pattern recognition, or by comparing to historical data.
- D The reported result is from a dilution.
- E The result is an estimated value.
- J The result is an estimated value.
- N The result is presumptive. The analyte was tentatively identified, but a confirmation analysis was not performed.
- P The GC or HPLC confirmation criteria was exceeded. The relative percent difference is greater than 40% between the two analytical results.
- U The analyte was analyzed for, but was not detected ("Non-detect") at or above the MRL/MDL.
DOD-QSM 4.2 definition : Analyte was not detected and is reported as less than the LOD or as defined by the project. The detection limit is adjusted for dilution.
- i The MRL/MDL or LOQ/LOD is elevated due to a chromatographic interference.
- X See case narrative.
- Q See case narrative. One or more quality control criteria was outside the limits.

Additional Petroleum Hydrocarbon Specific Qualifiers

- F The chromatographic fingerprint of the sample matches the elution pattern of the calibration standard.
- L The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of lighter molecular weight constituents than the calibration standard.
- H The chromatographic fingerprint of the sample resembles a petroleum product, but the elution pattern indicates the presence of a greater amount of heavier molecular weight constituents than the calibration standard.
- O The chromatographic fingerprint of the sample resembles an oil, but does not match the calibration standard.
- Y The chromatographic fingerprint of the sample resembles a petroleum product eluting in approximately the correct carbon range, but the elution pattern does not match the calibration standard.
- Z The chromatographic fingerprint does not resemble a petroleum product.

**ALS Group USA Corp. dba ALS Environmental (ALS) - Kelso
State Certifications, Accreditations, and Licenses**

Agency	Web Site	Number
Alaska DEH	http://dec.alaska.gov/eh/lab/cs/csapproval.htm	UST-040
Arizona DHS	http://www.azdhs.gov/lab/license/env.htm	AZ0339
Arkansas - DEQ	http://www.adeq.state.ar.us/techsvs/labcert.htm	88-0637
California DHS (ELAP)	http://www.cdph.ca.gov/certlic/labs/Pages/ELAP.aspx	2795
DOD ELAP	http://www.denix.osd.mil/edqw/Accreditation/AccreditedLabs.cfm	L16-58-R4
Florida DOH	http://www.doh.state.fl.us/lab/EnvLabCert/WaterCert.htm	E87412
Hawaii DOH	http://health.hawaii.gov/	-
ISO 17025	http://www.pjllabs.com/	L16-57
Louisiana DEQ	http://www.deq.louisiana.gov/page/la-lab-accreditation	03016
Maine DHS	http://www.maine.gov/dhhs/	WA01276
Minnesota DOH	http://www.health.state.mn.us/accreditation	053-999-457
Nevada DEP	http://ndep.nv.gov/bsdw/labservice.htm	WA01276
New Jersey DEP	http://www.nj.gov/dep/enforcement/oqa.html	WA005
New York - DOH	https://www.wadsworth.org/regulatory/elap	12060
North Carolina DEQ	https://deq.nc.gov/about/divisions/water-resources/water-resources-data/water-sciences-home-page/laboratory-certification-branch/non-field-lab-certification	605
Oklahoma DEQ	http://www.deq.state.ok.us/CSDnew/labcert.htm	9801
Oregon – DEQ (NELAP)	http://public.health.oregon.gov/LaboratoryServices/EnvironmentalLaboratoryAccreditation/Pages/index.aspx	WA100010
South Carolina DHEC	http://www.scdhec.gov/environment/EnvironmentalLabCertification/	61002
Texas CEQ	http://www.tceq.texas.gov/field/qa/env_lab_accreditation.html	T104704427
Washington DOE	http://www.ecy.wa.gov/programs/eap/labs/lab-accreditation.html	C544
Wyoming (EPA Region 8)	https://www.epa.gov/region8-waterops/epa-region-8-certified-drinking-water	-
Kelso Laboratory Website	www.alsglobal.com	NA

Analyses were performed according to our laboratory's NELAP-approved quality assurance program. A complete listing of specific NELAP-certified analytes, can be found in the certification section at www.ALSGlobal.com or at the accreditation bodies web site.

Please refer to the certification and/or accreditation body's web site if samples are submitted for compliance purposes. The states highlighted above, require the analysis be listed on the state certification if used for compliance purposes and if the method/analyte is offered by that state.



Case Narrative

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com



Client: ALS Environmental - US
Project: HS19031492
Sample Matrix: Water

Service Request: K1902766
Date Received: 03/30/2019

CASE NARRATIVE

All analyses were performed consistent with the quality assurance program of ALS Environmental. This report contains analytical results for samples for the Tier level IV requested by the client.

Sample Receipt:

One water sample was received for analysis at ALS Environmental on 03/30/2019. Any discrepancies upon initial sample inspection are annotated on the sample receipt and preservation form included within this report. The sample was stored at minimum in accordance with the analytical method requirements.

General Chemistry:

No significant anomalies were noted with this analysis.

Approved by Kelley Lovejoy

Date 04/05/2019



Chain of Custody

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com



K1902766

00935669

10450 Standliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 11011

SUBCONTRACT TO:

ALS Environmental Kelso
1317 S. 13th Avenue
Kelso, WA 98626

Phone: +1 360 501 3312

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Standliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Standliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19031492
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19031492-01	LH18/24-SP650_032719	Water	27 Mar 2019 14:00
TOC Analysis for DOD Level IV			05 Apr 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: [Signature]
Received By: [Signature]
Cooler ID(s): _____

Date/Time: 3/29/19 18:00
Date/Time: 3/30/19 1100
Temperature(s): _____

RIGHT SOLUTIONS | RIGHT PARTNER



PC KL

Cooler Receipt and Preservation Form

Client ALS/Houston Service Request K19 02766

Received: 3/30/19 Opened: 3/30/19 By: [Signature] Unloaded: 3/30/19 By: [Signature]

- Samples were received via? USPS Fed Ex UPS DHL PDX Courier Hand Delivered
- Samples were received in: (circle) Cooler Box Envelope Other NA
- Were custody seals on coolers? NA Y N If yes, how many and where? 2, front
If present, were custody seals intact? Y N If present, were they signed and dated? Y N

Raw Cooler Temp	Corrected Cooler Temp	Raw Temp Blank	Corrected Temp Blank	Corr. Factor	Thermometer ID	Cooler/COC ID	Tracking Number	NA	Filed
0.1	0.1	1.5	1.5	0	349	Various	4809 7832	2378	
0.7	1.0	0.2	0.5	+0.3	384		" "	2389	
						11011			

- Packing material: Inserts Baggies Bubble Wrap Gel Packs Wet Ice Dry Ice Sleeves
- Were custody papers properly filled out (ink, signed, etc.)? NA Y N
- Were samples received in good condition (temperature, unbroken)? *Indicate in the table below.* NA Y N
If applicable, tissue samples were received: Frozen Partially Thawed Thawed
- Were all sample labels complete (i.e analysis, preservation, etc.)? NA Y N
- Did all sample labels and tags agree with custody papers? *Indicate major discrepancies in the table on page 2.* NA Y N
- Were appropriate bottles/containers and volumes received for the tests indicated? NA Y N
- Were the pH-preserved bottles (see SMO GEN SOP) received at the appropriate pH? *Indicate in the table below* NA Y N
- Were VOA vials received without headspace? *Indicate in the table below.* NA Y N
- Was C12/Res negative? NA Y N

Sample ID on Bottle	Sample ID on COC	Identified by:

Sample ID	Bottle Count	Bottle Type	Out of Temp	Head-space	Broke	pH	Reagent	Volume added	Reagent Lot Number	Initials	Time

Notes, Discrepancies, & Resolutions: _____



General Chemistry

ALS Environmental—Kelso Laboratory
1317 South 13th Avenue, Kelso, WA 98626
Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Analytical Report

Client: ALS Environmental - US
Project: HS19031492
Sample Matrix: Water
Analysis Method: SM 5310 C
Prep Method: None

Service Request: K1902766
Date Collected: 03/27/19
Date Received: 03/30/19
Units: mg/L
Basis: NA

Carbon, Total Organic

Sample Name	Lab Code	Result	LOQ	LOD	MDL	Dil.	Date Analyzed	Q
LH18/24-SP650_032719	K1902766-001	2.29	0.50	0.20	0.07	1	04/04/19 08:25	
Method Blank	K1902766-MB	ND U	0.50	0.20	0.07	1	04/04/19 06:02	

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19031492
Sample Matrix: Water

Service Request: K1902766
Date Collected: N/A
Date Received: N/A
Date Analyzed: 04/4/19
Date Extracted: NA

Matrix Spike Summary
Carbon, Total Organic

Sample Name: Batch QC
Lab Code: K1902703-001
Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA

Matrix Spike
K1902703-001MS

<u>Analyte Name</u>	<u>Sample Result</u>	<u>Result</u>	<u>Spike Amount</u>	<u>% Rec</u>	<u>% Rec Limits</u>
Carbon, Total Organic	3.05	29.7	25.0	107	83-117

Results flagged with an asterisk (*) indicate values outside control criteria.

Results flagged with a pound (#) indicate the control criteria is not applicable.

Percent recoveries and relative percent differences (RPD) are determined by the software using values in the calculation which have not been rounded.

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19031492
Sample Matrix: Water

Service Request: K1902766
Date Analyzed: 04/04/19
Date Extracted: NA

Lab Control Sample Summary
Carbon, Total Organic

Analysis Method: SM 5310 C
Prep Method: None

Units: mg/L
Basis: NA
Analysis Lot: 630890

Sample Name	Lab Code	Result	Spike Amount	% Rec	% Rec Limits
Lab Control Sample	K1902766-LCS	26.4	25.0	106	83-117

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19031492

Service Request: K1902766

Continuing Calibration Verification (CCV) Summary

Carbon, Total Organic

Analysis Method: SM 5310 C

Units: mg/L

	Analysis		Date	True	Measured	Percent	Acceptance
	Lot	Lab Code	Analyzed	Value	Value	Recovery	Limits
CCV1	630890	KQ1904456-04	04/04/19 05:33	25.0	25.5	102	90-110
CCV2	630890	KQ1904456-05	04/04/19 09:21	25.0	25.3	101	90-110
CCV3	630890	KQ1904456-06	04/04/19 14:32	25.0	25.8	103	90-110

ALS Group USA, Corp.
dba ALS Environmental

QA/QC Report

Client: ALS Environmental - US
Project: HS19031492

Service Request:K1902766

Continuing Calibration Blank (CCB) Summary
Carbon, Total Organic

Analysis Method: SM 5310 C

Units:mg/L

	Analysis Lot	Lab Code	Date Analyzed	LOQ	LOD	MDL	Result	Q
CCB1	630890	KQ1904456-01	04/04/19 05:47	0.50	0.20	0.07	ND	U
CCB2	630890	KQ1904456-02	04/04/19 09:36	0.50	0.20	0.07	ND	U
CCB3	630890	KQ1904456-03	04/04/19 14:46	0.50	0.20	0.07	ND	U



Raw Data

ALS Environmental—Kelso Laboratory
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General Chemistry

ALS Environmental—Kelso Laboratory
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Phone (360)577- 7222 Fax (360)636-1 068
www.alsglobal.com

Work Request # (Original) K1902928, 2829, 2526, 2660, 2705, 2766, 2818, 2831, 2853
 Tier: IV IV II II II IV # II II
 Date Analyzed: 4/13/19 TOC:630888, 630890
 Analyst: BCP Run # DOC:630891
 Analysis: TOC/DOC

DATA QUALITY REPORT INORGANICS

Explain any "no" responses to questions below, and any corrective actions in the comments section below.

1. Is the method name and number correct and appropriate? yes/no/NA
2. Holding times met for all analyses and for all samples? yes/no/NA
3. Are calculations correct? yes/no/NA
4. Is the reporting basis correct? (Dry Weight) yes/no/NA
5. All quality control criteria met? yes/no
6. Is the calibration curve correlation coefficient ≥ 0.995 ? yes/no/NA
7. MBs, CCVs, CCBs, LCSs, Dups, and Spikes, analyzed at proper frequency? yes/no/NA
8. Are ICVs, CCVs, and CCBs all within acceptance limits? yes/no/NA
9. Are results for methods blanks all ND? yes/no/NA
10. Are all QC samples within acceptance criteria? (LCS % rec, MS/DMS % rec, DUP or MS/DMS RPDs, etc.) yes/no/NA
11. Are all exceptions explained? yes/no/NA
12. Have all applicable service requests been reviewed? yes/no/NA
13. Are all samples labeled correctly? yes/no/NA
14. Have all instructions on the service request been followed? (e.g. Special MRLs, QC on a specific sample, Form V) yes/no/NA
15. Are detection limits and units reported correctly? yes/no/NA
16. Is the unused space on the benchsheet crossed out? yes/no/NA
17. Was analysis turned in by the due date? (n-2) (If not record SR#) yes/no/NA

COMMENTS: K1902660-3/3d report a high %RSD. However, these samples are less than the MRL.

Final Approved by: Fannings Date: 04/05/19 DQREPORT

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 630888 Method/Testcode: 9060/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902828-003	Carbon, Total Organic	N/A		Ground Water	0.88 mg/L	10 ml	0.88 mg/L	1	0.07	0.50			4/3/19 16:30	N	IV
K1902828-004	Carbon, Total Organic	N/A		Ground Water	0.86 mg/L	10 ml	0.86 mg/L	1	0.07	0.50			4/3/19 17:26	Y	IV
K1902828-005	Carbon, Total Organic	N/A		Ground Water	0.90 mg/L	10 ml	0.90 mg/L	1	0.07	0.50			4/3/19 19:32	N	IV
K1902828-006	Carbon, Total Organic	N/A		Ground Water	1.47 mg/L	10 ml	1.47 mg/L	1	0.07	0.50			4/3/19 20:57	N	IV
K1902828-007	Carbon, Total Organic	N/A		Ground Water	1.51 mg/L	10 ml	1.51 mg/L	1	0.07	0.50			4/3/19 21:53	N	IV
K1902828-008	Carbon, Total Organic	N/A		Ground Water	0.01 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/3/19 22:49	N	IV
K1902829-003	Carbon, Total Organic	N/A		Ground Water	0.52 mg/L	10 ml	0.52 mg/L	1	0.07	0.50			4/3/19 23:44	N	IV
K1902829-004	Carbon, Total Organic	N/A		Ground Water	0.37 mg/L	10 ml	0.37 mg/L J	1	0.07	0.50			4/4/19 00:40	Y	IV
K1902829-005	Carbon, Total Organic	N/A		Ground Water	0.23 mg/L	10 ml	0.23 mg/L J	1	0.07	0.50			4/4/19 02:46	N	IV
K1902829-006	Carbon, Total Organic	N/A		Ground Water	0.21 mg/L	10 ml	0.21 mg/L J	1	0.07	0.50			4/4/19 03:41	N	IV
K1902829-007	Carbon, Total Organic	N/A		Ground Water	8.38 mg/L	10 ml	8.38 mg/L	1	0.07	0.50			4/4/19 04:37	N	IV
KQ1904455-01	Carbon, Total Organic	CCB		Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/3/19 12:17	N	IV
KQ1904455-02	Carbon, Total Organic	CCB		Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/3/19 20:42	N	IV
KQ1904455-03	Carbon, Total Organic	CCB		Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/4/19 05:47	N	IV
KQ1904455-04	Carbon, Total Organic	CCV		Ground Water	25.78 mg/L	10 ml	25.8 mg/L	1					4/3/19 12:02	N	IV
KQ1904455-05	Carbon, Total Organic	CCV		Ground Water	25.44 mg/L	10 ml	25.4 mg/L	1					4/3/19 20:28	N	IV
KQ1904455-06	Carbon, Total Organic	CCV		Ground Water	25.51 mg/L	10 ml	25.5 mg/L	1					4/4/19 05:33	N	IV
KQ1904455-07	Carbon, Total Organic	MB		Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/3/19 12:32	N	IV
KQ1904455-08	Carbon, Total Organic	MB		Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/3/19 12:32	N	IV
KQ1904455-09	Carbon, Total Organic	MB		Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/3/19 12:32	N	IV
KQ1904455-10	Carbon, Total Organic	MB		Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50			4/3/19 12:32	N	IV
KQ1904455-11	Carbon, Total Organic	LCS		Ground Water	26.27 mg/L	10 ml	26.3 mg/L	1	0.07	0.50	105		4/3/19 13:28	N	IV
KQ1904455-12	Carbon, Total Organic	LCS		Ground Water	26.32 mg/L	10 ml	26.3 mg/L	1	0.07	0.50	105		4/3/19 13:28	N	IV

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Printed 4/5/19 10:08

Results Summary

04/05/19
Hawley

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 630888 Method/Testcode: 9060/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1904455-13	Carbon, Total Organic	LCS		Ground Water	26.41 mg/L	10 ml	26.4 mg/L	1	0.07	0.50	106		4/3/19 13:28	N	IV
KQ1904455-14	Carbon, Total Organic	LCS		Ground Water	26.29 mg/L	10 ml	26.3 mg/L	1	0.07	0.50	105		4/3/19 13:28	N	IV
KQ1904455-15	Carbon, Total Organic	MS	K1902828-004	Ground Water	28.15 mg/L	10 ml	28.1 mg/L	1	0.07	0.50	109		4/3/19 18:22	N	IV
KQ1904455-16	Carbon, Total Organic	MS	K1902828-004	Ground Water	28.12 mg/L	10 ml	28.1 mg/L	1	0.07	0.50	109		4/3/19 18:22	N	IV
KQ1904455-17	Carbon, Total Organic	MS	K1902828-004	Ground Water	27.87 mg/L	10 ml	27.9 mg/L	1	0.07	0.50	108		4/3/19 18:22	N	IV
KQ1904455-18	Carbon, Total Organic	MS	K1902828-004	Ground Water	27.80 mg/L	10 ml	27.8 mg/L	1	0.07	0.50	108		4/3/19 18:22	N	IV
KQ1904455-19	Carbon, Total Organic	MS	K1902829-004	Ground Water	27.22 mg/L	10 ml	27.2 mg/L	1	0.07	0.50	107		4/4/19 01:36	N	IV
KQ1904455-20	Carbon, Total Organic	MS	K1902829-004	Ground Water	27.07 mg/L	10 ml	27.1 mg/L	1	0.07	0.50	107		4/4/19 01:36	N	IV
KQ1904455-21	Carbon, Total Organic	MS	K1902829-004	Ground Water	27.23 mg/L	10 ml	27.2 mg/L	1	0.07	0.50	107		4/4/19 01:36	N	IV
KQ1904455-22	Carbon, Total Organic	MS	K1902829-004	Ground Water	27.03 mg/L	10 ml	27.0 mg/L	1	0.07	0.50	107		4/4/19 01:36	N	IV
KQ1904455-23	Carbon, Total Organic	DUP	K1902828-003	Ground Water	0.87 mg/L	10 ml	0.87 mg/L	1	0.07	0.50		<1	4/3/19 16:30	N	IV
KQ1904455-24	Carbon, Total Organic	TRP	K1902828-003	Ground Water	0.89 mg/L	10 ml	0.89 mg/L	1	0.07	0.50		1	4/3/19 16:30	N	IV
KQ1904455-25	Carbon, Total Organic	QUAD	K1902828-003	Ground Water	0.86 mg/L	10 ml	0.86 mg/L	1	0.07	0.50		1	4/3/19 16:30	N	IV
KQ1904455-26	Carbon, Total Organic	DUP	K1902828-004	Ground Water	0.82 mg/L	10 ml	0.82 mg/L	1	0.07	0.50		5	4/3/19 17:26	N	IV
KQ1904455-27	Carbon, Total Organic	TRP	K1902828-004	Ground Water	0.83 mg/L	10 ml	0.83 mg/L	1	0.07	0.50		3	4/3/19 17:26	N	IV
KQ1904455-28	Carbon, Total Organic	QUAD	K1902828-004	Ground Water	0.83 mg/L	10 ml	0.83 mg/L	1	0.07	0.50		2	4/3/19 17:26	N	IV
KQ1904455-29	Carbon, Total Organic	DUP	K1902828-005	Ground Water	0.85 mg/L	10 ml	0.85 mg/L	1	0.07	0.50		6	4/3/19 19:32	N	IV
KQ1904455-30	Carbon, Total Organic	TRP	K1902828-005	Ground Water	0.84 mg/L	10 ml	0.84 mg/L	1	0.07	0.50		4	4/3/19 19:32	N	IV
KQ1904455-31	Carbon, Total Organic	QUAD	K1902828-005	Ground Water	0.82 mg/L	10 ml	0.82 mg/L	1	0.07	0.50		4	4/3/19 19:32	N	IV
KQ1904455-32	Carbon, Total Organic	DUP	K1902828-006	Ground Water	1.45 mg/L	10 ml	1.45 mg/L	1	0.07	0.50		1	4/3/19 20:57	N	IV
KQ1904455-33	Carbon, Total Organic	TRP	K1902828-006	Ground Water	1.41 mg/L	10 ml	1.41 mg/L	1	0.07	0.50		2	4/3/19 20:57	N	IV
KQ1904455-34	Carbon, Total Organic	QUAD	K1902828-006	Ground Water	1.44 mg/L	10 ml	1.44 mg/L	1	0.07	0.50		2	4/3/19 20:57	N	IV
KQ1904455-35	Carbon, Total Organic	DUP	K1902828-007	Ground Water	1.44 mg/L	10 ml	1.44 mg/L	1	0.07	0.50		5	4/3/19 21:53	N	IV

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 630888 Method/Testcode: 9060/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
KQ1904455-36	Carbon, Total Organic	TRP	K1902828-007	Ground Water	1.44 mg/L	10 ml	1.44 mg/L	1	0.07	0.50		3	4/3/19 21:53	N	IV
KQ1904455-37	Carbon, Total Organic	QUAD	K1902828-007	Ground Water	1.47 mg/L	10 ml	1.47 mg/L	1	0.07	0.50		2	4/3/19 21:53	N	IV
KQ1904455-38	Carbon, Total Organic	DUP	K1902828-008	Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50		NC	4/3/19 22:49	N	IV
KQ1904455-39	Carbon, Total Organic	TRP	K1902828-008	Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50		NC	4/3/19 22:49	N	IV
KQ1904455-40	Carbon, Total Organic	QUAD	K1902828-008	Ground Water	0.00 mg/L	10 ml	0.50 mg/L U	1	0.07	0.50		NC	4/3/19 22:49	N	IV
KQ1904455-41	Carbon, Total Organic	DUP	K1902829-003	Ground Water	0.52 mg/L	10 ml	0.52 mg/L	1	0.07	0.50		<1	4/3/19 23:44	N	IV
KQ1904455-42	Carbon, Total Organic	TRP	K1902829-003	Ground Water	0.51 mg/L	10 ml	0.51 mg/L	1	0.07	0.50		1	4/3/19 23:44	N	IV
KQ1904455-43	Carbon, Total Organic	QUAD	K1902829-003	Ground Water	0.53 mg/L	10 ml	0.53 mg/L	1	0.07	0.50		2	4/3/19 23:44	N	IV
KQ1904455-44	Carbon, Total Organic	DUP	K1902829-004	Ground Water	0.35 mg/L	10 ml	0.35 mg/L J	1	0.07	0.50		7	4/4/19 00:40	N	IV
KQ1904455-45	Carbon, Total Organic	TRP	K1902829-004	Ground Water	0.38 mg/L	10 ml	0.38 mg/L J	1	0.07	0.50		5	4/4/19 00:40	N	IV
KQ1904455-46	Carbon, Total Organic	QUAD	K1902829-004	Ground Water	0.35 mg/L	10 ml	0.35 mg/L J	1	0.07	0.50		4	4/4/19 00:40	N	IV
KQ1904455-47	Carbon, Total Organic	DUP	K1902829-005	Ground Water	0.25 mg/L	10 ml	0.25 mg/L J	1	0.07	0.50		12	4/4/19 02:46	N	IV
KQ1904455-48	Carbon, Total Organic	TRP	K1902829-005	Ground Water	0.18 mg/L	10 ml	0.18 mg/L J	1	0.07	0.50		18	4/4/19 02:46	N	IV
KQ1904455-49	Carbon, Total Organic	QUAD	K1902829-005	Ground Water	0.24 mg/L	10 ml	0.24 mg/L J	1	0.07	0.50		15	4/4/19 02:46	N	IV
KQ1904455-50	Carbon, Total Organic	DUP	K1902829-006	Ground Water	0.24 mg/L	10 ml	0.24 mg/L J	1	0.07	0.50		17	4/4/19 03:41	N	IV
KQ1904455-51	Carbon, Total Organic	TRP	K1902829-006	Ground Water	0.20 mg/L	10 ml	0.20 mg/L J	1	0.07	0.50		11	4/4/19 03:41	N	IV
KQ1904455-52	Carbon, Total Organic	QUAD	K1902829-006	Ground Water	0.19 mg/L	10 ml	0.19 mg/L J	1	0.07	0.50		11	4/4/19 03:41	N	IV
KQ1904455-53	Carbon, Total Organic	DUP	K1902829-007	Ground Water	8.27 mg/L	10 ml	8.27 mg/L	1	0.07	0.50		1	4/4/19 04:37	N	IV
KQ1904455-54	Carbon, Total Organic	TRP	K1902829-007	Ground Water	8.42 mg/L	10 ml	8.42 mg/L	1	0.07	0.50		<1	4/4/19 04:37	N	IV
KQ1904455-55	Carbon, Total Organic	QUAD	K1902829-007	Ground Water	8.37 mg/L	10 ml	8.37 mg/L	1	0.07	0.50		<1	4/4/19 04:37	N	IV

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 630890 Method/Testcode: SM 5310 C/TOC T

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Ti
K1902526-003	Carbon, Total Organic	N/A		Water	4.23 mg/L	10 ml	423 mg/L	100	7	50			4/4/19 09:51	N	I
K1902526-004	Carbon, Total Organic	N/A		Water	4.85 mg/L	10 ml	485 mg/L	100	7	50			4/4/19 10:19	N	I
K1902660-001	Carbon, Total Organic	N/A		Ground Water	2.00 mg/L	10 ml	2.00 mg/L	1	0.07	0.50			4/4/19 11:16	N	I
K1902660-002	Carbon, Total Organic	N/A		Ground Water	1.17 mg/L	10 ml	1.17 mg/L	1	0.07	0.50			4/4/19 12:12	N	II
K1902660-003	Carbon, Total Organic	N/A		Ground Water	0.12 mg/L	10 ml	0.12 mg/L	J	1	0.07	0.50		4/4/19 12:40	N	II
K1902703-001	Carbon, Total Organic	N/A		Water	3.05 mg/L	10 ml	3.05 mg/L	1	0.07	0.50			4/4/19 07:28	N	II
K1902766-001	Carbon, Total Organic	N/A		Water	2.29 mg/L	10 ml	2.29 mg/L	1	0.07	0.50			4/4/19 08:25	N	I
K1902818-001	Carbon, Total Organic	N/A		Water	3.29 mg/L	10 ml	3.29 mg/L	1	0.07	0.50			4/4/19 13:36	N	II
K1902818-003	Carbon, Total Organic	N/A		Water	2.47 mg/L	10 ml	2.47 mg/L	1	0.07	0.50			4/4/19 14:04	N	II
K1902831-001	Carbon, Total Organic	N/A		Water	5.74 mg/L	10 ml	574 mg/L	100	7	50			4/4/19 13:08	N	II
K1902853-001	Carbon, Total Organic	N/A		Water	4.86 mg/L	10 ml	4.86 mg/L	1	0.07	0.50			4/4/19 08:53	N	II
KQ1904456-01	Carbon, Total Organic	CCB		Water	0.00 mg/L	10 ml	0.50 mg/L	U	1	0.07	0.50		4/4/19 05:47	N	II
KQ1904456-02	Carbon, Total Organic	CCB		Water	0.00 mg/L	10 ml	0.50 mg/L	U	1	0.07	0.50		4/4/19 09:36	N	II
KQ1904456-03	Carbon, Total Organic	CCB		Water	0.00 mg/L	10 ml	0.50 mg/L	U	1	0.07	0.50		4/4/19 14:46	N	II
KQ1904456-04	Carbon, Total Organic	CCV		Water	25.51 mg/L	10 ml	25.5 mg/L	1					4/4/19 05:33	N	II
KQ1904456-05	Carbon, Total Organic	CCV		Water	25.25 mg/L	10 ml	25.3 mg/L	1					4/4/19 09:21	N	II
KQ1904456-06	Carbon, Total Organic	CCV		Water	25.75 mg/L	10 ml	25.8 mg/L	1					4/4/19 14:32	N	II
KQ1904456-07	Carbon, Total Organic	LCS		Water	26.39 mg/L	10 ml	26.4 mg/L	1	0.07	0.50			4/4/19 06:17	N	II
KQ1904456-08	Carbon, Total Organic	MB		Water	0.00 mg/L	10 ml	0.50 mg/L	U	1	0.07	0.50		4/4/19 06:02	N	II
KQ1904456-09	Carbon, Total Organic	MS	K1902703-001	Water	29.74 mg/L	10 ml	29.7 mg/L	1	0.07	0.50			4/4/19 07:56	N	II
KQ1904456-10	Carbon, Total Organic	DUP	K1902703-001	Water	3.01 mg/L	10 ml	3.01 mg/L	1	0.07	0.50			4/4/19 07:28	N	II
KQ1904456-11	Carbon, Total Organic	DUP	K1902766-001	Water	2.20 mg/L	10 ml	2.20 mg/L	1	0.07	0.50			4/4/19 08:25	N	IV
KQ1904456-12	Carbon, Total Organic	DUP	K1902853-001	Water	4.78 mg/L	10 ml	4.78 mg/L	1	0.07	0.50			4/4/19 08:53	N	II
KQ1904456-13	Carbon, Total Organic	DUP	K1902526-003	Water	4.07 mg/L	10 ml	407 mg/L	100	7	50			4/4/19 09:51	N	II
KQ1904456-14	Carbon, Total Organic	DUP	K1902526-004	Water	4.78 mg/L	10 ml	478 mg/L	100	7	50			4/4/19 10:19	N	II
KQ1904456-15	Carbon, Total Organic	DUP	K1902660-001	Ground Water	1.86 mg/L	10 ml	1.86 mg/L	1	0.07	0.50			4/4/19 11:16	N	II
KQ1904456-16	Carbon, Total Organic	DUP	K1902660-002	Ground Water	1.13 mg/L	10 ml	1.13 mg/L	1	0.07	0.50			4/4/19 12:12	N	II
KQ1904456-17	Carbon, Total Organic	DUP	K1902660-003	Ground Water	0.15 mg/L	10 ml	0.15 mg/L	J	1	0.07	0.50		4/4/19 12:40	N	II
KQ1904456-18	Carbon, Total Organic	DUP	K1902831-001	Water	5.83 mg/L	10 ml	583 mg/L	100	7	50			4/4/19 13:08	N	II
KQ1904456-19	Carbon, Total Organic	DUP	K1902818-001	Water	3.28 mg/L	10 ml	3.28 mg/L	1	0.07	0.50			4/4/19 13:36	N	II
KQ1904456-20	Carbon, Total Organic	DUP	K1902818-003	Water	2.46 mg/L	10 ml	2.46 mg/L	1	0.07	0.50			4/4/19 14:04	N	II

indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

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Analytical Results Summary

Instrument Name: K-TOC-03

Analyst: BDITZLER

Analysis Lot: 630891 Method/Testcode: SM 5310 C/TOC D

Lab Code	Target Analytes	QC	Parent Sample	Matrix	Raw Result	Sample Amt.	Final Result	Dil	MDL	PQL	% Rec	% RSD	Date Analyzed	QC?	Tier
K1902818-002	Carbon, Dissolved Organic (DOC)	N/A		Water	3.20 mg/L	10 ml	3.20 mg/L	1	0.07	0.50			4/4/19 15:31	N	II
K1902818-004	Carbon, Dissolved Organic (DOC)	N/A		Water	2.36 mg/L	10 ml	2.36 mg/L	1	0.07	0.50			4/4/19 16:28	N	II
KQ1904457-01	Carbon, Dissolved Organic (DOC)	CCB		Water	0.00 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			4/4/19 14:46	N	II
KQ1904457-02	Carbon, Dissolved Organic (DOC)	CCB		Water	0.00 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			4/4/19 17:11	N	II
KQ1904457-03	Carbon, Dissolved Organic (DOC)	CCV		Water	25.75 mg/L	10 ml	25.8 mg/L	1			103		4/4/19 14:32	N	II
KQ1904457-04	Carbon, Dissolved Organic (DOC)	CCV		Water	25.70 mg/L	10 ml	25.7 mg/L	1			103		4/4/19 16:56	N	II
KQ1904457-05	Carbon, Dissolved Organic (DOC)	LCS		Water	26.54 mg/L	10 ml	26.5 mg/L	1	0.07	0.50	106		4/4/19 15:16	N	II
KQ1904457-06	Carbon, Dissolved Organic (DOC)	MB		Water	0.00 mg/L	10 ml	0.50 mg/L	U 1	0.07	0.50			4/4/19 15:01	N	II
KQ1904457-07	Carbon, Dissolved Organic (DOC)	MS	K1902818-002	Water	30.57 mg/L	10 ml	30.6 mg/L	1	0.07	0.50	110		4/4/19 15:59	N	II
KQ1904457-08	Carbon, Dissolved Organic (DOC)	DUP	K1902818-002	Water	3.11 mg/L	10 ml	3.11 mg/L	1	0.07	0.50		3	4/4/19 15:31	N	II
KQ1904457-09	Carbon, Dissolved Organic (DOC)	DUP	K1902818-004	Water	2.28 mg/L	10 ml	2.28 mg/L	1	0.07	0.50		4	4/4/19 16:28	N	II

04/05/19
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indicates Final Result is not yet adjusted for Solids because it has not yet been determined.

TOC: 630888,
630890
DOC: 630891

Schedule: 04032019

Version: 4

Instrument: Fusion1

Last Saved by: Fusion1 (Fusion1)

Last Saved on: 2019/04/03 10:38 - Wednesday

Position	Sample Type	Sample ID	Method ID (Calibration ID)	Reps	Use	State
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Clean)	Clean	Clean		1	True	Ready
(Blank)	Blank	Reagent/Acid Blank		1	True	Ready
D	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
1	Sample	MB1	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
2	Sample	ICS	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
3	Sample	LOD	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
4	Sample	LOQ	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
5	Sample	K1902828-003.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
6	Sample	K1902828-004.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
7	Sample	K1902828-004.33 ms	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
8	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
9	Sample	K1902828-005.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
10	Sample	K1902828-006.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
11	Sample	K1902828-007.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
12	Sample	K1902828-008.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
13	Sample	K1902829-003.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
14	Sample	K1902829-004.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
15	Sample	K1902829-004.33 ms	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
16	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
17	Sample	K1902829-005.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
18	Sample	K1902829-006.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
19	Sample	K1902829-007.08	CAS_salt_010711 (CAS_salt_010711)	4	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
20	Sample	MB2	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
C	Check Standard	[TOC] LCS [24.0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
21	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
22	Sample	LOD	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
23	Sample	LOQ	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
24	Sample	K1902703-001.02	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
25	Sample	K1902703-001.02 ms	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
26	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
27	Sample	K1902766-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
28	Sample	K1902853-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
D	Check Standard	[TOC] CCB [0 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready
29	Sample	K1902526-003.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
30	Sample	K1902526-004.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
31	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
32	Sample	K1902660-001.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
33	Sample	RB	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
34	Sample	K1902660-002.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
35	Sample	K1902660-003.03	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
36	Sample	K1902831-001.01 100x	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
37	Sample	K1902818-001.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
38	Sample	K1902818-003.01	CAS_salt_010711 (CAS_salt_010711)	2	True	Ready
B	Check Standard	[TOC] CCV 25 ppm [25 ppm]	CAS_salt_010711 (CAS_salt_010711)	1	True	Ready

Printed on: April 5, 2019 09:08:21

04/05/19
H. Williams

Schedule: 04032019

Case ID	Sample Type	Sample ID	Method ID / Description ID	Page	Job	Status
41	Check Standard	TCC CCB (3 ppm)	CAS met 010711 (CAS met 010711)	1	True	Ready
42	Check Standard	TCC LUS (4.0 ppm)	CAS met 010711 (CAS met 010711)	1	True	Ready
43	Sample	N1002518-001 (2) max conc	CAS met 010711 (CAS met 010711)	1	True	Ready
44	Sample	N1002518-004 (2) max conc	CAS met 010711 (CAS met 010711)	2	True	Ready
45	Check Standard	TCC CCB (3 ppm)	CAS met 010711 (CAS met 010711)	1	True	Ready
					False	

Fusion Report - 04032019

Wednesday, April 03, 2019 09:54 AM

(View - Reps, Unused Reps, Meta-Data, Signature, History)
Printed on 2019/04/05 09:08 - Friday

Report Summary Information

Company Location:	Gen Chem Lab	Engine Version:	1.1.5.1
Schedule Name:	04032019	Firmware Version:	1.2.0696
Instrument Name:	Fusion1	Connection:	RS232 COM1
Report Version:	1 of 1		
Report Creation by Operators (schedule version):	Fusion1 (Fusion1) (v2) Fusion1 (Fusion1) (v3) Fusion1 (Fusion1) (v4)		
Comment:			

Report Results

04/05/19
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Sample Type: Clean From Schedule Version 2

Pos	Analysis Type	Sample ID	Start Time
◆ (clean)		Clean	2019/04/03 09:55

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	13.40	18.23	4.83	49.70	05:26
2	TC Clean	10.63	13.62	2.99	49.84	07:19
3	TC Clean	2.30	5.41	3.10	49.87	07:03
4	TC Clean	1.52	4.69	3.17	49.88	07:00

Sample Type: Clean From Schedule Version 3

Pos	Analysis Type	Sample ID	Start Time
◆ (clean)		Clean	2019/04/03 10:27

Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	13.30	16.37	3.07	49.71	05:24
2	TC Clean	4.02	7.14	3.11	49.86	07:16
3	TC Clean	1.69	4.73	3.04	50.02	03:49
4	TC Clean	1.42	4.51	3.09	50.04	03:56

Sample Type: Clean			From Schedule Version 4			
Pos	Analysis Type	Sample ID			Start Time	
◊ (clean)		Clean			2019/04/03 10:52	
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	0.89	3.96	3.07	49.77	05:23
2	TC Clean	3.44	6.46	3.03	50.01	04:03
3	TC Clean	1.68	4.69	3.01	50.05	03:48
4	TC Clean	1.69	4.50	2.81	50.03	03:49

Sample Type: Blank (Creating v1242)			From Schedule Version 4			
Pos	Analysis Type	Sample ID			Start Time	
◊ (blank)		Reagent/Acid Blank			2019/04/03 11:14	
Rep #	Base Analysis Type	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	IC Clean	0.73	3.76	3.03	49.76	05:26
2	TC Clean	3.88	6.85	2.97	49.99	04:04
3	TC Clean	2.13	5.12	2.99	50.00	03:55
4	TC Clean	1.72	4.93	3.22	49.99	03:56
5	Reagent Blank	3.55	6.56	3.01	50.01	05:06
6	Acid Blank	1.06	3.83	2.77	49.72	05:30

Sample Type: Sample			From Schedule Version 4					
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
◊ D	TOC	RB	0.4227 ppm	0.0000 ppm	0.0000%	2019/04/03 11:48		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4227	4.2266	11.62	14.73	3.11	50.09	10:34
Dilution		Blank Contribution		Method		Calibration		
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.7836 ppm (PASS)	0.0000 ppm	0%	2019/04/03 12:02

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.7836	257.8359	184.48	187.58	3.10	50.10	10:35

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/04/03 12:17

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	8.04	11.15	3.11	50.10	10:32

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 1	TOC	MB1	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/03 12:32

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	7.65	10.39	2.75	50.11	10:31
2	TOC	0.0000	0.0000	7.29	10.24	2.95	50.10	10:28
3	TOC	0.0000	0.0000	7.15	10.19	3.04	50.12	10:29
4	TOC	0.0000	0.0000	7.46	10.28	2.82	50.12	10:27

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> LCS

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.3222 ppm	0.0626 ppm	0.24%	2019/04/03 13:28

(PASS)										
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.2700	262.7005	187.78	190.71	2.93	50.10	10:27
C	TOC	25.0 ppm	2	26.3167	263.1675	188.10	191.08	2.98	50.09	10:30
C	TOC	25.0 ppm	3	26.4116	264.1162	188.74	191.81	3.07	50.09	10:29
C	TOC	25.0 ppm	4	26.2902	262.9023	187.92	190.77	2.85	50.07	10:24

<u>Completion State</u>	<u>Success Action</u>	<u>Method</u>	<u>Calibration</u>	<u>STD Conc - Pos C</u>
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	25 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
2	TOC	ICS	0.1226 ppm	0.0000 ppm	0.0000%	2019/04/03 14:24

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1226	1.2256	9.59	12.41	2.82	50.04	10:32

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 8.7530 (IC) (v1242)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
3	TOC	LOD	0.0166 ppm	0.0191 ppm	115.1300%	2019/04/03 14:38

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0368	0.3682	9.00	11.80	2.80	50.04	10:27
2	TOC	0.0006	0.0058	8.76	11.68	2.92	50.02	10:29
3	TOC	0.0289	0.2887	8.95	11.77	2.82	50.00	10:26
4	TOC	0.0000	0.0000	8.50	11.56	3.06	49.99	10:30

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 8.7530 (IC) (v1242)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
4	TOC	LOQ	0.3479 ppm	0.0297 ppm	8.5400%	2019/04/03 15:34

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3483	3.4826	11.12	14.03	2.91	49.96	10:28
2	TOC	0.3297	3.2970	10.99	14.12	3.13	49.95	10:28
3	TOC	0.3897	3.8966	11.40	14.30	2.90	49.94	10:28
4	TOC	0.3240	3.2395	10.95	13.90	2.95	49.93	10:28

<u>Dilution</u>	<u>Blank Contribution</u>	<u>Method</u>	<u>Calibration</u>
1:10	(TC) 8.7530 (IC)	CAS_salt_010711	CAS_salt_010711

(v1242)

(v4)

(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
5	TOC	K1902828-003.08	0.8751 ppm	0.0119 ppm	1.3500%	2019/04/03 16:30

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.8758	8.7581	14.70	17.55	2.85	49.93	10:27
2	TOC	0.8720	8.7198	14.67	17.44	2.77	49.91	10:26
3	TOC	0.8905	8.9054	14.80	17.61	2.81	49.90	10:28
4	TOC	0.8620	8.6196	14.60	17.69	3.08	49.90	10:27

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
6	TOC	K1902828-004.08	0.8332 ppm	0.0183 ppm	2.2000%	2019/04/03 17:26

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.8587	8.5872	14.58	17.57	2.99	49.89	10:30
2	TOC	0.8151	8.1512	14.29	17.42	3.14	49.89	10:29
3	TOC	0.8293	8.2926	14.38	17.47	3.09	49.91	10:30
4	TOC	0.8296	8.2955	14.38	17.33	2.95	49.90	10:26

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
7	TOC	K1902828-004.33 ms	27.9836 ppm	0.1749 ppm	0.6200%	2019/04/03 18:22

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	28.1474	281.4741	199.82	202.95	3.13	49.91	10:28
2	TOC	28.1177	281.1765	199.61	202.59	2.97	49.92	10:26
3	TOC	27.8707	278.7074	197.94	201.07	3.13	49.93	10:25
4	TOC	27.7987	277.9870	197.45	200.42	2.97	49.95	10:27

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
8	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/03 19:17

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.01	11.07	3.06	49.95	10:33

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.9003	9.0027	14.86	17.73	2.87	49.97	10:29
2	TOC	0.8471	8.4708	14.50	17.46	2.96	49.99	10:30
3	TOC	0.8365	8.3648	14.43	17.25	2.82	50.00	10:30
4	TOC	0.8248	8.2484	14.35	17.32	2.97	50.00	10:29

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.4415 ppm (PASS)	0.0000 ppm	0%	2019/04/03 20:28

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.4415	254.4152	182.16	185.05	2.89	50.00	10:33

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB

From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/04/03 20:42

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	7.17	10.24	3.07	50.03	10:30

Completion State Success - Criteria met.
Success Action Do Nothing
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)
STD Conc - Pos D 0 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 10	TOC	K1902828-006.08	1.4433 ppm	0.0273 ppm	1.8900%	2019/04/03 20:57

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.4705	14.7054	18.74	21.59	2.85	50.04	10:30
2	TOC	1.4535	14.5345	18.62	21.42	2.81	50.07	10:28
3	TOC	1.4060	14.0602	18.30	21.24	2.95	50.07	10:27
4	TOC	1.4431	14.4314	18.55	21.39	2.84	50.09	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
11	TOC	K1902828-007.08	1.4624 ppm	0.0336 ppm	2.2900%	2019/04/03 21:53

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.5053	15.0531	18.97	21.68	2.71	50.11	10:28
2	TOC	1.4350	14.3504	18.49	21.42	2.92	50.09	10:28
3	TOC	1.4364	14.3636	18.50	21.55	3.05	50.11	10:30
4	TOC	1.4729	14.7290	18.75	21.63	2.88	50.11	10:30

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
12	TOC	K1902828-008.08	0.0017 ppm	0.0033 ppm	200.0000%	2019/04/03 22:49

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0066	0.0662	8.80	11.64	2.84	50.09	10:31
2	TOC	0.0000	0.0000	8.36	11.40	3.04	50.10	10:28
3	TOC	0.0000	0.0000	8.59	11.58	2.98	50.10	10:30
4	TOC	0.0000	0.0000	8.63	11.52	2.89	50.10	10:28

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
13	TOC	K1902829-003.08	0.5158 ppm	0.0085 ppm	1.6500%	2019/04/03 23:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.5161	5.1606	12.26	15.37	3.11	50.07	10:26
2	TOC	0.5162	5.1620	12.26	15.28	3.02	50.07	10:26
3	TOC	0.5050	5.0501	12.18	15.29	3.10	50.07	10:27
4	TOC	0.5258	5.2578	12.32	15.39	3.06	50.06	10:27

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis	Sample ID	Result (ppmC)	Std. Dev.	RSD	Start Time
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Type	(ppmC)
14 TOC K1902829-004.08	0.3633 ppm 0.0157 ppm 4.3300%

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.3727	3.7271	11.28	14.41	3.12	50.07	10:30
2	TOC	0.3459	3.4590	11.10	14.18	3.08	50.07	10:30
3	TOC	0.3799	3.7993	11.33	14.36	3.03	50.06	10:29
4	TOC	0.3547	3.5474	11.16	14.28	3.12	50.05	10:29

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
15	TOC	K1902829-004.33 ms	27.1361 ppm	0.1030 ppm	0.3800%	2019/04/04 01:36

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	27.2163	272.1635	193.50	196.30	2.80	50.05	10:31
2	TOC	27.0690	270.6903	192.50	195.51	3.01	50.05	10:27
3	TOC	27.2314	272.3138	193.60	196.55	2.95	50.03	10:26
4	TOC	27.0275	270.2748	192.21	195.25	3.03	50.06	10:25

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
16	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 02:31

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	7.75	10.84	3.09	50.07	10:30

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
17	TOC	K1902829-005.08	0.2235 ppm	0.0326 ppm	14.5800%	2019/04/04 02:46

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2261	2.2613	10.29	13.32	3.03	50.06	10:29
2	TOC	0.2546	2.5456	10.48	13.37	2.89	50.05	10:29
3	TOC	0.1780	1.7796	9.96	13.10	3.13	50.06	10:29
4	TOC	0.2353	2.3526	10.35	13.48	3.13	50.06	10:29

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
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18	TOC	K1902829-006.08	0.2106 ppm	0.0234 ppm	11.1200%	2019/04/04 03:41			
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Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.2068	2.0683	10.16	13.22	3.06	50.06	10:28
2	TOC	0.2445	2.4455	10.41	13.40	2.99	50.07	10:31
3	TOC	0.1983	1.9829	10.10	13.15	3.05	50.08	10:26
4	TOC	0.1925	1.9254	10.06	13.01	2.95	50.06	10:23

Dilution 1:10
 Blank Contribution (TC) 8.7530 (IC) (v1242)
 Method CAS_salt_010711 (v4)
 Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
19	TOC	K1902829-007.08	8.3598 ppm	0.0640 ppm	0.7700%	2019/04/04 04:37

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	8.3757	83.7573	65.61	68.70	3.10	50.05	10:26
2	TOC	8.2698	82.6981	64.89	67.91	3.03	50.06	10:27
3	TOC	8.4213	84.2125	65.92	68.97	3.06	50.05	10:28
4	TOC	8.3725	83.7249	65.58	68.73	3.15	50.07	10:26

Dilution 1:10
 Blank Contribution (TC) 8.7530 (IC) (v1242)
 Method CAS_salt_010711 (v4)
 Calibration CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.5103 ppm (PASS)	0.0000 ppm	0%	2019/04/04 05:33

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.5103	255.1032	182.62	185.55	2.92	50.09	10:33

Completion State Success - Criteria met.
 Success Action Do Nothing
 Method CAS_salt_010711 (v4)
 Calibration CAS_salt_010711 (v30)
 STD Conc - Pos B 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/04/04 05:47

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time

D	TOC	0 ppm	1	0.0000	0.0000	7.88	10.92	3.04	50.06	10:31
Completion State		Success Action		Method		Calibration		STD Conc - Pos D		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		0 ppmC		

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
20	TOC	MB2	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 06:02

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	7.96	10.94	2.98	50.07	10:28

Dilution		Blank Contribution		Method		Calibration	
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)	

Sample Type: Check Standard --> LCS From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.3894 ppm (PASS)	0.0000 ppm	0%	2019/04/04 06:17

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.3894	263.8937	188.59	191.64	3.05	50.08	10:31

Completion State		Success Action		Method		Calibration		STD Conc - Pos C	
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		25 ppmC	

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
21	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 06:31

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.05	11.14	3.09	50.07	10:27
2	TOC	0.0000	0.0000	7.32	10.51	3.20	50.07	10:28

Dilution		Blank Contribution		Method		Calibration	
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)	

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
22	TOC	LOD	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 06:59

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	7.77	10.93	3.16	50.07	10:29

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
23	TOC	LOQ	0.4151 ppm	0.0000 ppm	0.0000%	2019/04/04 07:14

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.4151	4.1514	11.57	14.72	3.15	50.06	10:34

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
24	TOC	K1902703-001.02	3.0262 ppm	0.0276 ppm	0.9100%	2019/04/04 07:28

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.0457	30.4569	29.43	32.32	2.89	50.06	10:26
2	TOC	3.0066	30.0665	29.16	32.17	3.00	50.09	10:28

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
25	TOC	K1902703-001.02 ms	29.7416 ppm	0.0000 ppm	0.0000%	2019/04/04 07:56

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	29.7416	297.4156	210.64	213.66	3.03	50.05	10:34

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
26	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 08:11

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	8.20	11.27	3.07	50.07	10:28

Dilution 1:10
Blank Contribution (TC) 8.7530 (IC) (v1242)
Method CAS_salt_010711 (v4)
Calibration CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
27	TOC	K1902766-001.01	2.2477 ppm	0.0649 ppm	2.8900%	2019/04/04 08:25

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.2936	22.9362	24.32	27.22	2.90	50.08	10:29
2	TOC	2.2018	22.0184	23.70	26.64	2.94	50.09	10:29

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
28	TOC	K1902853-001.01	4.8178 ppm	0.0573 ppm	1.1900%	2019/04/04 08:53

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	4.8583	48.5832	41.73	44.66	2.93	50.09	10:25
2	TOC	4.7773	47.7729	41.18	44.24	3.05	50.08	10:25

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.2521 ppm (PASS)	0.0000 ppm	0%	2019/04/04 09:21

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.2521	252.5206	180.87	183.96	3.09	50.09	10:33

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30) **STD Conc - Pos B** 50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/04/04 09:36

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	7.73	10.83	3.10	50.08	10:30

Completion State Success - Criteria met. **Success Action** Do Nothing **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30) **STD Conc - Pos D** 0 ppmC

Sample Type: Sample

From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
29	TOC	K1902526-003.01 100x	4.1472 ppm	0.1115 ppm	2.6900%	2019/04/04 09:51

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	4.2260	42.2602	37.44	40.45	3.01	50.09	10:25
2	TOC	4.0684	40.6839	36.37	39.44	3.07	50.08	10:25

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
30	TOC	K1902526-004.01 100x	4.8155 ppm	0.0434 ppm	0.9000%	2019/04/04 10:19

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	4.8462	48.4624	41.65	44.75	3.10	50.07	10:32
2	TOC	4.7848	47.8480	41.23	44.26	3.03	50.06	10:25

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
31	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 10:47

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.0000	0.0000	7.40	10.46	3.06	50.05	10:28
2	TOC	0.0000	0.0000	7.44	10.45	3.01	50.04	10:26

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
32	TOC	K1902660-001.03	1.9294 ppm	0.1023 ppm	5.3000%	2019/04/04 11:16

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.0018	20.0178	22.34	25.41	3.07	50.05	10:29
2	TOC	1.8571	18.5711	21.36	24.29	2.93	50.04	10:25

Dilution

1:10

Blank Contribution(TC) 8.7530 (IC)
(v1242)MethodCAS_salt_010711
(v4)CalibrationCAS_salt_010711
(v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
33	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 11:44

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time

1	TOC	0.0000	0.0000	6.76	9.80	3.04	50.04	10:28
2	TOC	0.0000	0.0000	6.90	9.83	2.93	50.05	10:27

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
34	TOC	K1902660-002.03	1.1509 ppm	0.0306 ppm	2.6600%	2019/04/04 12:12

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	1.1725	11.7251	16.71	19.69	2.98	50.04	10:27
2	TOC	1.1292	11.2920	16.42	19.46	3.04	50.05	10:27

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
35	TOC	K1902660-003.03	0.1343 ppm	0.0257 ppm	19.1600%	2019/04/04 12:40

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	0.1161	1.1608	9.54	12.60	3.06	50.05	10:26
2	TOC	0.1525	1.5247	9.79	12.65	2.86	50.07	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
36	TOC	K1902831-001.01 100x	5.7811 ppm	0.0621 ppm	1.0700%	2019/04/04 13:08

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	5.7372	57.3723	47.70	50.65	2.95	50.05	10:29
2	TOC	5.8250	58.2503	48.29	51.20	2.91	50.06	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
37	TOC	K1902818-001.01	3.2864 ppm	0.0025 ppm	0.0800%	2019/04/04 13:36

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.2882	32.8818	31.07	34.09	3.02	50.03	10:26
2	TOC	3.2846	32.8464	31.05	33.92	2.87	50.03	10:26

Dilution 1:10 **Blank Contribution** (TC) 8.7530 (IC) (v1242) **Method** CAS_salt_010711 (v4) **Calibration** CAS_salt_010711 (v30)

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
38	TOC	K1902818-003.01	2.4624 ppm	0.0071 ppm	0.2900%	2019/04/04 14:04

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.4675	24.6746	25.50	28.37	2.87	50.01	10:30
2	TOC	2.4574	24.5744	25.43	28.32	2.89	50.00	10:26

Dilution	Blank Contribution	Method	Calibration
1:10	(TC) 8.7530 (IC) (v1242)	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.7540 ppm (PASS)	0.0000 ppm	0%	2019/04/04 14:32

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.7540	257.5398	184.28	187.15	2.87	49.97	10:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos B
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	50 ppmC

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/04/04 14:46

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	7.96	10.97	3.01	49.97	10:30

Completion State	Success Action	Method	Calibration	STD Conc - Pos D
Success - Criteria met.	Do Nothing	CAS_salt_010711 (v4)	CAS_salt_010711 (v30)	0 ppmC

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
39	TOC	MB3	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 15:01

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time

1	TOC	0.0000	0.0000	7.38	10.48	3.10	49.96	10:31
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Sample Type: Check Standard --> LCS From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time
◊ C	TOC	25.0000	1:1	[TOC] LCS [24.0 ppm]	0 / infinity (NA / NA)	26.5417 ppm (PASS)	0.0000 ppm	0%	2019/04/04 15:16

Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
C	TOC	25.0 ppm	1	26.5417	265.4170	189.63	192.72	3.09	49.94	10:31

<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>		<u>Calibration</u>		<u>STD Conc - Pos C</u>	
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		25 ppmC	

Sample Type: Sample From Schedule Version 4

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 40	TOC	K1902818-002.02 doc	3.1511 ppm	0.0641 ppm	2.0300%	2019/04/04 15:31

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	3.1964	31.9640	30.45	33.46	3.01	49.94	10:28
2	TOC	3.1058	31.0580	29.84	32.78	2.95	49.93	10:29

<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 41	TOC	K1902818-002.02 ms doc	30.5720 ppm	0.0000 ppm	0.0000%	2019/04/04 15:59

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	30.5720	305.7201	216.27	219.15	2.87	49.92	10:32

<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time
◊ 42	TOC	RB	0.0000 ppm	0.0000 ppm	0.0000%	2019/04/04 16:13

Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time

1	TOC	0.0000	0.0000	8.40	11.41	3.01	49.93	10:35
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		
Pos	Analysis Type	Sample ID	Result (ppmC)	Std. Dev. (ppmC)	RSD	Start Time		
43	TOC	K1902818-004.02 doc	2.3184 ppm	0.0584 ppm	2.5200%	2019/04/04 16:28		
Rep #	Base Analysis Type	ppm	µg	Adjusted (Abs)	NDIR (Abs)	Baseline (Abs)	Pressure (psig)	Run Time
1	TOC	2.3598	23.5977	24.77	27.75	2.98	49.93	10:30
2	TOC	2.2771	22.7712	24.21	27.20	2.99	49.93	10:31
<u>Dilution</u>		<u>Blank Contribution</u>		<u>Method</u>		<u>Calibration</u>		
1:10		(TC) 8.7530 (IC) (v1242)		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		

Sample Type: Check Standard --> CCV 25 ppm From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
B	TOC	25.0000	1:2	[TOC] CCV 25 ppm [25 ppm]	0 / infinity (NA / NA)	25.7046 ppm (PASS)	0.0000 ppm	0%	2019/04/04 16:56	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
B	TOC	25 ppm	1	25.7046	257.0463	183.94	186.96	3.02	49.91	10:31
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>		<u>Calibration</u>		<u>STD Conc - Pos B</u>		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		50 ppmC		

Sample Type: Check Standard --> CCB From Schedule Version 4

Pos	BAT	Concentration (ppm)	Dil	Sample ID	Min / Max (% dev)	Result	Std. Dev.	RSD	Start Time	
D	TOC	0.0000	1:1	[TOC] CCB [0 ppm]	0 / infinity (NA / NA)	0.0000 ppm (PASS)	0.0000 ppm	0%	2019/04/04 17:11	
Pos	Base Analysis Type	ID	Rep #	ppm	µg	Adjusted	NDIR	Baseline	Pressure	Run Time
D	TOC	0 ppm	1	0.0000	0.0000	7.93	10.80	2.87	49.91	10:31
<u>Completion State</u>		<u>Success Action</u>		<u>Method</u>		<u>Calibration</u>		<u>STD Conc - Pos D</u>		
Success - Criteria met.		Do Nothing		CAS_salt_010711 (v4)		CAS_salt_010711 (v30)		0 ppmC		

Meta Data Used in this Report

Blanks

Version	Reagent (Abs)	Acid (Abs)	DI IC (Abs)	DI TC (Abs)	DI TOC (Abs)	Save Time	Operator
v1241	1.6887	1.2060	0.0000	0.0000	0.0000	2019/04/01 15:29	Fusion1 (Fusion1)
v1242	1.1840	1.0590	0.0000	0.0000	0.0000	2019/04/03 11:48	Fusion1 (Fusion1)

Calibrations**Name: CAS_salt_010711 (TOC)**

Version: v30 Calibration curve formula: TOC: $y = 6.788x + 9.463$
 Ver Creation: 2019/03/05 17:42 r^2 value: TOC: $r^2 = 0.99963$
 Comment:
 Operator: Fusion1 (Fusion1)
 Basic Analysis Type: TOC

Basic Analysis Type: TOC

Sample ID	Y Raw Value	X Expected	Message	End Time
DI Water	7.8970	0.0000		2019/03/05 16:15
0.500 ppm	11.5280	0.5000		2019/03/05 16:29
1.0 ppm	14.9760	1.0000		2019/03/05 16:44
5.0 ppm	43.6500	5.0000		2019/03/05 16:58
10 ppm	79.6020	10.0000		2019/03/05 17:12
25 ppm	183.3580	25.0000		2019/03/05 17:26
50 ppm	346.3230	50.0000		2019/03/05 17:40

Methods**Name: CAS_salt_010711 (TOC)**

Version: v4 Operator: Fusion1 (Fusion1)
 Ver Creation: 2019/02/21 17:57
 Comment:

Parameter	Value	Advanced Parameter	Value
SampleVolume	10.0 mL	NeedleRinseVolume	5.0 ml
Dilution	1:10	VialPrimeVolume	2.0 ml
AcidVolume	0.5 ml	ICSamplePrimeVolume	2.0 ml
ReagentVolume	2.0 ml	ICSpurgeRinseVolume	12.0 ml
UVReactorPrerinse	Off	BaselineStabilizeTime	0.70 min
UVReactorPrerinseVolume	5.0	DetectorPressureFlow	150 ml/min
NumberOfUVReactorPrerinse	1	SyringeSpeedWaste	10
ICSpurgeTime	1.00 mins	SyringeSpeedAcid	7
DetectorSweepFlow	500 ml/min	SyringeSpeedReagent	7

PreSpargeTime	2.00 mins	SyringeSpeedDIWater	7
SystemFlow	500 ml/min	NDIRPressurization	60 psig
		SyringeSpeedSampleDispense	5
		SyringeSpeedSampleAspirate	4
		SyringeSpeedUVDispense	5
		SyringeSpeedUVAspirate	5
		SyringeSpeedICDispense	5
		SyringeSpeedICAspirate	5
		NDIRPressureStabilize	1.75 min
		SampleMixing	Off
		SampleMixingCycles	1
		SampleMixingVolume	10.0
		LowLevelFilterNDIR	Off

Acceptance / Approval

Electronic Signatures

Report Version	User Name	Acceptance	Reason	Date
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Report History

Report History

Report Version	User Name	System Reason	User Reason	Date
1	Fusion1 (Fusion1)	Schedule completed	Schedule completed	2019/04/04 17:28

ALS Environmental

StarLIMS Run: 630888, 630890, 630891
 Analysis: TOC/DOC
 Method: 9060, 415.1, SM 5310 C, 9060A

CCV: 11-GEN-05-77C 50 ppm LCS: 11-GEN-05-77D 25.0 ppm

ICAL Date: 3/6/19

ICAL ID: 11-GEN-05-76H

ICS ID: 11-GEN-05-74A

ICS TV: 25.0 ppm

ICS % R = 2

Spike ID: 11-GEN-05-700 0.05 ml of 5000 ppm stock ---> 10.0 ml = 25.0 ppm x dilution factor

Sodium Persulfate: 11-GEN-05-77E

21 % H3PO4: 11-GEN-05-77F

Equipment ID: K-TOC-03

PIPETTE ID: 124276B, 129001F, N11314F, Marge

FILTER ID: NA

CBA: 0.06875 ⁰

Analyzed By: <i>BCD</i>	Date Analyzed: <i>4/3/19</i>
Reviewed By: <i>Fauquier</i>	Date Reviewed: <i>04/05/19</i>

0 BCD 4/5/19



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1909152; 1909153; 1909154;
1909947; 1909949

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2233 (236356)

General Set Information: There were ten field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Method QC data: The method blank (LMB 647198) was less than 1/2 the CRDL. The recovery for the LCS (647199) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1909152001 (Client ID's: LH18/24-SP650_032719_BIX). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.µg/L. The MS/MSD – 647200/01 failed QC acceptance criteria for percent recoveries, biased high. This is due to the fact that the unspiked sample result of 2.574µg/L was not subtracted from the MS/MSD results. The Matrix Spike and Matrix Spike duplicate is reported for the clients' information only. The sample matrix may be inappropriate for the method selected. The MS/MSD relative percent difference (RPD) was within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in µg/L. Results were calculated in µg/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (µg/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 647196) is reported from the analysis of the Laboratory Control Sample (LCS – 647199) at a level of 4.0µg/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch April 11, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: April 11, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1909152**

Project ID: HS19031492

Purchase Order: HS19031492

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP650_032719_BIX	1909152001	03/27/19	03/30/19	



ANALYTICAL REPORT

Workorder: 34-1909152

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP650_032719_BIX	Sampling Site: NA	Collected: 03/27/2019				
Lab ID: 1909152001	Media: 125 mL Nalgene	Received: 03/30/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2233 (HBN: 236356) Analyzed: 04/10/2019 10:55	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	2.6	1.0	2.0	4.0	1	J

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 236356)

Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 04/11/2019 09:27	/S/ Stephen Brose 04/11/2019 15:00

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1909152

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935712

Analysis Information

Workorder: 1909152

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2233 (HBN: 236356)
Analyzed By: Thomas Bosch

Blank

LMB: 647198 Analyzed: 04/10/2019 10:42 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 647199 Analyzed: 04/10/2019 10:15 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	3.68	4.00	91.9	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1909152001 Analyzed: 04/10/2019 10:55 Dilution: 1 Units: ug/L		MS: 647200 Analyzed: 04/10/2019 11:08 Dilution: 1 Units: ug/L				MSD: 647201 Analyzed: 04/10/2019 11:22 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	2.60	6.18	4	# 154	78.8 123.8	6.36	# 159	2.92	0.0 20.0

Comments

Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 04/11/2019 13:48	/S/ Stephen Brose 04/11/2019 15:00

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



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Subcontract Chain of Custody

COC ID: 11012

1909152

18698/#2

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19031492
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19031492-02	LH18/24-SP650_032719_BIX	Water	27 Mar 2019 14:00
SUB_Perch-6850			05 Apr 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. M. M. M.
Received By: Meredith M. M.
Cooler ID(s): 927

Date/Time: 3/29/19 18:00
Date/Time: 3/29/19 8:57
Temperature(s): 2

CHAIN OF CUSTODY SOLUTIONS' RIGHT PARTNER



**ALS Environmental
CHAIN-OF-CUSTODY**

Project / Job / Task:		Split:		Workorder ID: 1909152		Level: ENV_LVL4		Requested Analysis:	
Client: ALS Environmental (Houston)		Account: 8101				Type: 129Poly			
Comments:						Preservatives:			
						Containers:			
						ID(s)			
						Count			
Item	Collect Date/Time	Sample ID	Lab ID	QC	Matrix				
1	03/27/2019 14:00	LH18/24-SP650_032719_BIX	1909152001		Water	A	1	A	
2									
3									
4									
5									
6									
7									
8									
9									
10									

PA 8850.D.D GSM

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY				SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY			
Relinquished By: (Signature)	Date / Time	Received By: (Signature)	Reason for Transfer / Storage Location	Sample Prep / Analysis for: Prepared / Analyzed by:	Lab Notebook No.:	Received By: (Signature)	Reason for Transfer / Storage Location
Warath, Julie	03/30/2019 08:58	ALS Sample Receiving	Sample Login				
<i>Julie Warath</i>	4/19 1330	<i>CRB</i>	<i>storage</i>				
<i>R.33.1</i>	4/8/19 10:00	<i>T. Bush</i>	<i>6850</i>				



Must Deliver Next Business Day
Time and Temperature Sensitive!

Part # 159489-434 RIT2 EXP 11/19

ORIGIN ID:SGRA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77089
UNITED STATES US

SHIP DATE: 29MAR19
ACTWGT: 9.25 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

TO **SAMPLE RECEIVING
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE**

SALT LAKE CITY UT 84123

(801) 266-7700

REF: HS19031492/1508/1511 RJ



**FedEx
Express**



JT8111806050104

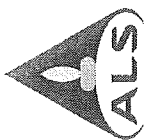
TRK# 4809 7832 2367
0201

**SATURDAY 12:00P
PRIORITY OVERNIGHT**

XO BTFA

**84123
UT-US SLC**





Batch Worklist

Batch: ELMS/ 2233 **Created:** 4/9/2019 08:42 **Instrument:** LCMS04 **HBN:** 236356
Rule: EPA 6850, DoD QSM Water **Analyst:** T. Bosch **Status:** WP



- Workorder:** 1909152 [ENV_LVL4]
- Workorder:** 1909153 [ENV_LVL4]
- Workorder:** 1909154 [ENV_LVL4]
- Workorder:** 1909947 [ENV_LVL4]
- Workorder:** 1909949 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	647195	CCV for HBN 236356 [ELMS/2233]				CCV	3		E685041C3Q	5311		4/11/2019	4/10/2019
2	647199	LCS for HBN 236356 [ELMS/2233]				LCS	3		E6850Q413Q	5311		4/11/2019	4/10/2019
3	647197	ICS for HBN 236356 [ELMS/2233]				ICS	3		E6850.D3Q	5311		4/11/2019	4/10/2019
4	647198	LMB for HBN 236356 [ELMS/2233]				LMB	3		E6850Q413Q	5311		4/11/2019	4/10/2019
5	1909152001	LH18/24-SP650_032719_BIX				SAMPLE	3	1909152001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
6	647200	LH18/24-SP650...(1909152001MS)				MS	3		E6850Q413Q	5311		4/11/2019	4/10/2019
7	647201	LH18/24-SP65...(1909152001MSD)				MSD	3		E6850Q413Q	5311		4/11/2019	4/10/2019
8	1909153001	LH18-24-SP140_032719				SAMPLE	3	1909153001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
9	1909154001	LH18/24-SP650_032719_BIX				SAMPLE	3	1909154001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
10	1909947001	LH18/25-SP650_040419_BIX				SAMPLE	3	1909947001-A	E6850Q41.3	5480	5/2/2019	4/18/2019	4/10/2019
11	1909949001	HBW7_040119				SAMPLE	3	1909949001-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
12	1909949002	HBW7_040119-a				SAMPLE	3	1909949002-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
13	1909949003	HBW10_040119				SAMPLE	3	1909949003-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
14	1909949004	HBW1_040119				SAMPLE	3	1909949004-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
15	1909949005	GPW1_040119				SAMPLE	3	1909949005-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
16	1909949006	GPW3_040119				SAMPLE	3	1909949006-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
17	647196	RLVS for HBN 236356 [ELMS/2233]				RLVS	3		E685041C3Q	5311		4/11/2019	4/10/2019
18	647202	CCV for HBN 236356 [ELMS/2233]				CCV	3		E685041C3Q	5311		4/11/2019	4/10/2019



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1909152 (001); 1909153 (001); 1909154 (001); 1909947 (001); 1909949 (001-06) ELMS Batch/HBN ID: 2233 (236356)
 Prep Date: 04/09/2019 Analysis Date: 04/10/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\28MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by **TNB**. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4) / 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 647199; Target = 4.0µg/L. ASTM type II water was used for LMB 647198.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1909152001 (Client ID's: LH18/24-SP650_032719_BIX). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- Results reported in µg/L. Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.
- All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The MS/MSD – 647200/01 failed QC acceptance criteria for percent recoveries, biased high. This is due to the fact that the unspiked sample result of 2.574µg/L was not subtracted from the MS/MSD results. The Matrix Spike and Matrix Spike duplicate is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\APR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- Notebook: \\slstws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\236356-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATA\REVIEW\HBN#
- The Reporting Limit Verification Standard (RLVS – 647196) is reported from the analysis of the Laboratory Control Sample (LCS – 647199) at a level 4.0µg/L.
- Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
<u>Batch(es)/SDG: E LMS: 2233 HBN: 236356</u> 1209947 / 1209949		
<u>Sample Set IDs if Applicable: 1909152/1909153/1909154</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK		Description - 6850 WKG Std 100.ug/L			
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK		Description - 6850 Stock AccStd 1,000ug/mL	
Standard: 43659	Created By: Thomas Bosch	Amount: 100 mL	
MFG: AccuStandard	Create Date: 09/17/2018 09:09AM	Expires: 07/25/2020	
MFG Lot: 218065075		Usable: No	
Part ID: IC-PER-10X-1		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT		Description - 6850 Intermdt AccStd 10.ug/mL			
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK		Description: 6850 QC WKG STD 100ug/L			
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided		Lab Lot: CLO4 QC WRK 100.ug/L			
Pos	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte:	Name:	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK		Description: 6850 QC Stock STD 1,000ug/mL	
Standard: 36748	Created By: Thomas Bosch	Amount: 100 mL	
MFG: Ultra Scientific	Create Date: 05/11/2017 01:05PM	Expires: 03/31/2020	
MFG Lot: CP-0860		Usable: Yes	
Part ID: ICC-013		Lab Lot: CLO4 QC STOCK	
Pos	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:

This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES
	perchlorate					

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis



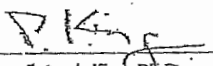
ISO Guide 34 Reference Material

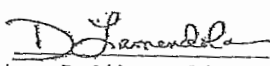
Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:

The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QAVRA



125 Market Street
New Haven, CT 06513
USA



Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is 10.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Melgan O'Leary

Melgan O'Leary, Inorganic QC Manager



Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:
 ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
 (Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$
 Labeled CAS Number: NA
 Unlabeled CAS Number: 7601-89-0
 MW*: 130.4
 Chemical Formula: NaCl*O4
 Storage: Store at room temperature away from light and moisture.
 Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 µg/mL
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 ± 2.8 µg/mL (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	647195	CCV@25	Vial 71	1	Control	1	1.06336e6	21.31933
#*	647199	QC@4.0	Vial 72	1	Control	2	1.97600e5	3.67714
#*	647197	ICS@4.0	Vial 73	1	Control	3	1.49399e5	3.21517
#*	647198	LMB	Vial 74	1	Control	4	0.00000	0.00000
#*	1909152001		Vial 75	1	Sample	5	9.15156e4	2.57433
#*	647200	91521MS	Vial 76	1	Sample	6	2.25049e5	6.17769
#*	647201	91521SD	Vial 77	1	Sample	7	2.49579e5	6.36091
#*	1909153001	1K	Vial 78	1	Sample	8	3.70174e5	6532.86203
#*	1909154001		Vial 79	1	Sample	9	7.79123e4	2.34023
#*	1909947001		Vial 80	1	Sample	10	0.00000	0.00000
#*	1909949001		Vial 81	1	Sample	11	0.00000	0.00000
#*	1909949002		Vial 82	1	Sample	12	0.00000	0.00000
#*	1909949003		Vial 83	1	Sample	13	0.00000	0.00000
#*	1909949004		Vial 84	1	Sample	14	0.00000	0.00000
#*	1909949005		Vial 85	1	Sample	15	0.00000	0.00000
#*	1909949006		Vial 86	1	Sample	16	0.00000	0.00000
*	647202	CCV@25	Vial 71	1	Control	17	1.08007e6	22.59518

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	647195	CCV@25	Vial 71	1	Control	1	3.43635e5	23.09461
#*	647199	QC@4.0	Vial 72	1	Control	2	7.01681e4	4.22460
#*	647197	ICS@4.0	Vial 73	1	Control	3	5.45965e4	3.77625
#*	647198	LMB	Vial 74	1	Control	4	0.00000	0.00000
#*	1909152001		Vial 75	1	Sample	5	3.18275e4	2.84107
#*	647200	91521MS	Vial 76	1	Sample	6	7.61114e4	6.88356
#*	647201	91521SD	Vial 77	1	Sample	7	8.42686e4	7.08137
#*	1909153001	1K	Vial 78	1	Sample	8	1.21054e5	7051.77497
#*	1909154001		Vial 79	1	Sample	9	2.76117e4	2.61418
#*	1909947001		Vial 80	1	Sample	10	0.00000	0.00000
#*	1909949001		Vial 81	1	Sample	11	0.00000	0.00000
#*	1909949002		Vial 82	1	Sample	12	0.00000	0.00000
#*	1909949003		Vial 83	1	Sample	13	0.00000	0.00000
#*	1909949004		Vial 84	1	Sample	14	0.00000	0.00000
#*	1909949005		Vial 85	1	Sample	15	0.00000	0.00000
#*	1909949006		Vial 86	1	Sample	16	0.00000	0.00000
*	647202	CCV@25	Vial 71	1	Control	17	3.36319e5	23.63294

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	647195	CCV@25	Vial 71	1	Control	1	1.53120e5	5.00000
#*	647199	QC@4.0	Vial 72	1	Control	2	1.77838e5	5.00000
#*	647197	ICS@4.0	Vial 73	1	Control	3	1.54931e5	5.00000
#*	647198	LMB	Vial 74	1	Control	4	1.76915e5	5.00000
#*	1909152001		Vial 75	1	Sample	5	1.20240e5	5.00000
#*	647200	91521MS	Vial 76	1	Sample	6	1.17757e5	5.00000
#*	647201	91521SD	Vial 77	1	Sample	7	1.26683e5	5.00000
#*	1909153001	1K	Vial 78	1	Sample	8	1.82758e5	5000.00000
#*	1909154001		Vial 79	1	Sample	9	1.13404e5	5.00000
#*	1909947001		Vial 80	1	Sample	10	1.15130e5	5.00000
#*	1909949001		Vial 81	1	Sample	11	1.14791e5	5.00000
#*	1909949002		Vial 82	1	Sample	12	1.24046e5	5.00000
#*	1909949003		Vial 83	1	Sample	13	1.23373e5	5.00000
#*	1909949004		Vial 84	1	Sample	14	1.20241e5	5.00000
#*	1909949005		Vial 85	1	Sample	15	1.27767e5	5.00000
#*	1909949006		Vial 86	1	Sample	16	1.23965e5	5.00000
*	647202	CCV@25	Vial 71	1	Control	17	1.46281e5	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	647195	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	647199	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	647197	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	647198	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1909152001		CLO4-AQN	1	Sample	
6	Vial 76	647200	91521MS	CLO4-AQN	1	Sample	
7	Vial 77	647201	91521SD	CLO4-AQN	1	Sample	
8	Vial 78	1909153001	1K	CLO4-AQN	1	Sample	
9	Vial 79	1909154001		CLO4-AQN	1	Sample	
10	Vial 80	1909947001		CLO4-AQN	1	Sample	
11	Vial 81	1909949001		CLO4-AQN	1	Sample	
12	Vial 82	1909949002		CLO4-AQN	1	Sample	
13	Vial 83	1909949003		CLO4-AQN	1	Sample	
14	Vial 84	1909949004		CLO4-AQN	1	Sample	
15	Vial 85	1909949005		CLO4-AQN	1	Sample	
16	Vial 86	1909949006		CLO4-AQN	1	Sample	
17	Vial 71	647202	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD01.D

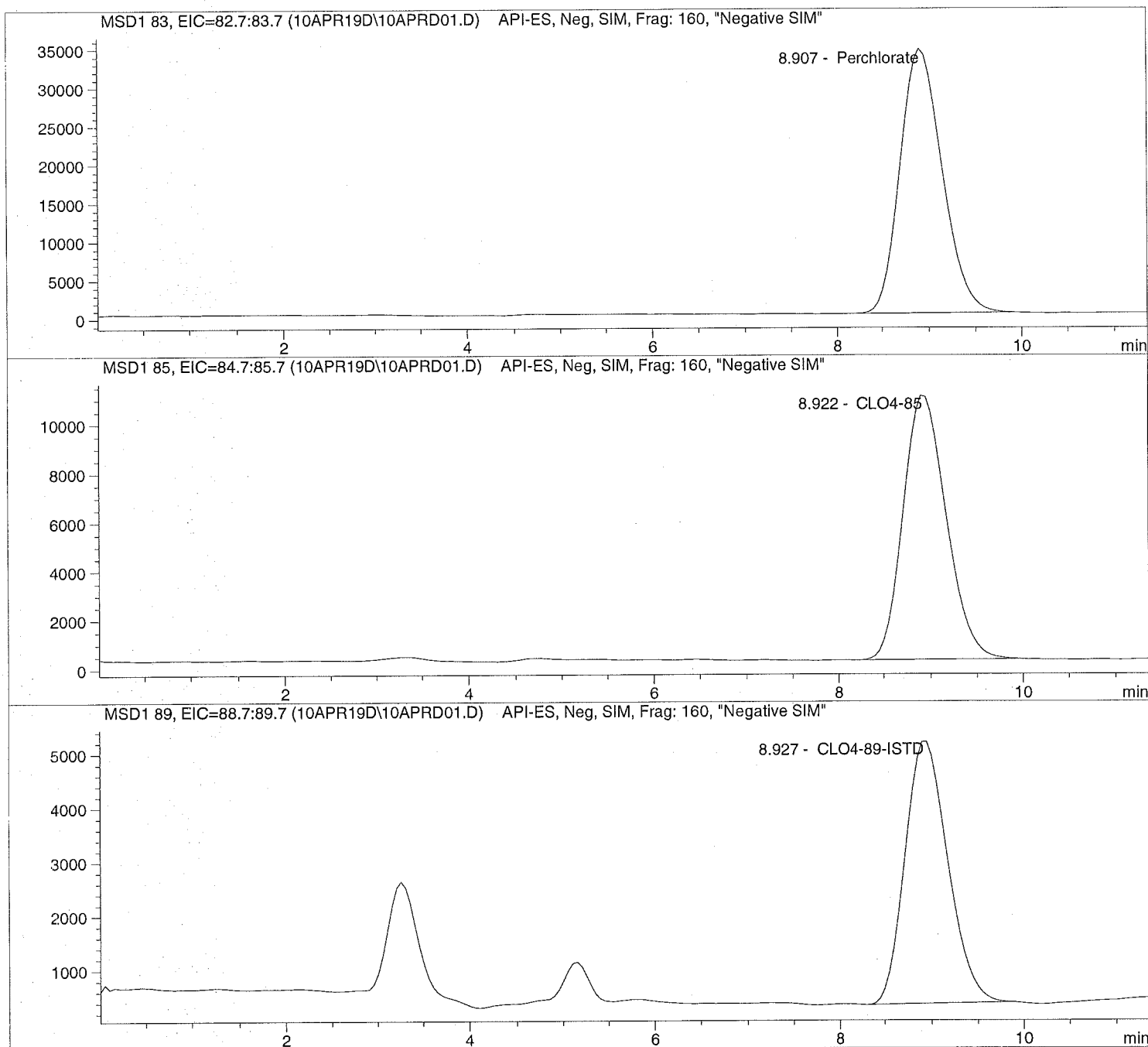
Sample Name: 647195 CCV@25

Injection Date: 4/10/2019 10:01:38
Sample Name: 647195 CCV@25
Acq Operator: TNB

Seq Line: 1
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD01.D Sample Name: 647195 CCV@25

```

=====
Injection Date: 4/10/2019 10:01:38      Seq Line: 1
Sample Name:    647195 CCV@25           Location:  Vial 71
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.907	PBA	1063358.2	21.3193	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.922	PBA	343635.4	23.0946	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.927	PBA	153120.0	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD02.D

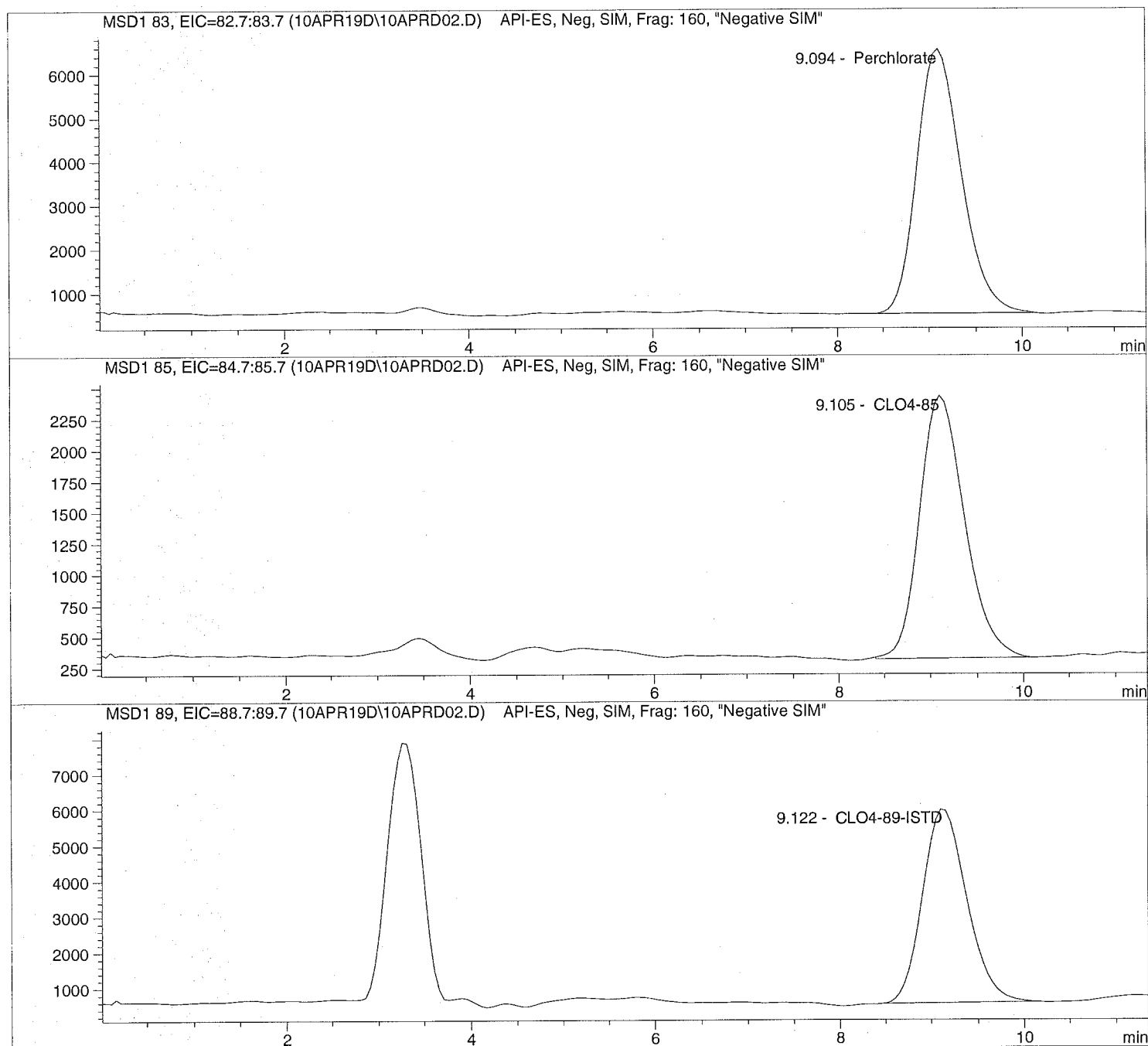
Sample Name: 647199 QC@4.0

=====
Injection Date: 4/10/2019 10:15:40
Sample Name: 647199 QC@4.0
Acq Operator: TNB

Seq Line: 2
Location: Vial 72
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD03.D

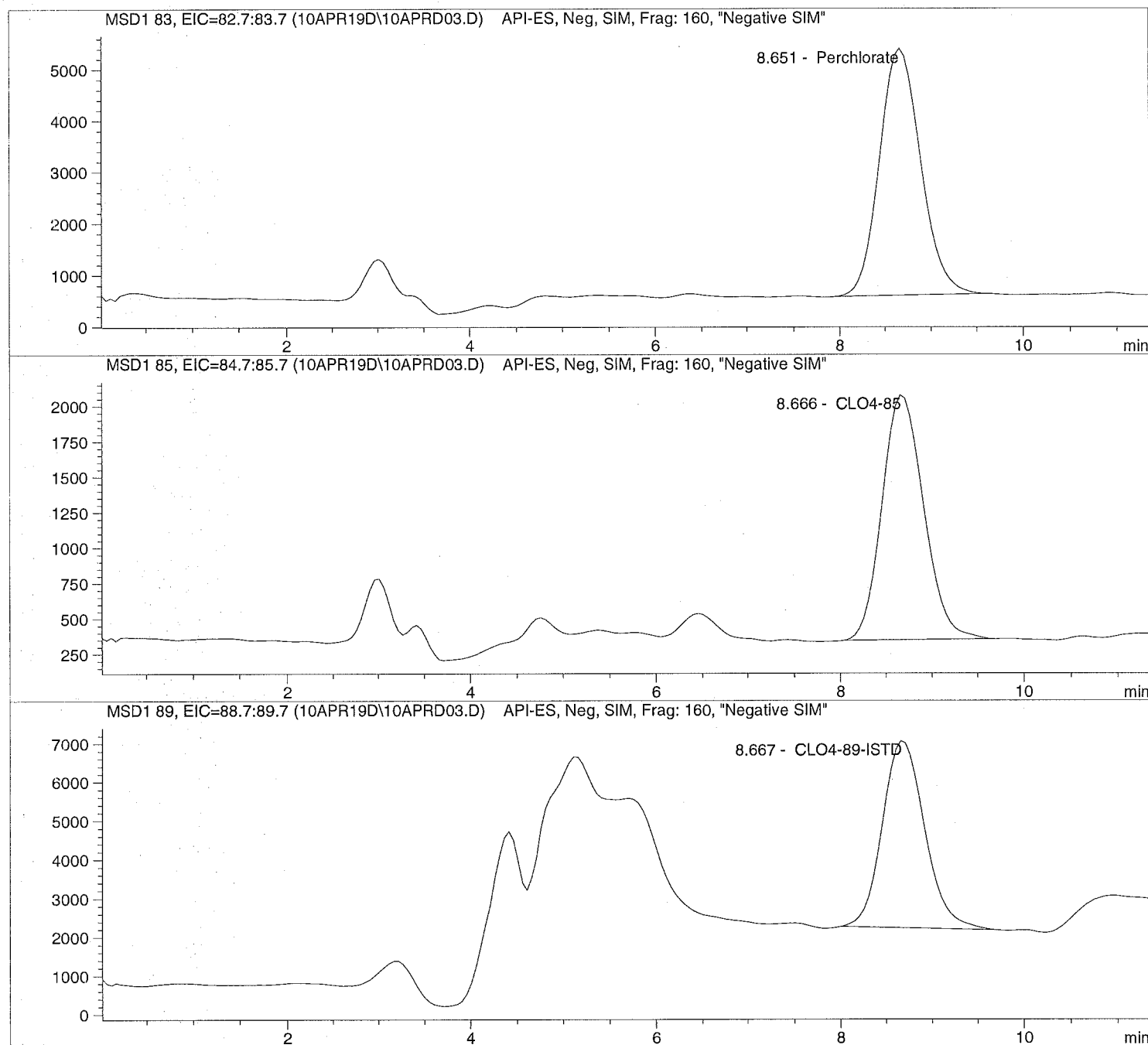
Sample Name: 647197 ICS@4.0

Injection Date: 4/10/2019 10:28:57
Sample Name: 647197 ICS@4.0
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD03.D Sample Name: 647197 ICS@4.0

```

=====
Injection Date: 4/10/2019 10:28:57      Seq Line: 3
Sample Name: 647197 ICS@4.0           Location: Vial 73
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 4.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.651	PBA	149398.5	3.2152	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.666	BBA	54596.5	3.7763	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.667	PBA	154930.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD04.D

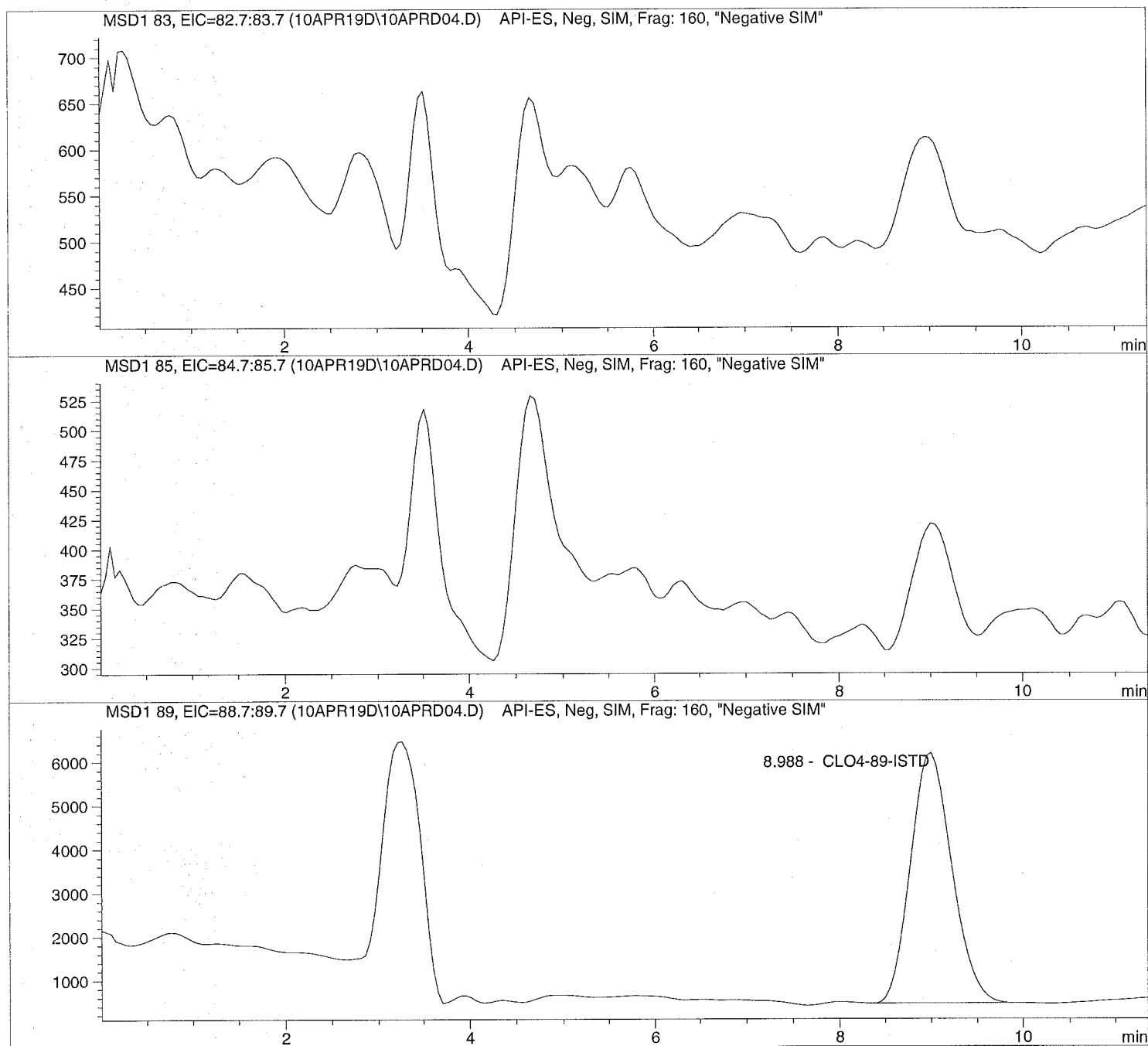
Sample Name: 647198 LMB

Injection Date: 4/10/2019 10:42:16
Sample Name: 647198 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD04.D Sample Name: 647198 LMB

```

=====
Injection Date: 4/10/2019 10:42:16 Seq Line: 4
Sample Name: 647198 LMB Location: Vial 74
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl
  
```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
  
```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
  
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.988	PBA	176915.0	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD05.D

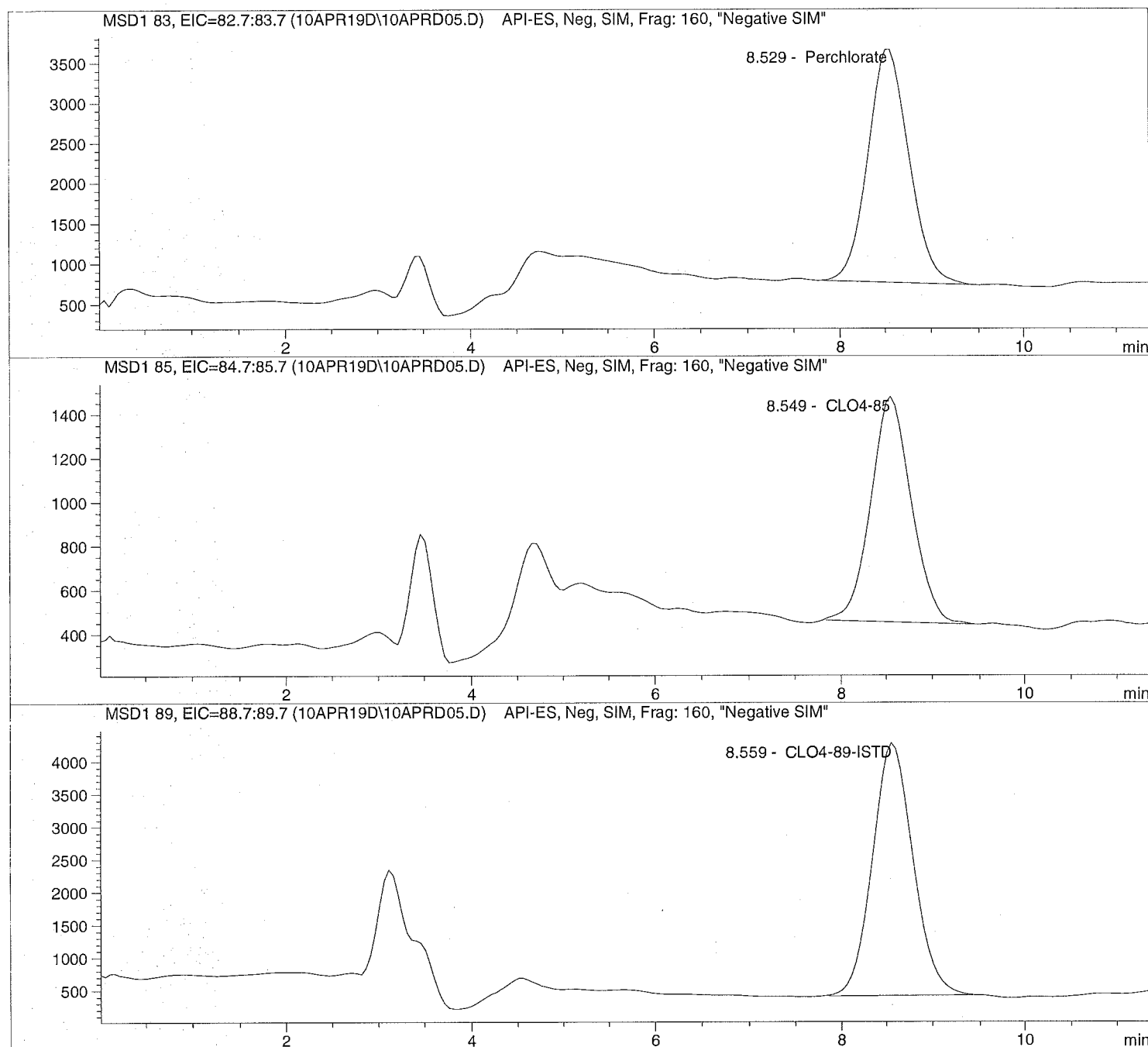
Sample Name: 1909152001

=====
Injection Date: 4/10/2019 10:55:33
Sample Name: 1909152001
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD05.D Sample Name: 1909152001

```

=====
Injection Date: 4/10/2019 10:55:33      Seq Line: 5
Sample Name: 1909152001                Location: Vial 75
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.529	BBA	91515.6	2.5743	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.549	BBA	31827.5	2.8411	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.559	BBA	120239.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD06.D

Sample Name: 647200 91521MS

Injection Date: 4/10/2019 11:08:49

Seq Line: 6

Sample Name: 647200 91521MS

Location: Vial 76

Acq Operator: TNB

Inj. No.: 1

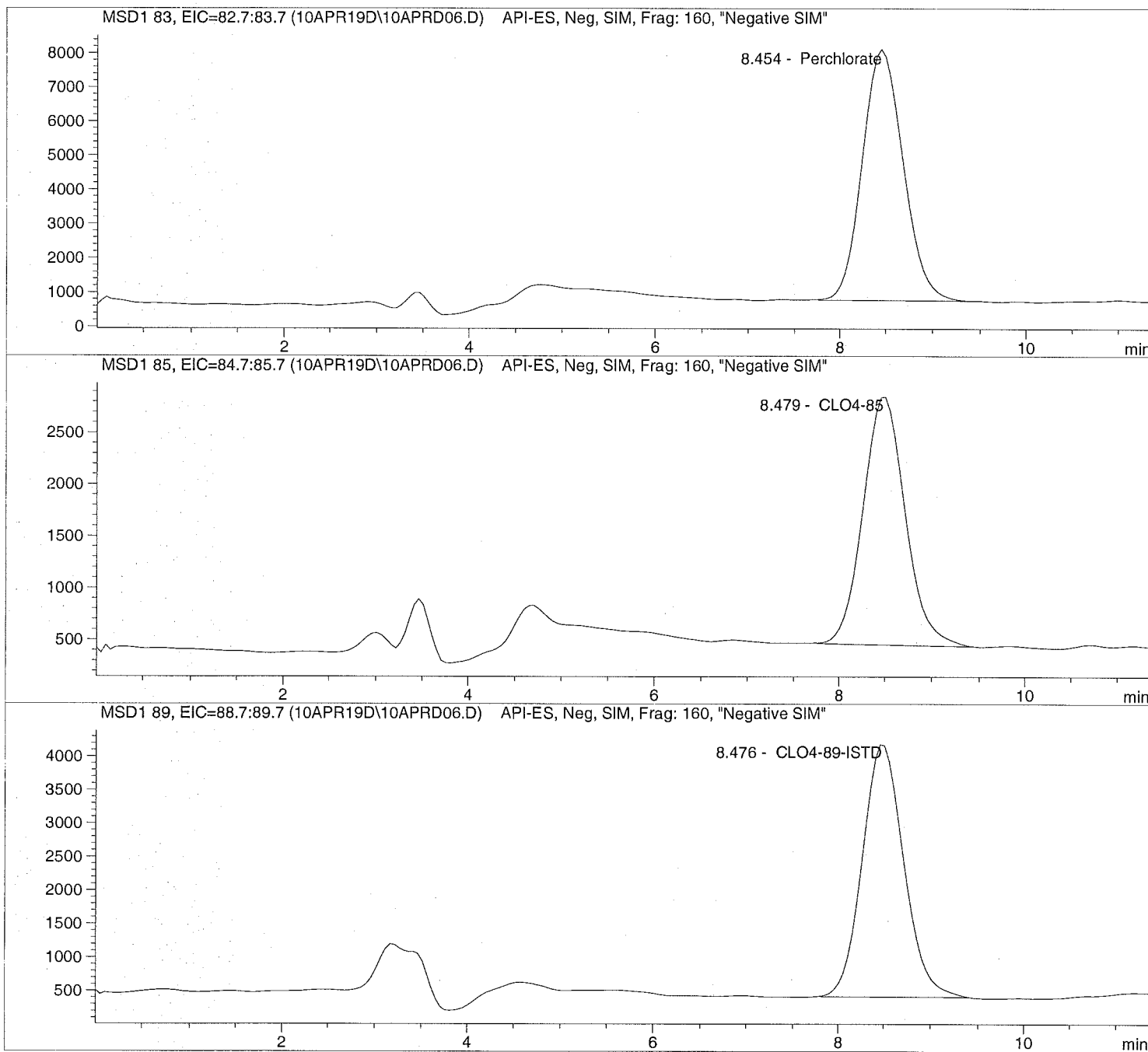
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD06.D Sample Name: 647200 91521MS

```

=====
Injection Date: 4/10/2019 11:08:49 Seq Line: 6
Sample Name: 647200 91521MS Location: Vial 76
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.454	BBA	225049.5	6.1777	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.479	BBA	76111.4	6.8836	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.476	BBA	117756.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

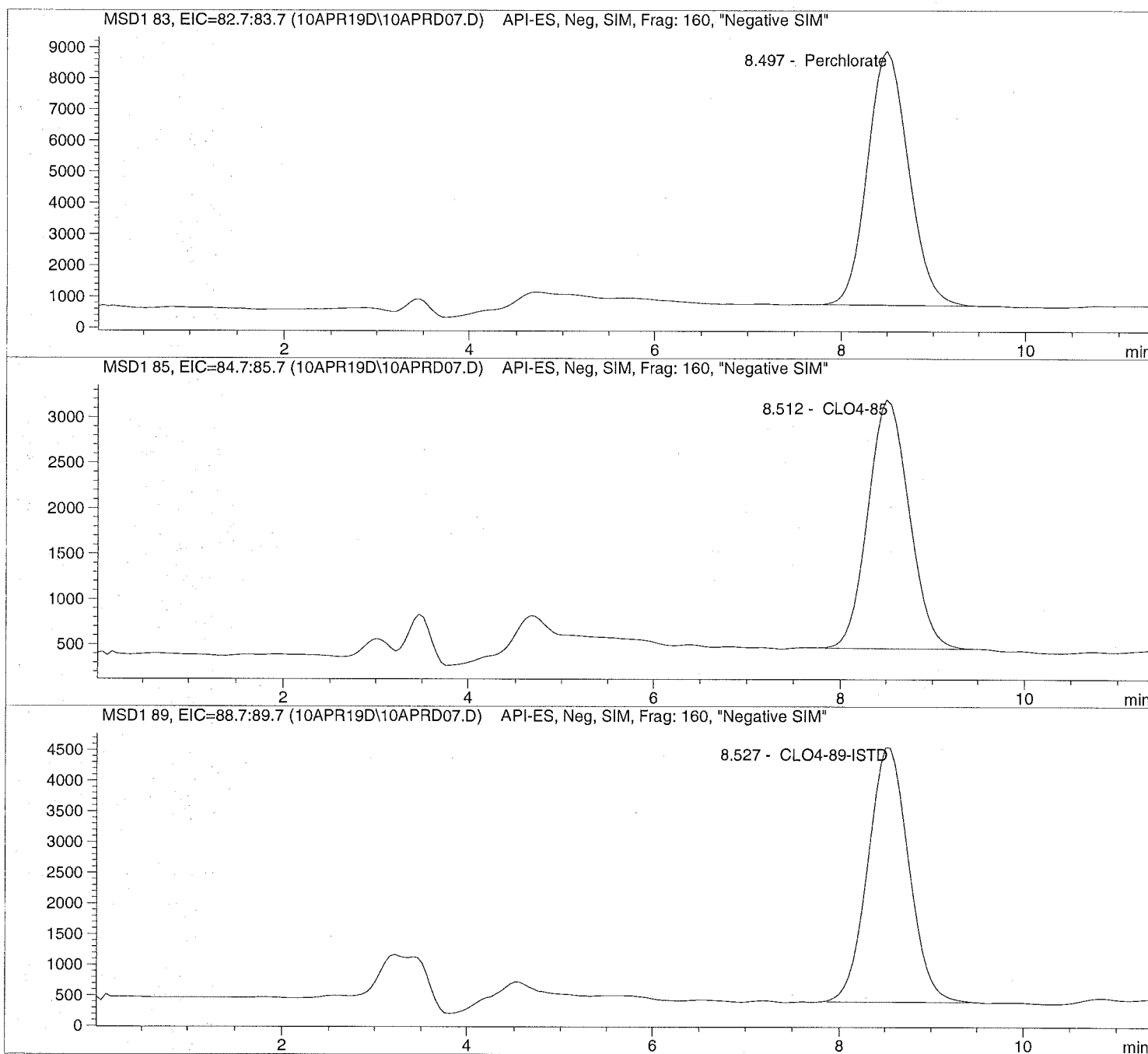
```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD07.D Sample Name: 647201 91521SD

=====
Injection Date: 4/10/2019 11:22:09 Seq Line: 7
Sample Name: 647201 91521SD Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD07.D Sample Name: 647201 91521SD

```

=====
Injection Date: 4/10/2019 11:22:09      Seq Line: 7
Sample Name: 647201 91521SD      Location: Vial 77
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.497	BBA	249579.0	6.3609	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.512	BBA	84268.6	7.0814	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.527	BBA	126683.1	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD08.D

Sample Name: 1909153001 1K

Injection Date: 4/10/2019 11:35:27

Seq Line: 8

Sample Name: 1909153001 1K

Location: Vial 78

Acq Operator: TNB

Inj. No.: 1

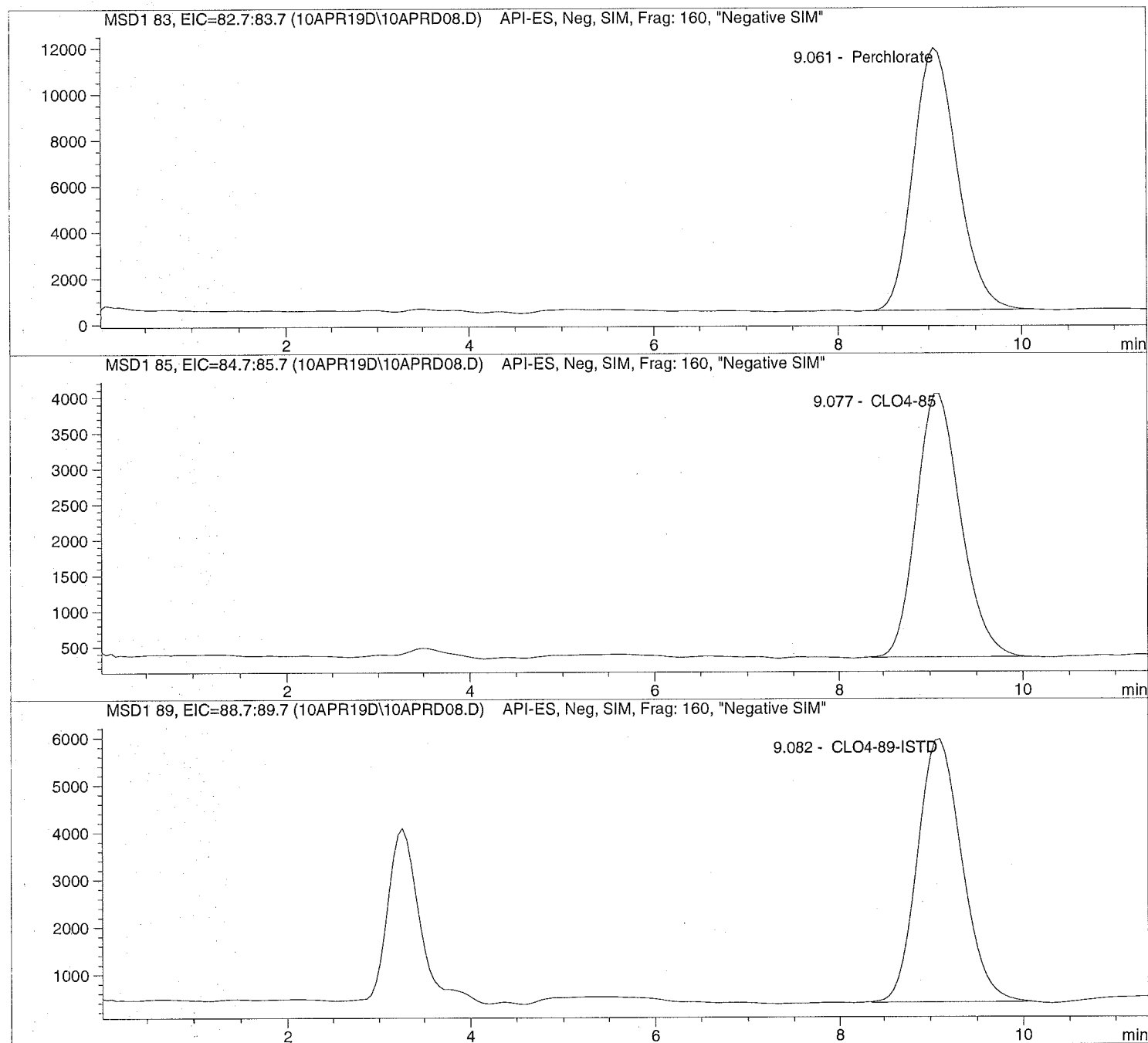
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD08.D Sample Name: 1909153001 1K

```

=====
Injection Date: 4/10/2019 11:35:27      Seq Line:      8
Sample Name:    1909153001 1K           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1000.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.061	PBA	370174.3	6532.8620	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.077	BBA	121053.5	7051.7750	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.082	BBA	182757.9	5000.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD09.D

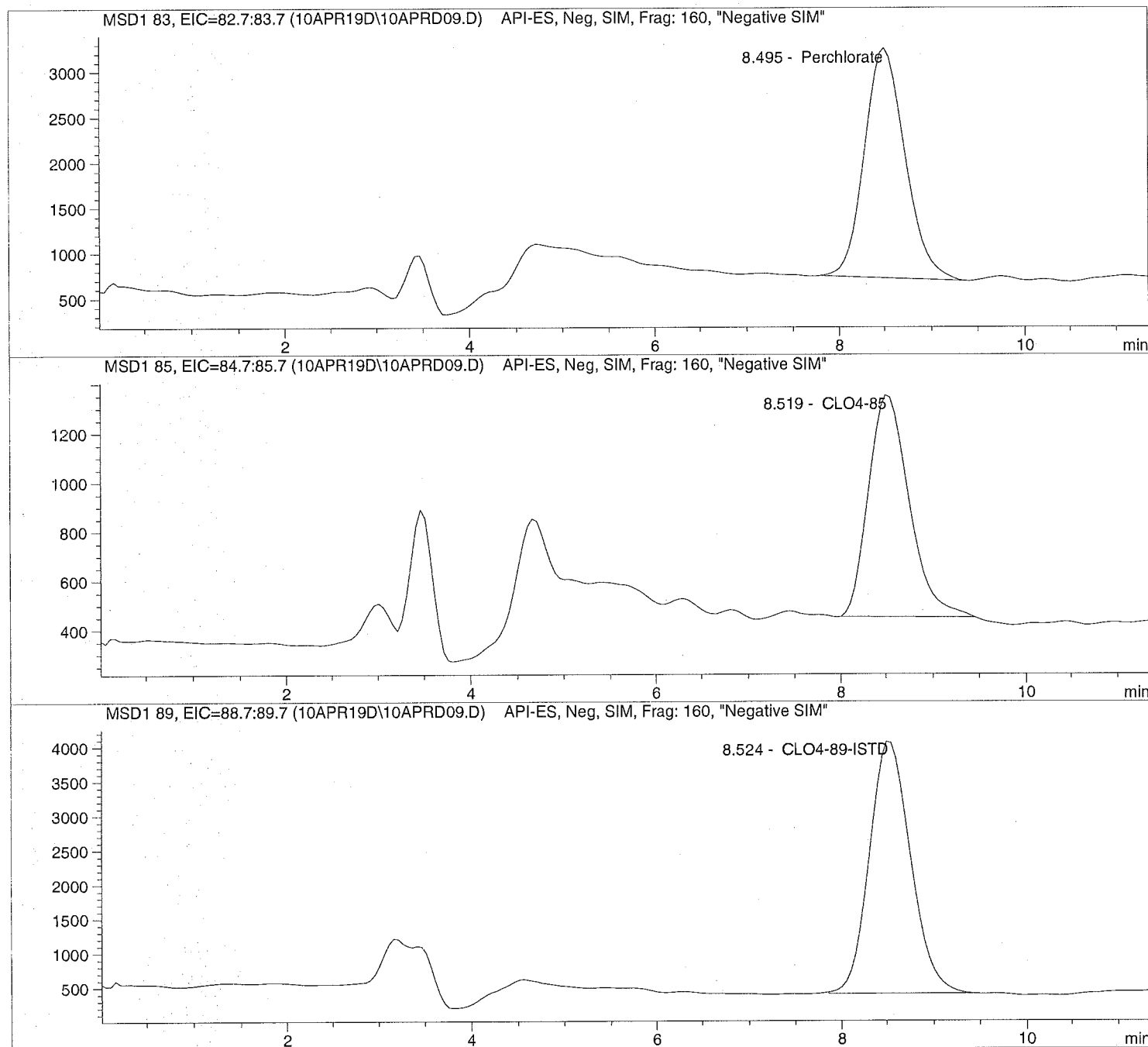
Sample Name: 1909154001

Injection Date: 4/10/2019 11:48:52
Sample Name: 1909154001
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD09.D Sample Name: 1909154001

```

=====
Injection Date: 4/10/2019 11:48:52 Seq Line: 9
Sample Name: 1909154001 Location: Vial 79
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.495	PBA	77912.3	2.3402	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.519	PBA	27611.7	2.6142	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.524	BBA	113404.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```


Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD10.D

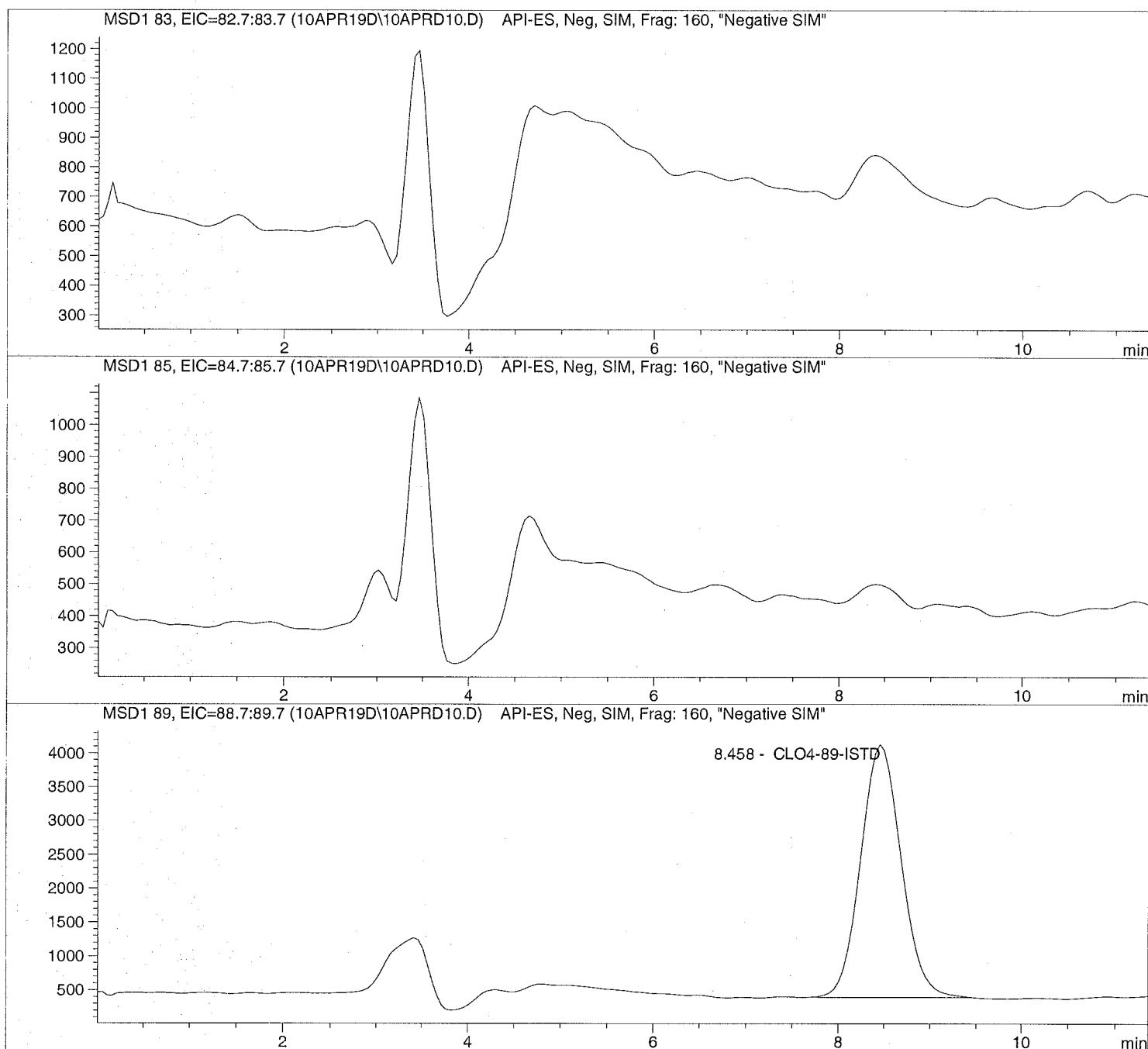
Sample Name: 1909947001

Injection Date: 4/10/2019 12:02:10
Sample Name: 1909947001
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD10.D

Sample Name: 1909947001

```

=====
Injection Date: 4/10/2019 12:02:10      Seq Line:          10
Sample Name:    1909947001              Location:          Vial 80
Acq Operator:   TNB                      Inj. No.:         1
                                           Inj. Vol.:        30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.458	BBA	115129.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

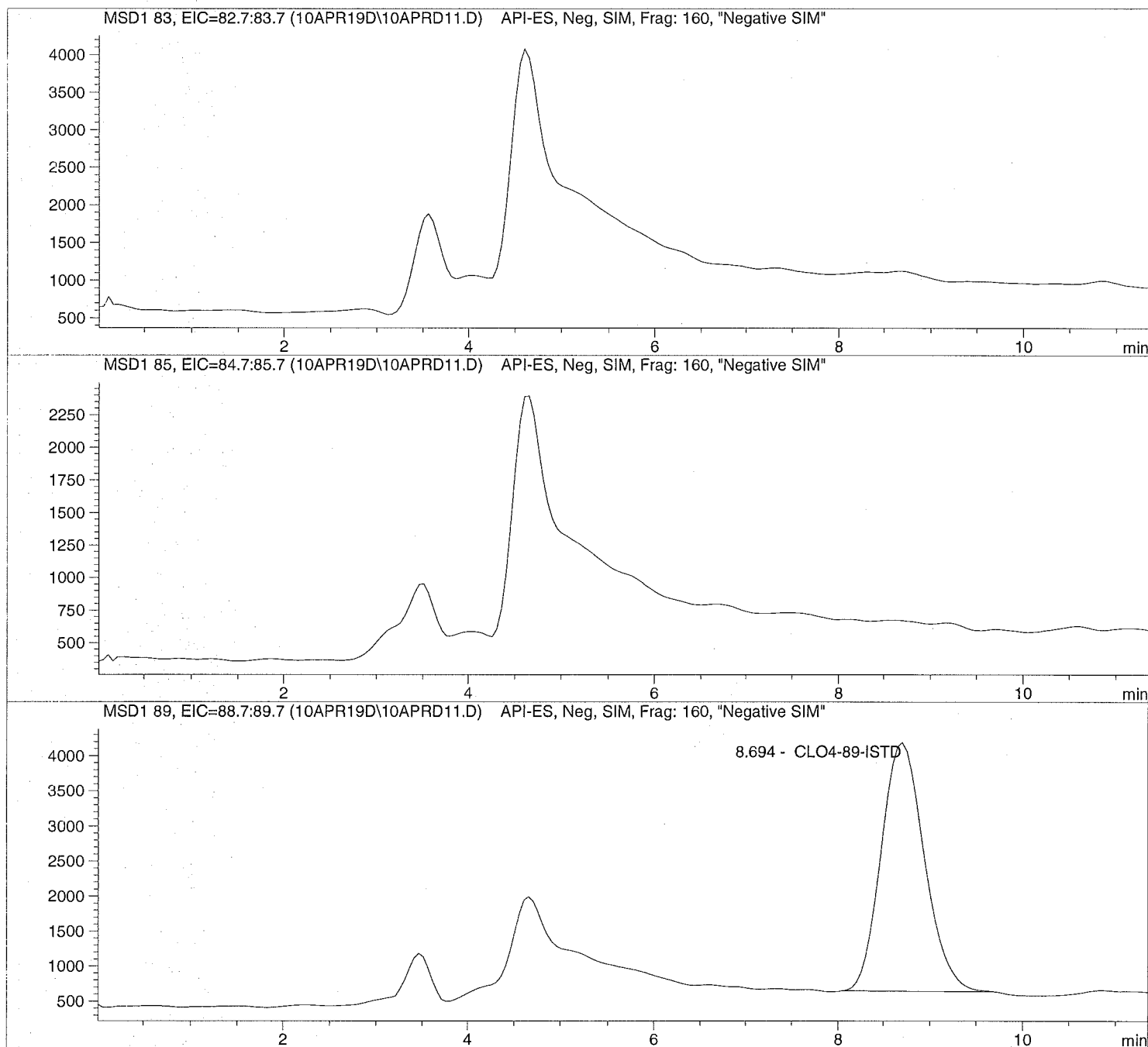
```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD11.D Sample Name: 1909949001

```
=====
Injection Date: 4/10/2019 12:15:27      Seq Line:      11
Sample Name:    1909949001              Location:      Vial 81
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD11.D Sample Name: 1909949001

```

=====
Injection Date: 4/10/2019 12:15:27      Seq Line:      11
Sample Name:    1909949001              Location:      Vial 81
Acq Operator:   TNB                      Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.694	PBA	114791.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD12.D

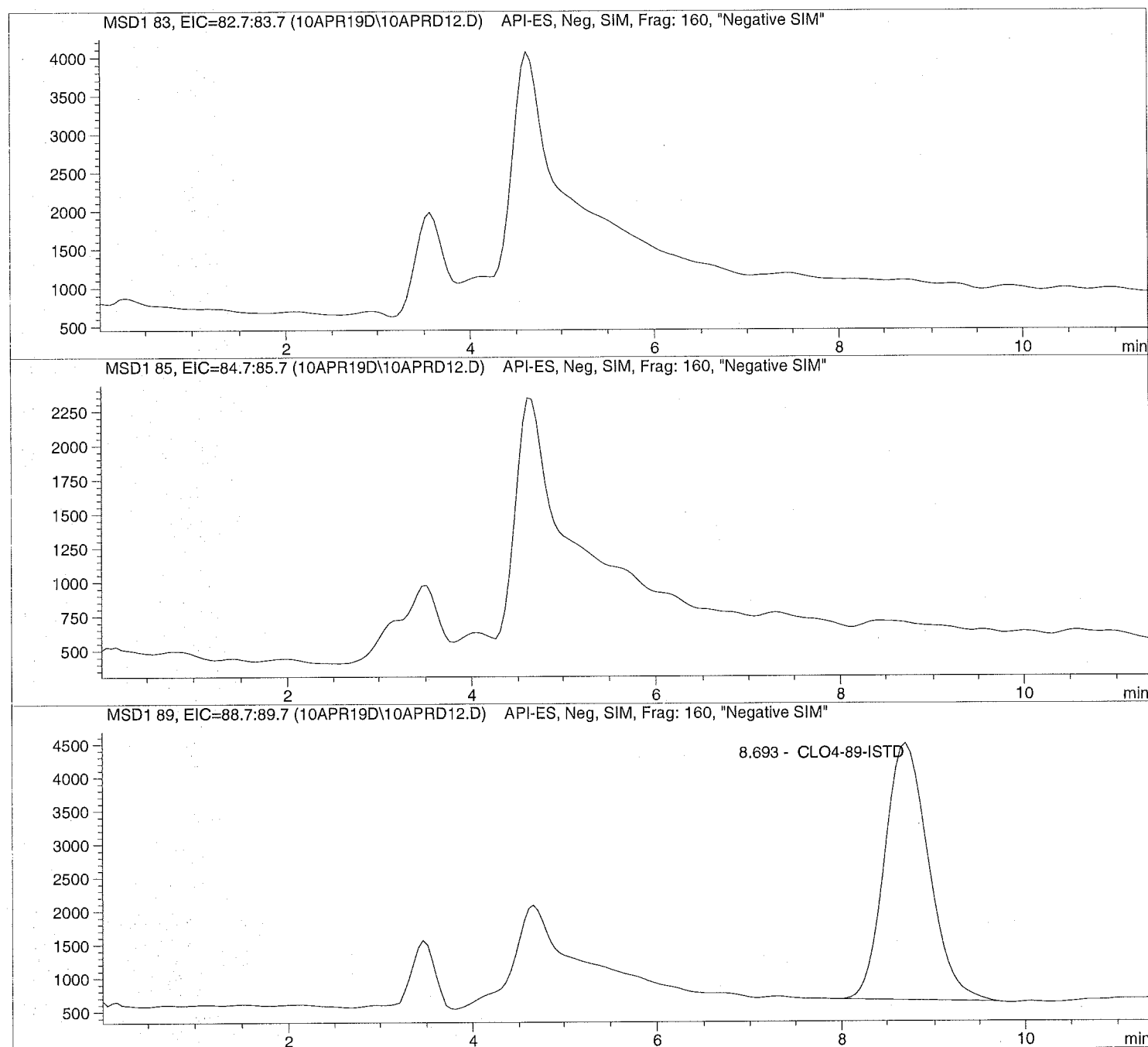
Sample Name: 1909949002

=====
Injection Date: 4/10/2019 12:28:41
Sample Name: 1909949002
Acq Operator: TNB

Seq Line: 12
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD12.D Sample Name: 1909949002

```

=====
Injection Date: 4/10/2019 12:28:41      Seq Line: 12
Sample Name: 1909949002      Location: Vial 82
Acq Operator: TNB      Inj. No.: 1
                                 Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

=====
Sample Information
=====

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.693	PBA	124045.6	5.0000	CLO4-89-ISTD

=====
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD13.D

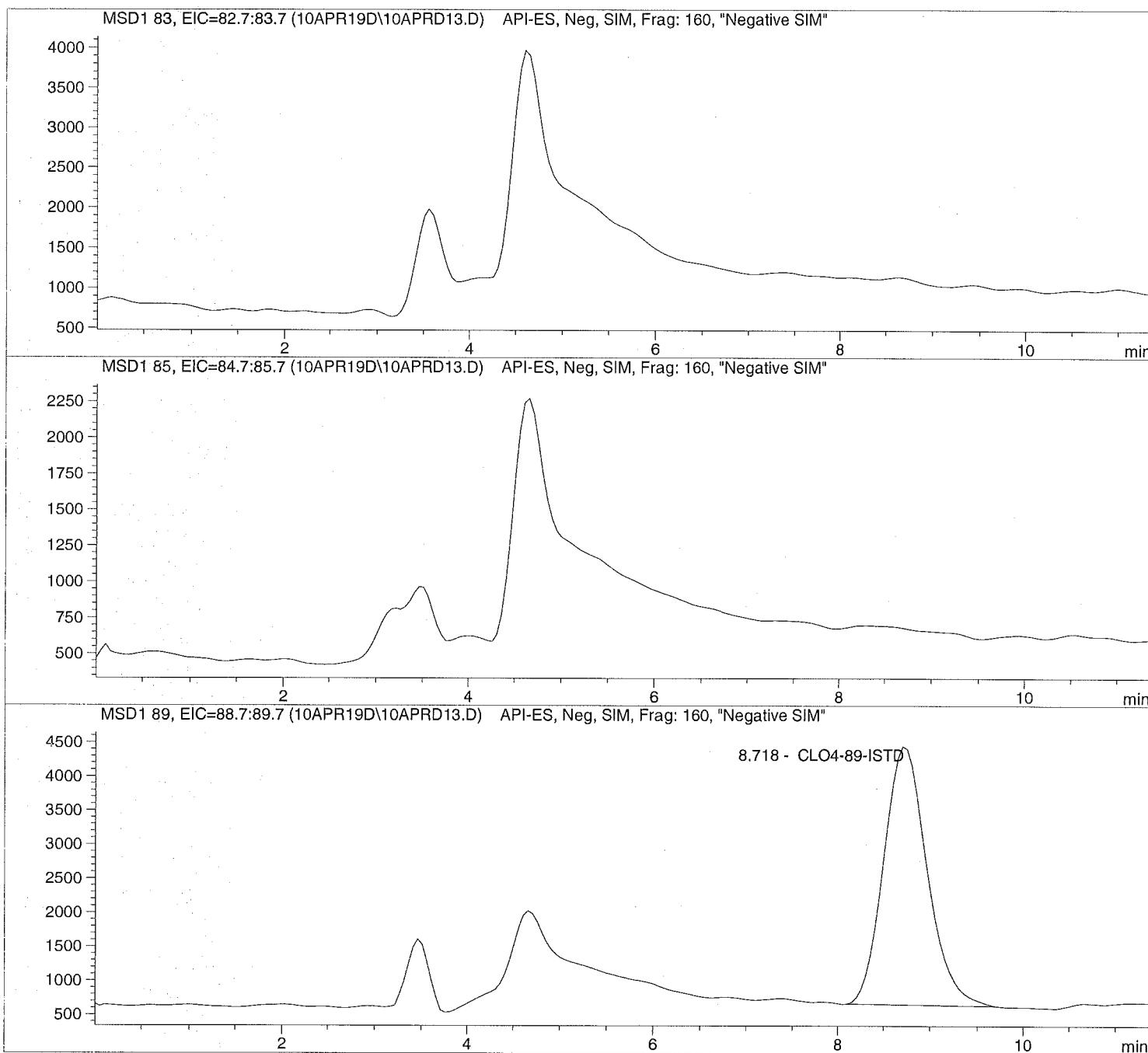
Sample Name: 1909949003

=====
Injection Date: 4/10/2019 12:41:59
Sample Name: 1909949003
Acq Operator: TNB

Seq Line: 13
Location: Vial 83
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD13.D Sample Name: 1909949003

```

=====
Injection Date: 4/10/2019 12:41:59 Seq Line: 13
Sample Name: 1909949003 Location: Vial 83
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl
  
```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
  
```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
  
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.718	PBA	123373.0	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD14.D

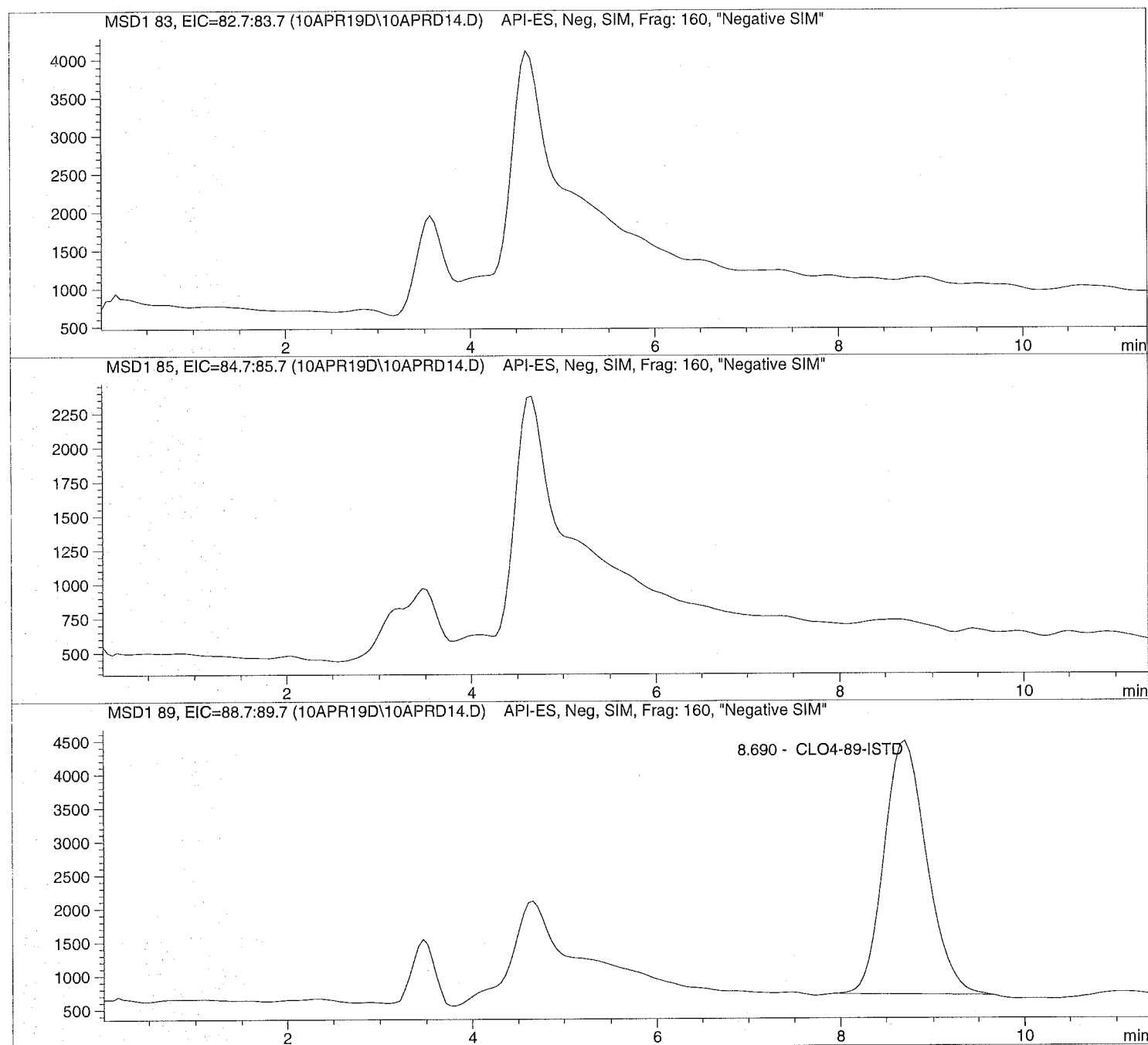
Sample Name: 1909949004

=====
Injection Date: 4/10/2019 12:55:15
Sample Name: 1909949004
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD14.D Sample Name: 1909949004

```

=====
Injection Date: 4/10/2019 12:55:15      Seq Line: 14
Sample Name: 1909949004                Location: Vial 84
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.690	BBA	120241.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD15.D

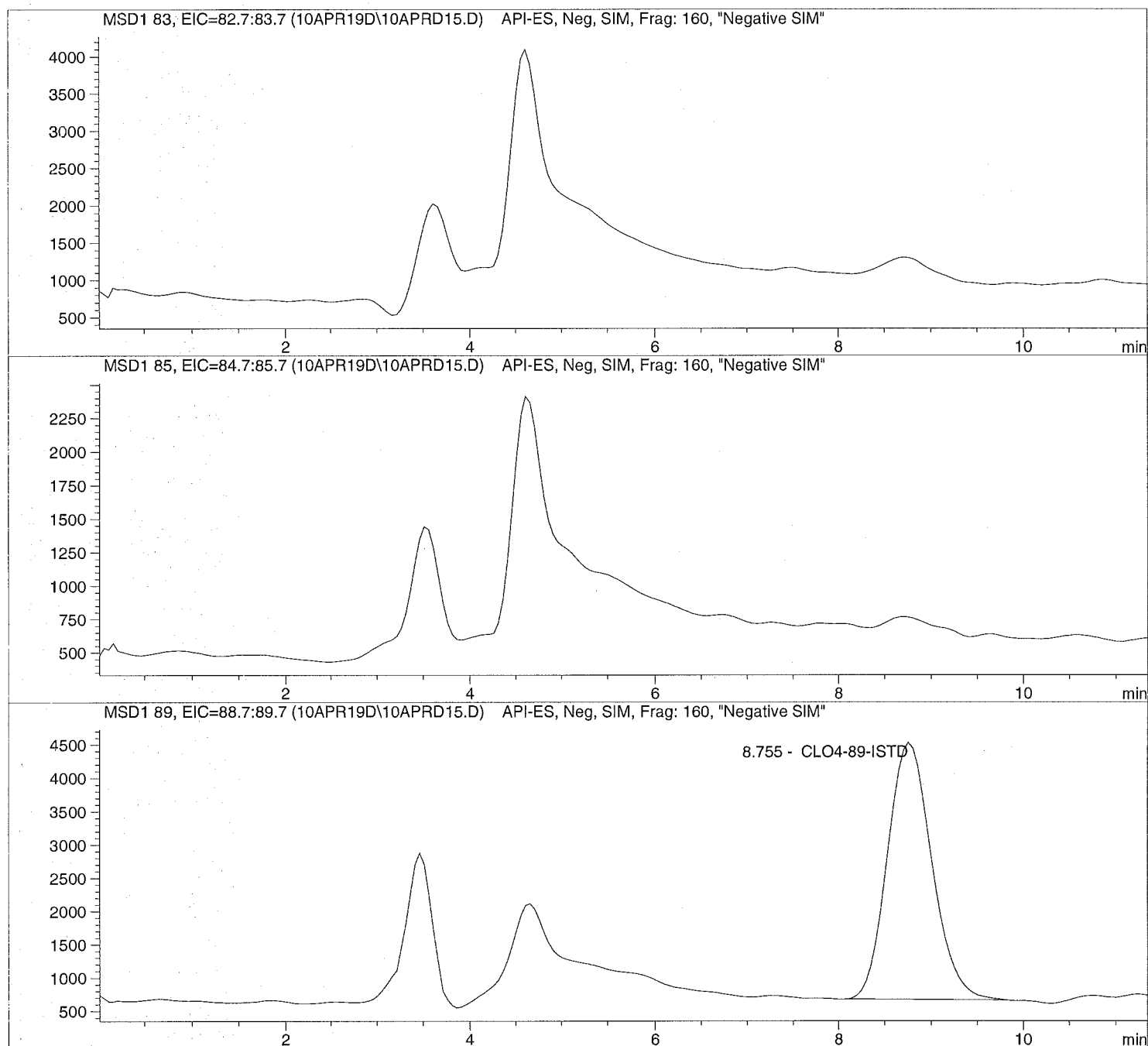
Sample Name: 1909949005

=====
Injection Date: 4/10/2019 13:08:30
Sample Name: 1909949005
Acq Operator: TNB

Seq Line: 15
Location: Vial 85
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD15.D

Sample Name: 1909949005

```

=====
Injection Date: 4/10/2019 13:08:30      Seq Line:      15
Sample Name:    1909949005              Location:      Vial 85
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	PBA	127767.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD16.D

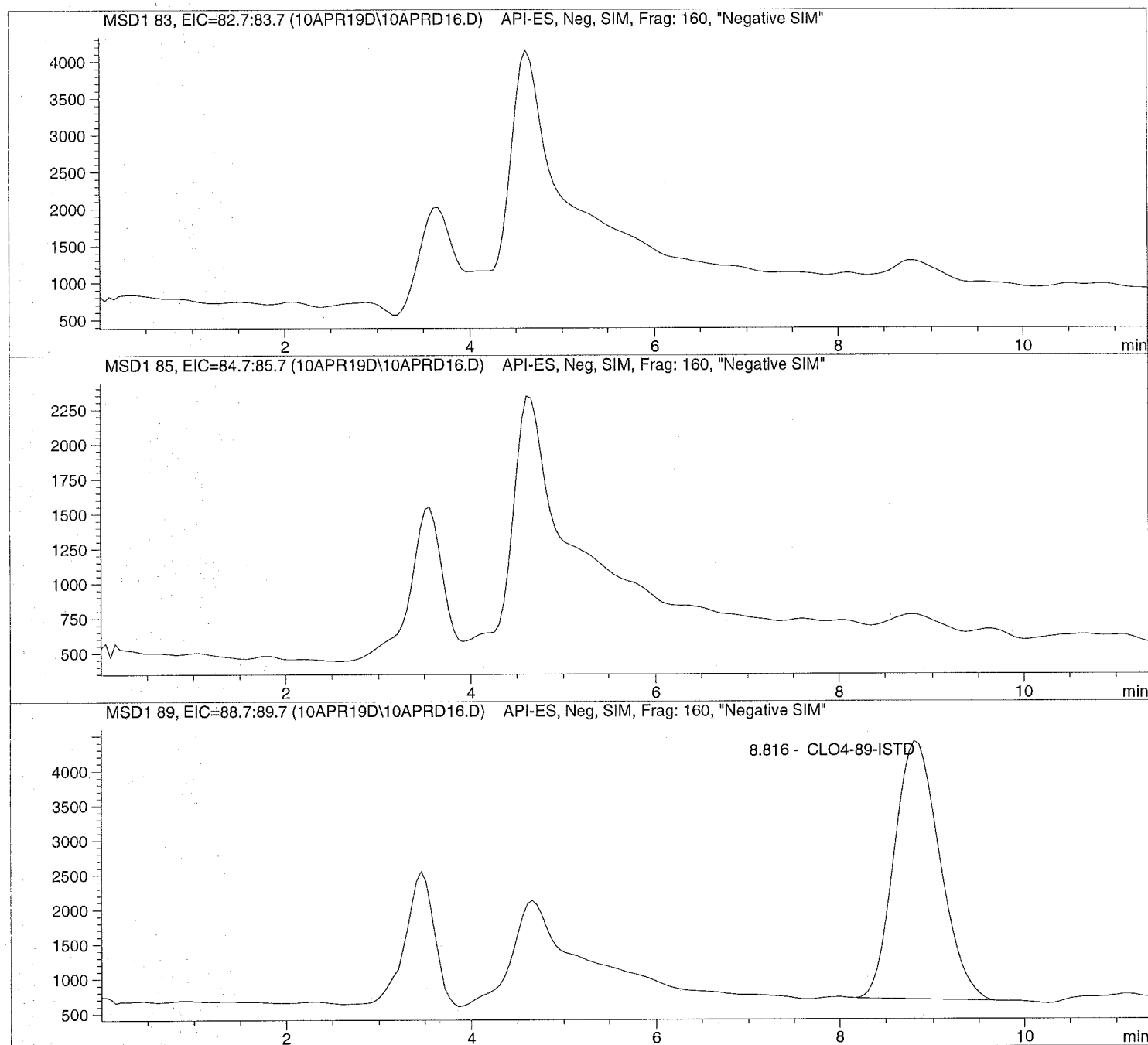
Sample Name: 1909949006

=====
Injection Date: 4/10/2019 13:21:46
Sample Name: 1909949006
Acq Operator: TNB

Seq Line: 16
Location: Vial 86
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD16.D

Sample Name: 1909949006

```

=====
Injection Date: 4/10/2019 13:21:46      Seq Line: 16
Sample Name: 1909949006                Location: Vial 86
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.816	BBA	123964.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD17.D

Sample Name: 647202 CCV@25

Injection Date: 4/10/2019 13:35:04

Seq Line: 17

Sample Name: 647202 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

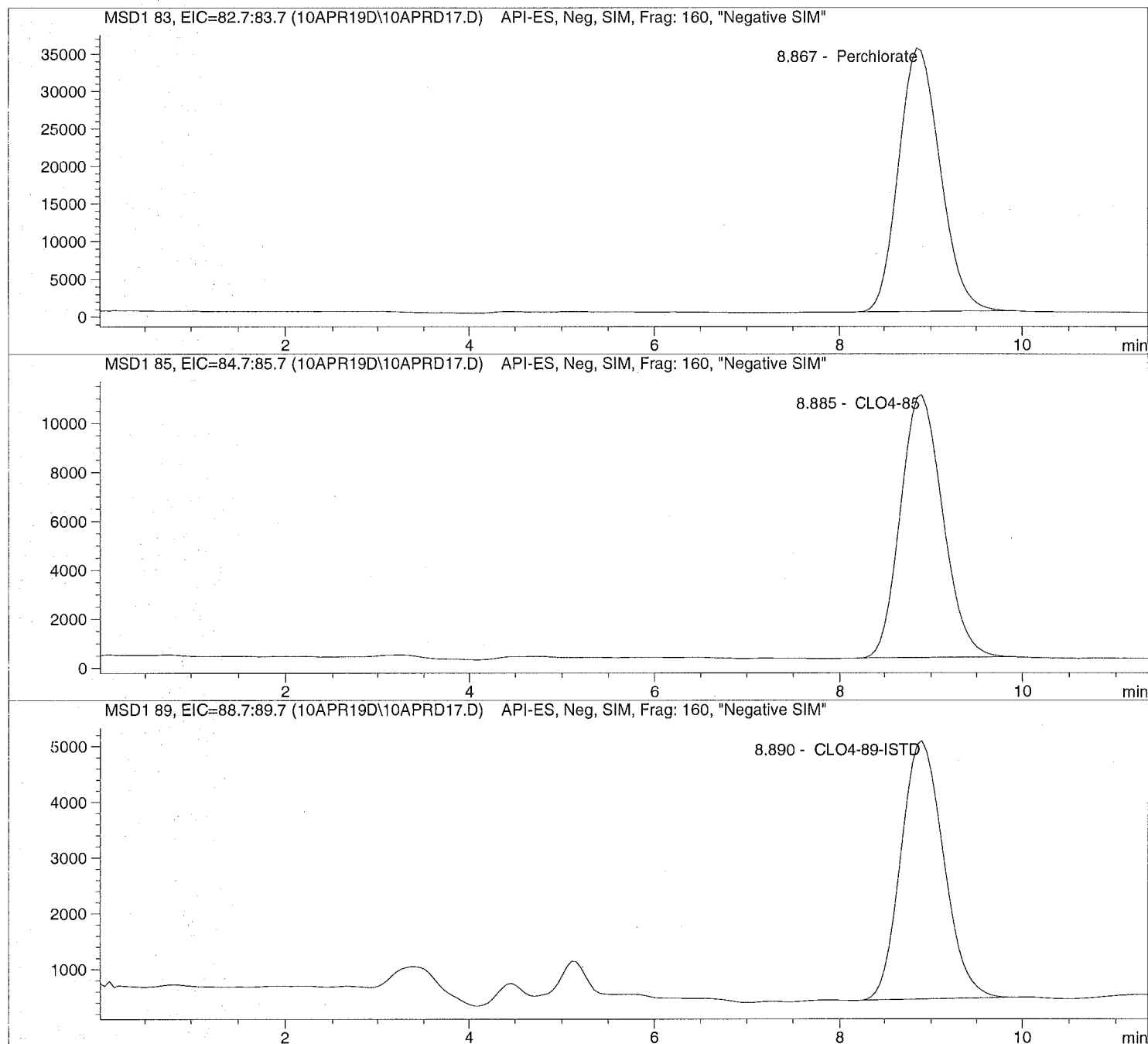
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis





ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
Calibration Table
=====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
Based on : Peak Area

Rel. Reference Window : 20.000 %
Abs. Reference Window : 0.000 min
Rel. Non-ref. Window : 20.000 %
Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
Uncalibrated Peaks : not reported
Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
Origin : Ignored (some peaks differ, see below)
Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
Average Response : Average all calibrations
Average Retention Time: Floating Average New 75%

Calibration Report Options :
Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
Signal 2: MSD1 85, EIC=84.7:85.7
Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl Sig	Amount	Area	Amt/Area	Ref	Grp Name
8.744	1 1	1.00000	7.76074e4	1.28854e-5	1	Perchlorate
	2	2.00000	1.35273e5	1.47849e-5		
	3	5.00000	3.37764e5	1.48033e-5		
	4	10.00000	6.83454e5	1.46316e-5		
	5	25.00000	2.08433e6	1.19943e-5		
	6	50.00000	4.13334e6	1.20968e-5		
	7	75.00000	5.99313e6	1.25143e-5		
8.755	2 1	1.00000	2.36780e4	4.22333e-5	1	CLO4-85
	2	2.00000	4.69486e4	4.25998e-5		
	3	5.00000	1.06124e5	4.71147e-5		
	4	10.00000	2.13523e5	4.68335e-5		
	5	25.00000	6.14295e5	4.06971e-5		
	6	50.00000	1.19814e6	4.17315e-5		
	7	75.00000	1.78355e6	4.20509e-5		
8.766	3 1	5.00000	2.73208e5	1.83011e-5	+I1	CLO4-89-ISTD
	2	5.00000	2.24886e5	2.22335e-5		
	3	5.00000	2.33196e5	2.14412e-5		
	4	5.00000	2.34454e5	2.13262e-5		
	5	5.00000	2.50568e5	1.99547e-5		
	6	5.00000	2.30977e5	2.16472e-5		

RetTime	Lvl	Amount	Area	Amt/Area	Ref Grp Name
[min]	Sig				
----- ---	-----	-----	-----	-----	-----
7		5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-85

Time Window : From 6.650 min To 12.505 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

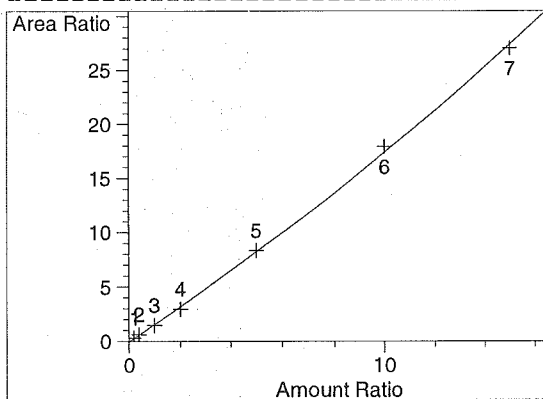
Compound: CLO4-89-ISTD

Time Window : From 6.659 min To 12.466 min
Curve Type : Linear
Origin : Included
Calibration Level Weights:/
Level 1 : 1
Level 2 : 1
Level 3 : 1
Level 4 : 1
Level 5 : 1
Level 6 : 1
Level 7 : 1

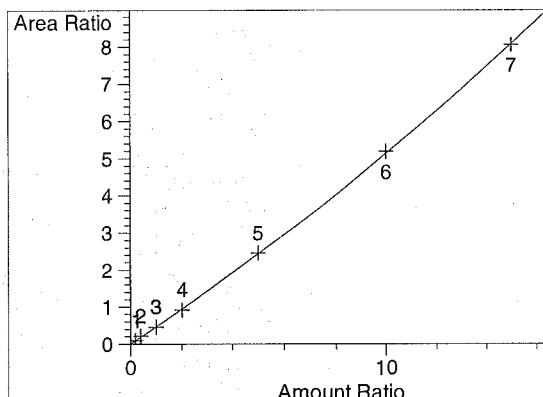
=====
Peak Sum Table
=====

No Entries in table
=====

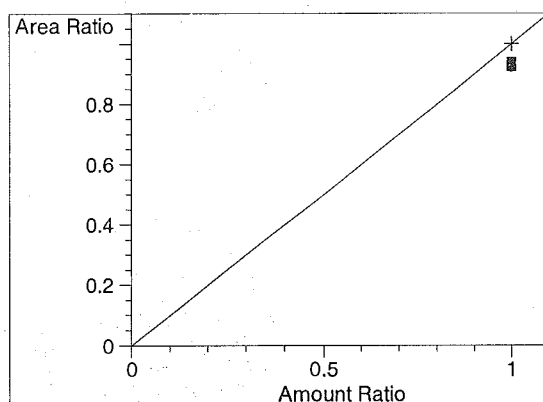
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

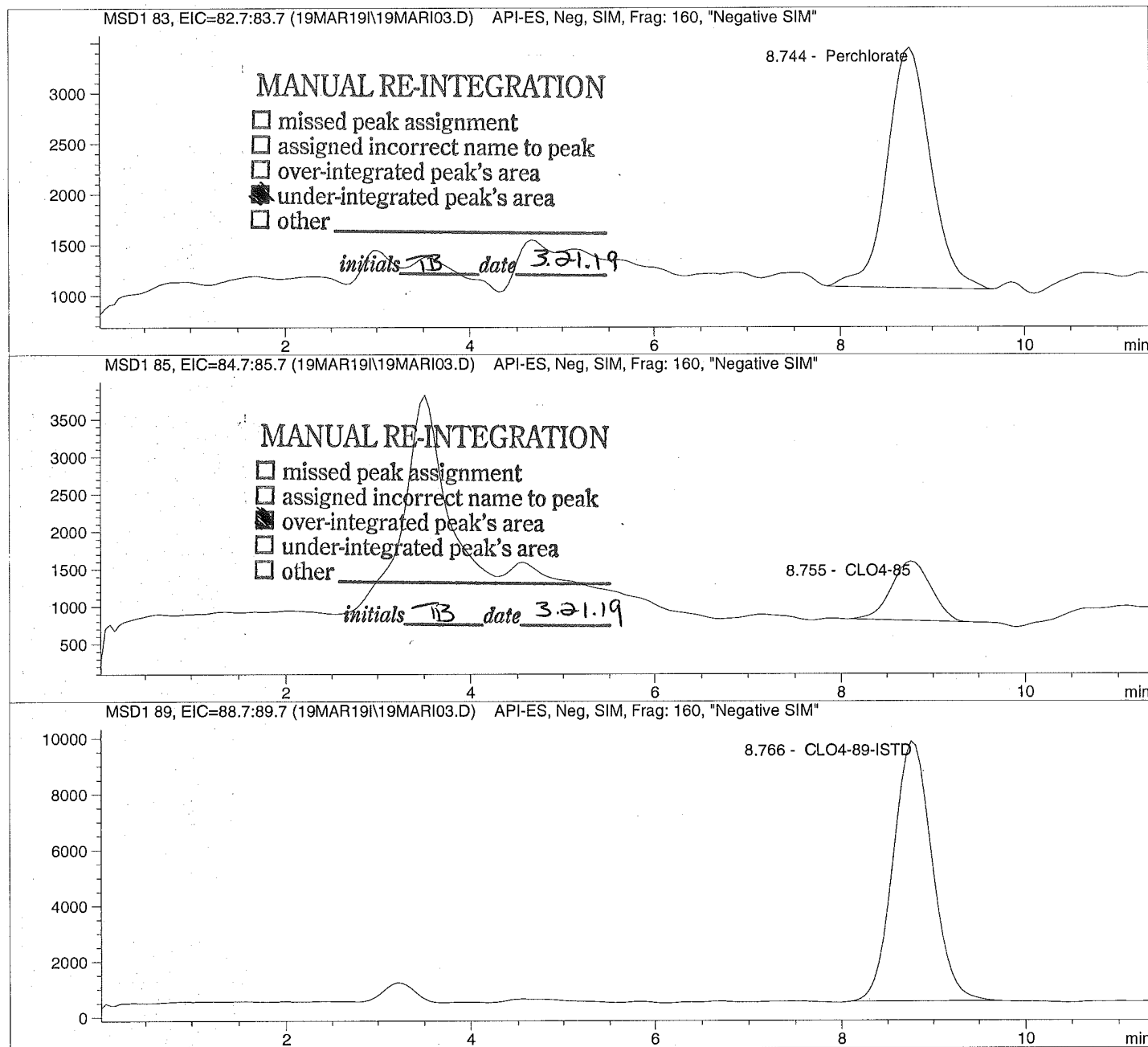
Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line: 3
Sample Name: CLO4@ 1.0ug/L              Location: Vial 73
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 1.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

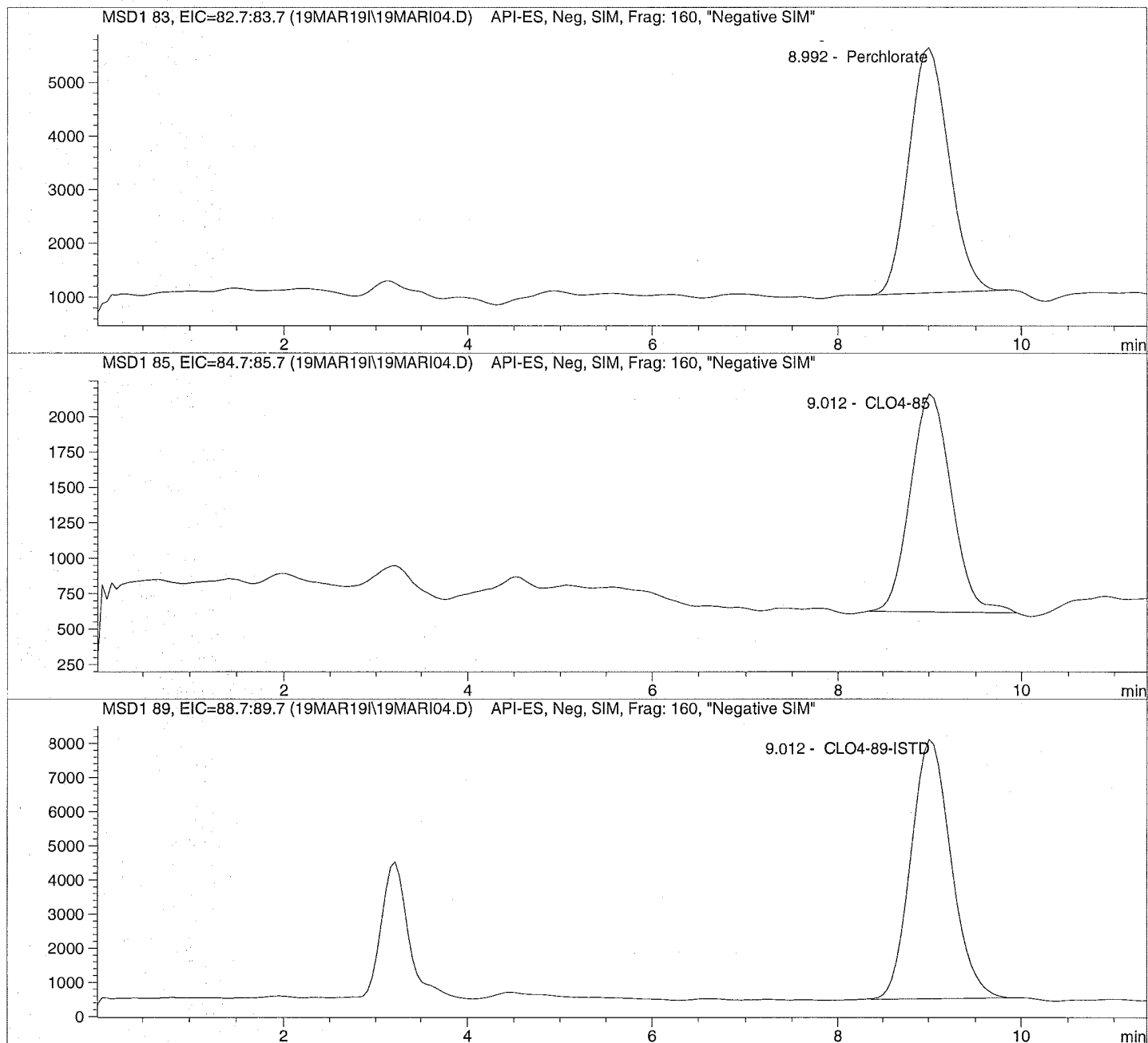
```


Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```
=====
Injection Date: 3/19/2019 09:53:00      Seq Line: 4
Sample Name:    CLO4@ 2.0ug/L           Location:  Vial 74
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line: 4
Sample Name:    CLO4@ 2.0ug/L           Location:  Vial 74
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D

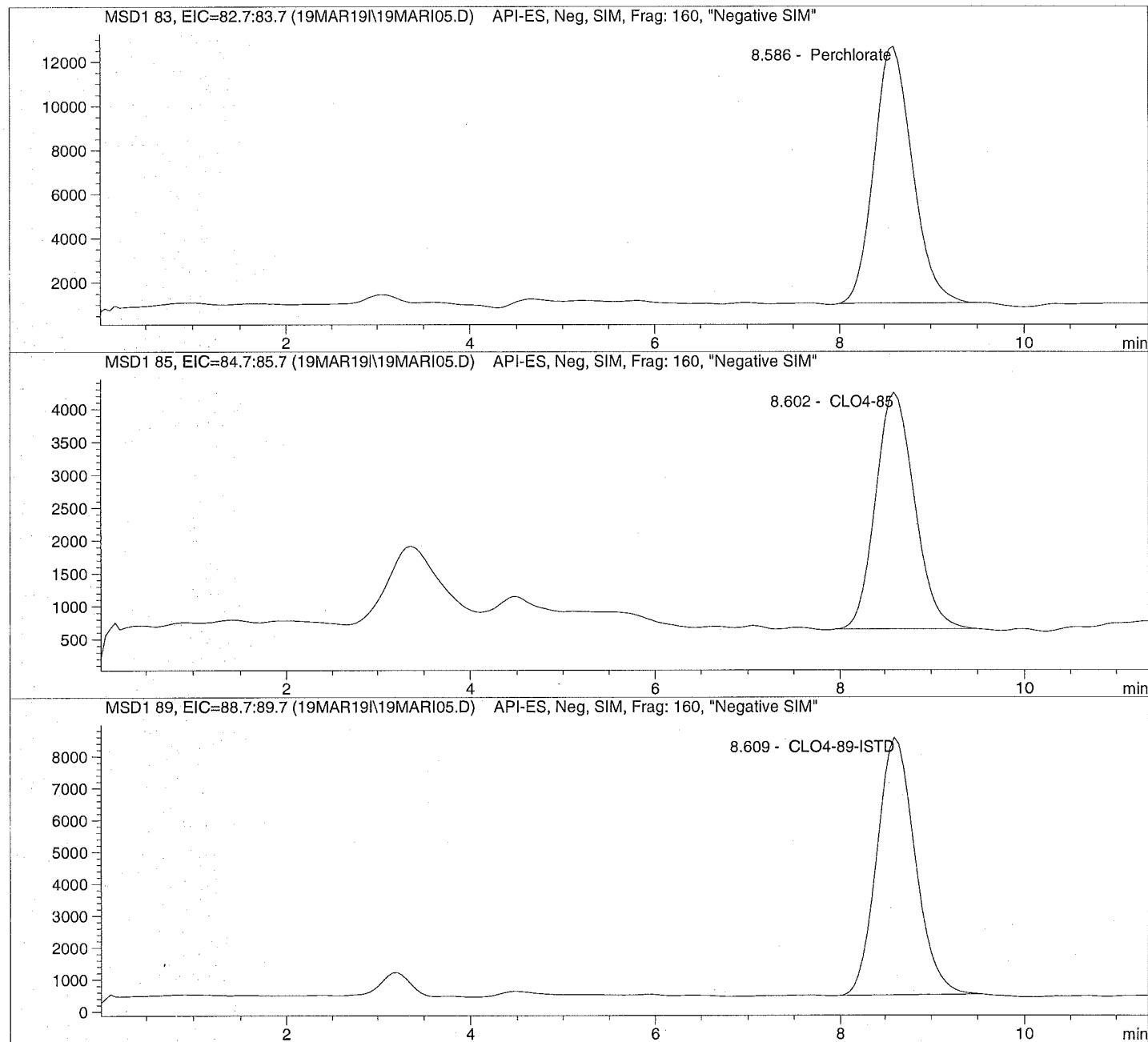
Sample Name: CLO4@ 5.0ug/L

=====
Injection Date: 3/19/2019 10:06:16
Sample Name: CLO4@ 5.0ug/L
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line: 5
Sample Name:    CLO4@ 5.0ug/L           Location:  Vial 75
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D

Sample Name: CLO4@ 10.ug/L

Injection Date: 3/19/2019 10:19:32

Seq Line: 6

Sample Name: CLO4@ 10.ug/L

Location: Vial 76

Acq Operator: TNB

Inj. No.: 1

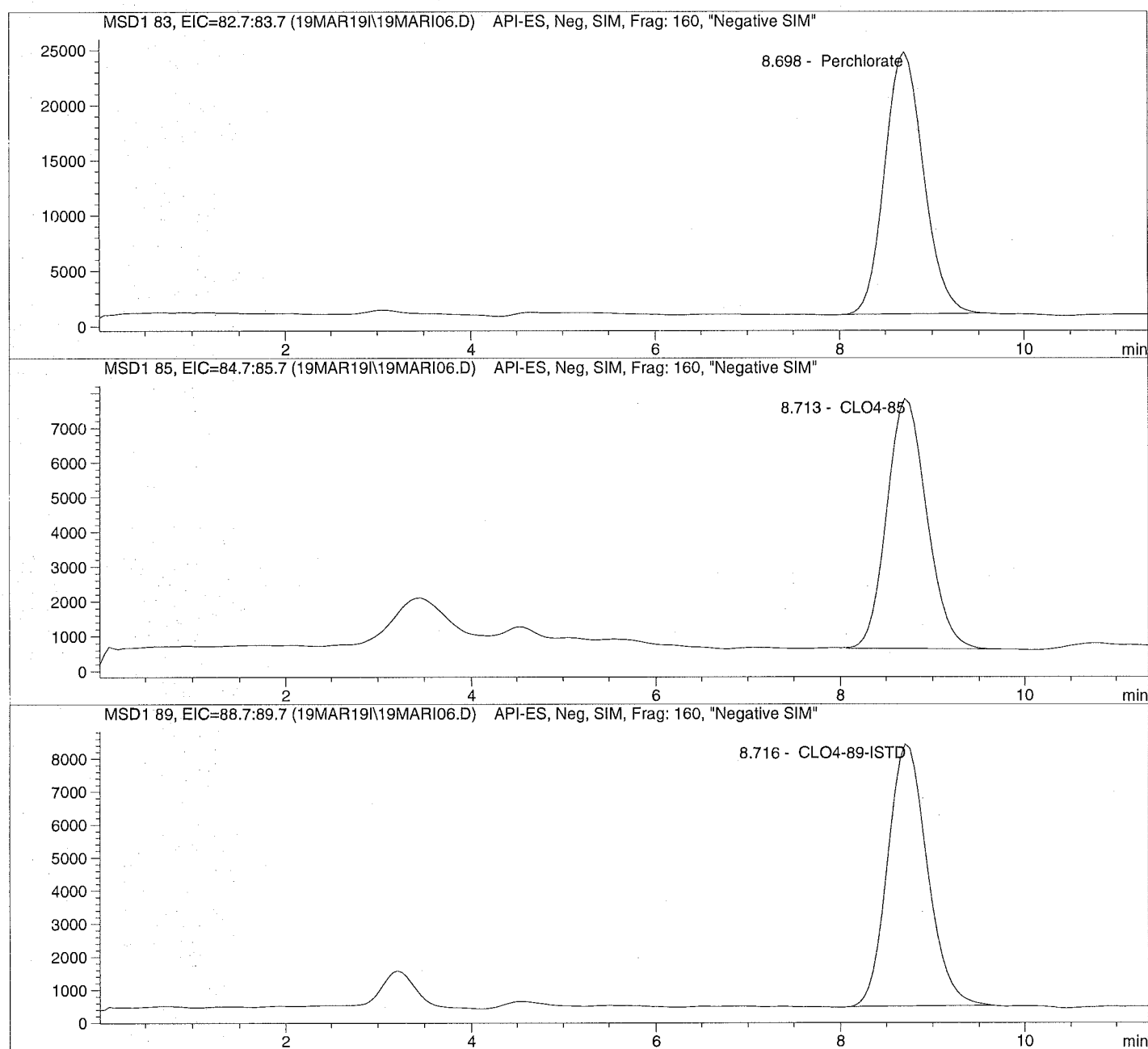
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

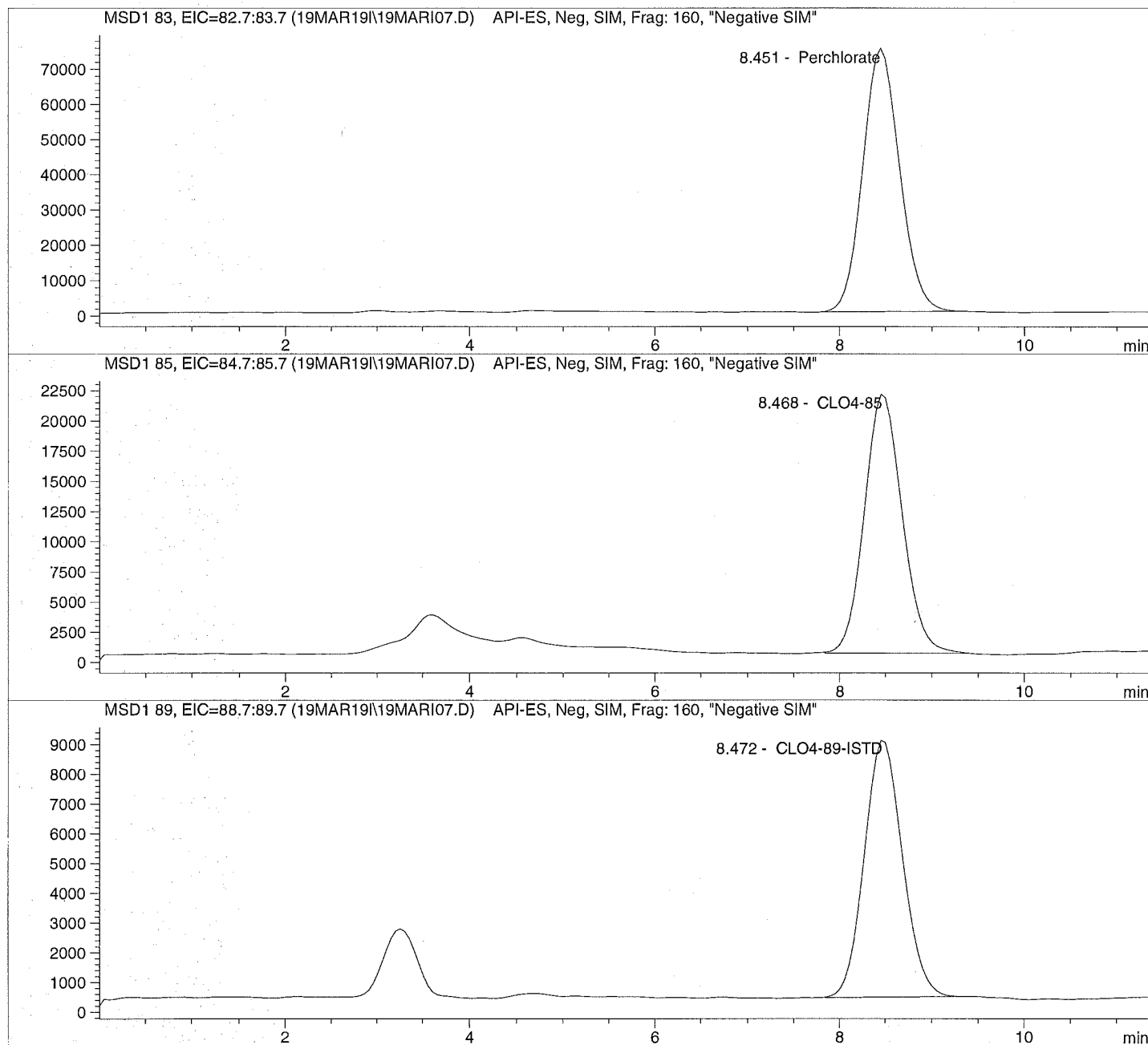
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```
=====
Injection Date: 3/19/2019 10:32:49      Seq Line: 7
Sample Name:    CLO4@ 25.ug/L           Location:  Vial 77
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 3/19/2019 10:32:49      Seq Line: 7
Sample Name:    CLO4@ 25.ug/L           Location:  Vial 77
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

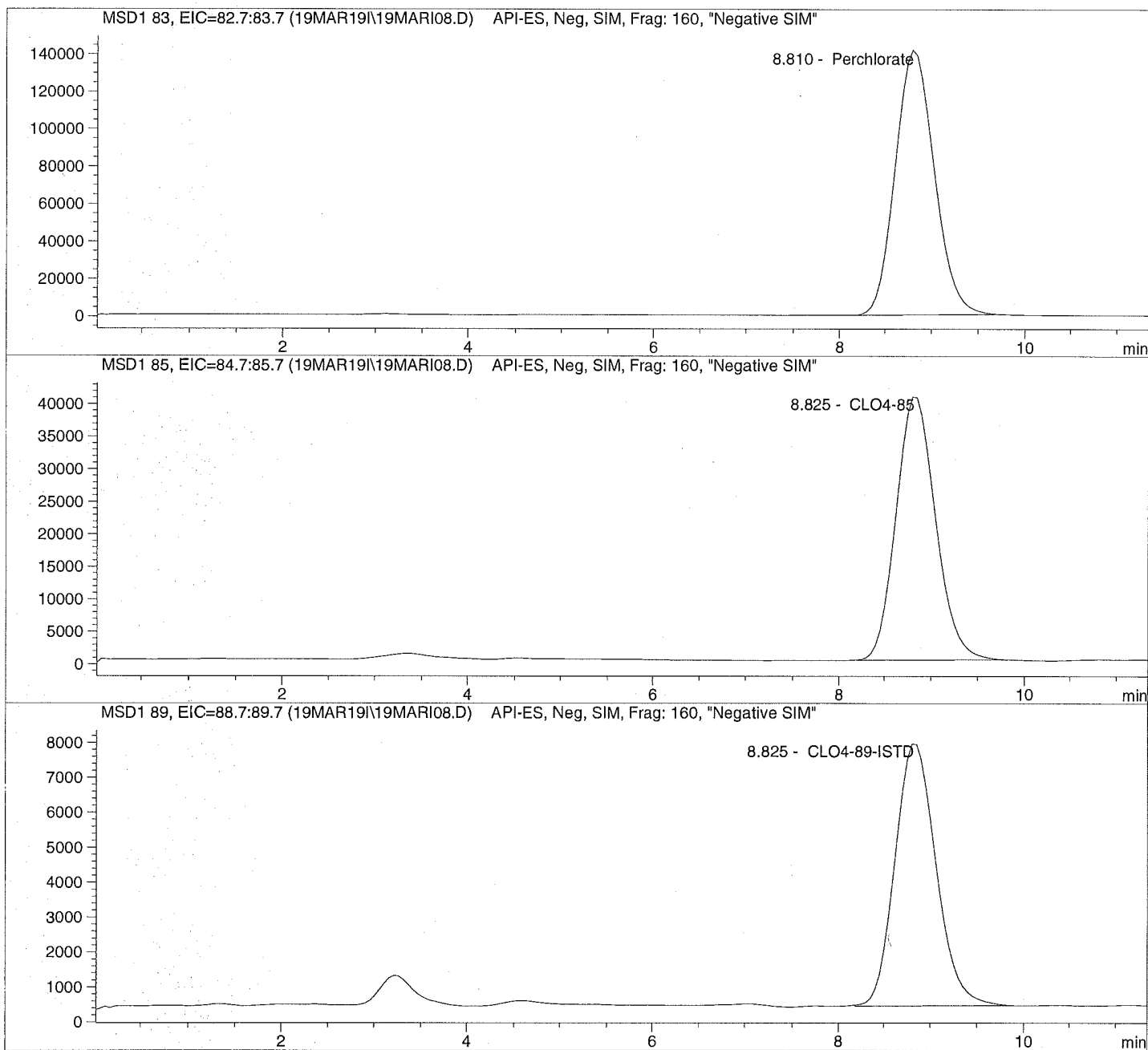
```


Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```
=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  50.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

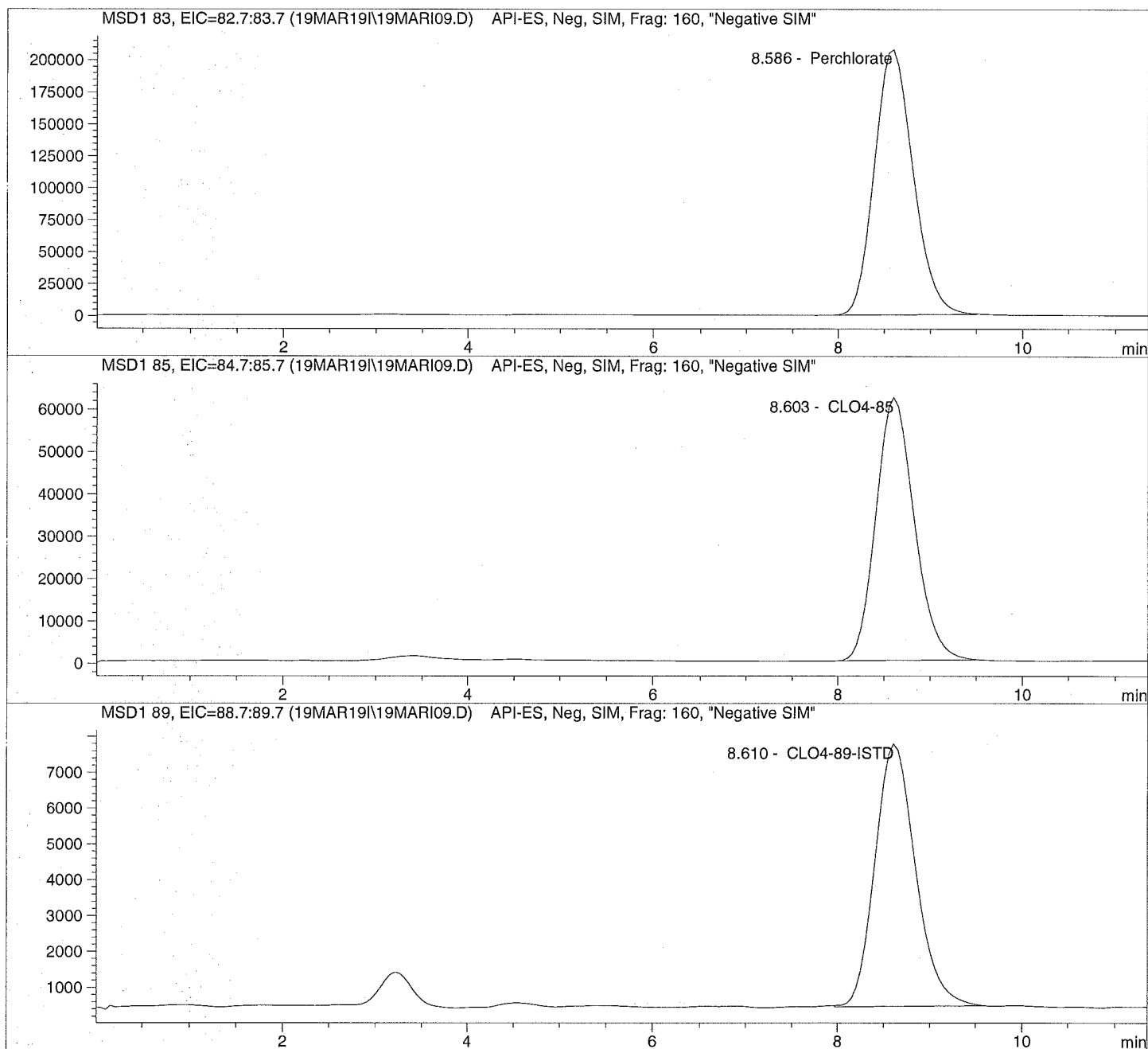
```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```
=====
Injection Date: 3/19/2019 10:59:22      Seq Line:          9
Sample Name:    CLO4@ 75.ug/L           Location:           Vial 79
Acq Operator:   TNB                     Inj. No.:          1
                                           Inj. Vol.:         30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date: 3/19/2019 10:59:22      Seq Line: 9
Sample Name: CLO4@ 75.ug/L      Location: Vial 79
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 75.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI10.D

Sample Name: ICAL Verf@10ug/L

Injection Date: 3/19/2019 11:12:42

Seq Line: 10

Sample Name: ICAL Verf@10ug/L

Location: Vial 80

Acq Operator: TNB

Inj. No.: 1

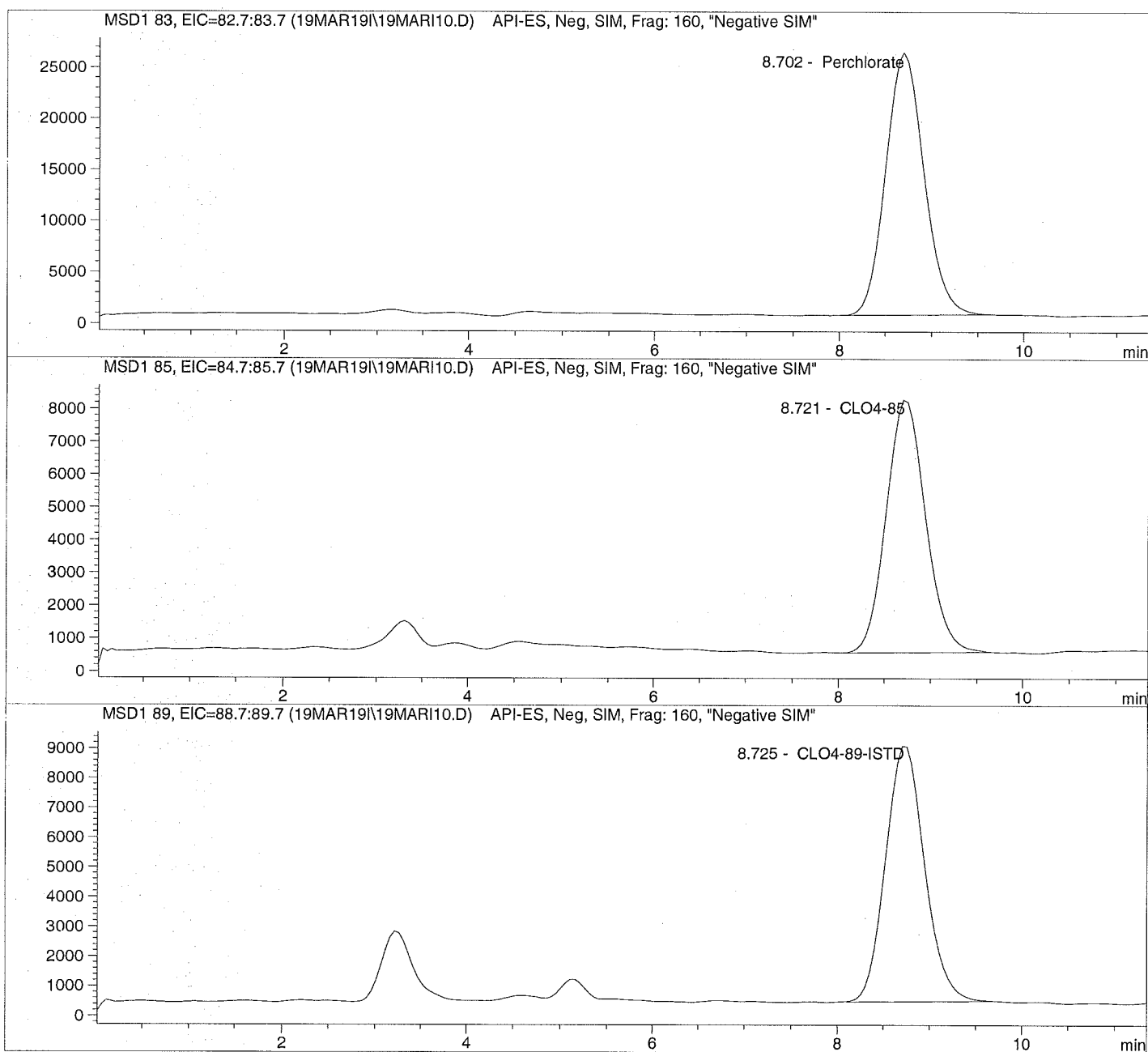
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 3/19/2019 11:12:42      Seq Line:            10
Sample Name:    ICAL Verf@10ug/L        Location:            Vial 80
Acq Operator:   TNB                      Inj. No.:            1
                                         Inj. Vol.:            30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22

```

Perchlorate analysis

=====
Sample Information
=====

```

Sorted By:                    Signal
Calib. Data Modified:    Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                1.000000
Dilution:                  1.000000
Sample Amount:             10.000

```

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

=====
*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

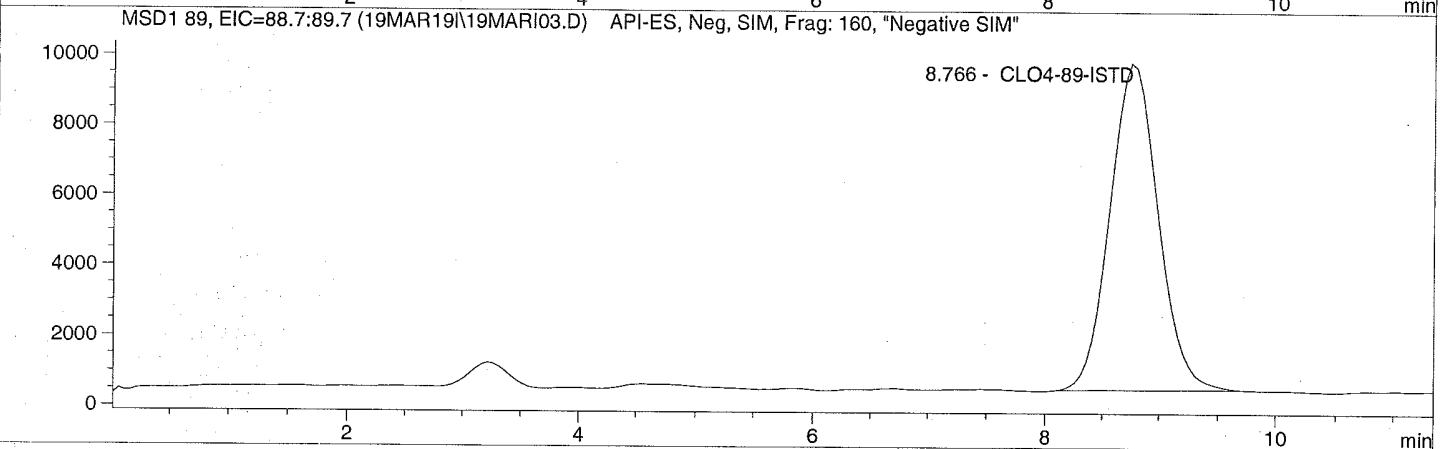
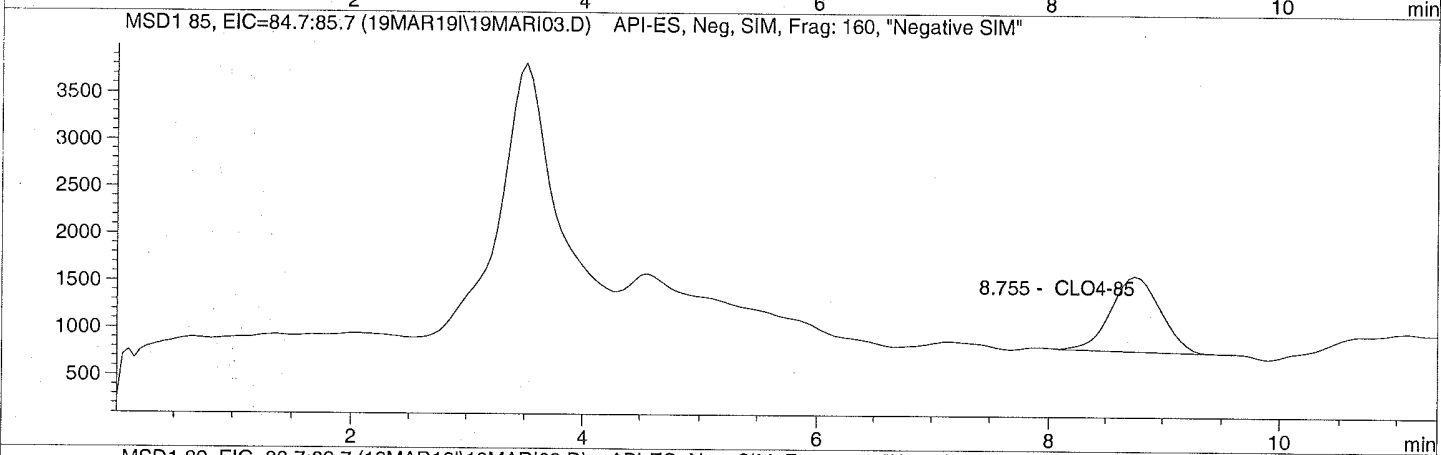
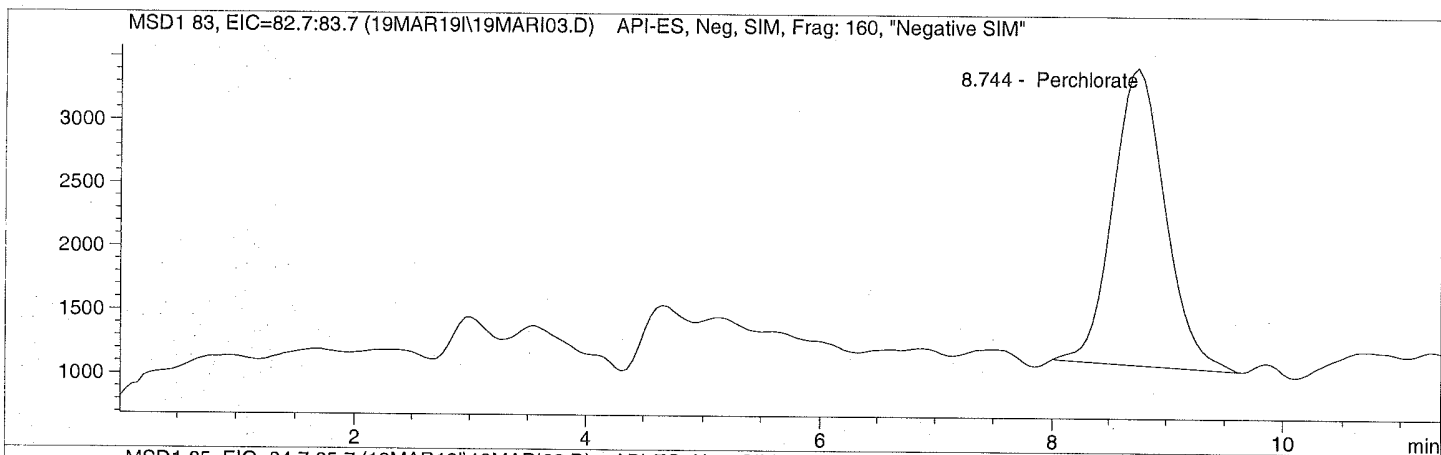
Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

=====
Injection Date: 3/19/2019 09:39:40 Seq Line: 3
Sample Name: CLO4@ 1.0ug/L Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L          Location:  Vial 73
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:38:25
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

*** End of Report ***



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Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

April 12, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19031508**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 2 sample(s) on Mar 28, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031508

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19031508-01	LH18/24-SP140_032719	Water		27-Mar-2019 14:00	28-Mar-2019 08:30	<input type="checkbox"/>
HS19031508-02	Trip Blank -ALS-020119-59	Water		27-Mar-2019 00:00	28-Mar-2019 08:30	<input type="checkbox"/>

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.

GCMS Semivolatiles by Method SW8270SIM**Batch ID: 139415****Sample ID: LH18/24-SP140_032719 (HS19031508-01)**

- The surrogate recoveries could not be determined due to dilution below the calibration range.

GCMS Volatiles by Method SW8260**Batch ID: R335882****Sample ID: CCV**

- 1,2-Dibromoethane, Bromoform and Dibromochloromethane exceeded %D limits for CCV. Samples are ND for these compounds.

Sample ID: HS19040036-36MS

- MS and MSD are for an unrelated sample

Metals by Method SW6020**Batch ID: 139323****Sample ID: HS19031584-01MS**

- MS/MSD and DUPs are for an unrelated sample

Metals by Method SW7470**Batch ID: 139318**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

WetChemistry by Method E1664A**Batch ID: R336080**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

WetChemistry by Method SW9056**Batch ID: R335875**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

WetChemistry by Method E410.4**Batch ID: R335804**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
-

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP140_032719
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031508
 Lab ID:HS19031508-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,1,1-Trichloroethane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,1,2,2-Tetrachloroethane	5.0	U	5.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,1,2-Trichloroethane	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,1-Dichloroethane	6.4	J	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,1-Dichloroethene	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,1-Dichloropropene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2,3-Trichlorobenzene	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2,3-Trichloropropane	5.0	U	5.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2,4-Trichlorobenzene	5.0	U	5.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2,4-Trimethylbenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2-Dibromo-3-chloropropane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2-Dibromoethane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2-Dichlorobenzene	5.0	U	5.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2-Dichloroethane	58		2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,2-Dichloropropane	5.0	U	5.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,3,5-Trimethylbenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,3-Dichlorobenzene	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,3-Dichloropropane	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
1,4-Dichlorobenzene	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
2,2-Dichloropropane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
2-Butanone	10	U	5.0	10	20	UG/L	10	03-Apr-2019 14:28	
2-Chlorotoluene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
2-Hexanone	10	U	10	10	20	UG/L	10	03-Apr-2019 14:28	
4-Chlorotoluene	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
4-Isopropyltoluene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
4-Methyl-2-pentanone	10	U	7.0	10	20	UG/L	10	03-Apr-2019 14:28	
Acetone	20	U	4.0	20	20	UG/L	10	03-Apr-2019 14:28	
Benzene	7.8	J	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Bromobenzene	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Bromochloromethane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Bromodichloromethane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Bromoform	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Bromomethane	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Carbon disulfide	10	U	6.0	10	20	UG/L	10	03-Apr-2019 14:28	
Carbon tetrachloride	25		5.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Chlorobenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Chloroethane	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Chloroform	9.1	J	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP140_032719
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031508
 Lab ID:HS19031508-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
cis-1,2-Dichloroethene	1,900		2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
cis-1,3-Dichloropropene	5.0	U	1.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Dibromochloromethane	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Dibromomethane	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Dichlorodifluoromethane	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Ethylbenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Hexachlorobutadiene	10	U	10	10	10	UG/L	10	03-Apr-2019 14:28	
Isopropylbenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
m,p-Xylene	10	U	5.0	10	20	UG/L	10	03-Apr-2019 14:28	
Methylene chloride	110		4.0	5.0	20	UG/L	10	03-Apr-2019 14:28	
n-Butylbenzene	5.0	U	4.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
n-Propylbenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Naphthalene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
o-Xylene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
sec-Butylbenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Styrene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
tert-Butylbenzene	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Tetrachloroethene	15		3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Toluene	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
trans-1,2-Dichloroethene	5.1	J	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
trans-1,3-Dichloropropene	5.0	U	2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Trichloroethene	5,600		20	50	100	UG/L	100	03-Apr-2019 15:40	
Trichlorofluoromethane	5.0	U	3.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Vinyl chloride	41		2.0	5.0	10	UG/L	10	03-Apr-2019 14:28	
Surr: 1,2-Dichloroethane-d4	88.0			0	81-118	%REC	10	03-Apr-2019 14:28	
Surr: 1,2-Dichloroethane-d4	88.1			0	81-118	%REC	100	03-Apr-2019 15:40	
Surr: 4-Bromofluorobenzene	99.4			0	85-114	%REC	10	03-Apr-2019 14:28	
Surr: 4-Bromofluorobenzene	97.7			0	85-114	%REC	100	03-Apr-2019 15:40	
Surr: Dibromofluoromethane	90.6			0	80-119	%REC	100	03-Apr-2019 15:40	
Surr: Dibromofluoromethane	90.2			0	80-119	%REC	10	03-Apr-2019 14:28	
Surr: Toluene-d8	106			0	89-112	%REC	100	03-Apr-2019 15:40	
Surr: Toluene-d8	105			0	89-112	%REC	10	03-Apr-2019 14:28	
SEMIVOLATILES SIM		Method:SW8270SIM						Prep:SW3510 / 03-Apr-2019 Analyst: QX	
1,4-Dioxane	16		1.0	1.0	1.0	ug/L	100	03-Apr-2019 14:32	
Surr: 2-Fluorobiphenyl	0	S		0	40-140	%REC	100	03-Apr-2019 14:32	
Surr: 4-Terphenyl-d14	0	S		0	40-140	%REC	100	03-Apr-2019 14:32	
Surr: Nitrobenzene-d5	0	S		0	40-140	%REC	100	03-Apr-2019 14:32	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP140_032719
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031508
 Lab ID:HS19031508-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
ICP-MS METALS BY SW6020A			Method:SW6020			Prep:SW3010A / 01-Apr-2019		Analyst: JHD
Aluminum	0.0643		0.00180	0.00500	0.0100	mg/L	1	03-Apr-2019 13:22
Antimony	0.00216		0.000400	0.000500	0.00200	mg/L	1	03-Apr-2019 13:22
Arsenic	0.000971	J	0.000400	0.000500	0.00200	mg/L	1	03-Apr-2019 13:22
Barium	0.223		0.00190	0.00250	0.00400	mg/L	1	03-Apr-2019 13:22
Beryllium	0.00250	U	0.000200	0.00250	0.00200	mg/L	1	03-Apr-2019 13:22
Cadmium	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	03-Apr-2019 13:22
Calcium	16.8		0.0340	0.0500	0.500	mg/L	1	03-Apr-2019 13:22
Chromium	0.00307	J	0.000400	0.000500	0.00400	mg/L	1	03-Apr-2019 13:22
Cobalt	0.00548		0.000200	0.000500	0.00500	mg/L	1	03-Apr-2019 13:22
Iron	0.317		0.0120	0.0500	0.200	mg/L	1	03-Apr-2019 13:22
Lead	0.00100	U	0.000600	0.00100	0.00200	mg/L	1	03-Apr-2019 13:22
Magnesium	13.8		0.0100	0.0500	0.200	mg/L	1	03-Apr-2019 13:22
Manganese	0.214		0.000700	0.00100	0.00500	mg/L	1	03-Apr-2019 13:22
Nickel	0.0114		0.000600	0.00100	0.00200	mg/L	1	03-Apr-2019 13:22
Potassium	1.30		0.0180	0.0500	0.200	mg/L	1	03-Apr-2019 13:22
Selenium	0.00250	U	0.00110	0.00250	0.00200	mg/L	1	03-Apr-2019 13:22
Silver	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	03-Apr-2019 13:22
Sodium	196		0.140	0.500	2.00	mg/L	10	02-Apr-2019 16:00
Thallium	0.00100	U	0.000200	0.00100	0.00200	mg/L	1	03-Apr-2019 13:22
Vanadium	0.00237	J	0.000600	0.00100	0.00500	mg/L	1	03-Apr-2019 13:22
Zinc	0.119		0.00200	0.00250	0.00400	mg/L	1	03-Apr-2019 13:22
MERCURY BY SW7470A			Method:SW7470			Prep:SW7470 / 01-Apr-2019		Analyst: FO
Mercury	0.000100	U	0.0000300	0.000100	0.000200	mg/L	1	01-Apr-2019 17:52
OIL & GREASE (HEM) BY E1664A			Method:E1664A					Analyst: KAH
Oil and Grease	0.729	J	0.610	1.00	2.00	mg/L	1	05-Apr-2019 14:45
CHEMICAL OXYGEN DEMAND BY E410.4			Method:E410.4					Analyst: AJH
Chemical Oxygen Demand	6.00	J	5.00	5.00	15.0	mg/L	1	02-Apr-2019 16:45
ANIONS BY SW9056A			Method:SW9056					Analyst: KMU
Chloride	197		2.00	5.00	5.00	mg/L	10	03-Apr-2019 05:42
Sulfate	32.0		2.00	5.00	5.00	mg/L	10	03-Apr-2019 05:42
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)			Method:NA					Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	12-Apr-2019 18:22

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: Trip Blank -ALS-020119-59
 Collection Date: 27-Mar-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19031508
 Lab ID:HS19031508-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	03-Apr-2019 16:53	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	03-Apr-2019 16:53	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	03-Apr-2019 16:53	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	03-Apr-2019 16:53	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	03-Apr-2019 16:53	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: Trip Blank -ALS-020119-59
 Collection Date: 27-Mar-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19031508
 Lab ID:HS19031508-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD		Method:SW8260							Analyst: PC
8260C									
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	03-Apr-2019 16:53	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	03-Apr-2019 16:53	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	03-Apr-2019 16:53	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Trichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 16:53	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>88.6</i>			<i>0</i>	<i>81-118</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 16:53</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>99.3</i>			<i>0</i>	<i>85-114</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 16:53</i>	
<i>Surr: Dibromofluoromethane</i>	<i>90.0</i>			<i>0</i>	<i>80-119</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 16:53</i>	
<i>Surr: Toluene-d8</i>	<i>104</i>			<i>0</i>	<i>89-112</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 16:53</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

WEIGHT LOG

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

Batch ID: 139318 **Method:** MERCURY BY SW7470A **Prep:** HG_WPR

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19031508-01	1	10 (mL)	10 (mL)	1

Batch ID: 139323 **Method:** ICP-MS METALS BY SW6020A **Prep:** 3010A

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19031508-01	1	10	10 (mL)	1

Batch ID: 139415 **Method:** SEMIVOLATILES SIM **Prep:** 3510_B_SIM

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19031508-01	1	1000	1 (mL)	0.001

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID 139318	Test Name : MERCURY BY SW7470A				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00		01 Apr 2019 11:00	01 Apr 2019 17:52	1
Batch ID 139323	Test Name : ICP-MS METALS BY SW6020A				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00		01 Apr 2019 12:30	03 Apr 2019 13:22	1
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00		01 Apr 2019 12:30	02 Apr 2019 16:00	10
Batch ID 139415	Test Name : SEMIVOLATILES SIM				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00		03 Apr 2019 10:00	03 Apr 2019 14:32	100
Batch ID R335804	Test Name : CHEMICAL OXYGEN DEMAND BY E410.4				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00			02 Apr 2019 16:45	1
Batch ID R335875	Test Name : ANIONS BY SW9056A				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00			03 Apr 2019 05:42	10
Batch ID R335882	Test Name : VOLATILES ORGANICS BY METHOD 8260C				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00			03 Apr 2019 15:40	100
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00			03 Apr 2019 14:28	10
HS19031508-02	Trip Blank -ALS-020119-59	27 Mar 2019 00:00			03 Apr 2019 16:53	1
Batch ID R336080	Test Name : OIL & GREASE (HEM) BY E1664A				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00			05 Apr 2019 14:45	1
Batch ID R336536	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)				Matrix: Water	
HS19031508-01	LH18/24-SP140_032719	27 Mar 2019 14:00			12 Apr 2019 18:22	1

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139318 (0)		Instrument: HG03		Method: MERCURY BY SW7470A						
MBLK	Sample ID: MBLK-139318	Units: mg/L		Analysis Date: 01-Apr-2019 17:33						
Client ID:	Run ID: HG03_335717	SeqNo: 5016992		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.000100	0.000200							U	
LCS	Sample ID: LCS-139318	Units: mg/L		Analysis Date: 01-Apr-2019 17:35						
Client ID:	Run ID: HG03_335717	SeqNo: 5016993		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.00509	0.000200	0.005	0	102	80 - 120				
MS	Sample ID: HS19031507-01MS	Units: mg/L		Analysis Date: 01-Apr-2019 17:38						
Client ID:	Run ID: HG03_335717	SeqNo: 5016995		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.00514	0.000200	0.005	-0.000029	103	75 - 125				
MSD	Sample ID: HS19031507-01MSD	Units: mg/L		Analysis Date: 01-Apr-2019 17:40						
Client ID:	Run ID: HG03_335717	SeqNo: 5016996		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.00518	0.000200	0.005	-0.000029	104	75 - 125	0.00514	0.775	20	

The following samples were analyzed in this batch: HS19031508-01

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
MBLK	Sample ID: MBLK-139323	Units: mg/L			Analysis Date: 03-Apr-2019 13:13					
Client ID:	Run ID: ICPMS05_335865	SeqNo: 5020436	PrepDate: 01-Apr-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.00500	0.0100								U
Antimony	0.000500	0.00200								U
Arsenic	0.000500	0.00200								U
Barium	0.00250	0.00400								U
Beryllium	0.00250	0.00200								U
Cadmium	0.000500	0.00200								U
Calcium	0.0500	0.500								U
Chromium	0.000500	0.00400								U
Cobalt	0.000500	0.00500								U
Iron	0.0500	0.200								U
Lead	0.00100	0.00200								U
Magnesium	0.0500	0.200								U
Manganese	0.00100	0.00500								U
Nickel	0.00100	0.00200								U
Potassium	0.0500	0.200								U
Selenium	0.00250	0.00200								U
Silver	0.000500	0.00200								U
Sodium	0.0500	0.200								U
Thallium	0.00100	0.00200								U
Vanadium	0.001347	0.00500								J
Zinc	0.00250	0.00400								U

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
LCS	Sample ID: LCS-139323	Units: mg/L			Analysis Date: 02-Apr-2019 15:47					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018925	PrepDate: 01-Apr-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.1079	0.0100	0.1	0	108	80 - 120				
Antimony	0.05106	0.00200	0.05	0	102	80 - 120				
Arsenic	0.05071	0.00200	0.05	0	101	80 - 120				
Barium	0.05062	0.00400	0.05	0	101	80 - 120				
Beryllium	0.04871	0.00200	0.05	0	97.4	80 - 120				
Cadmium	0.05226	0.00200	0.05	0	105	80 - 120				
Calcium	5.088	0.500	5	0	102	80 - 120				
Chromium	0.05165	0.00400	0.05	0	103	80 - 120				
Cobalt	0.05122	0.00500	0.05	0	102	80 - 120				
Iron	5.174	0.200	5	0	103	80 - 120				
Lead	0.04996	0.00200	0.05	0	99.9	80 - 120				
Magnesium	4.996	0.200	5	0	99.9	80 - 120				
Manganese	0.05141	0.00500	0.05	0	103	80 - 120				
Nickel	0.05127	0.00200	0.05	0	103	80 - 120				
Potassium	4.971	0.200	5	0	99.4	80 - 120				
Selenium	0.05018	0.00200	0.05	0	100	80 - 120				
Silver	0.05235	0.00200	0.05	0	105	80 - 120				
Sodium	5.458	0.200	5	0	109	80 - 120				
Thallium	0.04811	0.00200	0.05	0	96.2	80 - 120				
Vanadium	0.05077	0.00500	0.05	0	102	80 - 120				
Zinc	0.05274	0.00400	0.05	0	105	80 - 120				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
MS	Sample ID: HS19031584-01MS	Units: mg/L			Analysis Date: 02-Apr-2019 15:53					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018928	PrepDate: 01-Apr-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.106	0.0100	0.1	0.004897	101	80 - 120				
Antimony	0.05139	0.00200	0.05	0.000336	102	80 - 120				
Arsenic	0.05303	0.00200	0.05	0.000014	106	80 - 120				
Barium	1.01	0.00400	0.05	0.9881	44.4	80 - 120				SO
Beryllium	0.04853	0.00200	0.05	0.000012	97.0	80 - 120				
Cadmium	0.04927	0.00200	0.05	0.000018	98.5	80 - 120				
Calcium	81.44	0.500	5	74.56	138	80 - 120				SO
Chromium	0.05181	0.00400	0.05	-0.00017	104	80 - 120				
Cobalt	0.0509	0.00500	0.05	0.000005	102	80 - 120				
Iron	5.334	0.200	5	0.2078	103	80 - 120				
Lead	0.04855	0.00200	0.05	0.000157	96.8	80 - 120				
Magnesium	28.82	0.200	5	23.36	109	80 - 120				O
Manganese	0.1757	0.00500	0.05	0.1227	106	80 - 120				
Nickel	0.05074	0.00200	0.05	0.000118	101	80 - 120				
Potassium	7.73	0.200	5	2.581	103	80 - 120				
Selenium	0.0507	0.00200	0.05	0.00072	100.0	80 - 120				
Silver	0.04762	0.00200	0.05	0.000047	95.1	80 - 120				
Sodium	360.5	0.200	5	354.5	119	80 - 120				EO
Thallium	0.04538	0.00200	0.05	0.000243	90.3	80 - 120				
Vanadium	0.05283	0.00500	0.05	0.000556	105	80 - 120				
Zinc	0.1307	0.00400	0.05	0.07703	107	80 - 120				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05			Method: ICP-MS METALS BY SW6020A					
MSD	Sample ID: HS19031584-01MSD	Units: mg/L			Analysis Date: 02-Apr-2019 15:56					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018929			PrepDate: 01-Apr-2019		DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.1071	0.0100	0.1	0.004897	102	80 - 120	0.106	1.01	20	
Antimony	0.05105	0.00200	0.05	0.000336	101	80 - 120	0.05139	0.658	20	
Arsenic	0.05169	0.00200	0.05	0.000014	103	80 - 120	0.05303	2.56	20	
Barium	1.007	0.00400	0.05	0.9881	37.2	80 - 120	1.01	0.356	20	SO
Beryllium	0.04992	0.00200	0.05	0.000012	99.8	80 - 120	0.04853	2.83	20	
Cadmium	0.05007	0.00200	0.05	0.000018	100	80 - 120	0.04927	1.62	20	
Calcium	78.73	0.500	5	74.56	83.3	80 - 120	81.44	3.38	20	O
Chromium	0.05081	0.00400	0.05	-0.00017	102	80 - 120	0.05181	1.95	20	
Cobalt	0.04989	0.00500	0.05	0.000005	99.8	80 - 120	0.0509	2	20	
Iron	5.239	0.200	5	0.2078	101	80 - 120	5.334	1.78	20	
Lead	0.04939	0.00200	0.05	0.000157	98.5	80 - 120	0.04855	1.72	20	
Magnesium	28.59	0.200	5	23.36	105	80 - 120	28.82	0.802	20	O
Manganese	0.1711	0.00500	0.05	0.1227	96.7	80 - 120	0.1757	2.64	20	
Nickel	0.04944	0.00200	0.05	0.000118	98.6	80 - 120	0.05074	2.59	20	
Potassium	7.883	0.200	5	2.581	106	80 - 120	7.73	1.95	20	
Selenium	0.05281	0.00200	0.05	0.00072	104	80 - 120	0.0507	4.08	20	
Silver	0.04711	0.00200	0.05	0.000047	94.1	80 - 120	0.04762	1.07	20	
Sodium	357.6	0.200	5	354.5	60.2	80 - 120	360.5	0.816	20	SEO
Thallium	0.04656	0.00200	0.05	0.000243	92.6	80 - 120	0.04538	2.57	20	
Vanadium	0.05292	0.00500	0.05	0.000556	105	80 - 120	0.05283	0.161	20	
Zinc	0.1301	0.00400	0.05	0.07703	106	80 - 120	0.1307	0.456	20	

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05			Method: ICP-MS METALS BY SW6020A					
PDS	Sample ID: HS19031584-01PDS	Units: mg/L			Analysis Date: 02-Apr-2019 15:58					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018930		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.09986	0.0100	0.1	0.004897	95.0	75 - 125				
Antimony	0.08829	0.00200	0.1	0.000336	88.0	75 - 125				
Arsenic	0.0993	0.00200	0.1	0.000014	99.3	75 - 125				
Barium	1.062	0.00400	0.1	0.9881	74.1	75 - 125				SO
Beryllium	0.1001	0.00200	0.1	0.000012	100	75 - 125				
Cadmium	0.09637	0.00200	0.1	0.000018	96.4	75 - 125				
Calcium	80.82	0.500	10	74.56	62.6	75 - 125				SO
Chromium	0.09674	0.00400	0.1	-0.00017	96.9	75 - 125				
Cobalt	0.09603	0.00500	0.1	0.000005	96.0	75 - 125				
Iron	9.848	0.200	10	0.2078	96.4	75 - 125				
Lead	0.0916	0.00200	0.1	0.000157	91.4	75 - 125				
Magnesium	31.73	0.200	10	23.36	83.7	75 - 125				
Manganese	0.2125	0.00500	0.1	0.1227	89.8	75 - 125				
Nickel	0.09352	0.00200	0.1	0.000118	93.4	75 - 125				
Potassium	12.44	0.200	10	2.581	98.6	75 - 125				
Selenium	0.09928	0.00200	0.1	0.00072	98.6	75 - 125				
Silver	0.08785	0.00200	0.1	0.000047	87.8	75 - 125				
Thallium	0.08932	0.00200	0.1	0.000243	89.1	75 - 125				
Vanadium	0.09766	0.00500	0.1	0.000556	97.1	75 - 125				
Zinc	0.1737	0.00400	0.1	0.07703	96.7	75 - 125				
PDS	Sample ID: HS19031584-01PDS	Units: mg/L			Analysis Date: 03-Apr-2019 13:20					
Client ID:	Run ID: ICPMS05_335865	SeqNo: 5020439		PrepDate: 01-Apr-2019		DF: 10				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Sodium	430.6	2.00	100	334.6	96.0	75 - 125				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
SD	Sample ID: HS19031584-01SD	Units: mg/L			Analysis Date: 02-Apr-2019 15:51					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018927	PrepDate: 01-Apr-2019	DF: 5						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	%D Limit	Qual
Aluminum	0.009832	0.0500					0.004897	0	10	J
Antimony	0.00250	0.0100					0.000336	0	10	U
Arsenic	0.00250	0.0100					0.000014	0	10	U
Barium	0.9199	0.0200					0.9881	6.9	10	
Beryllium	0.0125	0.0100					0.000012	0	10	U
Cadmium	0.00250	0.0100					0.000018	0	10	U
Calcium	70.09	2.50					74.56	5.99	10	
Chromium	0.00250	0.0200					-0.00017	0	10	U
Cobalt	0.00250	0.0250					0.000005	0	10	U
Iron	0.2043	1.00					0.2078	0	10	J
Lead	0.00500	0.0100					0.000157	0	10	U
Magnesium	21.3	1.00					23.36	8.79	10	
Manganese	0.1188	0.0250					0.1227	3.19	10	
Nickel	0.00500	0.0100					0.000118	0	10	U
Potassium	2.471	1.00					2.581	4.24	10	
Selenium	0.0125	0.0100					0.00072	0	10	U
Silver	0.00250	0.0100					0.000047	0	10	U
Thallium	0.00500	0.0100					0.000243	0	10	U
Vanadium	0.005215	0.0250					0.000556	0	10	J
Zinc	0.07308	0.0200					0.07703	5.12	10	
SD	Sample ID: HS19031584-01SD	Units: mg/L			Analysis Date: 03-Apr-2019 13:18					
Client ID:	Run ID: ICPMS05_335865	SeqNo: 5020438	PrepDate: 01-Apr-2019	DF: 50						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	%D Limit	Qual
Sodium	340.7	10.0					334.6	1.83	10	

The following samples were analyzed in this batch: HS19031508-01

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: 139415 (0)		Instrument: SV-5		Method: SEMIVOLATILES SIM						
MBLK	Sample ID: MBLK-139415	Units: ug/L			Analysis Date: 03-Apr-2019 12:47					
Client ID:	Run ID: SV-5_335930	SeqNo: 5021779		PrepDate: 03-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
1,4-Dioxane	0.010	0.010							U	
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.09201</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>115</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.07352</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>91.9</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.08203</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>103</i>	<i>40 - 140</i>				
LCS	Sample ID: LCS-139415	Units: ug/L			Analysis Date: 03-Apr-2019 13:08					
Client ID:	Run ID: SV-5_335930	SeqNo: 5021780		PrepDate: 03-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
1,4-Dioxane	0.07995	0.010	0.08	0	99.9	40 - 140				
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.08927</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>112</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.07072</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>88.4</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.07169</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>89.6</i>	<i>40 - 140</i>				
LCSD	Sample ID: LCSD-139415	Units: ug/L			Analysis Date: 03-Apr-2019 13:29					
Client ID:	Run ID: SV-5_335930	SeqNo: 5021781		PrepDate: 03-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
1,4-Dioxane	0.08058	0.010	0.08	0	101	40 - 140	0.07995	0.787	20	
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.08169</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>102</i>	<i>40 - 140</i>	<i>0.08927</i>	<i>8.86</i>	<i>20</i>	
<i>Surr: 4-Terphenyl-d14</i>	<i>0.07017</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>87.7</i>	<i>40 - 140</i>	<i>0.07072</i>	<i>0.788</i>	<i>20</i>	
<i>Surr: Nitrobenzene-d5</i>	<i>0.07503</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>93.8</i>	<i>40 - 140</i>	<i>0.07169</i>	<i>4.56</i>	<i>20</i>	
The following samples were analyzed in this batch: HS19031508-01										

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 12:28					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020560	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	0.50	1.0								U
1,1,1-Trichloroethane	0.50	1.0								U
1,1,2,2-Tetrachloroethane	0.50	1.0								U
1,1,2-Trichloroethane	0.50	1.0								U
1,1-Dichloroethane	0.50	1.0								U
1,1-Dichloroethene	0.50	1.0								U
1,1-Dichloropropene	0.50	1.0								U
1,2,3-Trichlorobenzene	0.50	1.0								U
1,2,3-Trichloropropane	0.50	1.0								U
1,2,4-Trichlorobenzene	0.50	1.0								U
1,2,4-Trimethylbenzene	0.50	1.0								U
1,2-Dibromo-3-chloropropane	0.50	1.0								U
1,2-Dibromoethane	0.50	1.0								U
1,2-Dichlorobenzene	0.50	1.0								U
1,2-Dichloroethane	0.50	1.0								U
1,2-Dichloropropane	0.50	1.0								U
1,3,5-Trimethylbenzene	0.50	1.0								U
1,3-Dichlorobenzene	0.50	1.0								U
1,3-Dichloropropane	0.50	1.0								U
1,4-Dichlorobenzene	0.50	1.0								U
2,2-Dichloropropane	0.50	1.0								U
2-Butanone	1.0	2.0								U
2-Chlorotoluene	0.50	1.0								U
2-Hexanone	1.0	2.0								U
4-Chlorotoluene	0.50	1.0								U
4-Isopropyltoluene	0.50	1.0								U
4-Methyl-2-pentanone	1.0	2.0								U
Acetone	2.0	2.0								U
Benzene	0.50	1.0								U
Bromobenzene	0.50	1.0								U
Bromochloromethane	0.50	1.0								U
Bromodichloromethane	0.50	1.0								U
Bromoform	0.50	1.0								U
Bromomethane	0.50	1.0								U

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 12:28					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020560	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	43.62	1.0	50	0	87.2	81 - 118				
Surr: 4-Bromofluorobenzene	49.43	1.0	50	0	98.9	85 - 114				
Surr: Dibromofluoromethane	45.1	1.0	50	0	90.2	80 - 119				
Surr: Toluene-d8	53	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 11:40					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020559	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	23.34	1.0	20	0	117	78 - 124				
1,1,1-Trichloroethane	19.93	1.0	20	0	99.7	74 - 131				
1,1,2,2-Tetrachloroethane	20.44	1.0	20	0	102	71 - 121				
1,1,2-Trichloroethane	22.06	1.0	20	0	110	80 - 119				
1,1-Dichloroethane	19.85	1.0	20	0	99.2	77 - 125				
1,1-Dichloroethene	19.49	1.0	20	0	97.4	71 - 131				
1,1-Dichloropropene	19.9	1.0	20	0	99.5	78 - 125				
1,2,3-Trichlorobenzene	21.21	1.0	20	0	106	69 - 129				
1,2,3-Trichloropropane	21.32	1.0	20	0	107	73 - 122				
1,2,4-Trichlorobenzene	21.68	1.0	20	0	108	69 - 130				
1,2,4-Trimethylbenzene	23.62	1.0	20	0	118	76 - 124				
1,2-Dibromo-3-chloropropane	21.63	1.0	20	0	108	62 - 128				
1,2-Dibromoethane	22.93	1.0	20	0	115	77 - 121				
1,2-Dichlorobenzene	21.63	1.0	20	0	108	80 - 119				
1,2-Dichloroethane	21.35	1.0	20	0	107	73 - 128				
1,2-Dichloropropane	20.9	1.0	20	0	104	78 - 122				
1,3,5-Trimethylbenzene	22.82	1.0	20	0	114	75 - 124				
1,3-Dichlorobenzene	21.78	1.0	20	0	109	80 - 119				
1,3-Dichloropropane	22.02	1.0	20	0	110	80 - 119				
1,4-Dichlorobenzene	21.6	1.0	20	0	108	79 - 118				
2,2-Dichloropropane	20.72	1.0	20	0	104	60 - 139				
2-Butanone	39.59	2.0	40	0	99.0	56 - 143				
2-Chlorotoluene	21.51	1.0	20	0	108	79 - 122				
2-Hexanone	42.13	2.0	40	0	105	57 - 139				
4-Chlorotoluene	21.85	1.0	20	0	109	78 - 122				
4-Isopropyltoluene	22.25	1.0	20	0	111	77 - 127				
4-Methyl-2-pentanone	41.02	2.0	40	0	103	67 - 130				
Acetone	36.89	2.0	40	0	92.2	39 - 160				
Benzene	21.72	1.0	20	0	109	79 - 120				
Bromobenzene	22.97	1.0	20	0	115	80 - 120				
Bromochloromethane	20.97	1.0	20	0	105	78 - 123				
Bromodichloromethane	22.16	1.0	20	0	111	79 - 125				
Bromoform	23.37	1.0	20	0	117	66 - 130				
Bromomethane	24.03	1.0	20	0	120	53 - 141				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 11:40					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020559	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	37.4	2.0	40	0	93.5	64 - 133				
Carbon tetrachloride	22.65	1.0	20	0	113	72 - 136				
Chlorobenzene	21.88	1.0	20	0	109	82 - 118				
Chloroethane	18.95	1.0	20	0	94.7	60 - 138				
Chloroform	20.52	1.0	20	0	103	79 - 124				
Chloromethane	19.74	1.0	20	0	98.7	50 - 139				
cis-1,2-Dichloroethene	20.43	1.0	20	0	102	78 - 123				
cis-1,3-Dichloropropene	20.84	1.0	20	0	104	75 - 124				
Dibromochloromethane	22.56	1.0	20	0	113	74 - 126				
Dibromomethane	21.6	1.0	20	0	108	79 - 123				
Dichlorodifluoromethane	21.85	1.0	20	0	109	32 - 152				
Ethylbenzene	23.41	1.0	20	0	117	79 - 121				
Hexachlorobutadiene	23.54	1.0	20	0	118	66 - 134				
Isopropylbenzene	22.97	1.0	20	0	115	72 - 131				
m,p-Xylene	45.47	2.0	40	0	114	80 - 121				
Methylene chloride	20.82	2.0	20	0	104	74 - 124				
Naphthalene	21.06	1.0	20	0	105	61 - 128				
n-Butylbenzene	22.89	1.0	20	0	114	75 - 128				
n-Propylbenzene	21.9	1.0	20	0	109	76 - 126				
o-Xylene	22.22	1.0	20	0	111	78 - 122				
sec-Butylbenzene	21.56	1.0	20	0	108	77 - 126				
Styrene	23.58	1.0	20	0	118	78 - 123				
tert-Butylbenzene	21.9	1.0	20	0	109	78 - 124				
Tetrachloroethene	22.75	1.0	20	0	114	74 - 129				
Toluene	22.69	1.0	20	0	113	80 - 121				
trans-1,2-Dichloroethene	20.31	1.0	20	0	102	75 - 124				
trans-1,3-Dichloropropene	22.21	1.0	20	0	111	73 - 127				
Trichloroethene	20.76	1.0	20	0	104	79 - 123				
Trichlorofluoromethane	19.87	1.0	20	0	99.3	65 - 141				
Vinyl chloride	18.34	1.0	20	0	91.7	58 - 137				
Surr: 1,2-Dichloroethane-d4	44.85	1.0	50	0	89.7	81 - 118				
Surr: 4-Bromofluorobenzene	49.44	1.0	50	0	98.9	85 - 114				
Surr: Dibromofluoromethane	46.66	1.0	50	0	93.3	80 - 119				
Surr: Toluene-d8	52.9	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19040036-36MS	Units: UG/L			Analysis Date: 03-Apr-2019 14:52					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020790	PrepDate:	DF: 1000						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	24370	1000	20000	0	122	78 - 124				
1,1,1-Trichloroethane	22280	1000	20000	0	111	74 - 131				
1,1,2,2-Tetrachloroethane	22750	1000	20000	0	114	71 - 121				
1,1,2-Trichloroethane	23590	1000	20000	0	118	80 - 119				
1,1-Dichloroethane	21090	1000	20000	0	105	77 - 125				
1,1-Dichloroethene	22220	1000	20000	0	111	71 - 131				
1,1-Dichloropropene	22920	1000	20000	0	115	78 - 125				
1,2,3-Trichlorobenzene	21690	1000	20000	0	108	69 - 129				
1,2,3-Trichloropropane	22780	1000	20000	0	114	73 - 122				
1,2,4-Trichlorobenzene	22700	1000	20000	0	114	69 - 130				
1,2,4-Trimethylbenzene	24950	1000	20000	0	125	76 - 124				S
1,2-Dibromo-3-chloropropane	23950	1000	20000	0	120	62 - 128				
1,2-Dibromoethane	23610	1000	20000	0	118	77 - 121				
1,2-Dichlorobenzene	23700	1000	20000	0	118	80 - 119				
1,2-Dichloroethane	23200	1000	20000	0	116	73 - 128				
1,2-Dichloropropane	21950	1000	20000	0	110	78 - 122				
1,3,5-Trimethylbenzene	24840	1000	20000	0	124	75 - 124				S
1,3-Dichlorobenzene	24090	1000	20000	0	120	80 - 119				S
1,3-Dichloropropane	23000	1000	20000	0	115	80 - 119				
1,4-Dichlorobenzene	23350	1000	20000	0	117	79 - 118				
2,2-Dichloropropane	22210	1000	20000	0	111	60 - 139				
2-Butanone	41410	2000	40000	0	104	56 - 143				
2-Chlorotoluene	23870	1000	20000	0	119	79 - 122				
2-Hexanone	44400	2000	40000	0	111	57 - 139				
4-Chlorotoluene	23820	1000	20000	0	119	78 - 122				
4-Isopropyltoluene	25150	1000	20000	0	126	77 - 127				
4-Methyl-2-pentanone	44310	2000	40000	0	111	67 - 130				
Acetone	39450	2000	40000	0	98.6	39 - 160				
Benzene	163700	1000	20000	136400	137	79 - 120				SO
Bromobenzene	24010	1000	20000	0	120	80 - 120				S
Bromochloromethane	20610	1000	20000	0	103	78 - 123				
Bromodichloromethane	22820	1000	20000	0	114	79 - 125				
Bromoform	23650	1000	20000	0	118	66 - 130				
Bromomethane	24600	1000	20000	0	123	53 - 141				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19040036-36MS	Units: UG/L			Analysis Date: 03-Apr-2019 14:52					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020790	PrepDate:	DF: 1000						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	43460	2000	40000	0	109	64 - 133				
Carbon tetrachloride	25400	1000	20000	0	127	72 - 136				
Chlorobenzene	23430	1000	20000	0	117	82 - 118				
Chloroethane	20830	1000	20000	0	104	60 - 138				
Chloroform	21220	1000	20000	0	106	79 - 124				
Chloromethane	23190	1000	20000	0	116	50 - 139				
cis-1,2-Dichloroethene	21440	1000	20000	0	107	78 - 123				
cis-1,3-Dichloropropene	22680	1000	20000	0	113	75 - 124				
Dibromochloromethane	24100	1000	20000	0	120	74 - 126				
Dibromomethane	22210	1000	20000	0	111	79 - 123				
Dichlorodifluoromethane	22700	1000	20000	0	113	32 - 152				
Ethylbenzene	25690	1000	20000	1292	122	79 - 121				S
Hexachlorobutadiene	26030	1000	20000	0	130	66 - 134				
Isopropylbenzene	24720	1000	20000	0	124	72 - 131				
m,p-Xylene	53610	2000	40000	4701	122	80 - 121				S
Methylene chloride	22110	2000	20000	0	111	74 - 124				
Naphthalene	21950	1000	20000	0	110	61 - 128				
n-Butylbenzene	25170	1000	20000	0	126	75 - 128				
n-Propylbenzene	25030	1000	20000	0	125	76 - 126				
o-Xylene	26700	1000	20000	2345	122	78 - 122				
sec-Butylbenzene	24850	1000	20000	0	124	77 - 126				
Styrene	26250	1000	20000	1811	122	78 - 123				
tert-Butylbenzene	24760	1000	20000	0	124	78 - 124				
Tetrachloroethene	25340	1000	20000	0	127	74 - 129				
Toluene	52760	1000	20000	28020	124	80 - 121				S
trans-1,2-Dichloroethene	22170	1000	20000	0	111	75 - 124				
trans-1,3-Dichloropropene	22600	1000	20000	0	113	73 - 127				
Trichloroethene	24510	1000	20000	0	123	79 - 123				
Trichlorofluoromethane	23470	1000	20000	0	117	65 - 141				
Vinyl chloride	20510	1000	20000	0	103	58 - 137				
Surr: 1,2-Dichloroethane-d4	43910	1000	50000	0	87.8	81 - 118				
Surr: 4-Bromofluorobenzene	49050	1000	50000	0	98.1	85 - 114				
Surr: Dibromofluoromethane	45590	1000	50000	0	91.2	80 - 119				
Surr: Toluene-d8	52160	1000	50000	0	104	89 - 112				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19040036-36MSD	Units: UG/L			Analysis Date: 03-Apr-2019 15:16					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020791	PrepDate:	DF: 1000						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	24320	1000	20000	0	122	78 - 124	24370	0.228	20	
1,1,1-Trichloroethane	21690	1000	20000	0	108	74 - 131	22280	2.69	20	
1,1,2,2-Tetrachloroethane	22810	1000	20000	0	114	71 - 121	22750	0.276	20	
1,1,2-Trichloroethane	23280	1000	20000	0	116	80 - 119	23590	1.3	20	
1,1-Dichloroethane	20610	1000	20000	0	103	77 - 125	21090	2.28	20	
1,1-Dichloroethene	21230	1000	20000	0	106	71 - 131	22220	4.59	20	
1,1-Dichloropropene	22490	1000	20000	0	112	78 - 125	22920	1.87	20	
1,2,3-Trichlorobenzene	22690	1000	20000	0	113	69 - 129	21690	4.52	20	
1,2,3-Trichloropropane	23960	1000	20000	0	120	73 - 122	22780	5.05	20	
1,2,4-Trichlorobenzene	22750	1000	20000	0	114	69 - 130	22700	0.208	20	
1,2,4-Trimethylbenzene	24690	1000	20000	0	123	76 - 124	24950	1.03	20	
1,2-Dibromo-3-chloropropane	24840	1000	20000	0	124	62 - 128	23950	3.68	20	
1,2-Dibromoethane	23750	1000	20000	0	119	77 - 121	23610	0.569	20	
1,2-Dichlorobenzene	23560	1000	20000	0	118	80 - 119	23700	0.565	20	
1,2-Dichloroethane	23300	1000	20000	0	116	73 - 128	23200	0.421	20	
1,2-Dichloropropane	21950	1000	20000	0	110	78 - 122	21950	0.0278	20	
1,3,5-Trimethylbenzene	24400	1000	20000	0	122	75 - 124	24840	1.78	20	
1,3-Dichlorobenzene	23770	1000	20000	0	119	80 - 119	24090	1.35	20	
1,3-Dichloropropane	23040	1000	20000	0	115	80 - 119	23000	0.159	20	
1,4-Dichlorobenzene	23460	1000	20000	0	117	79 - 118	23350	0.488	20	
2,2-Dichloropropane	21530	1000	20000	0	108	60 - 139	22210	3.12	20	
2-Butanone	42690	2000	40000	0	107	56 - 143	41410	3.06	20	
2-Chlorotoluene	23450	1000	20000	0	117	79 - 122	23870	1.75	20	
2-Hexanone	45910	2000	40000	0	115	57 - 139	44400	3.33	20	
4-Chlorotoluene	23550	1000	20000	0	118	78 - 122	23820	1.15	20	
4-Isopropyltoluene	24580	1000	20000	0	123	77 - 127	25150	2.31	20	
4-Methyl-2-pentanone	45920	2000	40000	0	115	67 - 130	44310	3.58	20	
Acetone	43390	2000	40000	0	108	39 - 160	39450	9.5	20	
Benzene	161100	1000	20000	136400	123	79 - 120	163700	1.64	20	SO
Bromobenzene	23980	1000	20000	0	120	80 - 120	24010	0.139	20	
Bromochloromethane	20980	1000	20000	0	105	78 - 123	20610	1.77	20	
Bromodichloromethane	22510	1000	20000	0	113	79 - 125	22820	1.39	20	
Bromoform	24000	1000	20000	0	120	66 - 130	23650	1.45	20	
Bromomethane	22690	1000	20000	0	113	53 - 141	24600	8.06	20	

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19040036-36MSD	Units: UG/L			Analysis Date: 03-Apr-2019 15:16					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020791		PrepDate:		DF: 1000				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	41860	2000	40000	0	105	64 - 133	43460	3.75	20	
Carbon tetrachloride	24790	1000	20000	0	124	72 - 136	25400	2.42	20	
Chlorobenzene	23190	1000	20000	0	116	82 - 118	23430	1.07	20	
Chloroethane	19980	1000	20000	0	99.9	60 - 138	20830	4.15	20	
Chloroform	20990	1000	20000	0	105	79 - 124	21220	1.06	20	
Chloromethane	21200	1000	20000	0	106	50 - 139	23190	8.96	20	
cis-1,2-Dichloroethene	20790	1000	20000	0	104	78 - 123	21440	3.08	20	
cis-1,3-Dichloropropene	22920	1000	20000	0	115	75 - 124	22680	1.02	20	
Dibromochloromethane	24120	1000	20000	0	121	74 - 126	24100	0.0892	20	
Dibromomethane	22680	1000	20000	0	113	79 - 123	22210	2.1	20	
Dichlorodifluoromethane	21730	1000	20000	0	109	32 - 152	22700	4.35	20	
Ethylbenzene	25210	1000	20000	1292	120	79 - 121	25690	1.92	20	
Hexachlorobutadiene	26760	1000	20000	0	134	66 - 134	26030	2.75	20	
Isopropylbenzene	24210	1000	20000	0	121	72 - 131	24720	2.07	20	
m,p-Xylene	52410	2000	40000	4701	119	80 - 121	53610	2.27	20	
Methylene chloride	21630	2000	20000	0	108	74 - 124	22110	2.19	20	
Naphthalene	23380	1000	20000	0	117	61 - 128	21950	6.3	20	
n-Butylbenzene	24790	1000	20000	0	124	75 - 128	25170	1.55	20	
n-Propylbenzene	24480	1000	20000	0	122	76 - 126	25030	2.2	20	
o-Xylene	26170	1000	20000	2345	119	78 - 122	26700	2.01	20	
sec-Butylbenzene	24340	1000	20000	0	122	77 - 126	24850	2.07	20	
Styrene	26210	1000	20000	1811	122	78 - 123	26250	0.147	20	
tert-Butylbenzene	24250	1000	20000	0	121	78 - 124	24760	2.09	20	
Tetrachloroethene	24480	1000	20000	0	122	74 - 129	25340	3.45	20	
Toluene	51510	1000	20000	28020	117	80 - 121	52760	2.38	20	
trans-1,2-Dichloroethene	21410	1000	20000	0	107	75 - 124	22170	3.5	20	
trans-1,3-Dichloropropene	23180	1000	20000	0	116	73 - 127	22600	2.52	20	
Trichloroethene	23240	1000	20000	0	116	79 - 123	24510	5.32	20	
Trichlorofluoromethane	23010	1000	20000	0	115	65 - 141	23470	1.98	20	
Vinyl chloride	19890	1000	20000	0	99.5	58 - 137	20510	3.06	20	
Surr: 1,2-Dichloroethane-d4	43970	1000	50000	0	87.9	81 - 118	43910	0.129	20	
Surr: 4-Bromofluorobenzene	49970	1000	50000	0	99.9	85 - 114	49050	1.86	20	
Surr: Dibromofluoromethane	45550	1000	50000	0	91.1	80 - 119	45590	0.0906	20	
Surr: Toluene-d8	51790	1000	50000	0	104	89 - 112	52160	0.709	20	

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335882 (0)	Instrument: VOA6	Method: VOLATILES ORGANICS BY METHOD 8260C
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The following samples were analyzed in this batch:

HS19031508-01	HS19031508-02
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ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID:	R335804 (0)	Instrument:	WetChem_HS	Method:	CHEMICAL OXYGEN DEMAND BY E410.4					
MBLK	Sample ID: MBLK-R335804	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018767	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	5.00	15.0								U
LCS	Sample ID: LCS-R335804	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018766	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	97	15.0	100	0	97.0	85 - 115				
MS	Sample ID: HS19040002-01MS	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5020059	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	60	15.0	50	7	106	80 - 120				
MS	Sample ID: HS19031461-01MS	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018770	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	94	15.0	50	37	114	80 - 120				
DUP	Sample ID: HS19040002-01DUP	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5020060	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	8	15.0					7	0	20	J
DUP	Sample ID: HS19031461-01DUP	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018771	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	37	15.0					37	0	20	

The following samples were analyzed in this batch: HS19031508-01

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R335875 (0)		Instrument: ICS2100		Method: ANIONS BY SW9056A						
MBLK	Sample ID: WBLKW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:16					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020414		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	0.500	0.500							U	
Sulfate	0.500	0.500							U	
LCS	Sample ID: WLCSW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:31					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020415		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	20.22	0.500	20	0	101	80 - 120				
Sulfate	19.98	0.500	20	0	99.9	80 - 120				
LCSD	Sample ID: WLCSDW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:46					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020416		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	18.95	0.500	20	0	94.8	80 - 120	20.22	6.51	20	
Sulfate	18.67	0.500	20	0	93.3	80 - 120	19.98	6.79	20	
MS	Sample ID: HS19031013-07MS	Units: mg/L			Analysis Date: 03-Apr-2019 04:28					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020427		PrepDate:			DF: 50			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	1535	25.0	500	1090	89.1	80 - 120				
Sulfate	1565	25.0	500	1119	89.2	80 - 120				
MSD	Sample ID: HS19031013-07MSD	Units: mg/L			Analysis Date: 03-Apr-2019 04:43					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020428		PrepDate:			DF: 50			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	1605	25.0	500	1090	103	80 - 120	1535	4.46	20	
Sulfate	1638	25.0	500	1119	104	80 - 120	1565	4.57	20	

The following samples were analyzed in this batch:

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031508

QC BATCH REPORT NEW

Batch ID: R336080 (0)		Instrument: Balance1		Method: OIL & GREASE (HEM) BY E1664A						
MBLK	Sample ID: WBLKW-040519	Units: mg/L		Analysis Date: 05-Apr-2019 14:45						
Client ID:	Run ID: Balance1_336080	SeqNo: 5024526		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	1.00	2.00							U	
LCS	Sample ID: WLCSW-040519	Units: mg/L		Analysis Date: 05-Apr-2019 14:45						
Client ID:	Run ID: Balance1_336080	SeqNo: 5024528		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	38.8	2.00	40	0	97.0	78 - 114				
LCSD	Sample ID: WLCSDW-040519	Units: mg/L		Analysis Date: 05-Apr-2019 14:45						
Client ID:	Run ID: Balance1_336080	SeqNo: 5024527		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	38.8	2.00	40	0	97.0	78 - 114	38.8	0	18	
MS	Sample ID: HS19031453-01MS	Units: mg/L		Analysis Date: 05-Apr-2019 14:45						
Client ID:	Run ID: Balance1_336080	SeqNo: 5024506		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	39.58	2.00	40	1.25	95.8	78 - 114				

The following samples were analyzed in this batch: HS19031508-01

ALS Houston, US

Date: 12-Apr-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19031508	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

Unit Reported	Description
mg/L	Milligrams per Liter

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

ALS Houston, US

Date: 12-Apr-19

Client: Bhat Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031508

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19031508-01	LH18/24-SP140_032719	Login	3/29/2019 10:12:37 AM	NDR	WET094
HS19031508-01	LH18/24-SP140_032719	Login	3/29/2019 10:12:37 AM	NDR	WET094
HS19031508-01	LH18/24-SP140_032719	Login	3/29/2019 10:12:37 AM	NDR	Sub

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19031508

Date/Time Received: **28-Mar-2019 08:30**
 Received by: **PMG**

Checklist completed by: Nilesh D. Ranchod 29-Mar-2019
 eSignature Date

Reviewed by: RJ Modashia 29-Mar-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.5c UC/C IR25
 Cooler(s)/Kit(s): 44581
 Date/Time sample(s) sent to storage: 03/28/2019 18:00

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

HS19031508

Bhate Environmental Associates, Inc.
118/24 Longhorn GW Treatment Plant Weekly Sample

Name Of Lab Shipping To: ALS 10450 Stancliff Rd., Suite 210 Houston, Tx. 77099 ATTN: RJ Modashia


CHAIN OF CUSTODY

Project: BHATE LONGHORN ARMY AMMN. PLANT (LHAAP) GROUNDWATER TREATMENT PLANT (GWTP) KARNACK, TEXAS			Project No. NWO1312.0150.0 16.0001		Analyses										
Job: GROUNDWATER TREATMENT PLANT QUARTERLY INFLUENT SAMPLES					MS / MSD	No. OF CONTAINERS	ROD Volatiles	Total Metals	Oil & Grease	Chemical Oxygen Demand	Chloride & Sulfate	1, 4 - DIOXANE	Perchlorate	Remarks (Preservatives, etc.)	Lab I.D.#
Prepared By: Scott Beesinger			P. O. Number												
Field Sample I.D.	Sample Matrix	Date / Time													
LH18/24-SP140_032719	Water	03/27/19 / 14:00	4	3		1								HCL	
LH18/24-SP140_032719	Water	03/27/19 / 14:00	1		1									HNO3	
LH18/24-SP140_032719	Water	03/27/19 / 14:00	2						1	1				NONE	
LH18/24-SP140_032719	Water	03/27/19 / 14:00	1					1						H2SO4	
LH18/24-SP140_032719	Water	03/27/19 / 14:00	1								1			NONE	
Trip Blank	Water	03/27/19	2	2										HCL	
Additional Remarks: STANDARD TURN AROUND TIME															
Relinquished By: <i>Scott Beesinger</i>		Date: 03/27/19	Time: 14:30	Received By: <i>[Signature]</i>		Date: 3/28/19	Time: 08:30	Relinquished By:		Date:	Time:	Received By:		Date:	Time:



Received At Lab By:										For Lab Use Only				
			Date	Time	Albill No.	Opened By:			Date	Time	Temp of Container	Seal No.	Condition	
Remarks:														

44581
1-50
4-25
6/10/00

 ALS Environmental 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CUSTODY SEAL		Seal Broken By: <i>SM</i>
	Date: <i>3/27/19</i>	Time: <i>1130</i>	Date: <i>03/28/19</i>
	Name: <i>Scott Deisinger</i>		Company: <i>DAVE</i>

44581 MAR 28 2019



Must Deliver Next Business Day
Time and Temperature Sensitive!

44581

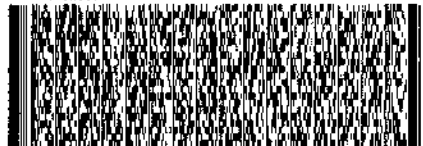
ORIGIN ID: SGRA (303) 597-2450
SCOTT BEES INER
BHAITE ENVIRONMENTAL ASSOCIATES
1203-B EAST GRAND AVE. PMB202
MARSHALL, TX 75670
UNITED STATES US

SHIP DATE: 06MAR19
ACTWT: 1.00 LB MAN
CRO: 300130/CAF3211
DIM: 26x14x14 IN

TO **CLIENT SERVICES**
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON TX 77099

(281) 530-5666
REF: LHAAP-58-80 64113-RJ

RMA: 01191100

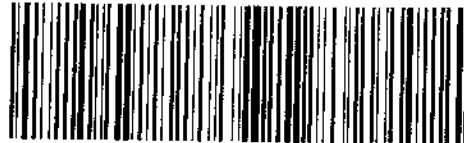


FedEx
TRK#
0221 4809 7831 3421

THU - 28 MAR 10:30A
PRIORITY OVERNIGHT

AB SGRA

77099
TX-US
IAH



13D 162785 27MAR19 666A 553C1J4685/RCSA



Case Narrative

Method: 6850
Analysis: Perchlorate
Analysis SOP: LC-MS-CLO4
ALS WO ID(s): 1909152; 1909153; 1909154;
1909947; 1909949

Client: ALS Laboratories (Houston, TX)
Matrix: Water
ELMS Batch (HBN): 2233 (236356)

General Set Information: There were ten field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Method QC data: The method blank (LMB 647198) was less than 1/2 the CRDL. The recovery for the LCS (647199) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1909152001 (Client ID's: LH18/24-SP650_032719_BIX). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.µg/L. The MS/MSD – 647200/01 failed QC acceptance criteria for percent recoveries, biased high. This is due to the fact that the unspiked sample result of 2.574µg/L was not subtracted from the MS/MSD results. The Matrix Spike and Matrix Spike duplicate is reported for the clients' information only. The sample matrix may be inappropriate for the method selected. The MS/MSD relative percent difference (RPD) was within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in µg/L. Results were calculated in µg/L by the equation $(A) \times (B)$,

where: A = Analyte concentration from the standard curve (µg/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 647196) is reported from the analysis of the Laboratory Control Sample (LCS – 647199) at a level of 4.0µg/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch April 11, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: April 11, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1909153**

Project ID: HS19031508

Purchase Order: HS19031508

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18-24-SP140_032719	1909153001	03/27/19	03/30/19	



ANALYTICAL REPORT

Workorder: 34-1909153

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18-24-SP140_032719	Sampling Site: NA	Collected: 03/27/2019				
Lab ID: 1909153001	Media: 125 mL Nalgene	Received: 03/30/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2233 (HBN: 236356) Analyzed: 04/10/2019 11:35	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	6500	1000	2000	4000	1000	

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 236356)

Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 04/11/2019 09:27	/S/ Stephen Brose 04/11/2019 15:00

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1909153

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935838

Analysis Information

Workorder: 1909153

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2233 (HBN: 236356)
Analyzed By: Thomas Bosch

Blank

LMB: 647198 Analyzed: 04/10/2019 10:42 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 647199 Analyzed: 04/10/2019 10:15 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	3.68	4.00	91.9	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1909152001 Analyzed: 04/10/2019 10:55 Dilution: 1 Units: ug/L		MS: 647200 Analyzed: 04/10/2019 11:08 Dilution: 1 Units: ug/L				MSD: 647201 Analyzed: 04/10/2019 11:22 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	2.60	6.18	4 #	154	78.8 123.8	6.36 #	159	2.92	0.0 20.0

Comments

Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 04/11/2019 13:48	/S/ Stephen Brose 04/11/2019 15:00

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

18698/#2

COC ID: 11013

1909153

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19031508
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19031508-01	LH18/24-SP140_032719	Water	27 Mar 2019 14:00
SUB_Perch-6850			05 Apr 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. Miskany
Received By: Meredith J. Lawal
Cooler ID(s): 9276

Date/Time: 3/29/19 18:00
Date/Time: 3/30/19 8:58
Temperature(s): 2°

RIGHT SOLUTIONS | RIGHT PARTNER



ALS Environmental CHAIN-OF-CUSTODY

00935840

Project / Job / Task: HS19031508		Split:		Workorder ID: 1909153	Level: ENV_LVL4		Requested Analysis								
Client: ALS Environmental (Houston)		Account: 8101				Type: 125Poly									
Comments:						Preservatives									
						Containers									
Item	Collect Date/Time	Sample ID	Lab ID	QC	Matrix	ID(s)	Count								
1	03/27/2019 14:00	LH18-24-SP140_032719	1909153001		Water	A	1								
2															
3															
4															
5															
6															
7															
8															
9															
10															
44															

EP A 6850, DOD QSM

ORIGINAL FIELD SAMPLE CHAIN-OF-CUSTODY				SAMPLE PREPARATION / ANALYSIS CHAIN-OF-CUSTODY			
Relinquished By: (Signature)		Received By: (Signature)		Relinquished By: (Signature)		Received By: (Signature)	
Date / Time		Date / Time		Date / Time		Date / Time	
Reason for Transfer / Storage Location		Reason for Transfer / Storage Location		Reason for Transfer / Storage Location		Reason for Transfer / Storage Location	
Sample LogIn		Sample LogIn		Sample LogIn		Sample LogIn	
ALS Sample Receiving		ALS Sample Receiving		ALS Sample Receiving		ALS Sample Receiving	
Signature		Signature		Signature		Signature	
Date / Time		Date / Time		Date / Time		Date / Time	
Lab Notebook No.:		Lab Notebook No.:		Lab Notebook No.:		Lab Notebook No.:	
Prepared / Analyzed by:		Prepared / Analyzed by:		Prepared / Analyzed by:		Prepared / Analyzed by:	
Date / Time		Date / Time		Date / Time		Date / Time	

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: ALS HOLDING Project/Task/Site: 1909153
 Date/Time of Receipt: 3/20/19 858 Number of Coolers Received: 1

Condition of Coolers:	Acceptable/Unacceptable	Temperature Control:	Present/Not Included
Cooler Custody Seals:	Present/Absent/NA	Location Temp Taken:	Control/Between Samples
Container Custody Seals:	Present/Absent/NA	Are all temperatures within project specific guidelines?	Yes/No/NA
Ice Present:	Yes/No/NA	VOA Headspace Present?	Yes/No/NA
	Frozen/Melted/NA		

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9276</u>	2 °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: Meredith Edrington Signature Meredith Edrington Printed Name 3/20/19 Date

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Must Deliver Next Business Day
Time and Temperature Sensitive!

Part # 159469-434 RITZ EXP 11/19

ORIGIN ID:SGRA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

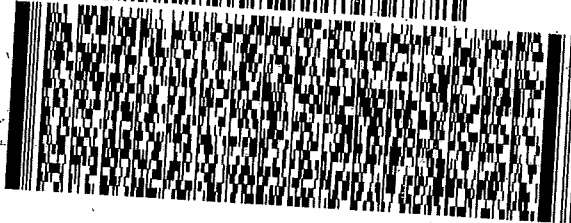
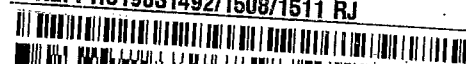
SHIP DATE: 29MAR19
ACTWGT: 9.25 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN
BILL THIRD PARTY

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(801) 266-7700

REF: HS19031492/1508/1511 RJ



FedEx
Express

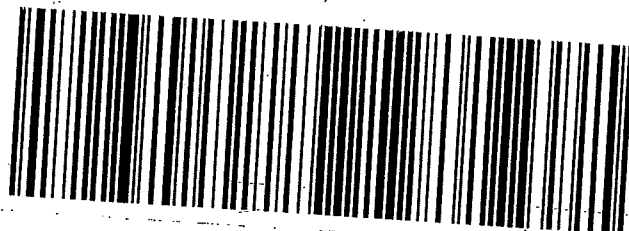


TRK# 4809 7832 2367
0201

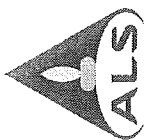
SATURDAY 12:00P
PRIORITY OVERNIGHT

XO BTFA

84123
UT-US SLC



551C1/4E93/413155



Batch Worklist

Batch: ELMS/ 2233 **Created:** 4/9/2019 08:42 **Instrument:** LCMS04 **HBN:** 236356
Rule: EPA 6850, DoD QSM Water **Analyst:** T. Bosch **Status:** WP



- Workorder:** 1909152 [ENV_LVL4]
- Workorder:** 1909153 [ENV_LVL4]
- Workorder:** 1909154 [ENV_LVL4]
- Workorder:** 1909947 [ENV_LVL4]
- Workorder:** 1909949 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	647195	CCV for HBN 236356 [ELMS/2233]				CCV	3		E685041C3Q	5311		4/11/2019	4/10/2019
2	647199	LCS for HBN 236356 [ELMS/2233]				LCS	3		E6850Q413Q	5311		4/11/2019	4/10/2019
3	647197	ICS for HBN 236356 [ELMS/2233]				ICS	3		E6850.D3Q	5311		4/11/2019	4/10/2019
4	647198	LMB for HBN 236356 [ELMS/2233]				LMB	3		E6850Q413Q	5311		4/11/2019	4/10/2019
5	1909152001	LH18/24-SP650_032719_BIX				SAMPLE	3	1909152001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
6	647200	LH18/24-SP650...(1909152001MS)				MS	3		E6850Q413Q	5311		4/11/2019	4/10/2019
7	647201	LH18/24-SP65...(1909152001MSD)				MSD	3		E6850Q413Q	5311		4/11/2019	4/10/2019
8	1909153001	LH18-24-SP140_032719				SAMPLE	3	1909153001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
9	1909154001	LH18/24-SP650_032719_BIX				SAMPLE	3	1909154001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
10	1909947001	LH18/25-SP650_040419_BIX				SAMPLE	3	1909947001-A	E6850Q41.3	5480	5/2/2019	4/18/2019	4/10/2019
11	1909949001	HBW7_040119				SAMPLE	3	1909949001-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
12	1909949002	HBW7_040119-a				SAMPLE	3	1909949002-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
13	1909949003	HBW10_040119				SAMPLE	3	1909949003-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
14	1909949004	HBW1_040119				SAMPLE	3	1909949004-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
15	1909949005	GPW1_040119				SAMPLE	3	1909949005-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
16	1909949006	GPW3_040119				SAMPLE	3	1909949006-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
17	647196	RLVS for HBN 236356 [ELMS/2233]				RLVS	3		E685041C3Q	5311		4/11/2019	4/10/2019
18	647202	CCV for HBN 236356 [ELMS/2233]				CCV	3		E685041C3Q	5311		4/11/2019	4/10/2019



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1909152 (001); 1909153 (001); 1909154 (001); 1909947 (001); 1909949 (001-06) ELMS Batch/HBN ID: 2233 (236356)
 Prep Date: 04/09/2019 Analysis Date: 04/10/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\28MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by **TNB**. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4) / 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 647199; Target = 4.0µg/L. ASTM type II water was used for LMB 647198.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1909152001 (Client ID's: LH18/24-SP650_032719_BIX). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The MS/MSD – 647200/01 failed QC acceptance criteria for percent recoveries, biased high. This is due to the fact that the unspiked sample result of 2.574µg/L was not subtracted from the MS/MSD results. The Matrix Spike and Matrix Spike duplicate is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\APR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\slstws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\236356-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATA\REVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 647196) is reported from the analysis of the Laboratory Control Sample (LCS – 647199) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
<u>Batch(es)/SDG: E LMS: 2233 HBN: 236356</u> 1209947 / 1209949		
<u>Sample Set IDs if Applicable: 1909152/1909153/1909154</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR#</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK		Description - 6850 WKG Std 100.ug/L			
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK		Description - 6850 Stock AccStd 1,000ug/mL	
Standard: 43659	Created By: Thomas Bosch	Amount: 100 mL	
MFG: AccuStandard	Create Date: 09/17/2018 09:09AM	Expires: 07/25/2020	
MFG Lot: 218065075		Usable: No	
Part ID: IC-PER-10X-1		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT		Description - 6850 Intermdt AccStd 10.ug/mL			
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK		Description - 6850 QC WKG STD 100ug/L			
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK		Description - 6850 QC Stock STD 1,000ug/mL	
Standard: 36748		Created By: Thomas Bosch	Amount: 100 mL
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	Expires: 03/31/2020
MFG Lot: CP-0860			Usable: Yes
Part ID: ICC-013			Lab Lot: CLO4 QC STOCK
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

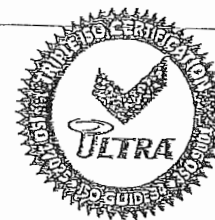
Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:

This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis



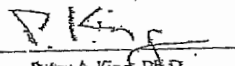
ISO Guide 34 Reference Material

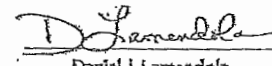
Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:

The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QA/RA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275B72-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:
ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$
Labeled CAS Number: NA
Unlabeled CAS Number: 7601-89-0
MW*: 130.4
Chemical Formula: NaCl^*O_4
Storage: Store at room temperature away from light and moisture.
Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 $\mu\text{g/mL}$
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 \pm 2.8 $\mu\text{g/mL}$ (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	647195	CCV@25	Vial 71	1	Control	1	1.06336e6	21.31933
#*	647199	QC@4.0	Vial 72	1	Control	2	1.97600e5	3.67714
#*	647197	ICS@4.0	Vial 73	1	Control	3	1.49399e5	3.21517
#*	647198	LMB	Vial 74	1	Control	4	0.00000	0.00000
#*	1909152001		Vial 75	1	Sample	5	9.15156e4	2.57433
#*	647200	91521MS	Vial 76	1	Sample	6	2.25049e5	6.17769
#*	647201	91521SD	Vial 77	1	Sample	7	2.49579e5	6.36091
#*	1909153001	1K	Vial 78	1	Sample	8	3.70174e5	6532.86203
#*	1909154001		Vial 79	1	Sample	9	7.79123e4	2.34023
#*	1909947001		Vial 80	1	Sample	10	0.00000	0.00000
#*	1909949001		Vial 81	1	Sample	11	0.00000	0.00000
#*	1909949002		Vial 82	1	Sample	12	0.00000	0.00000
#*	1909949003		Vial 83	1	Sample	13	0.00000	0.00000
#*	1909949004		Vial 84	1	Sample	14	0.00000	0.00000
#*	1909949005		Vial 85	1	Sample	15	0.00000	0.00000
#*	1909949006		Vial 86	1	Sample	16	0.00000	0.00000
*	647202	CCV@25	Vial 71	1	Control	17	1.08007e6	22.59518

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	647195	CCV@25	Vial 71	1	Control	1	3.43635e5	23.09461
#*	647199	QC@4.0	Vial 72	1	Control	2	7.01681e4	4.22460
#*	647197	ICS@4.0	Vial 73	1	Control	3	5.45965e4	3.77625
#*	647198	LMB	Vial 74	1	Control	4	0.00000	0.00000
#*	1909152001		Vial 75	1	Sample	5	3.18275e4	2.84107
#*	647200	91521MS	Vial 76	1	Sample	6	7.61114e4	6.88356
#*	647201	91521SD	Vial 77	1	Sample	7	8.42686e4	7.08137
#*	1909153001	1K	Vial 78	1	Sample	8	1.21054e5	7051.77497
#*	1909154001		Vial 79	1	Sample	9	2.76117e4	2.61418
#*	1909947001		Vial 80	1	Sample	10	0.00000	0.00000
#*	1909949001		Vial 81	1	Sample	11	0.00000	0.00000
#*	1909949002		Vial 82	1	Sample	12	0.00000	0.00000
#*	1909949003		Vial 83	1	Sample	13	0.00000	0.00000
#*	1909949004		Vial 84	1	Sample	14	0.00000	0.00000
#*	1909949005		Vial 85	1	Sample	15	0.00000	0.00000
#*	1909949006		Vial 86	1	Sample	16	0.00000	0.00000
*	647202	CCV@25	Vial 71	1	Control	17	3.36319e5	23.63294

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	647195	CCV@25	Vial 71	1	Control	1	1.53120e5	5.00000
#*	647199	QC@4.0	Vial 72	1	Control	2	1.77838e5	5.00000
#*	647197	ICS@4.0	Vial 73	1	Control	3	1.54931e5	5.00000
#*	647198	LMB	Vial 74	1	Control	4	1.76915e5	5.00000
#*	1909152001		Vial 75	1	Sample	5	1.20240e5	5.00000
#*	647200	91521MS	Vial 76	1	Sample	6	1.17757e5	5.00000
#*	647201	91521SD	Vial 77	1	Sample	7	1.26683e5	5.00000
#*	1909153001	1K	Vial 78	1	Sample	8	1.82758e5	5000.00000
#*	1909154001		Vial 79	1	Sample	9	1.13404e5	5.00000
#*	1909947001		Vial 80	1	Sample	10	1.15130e5	5.00000
#*	1909949001		Vial 81	1	Sample	11	1.14791e5	5.00000
#*	1909949002		Vial 82	1	Sample	12	1.24046e5	5.00000
#*	1909949003		Vial 83	1	Sample	13	1.23373e5	5.00000
#*	1909949004		Vial 84	1	Sample	14	1.20241e5	5.00000
#*	1909949005		Vial 85	1	Sample	15	1.27767e5	5.00000
#*	1909949006		Vial 86	1	Sample	16	1.23965e5	5.00000
*	647202	CCV@25	Vial 71	1	Control	17	1.46281e5	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

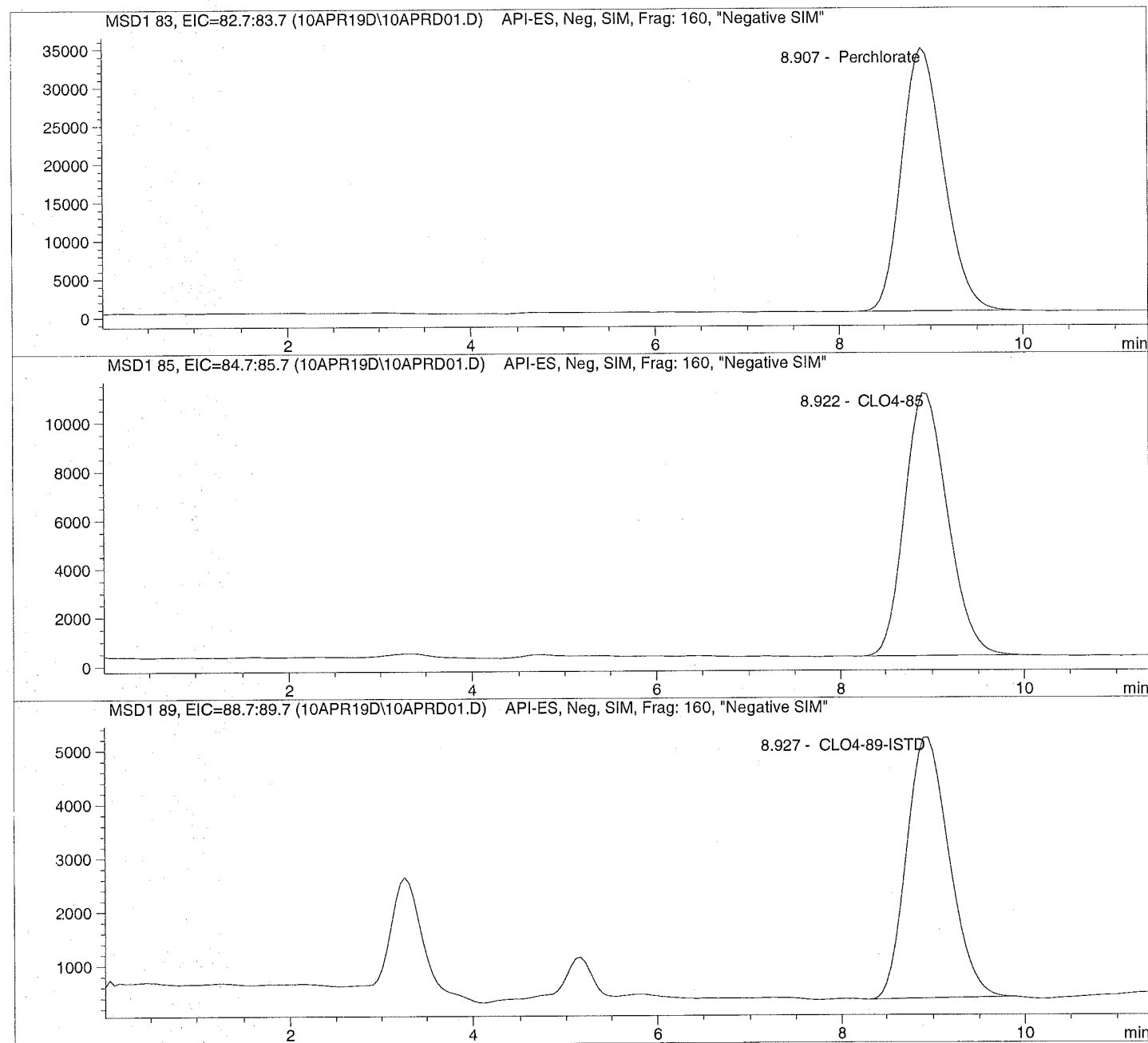
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
1	Vial 71	647195	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	647199	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	647197	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	647198	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1909152001		CLO4-AQN	1	Sample	
6	Vial 76	647200	91521MS	CLO4-AQN	1	Sample	
7	Vial 77	647201	91521SD	CLO4-AQN	1	Sample	
8	Vial 78	1909153001	1K	CLO4-AQN	1	Sample	
9	Vial 79	1909154001		CLO4-AQN	1	Sample	
10	Vial 80	1909947001		CLO4-AQN	1	Sample	
11	Vial 81	1909949001		CLO4-AQN	1	Sample	
12	Vial 82	1909949002		CLO4-AQN	1	Sample	
13	Vial 83	1909949003		CLO4-AQN	1	Sample	
14	Vial 84	1909949004		CLO4-AQN	1	Sample	
15	Vial 85	1909949005		CLO4-AQN	1	Sample	
16	Vial 86	1909949006		CLO4-AQN	1	Sample	
17	Vial 71	647202	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD01.D Sample Name: 647195 CCV@25

```
=====
Injection Date: 4/10/2019 10:01:38      Seq Line: 1
Sample Name: 647195 CCV@25             Location: Vial 71
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD01.D Sample Name: 647195 CCV@25

```

=====
Injection Date: 4/10/2019 10:01:38      Seq Line: 1
Sample Name:    647195  CCV@25          Location:  Vial 71
Acq Operator:   TNB                    Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.907	PBA	1063358.2	21.3193	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.922	PBA	343635.4	23.0946	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.927	PBA	153120.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

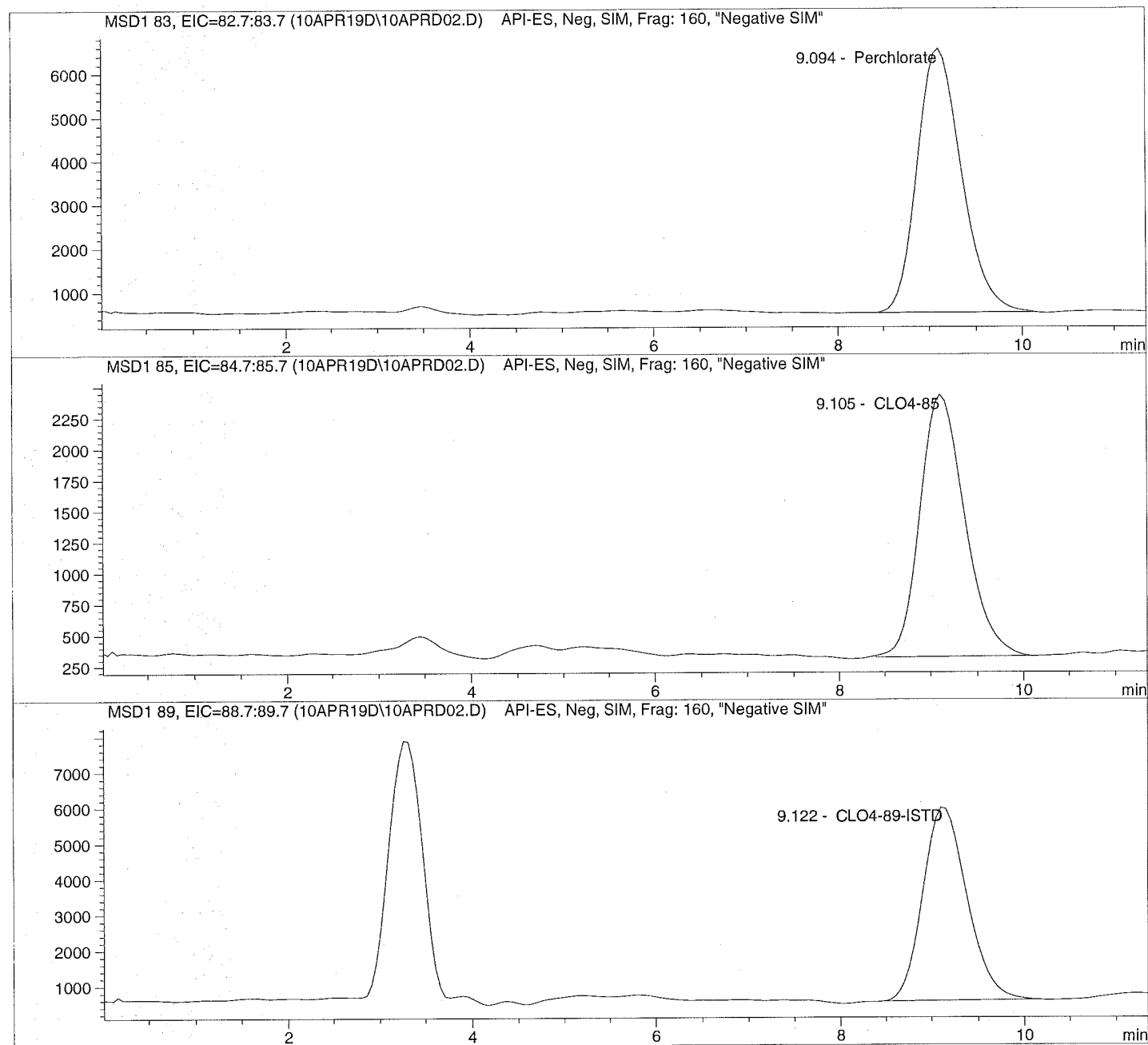
```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD02.D Sample Name: 647199 QC@4.0

=====
Injection Date: 4/10/2019 10:15:40 Seq Line: 2
Sample Name: 647199 QC@4.0 Location: Vial 72
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD02.D Sample Name: 647199 QC@4.0

```

=====
Injection Date: 4/10/2019 10:15:40      Seq Line:      2
Sample Name:    647199 QC@4.0           Location:      Vial 72
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  4.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.094	BBA	197600.1	3.6771	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.105	BBA	70168.1	4.2246	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.122	BBA	177837.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

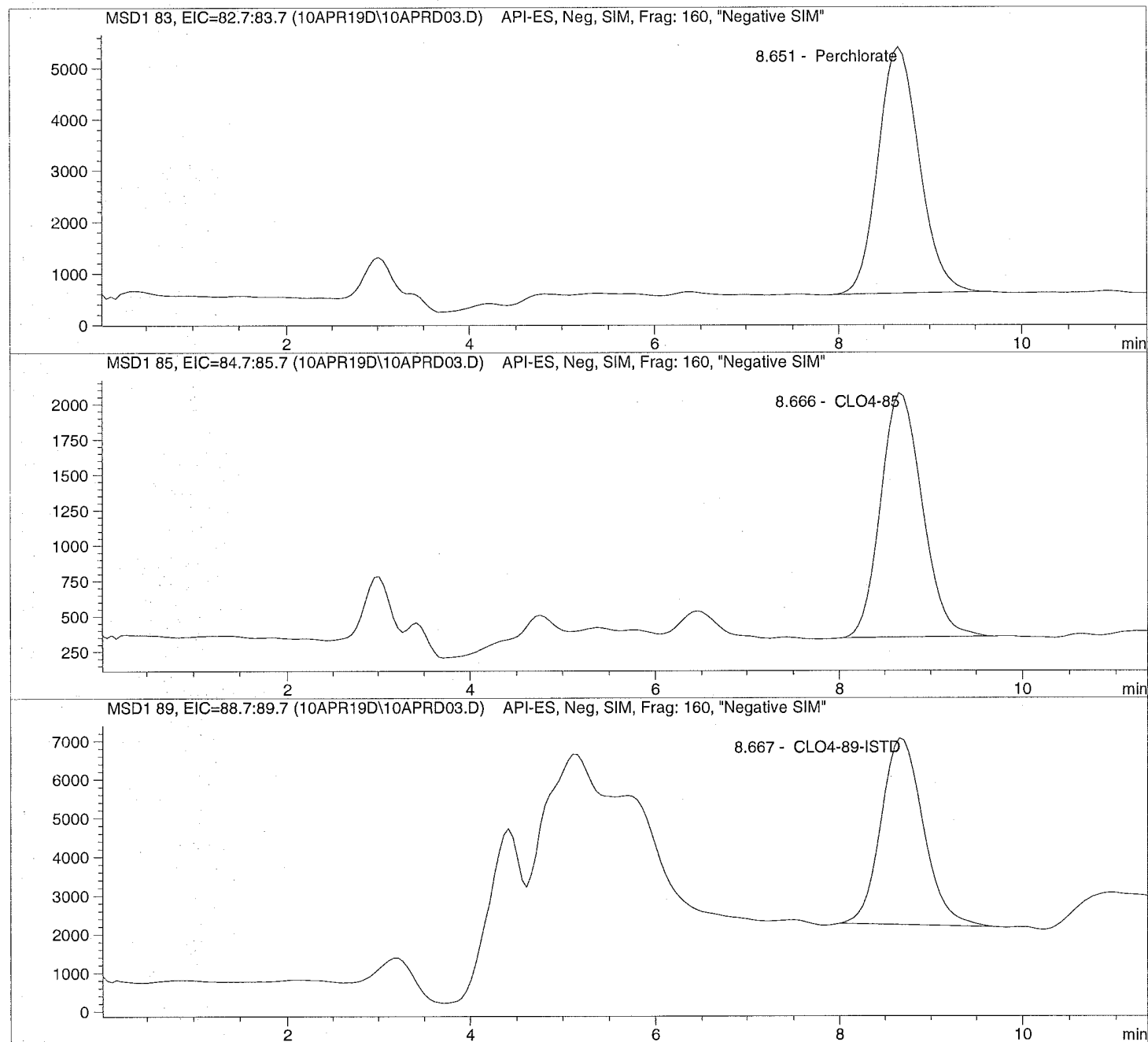
```


Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD03.D Sample Name: 647197 ICS@4.0

```
=====
Injection Date: 4/10/2019 10:28:57      Seq Line: 3
Sample Name: 647197 ICS@4.0             Location: Vial 73
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD03.D Sample Name: 647197 ICS@4.0

```

=====
Injection Date: 4/10/2019 10:28:57      Seq Line:      3
Sample Name:   647197 ICS@4.0          Location:      Vial 73
Acq Operator:  TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 4.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.651	PBA	149398.5	3.2152	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.666	BBA	54596.5	3.7763	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.667	PBA	154930.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

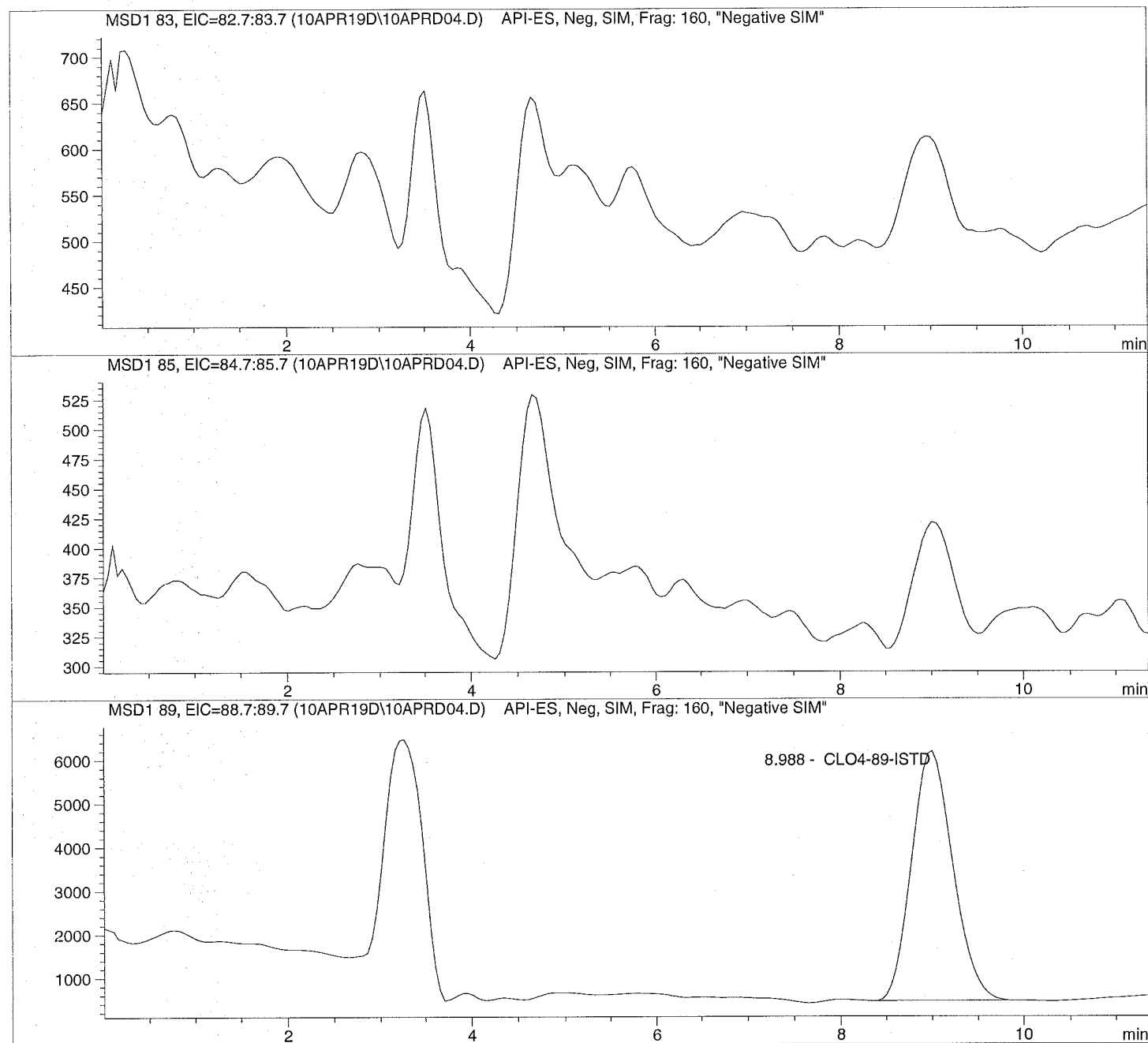
```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD04.D Sample Name: 647198 LMB

```
=====
Injection Date: 4/10/2019 10:42:16      Seq Line: 4
Sample Name: 647198 LMB                  Location: Vial 74
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD04.D Sample Name: 647198 LMB

```

=====
Injection Date: 4/10/2019 10:42:16      Seq Line: 4
Sample Name: 647198 LMB                  Location: Vial 74
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.988	PBA	176915.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD05.D

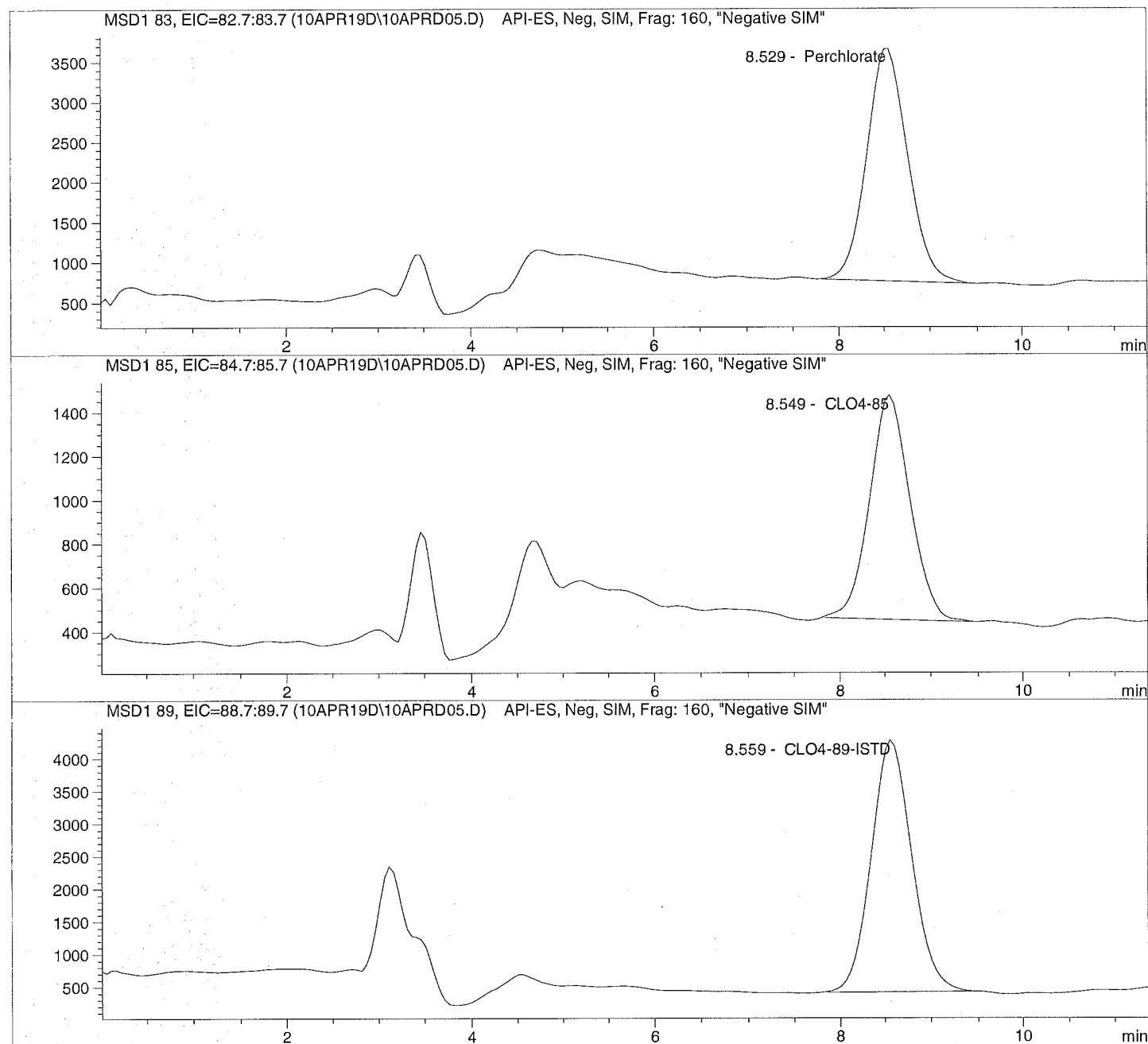
Sample Name: 1909152001

=====
Injection Date: 4/10/2019 10:55:33
Sample Name: 1909152001
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD05.D

Sample Name: 1909152001

```

=====
Injection Date: 4/10/2019 10:55:33      Seq Line:      5
Sample Name:    1909152001              Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.529	BBA	91515.6	2.5743	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.549	BBA	31827.5	2.8411	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.559	BBA	120239.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

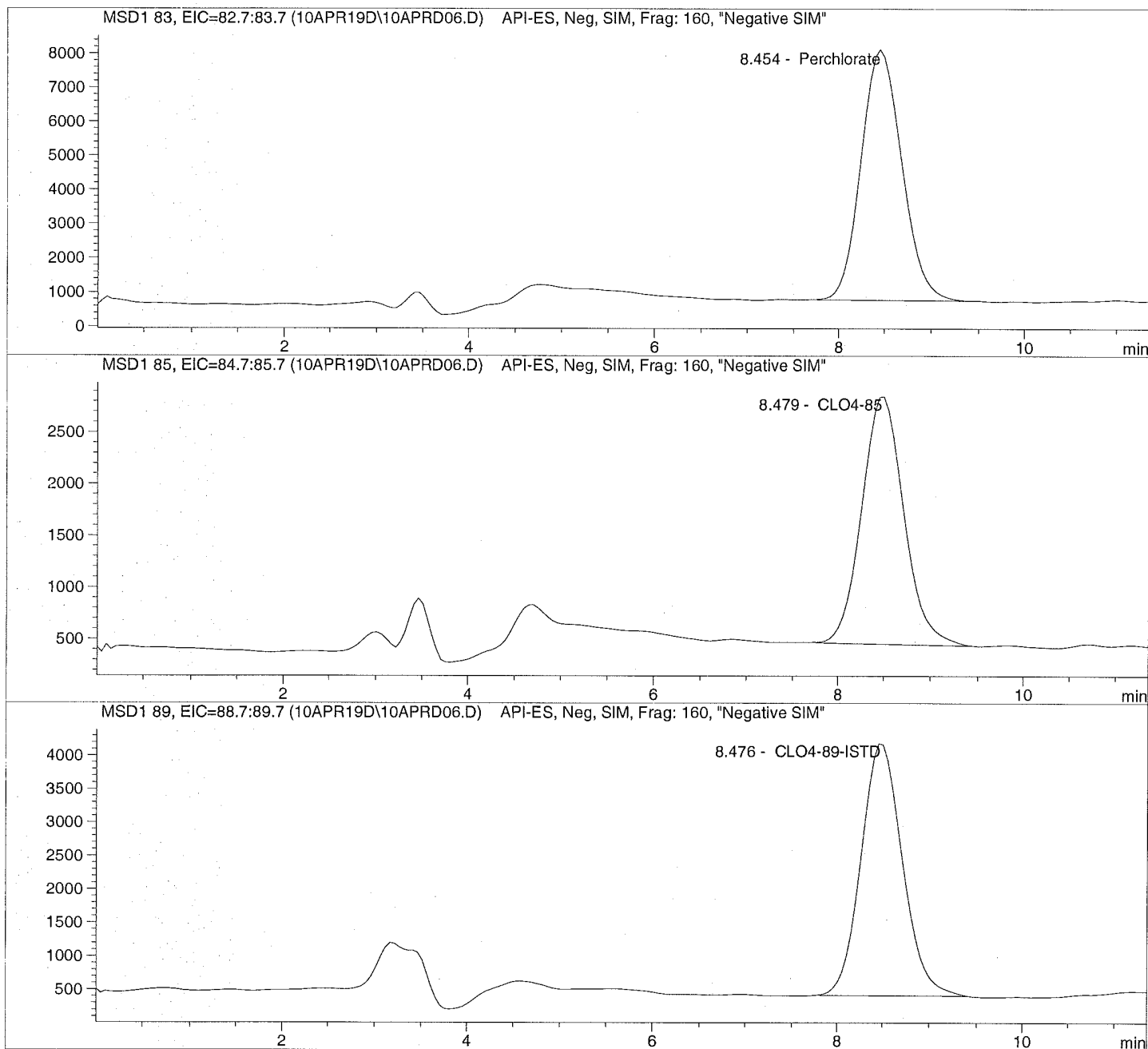
```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD06.D Sample Name: 647200 91521MS

=====
Injection Date: 4/10/2019 11:08:49 Seq Line: 6
Sample Name: 647200 91521MS Location: Vial 76
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD06.D Sample Name: 647200 91521MS

```

=====
Injection Date: 4/10/2019 11:08:49      Seq Line: 6
Sample Name: 647200 91521MS             Location: Vial 76
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.454	BBA	225049.5	6.1777	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.479	BBA	76111.4	6.8836	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.476	BBA	117756.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

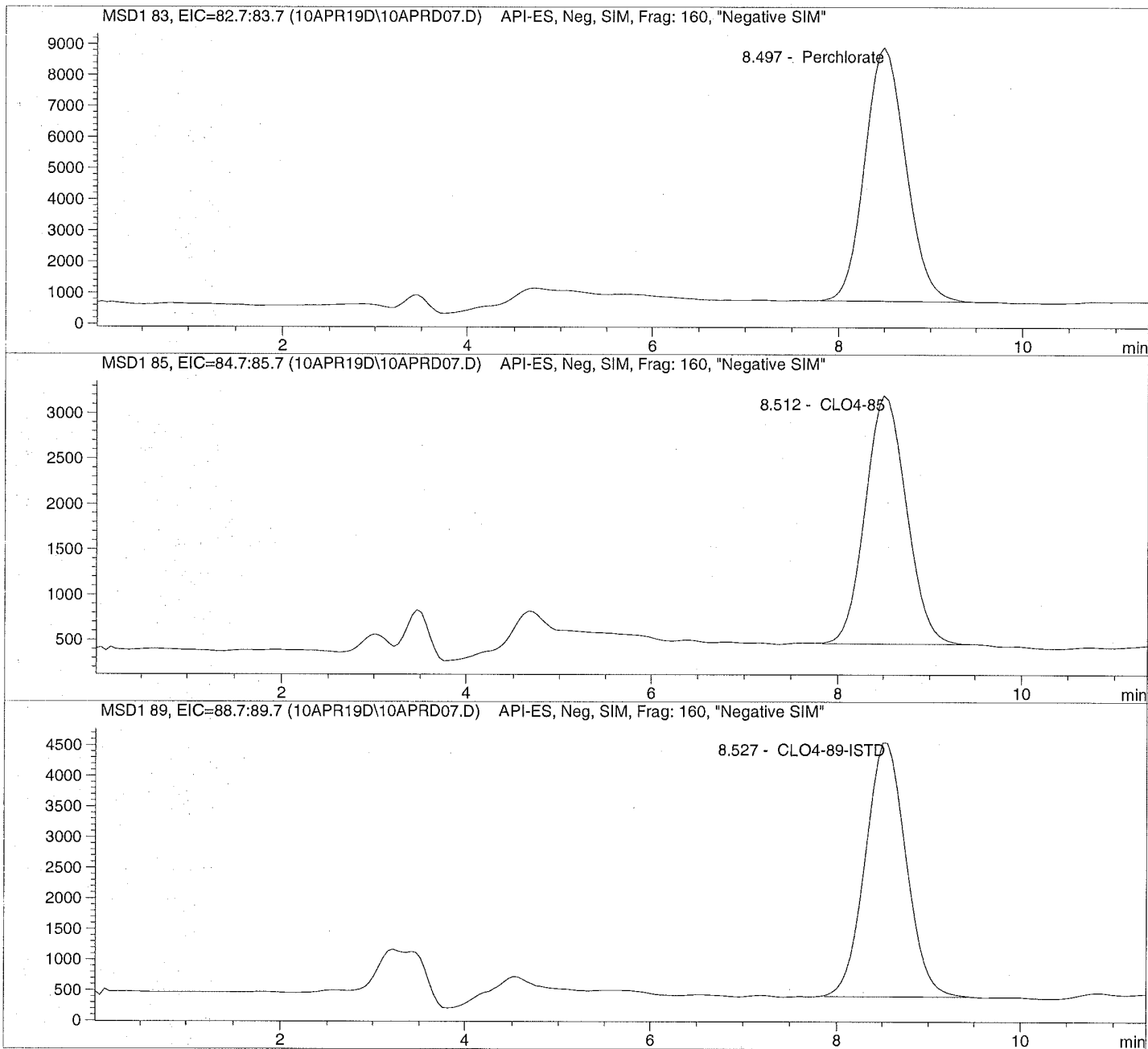
```


Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD07.D Sample Name: 647201 91521SD

```
=====
Injection Date: 4/10/2019 11:22:09 Seq Line: 7
Sample Name: 647201 91521SD Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD07.D Sample Name: 647201 91521SD

```

=====
Injection Date:  4/10/2019  11:22:09      Seq Line:      7
Sample Name:    647201    91521SD        Location:      Vial 77
Acq Operator:   TNB                Inj. No.:     1
                                           Inj. Vol.:    30 µl
  
```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
  
```

Perchlorate analysis

=====

Sample Information

=====

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount:  0.000
  
```

=====

LCMS Results

=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.497	BBA	249579.0	6.3609	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.512	BBA	84268.6	7.0814	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.527	BBA	126683.1	5.0000	CLO4-89-ISTD

=====

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD08.D

Sample Name: 1909153001 1K

Injection Date: 4/10/2019 11:35:27

Seq Line: 8

Sample Name: 1909153001 1K

Location: Vial 78

Acq Operator: TNB

Inj. No.: 1

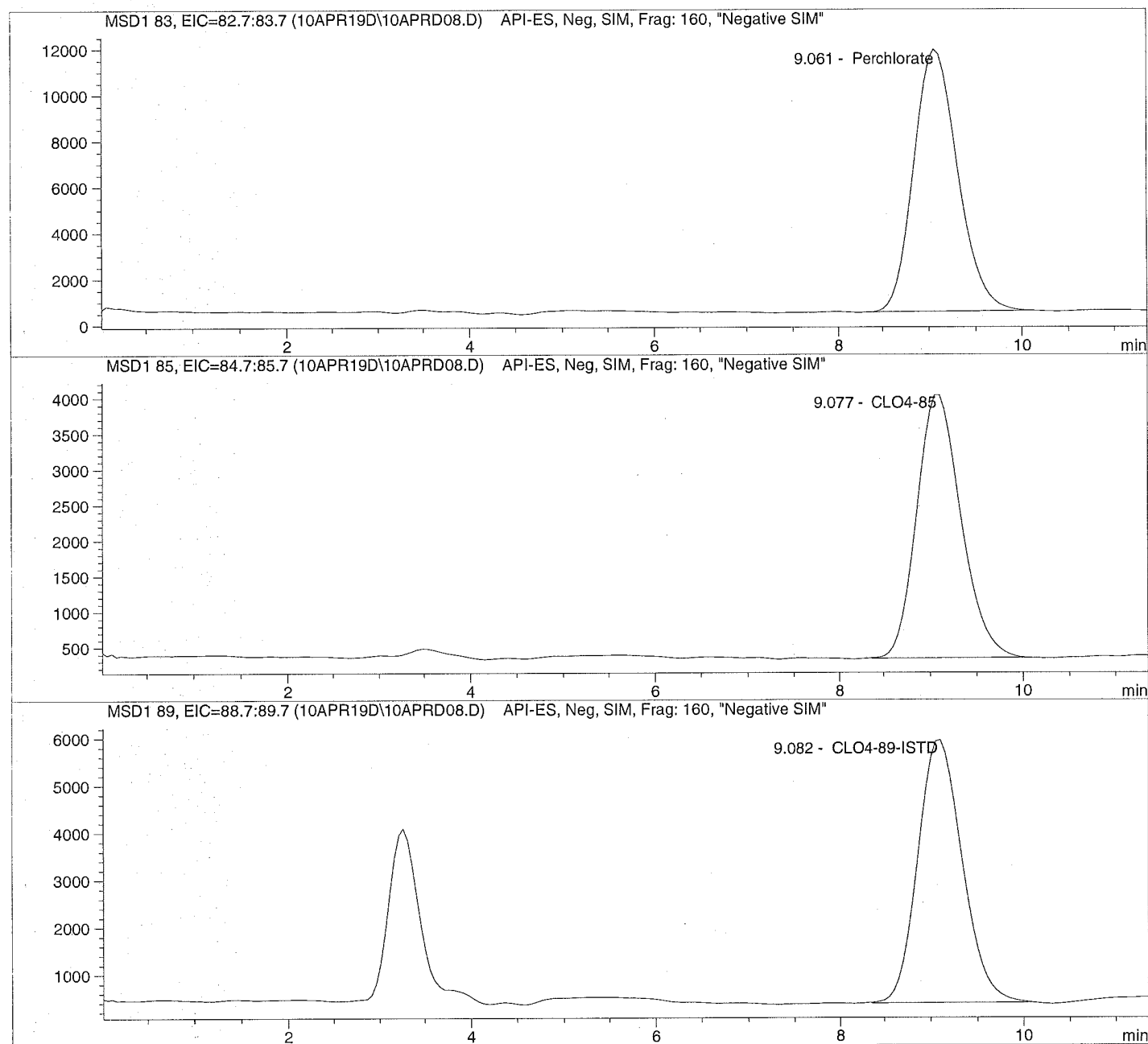
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD08.D Sample Name: 1909153001 1K

```

=====
Injection Date: 4/10/2019 11:35:27      Seq Line:      8
Sample Name:   1909153001 1K           Location:      Vial 78
Acq Operator:  TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1000.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.061	PBA	370174.3	6532.8620	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.077	BBA	121053.5	7051.7750	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.082	BBA	182757.9	5000.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD09.D

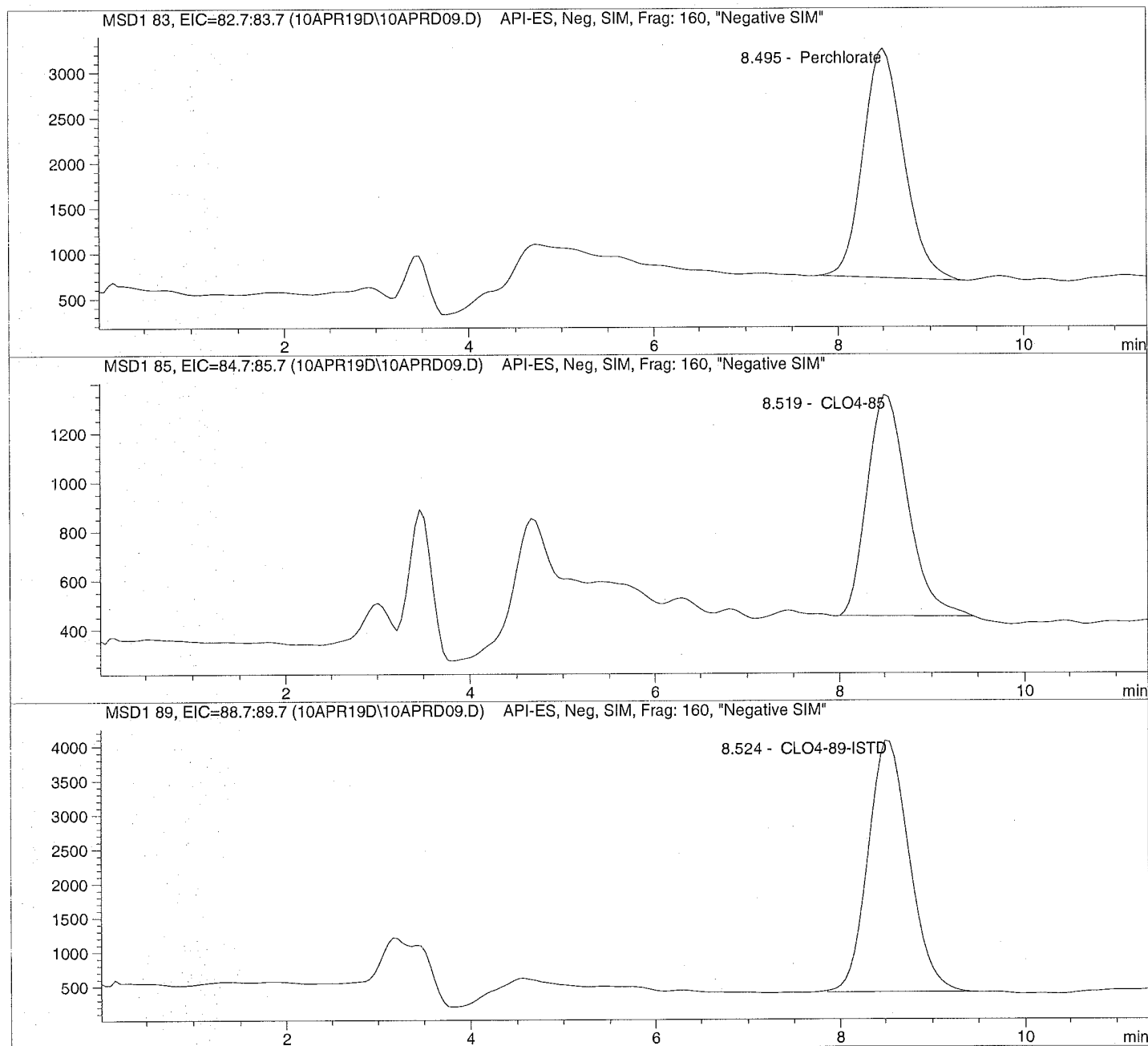
Sample Name: 1909154001

=====
Injection Date: 4/10/2019 11:48:52
Sample Name: 1909154001
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD09.D

Sample Name: 1909154001

```

=====
Injection Date: 4/10/2019 11:48:52      Seq Line:          9
Sample Name:   1909154001                Location:          Vial 79
Acq Operator:  TNB                       Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:          Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:         1.000000
Dilution:           1.000000
Sample Amount:      0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.495	PBA	77912.3	2.3402	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.519	PBA	27611.7	2.6142	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.524	BBA	113404.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD10.D

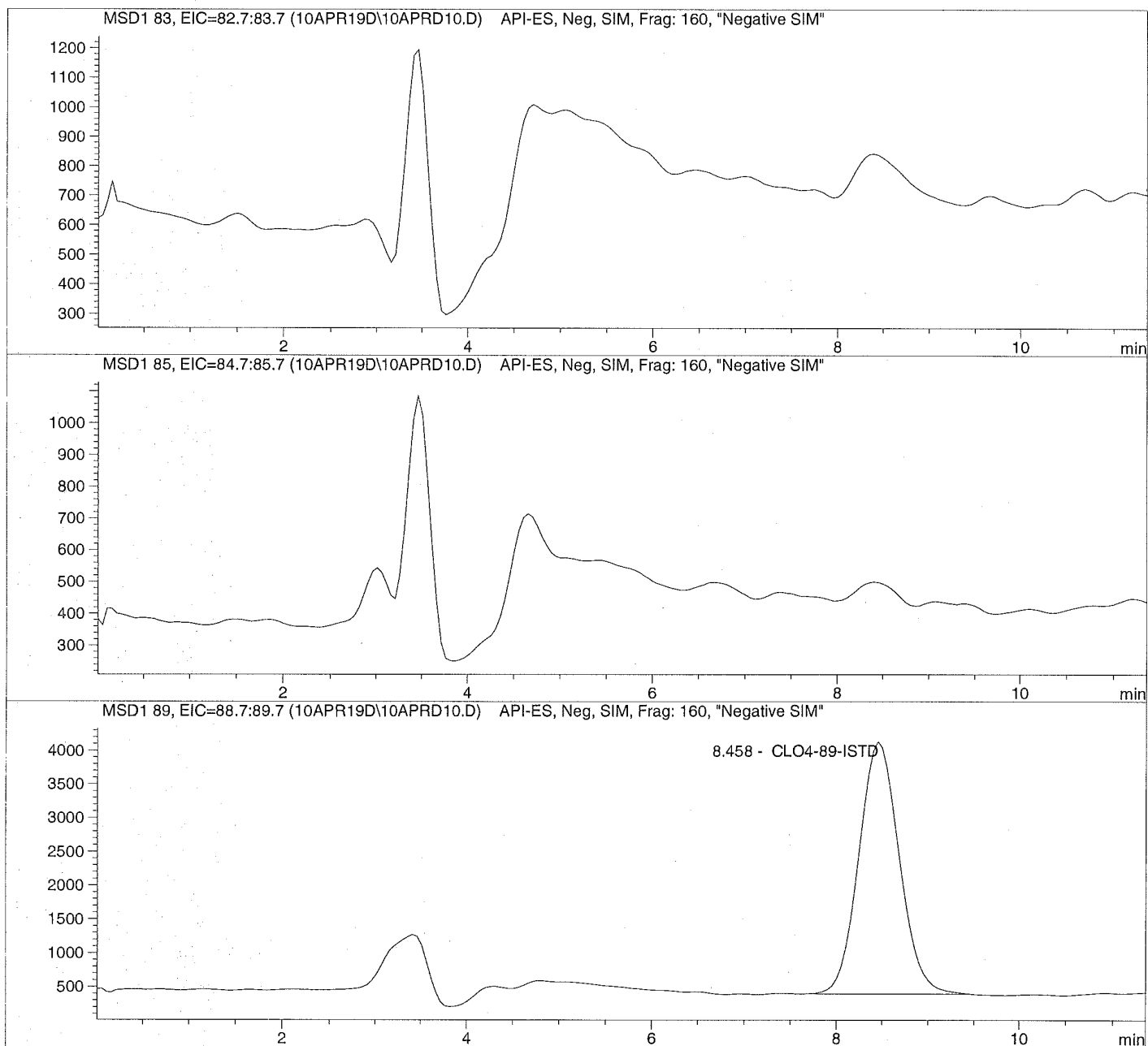
Sample Name: 1909947001

Injection Date: 4/10/2019 12:02:10
Sample Name: 1909947001
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD10.D Sample Name: 1909947001

```

=====
Injection Date: 4/10/2019 12:02:10      Seq Line:      10
Sample Name:   1909947001              Location:      Vial 80
Acq Operator:  TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount:  0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.458	BBA	115129.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```


Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD11.D

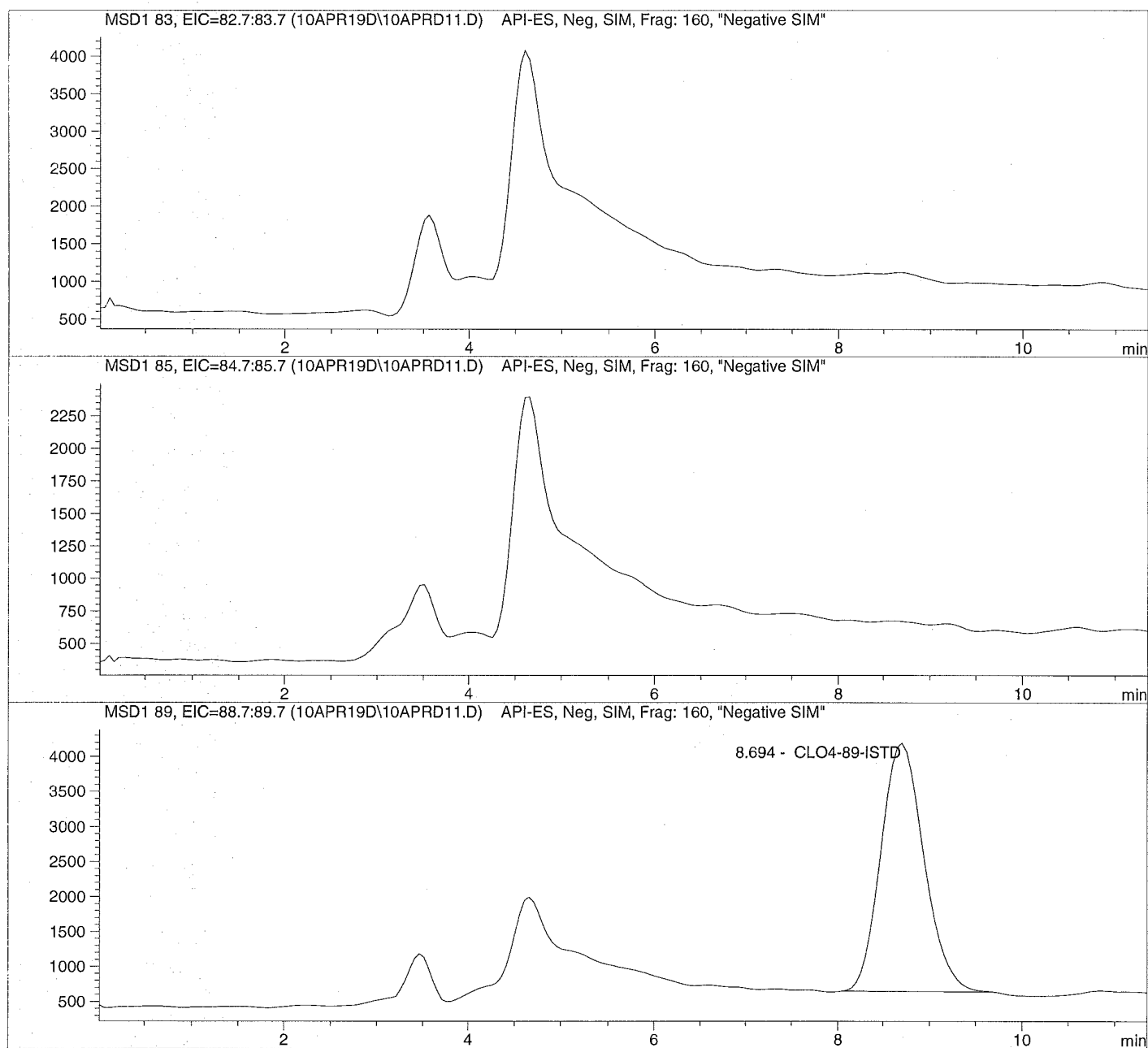
Sample Name: 1909949001

=====
Injection Date: 4/10/2019 12:15:27
Sample Name: 1909949001
Acq Operator: TNB

Seq Line: 11
Location: Vial 81
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD11.D Sample Name: 1909949001

```

=====
Injection Date: 4/10/2019 12:15:27      Seq Line:          11
Sample Name:    1909949001              Location:         Vial 81
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.694	PBA	114791.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD12.D

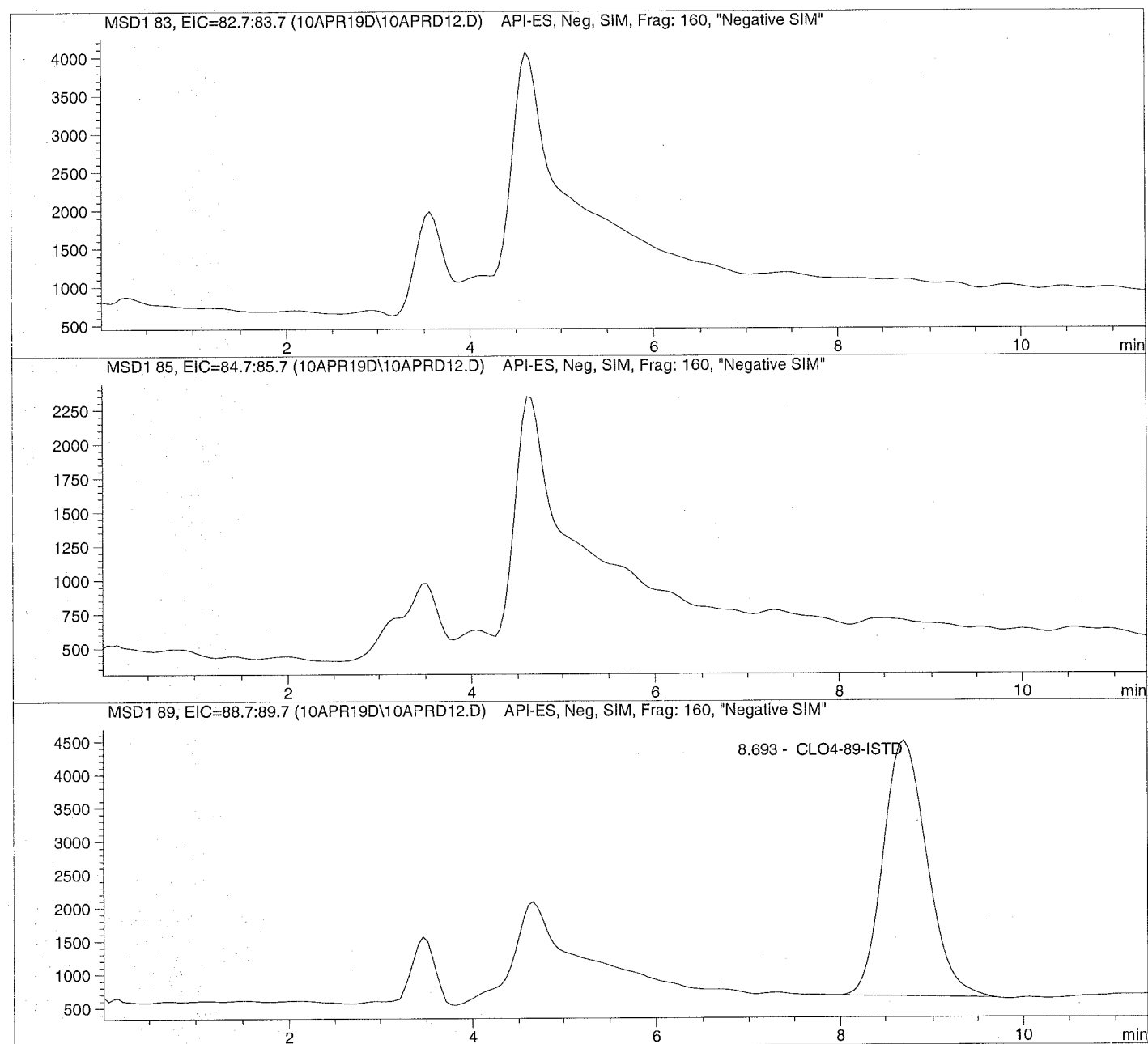
Sample Name: 1909949002

=====
Injection Date: 4/10/2019 12:28:41
Sample Name: 1909949002
Acq Operator: TNB

Seq Line: 12
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD12.D Sample Name: 1909949002

```

=====
Injection Date: 4/10/2019 12:28:41      Seq Line: 12
Sample Name: 1909949002                Location: Vial 82
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.693	PBA	124045.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD13.D

Sample Name: 1909949003

Injection Date: 4/10/2019 12:41:59

Seq Line: 13

Sample Name: 1909949003

Location: Vial 83

Acq Operator: TNB

Inj. No.: 1

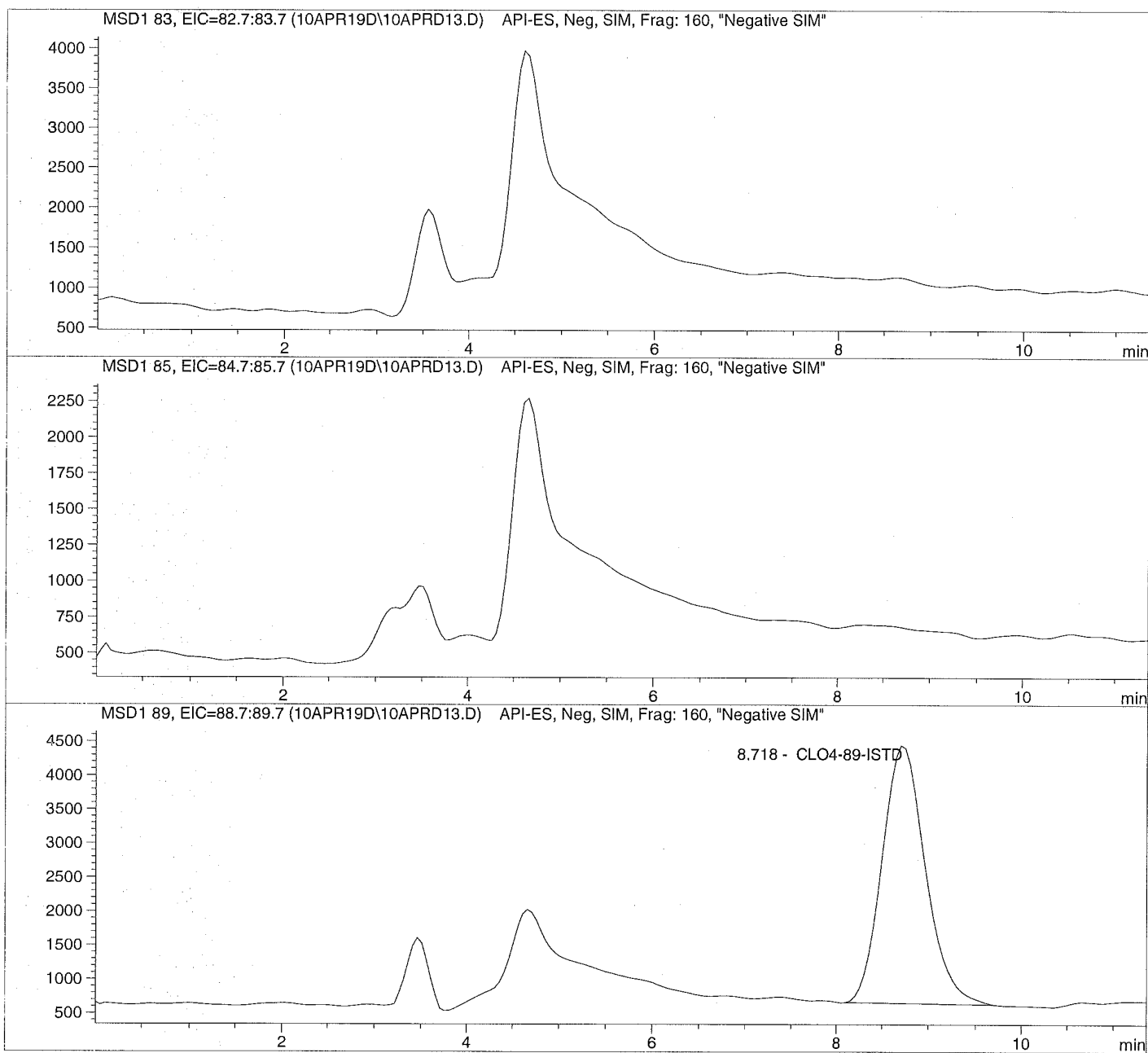
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD13.D Sample Name: 1909949003

```

=====
Injection Date:  4/10/2019  12:41:59      Seq Line:      13
Sample Name:    1909949003      Location:      Vial 83
Acq Operator:   TNB              Inj. No.:     1
                                      Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.718	PBA	123373.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD14.D

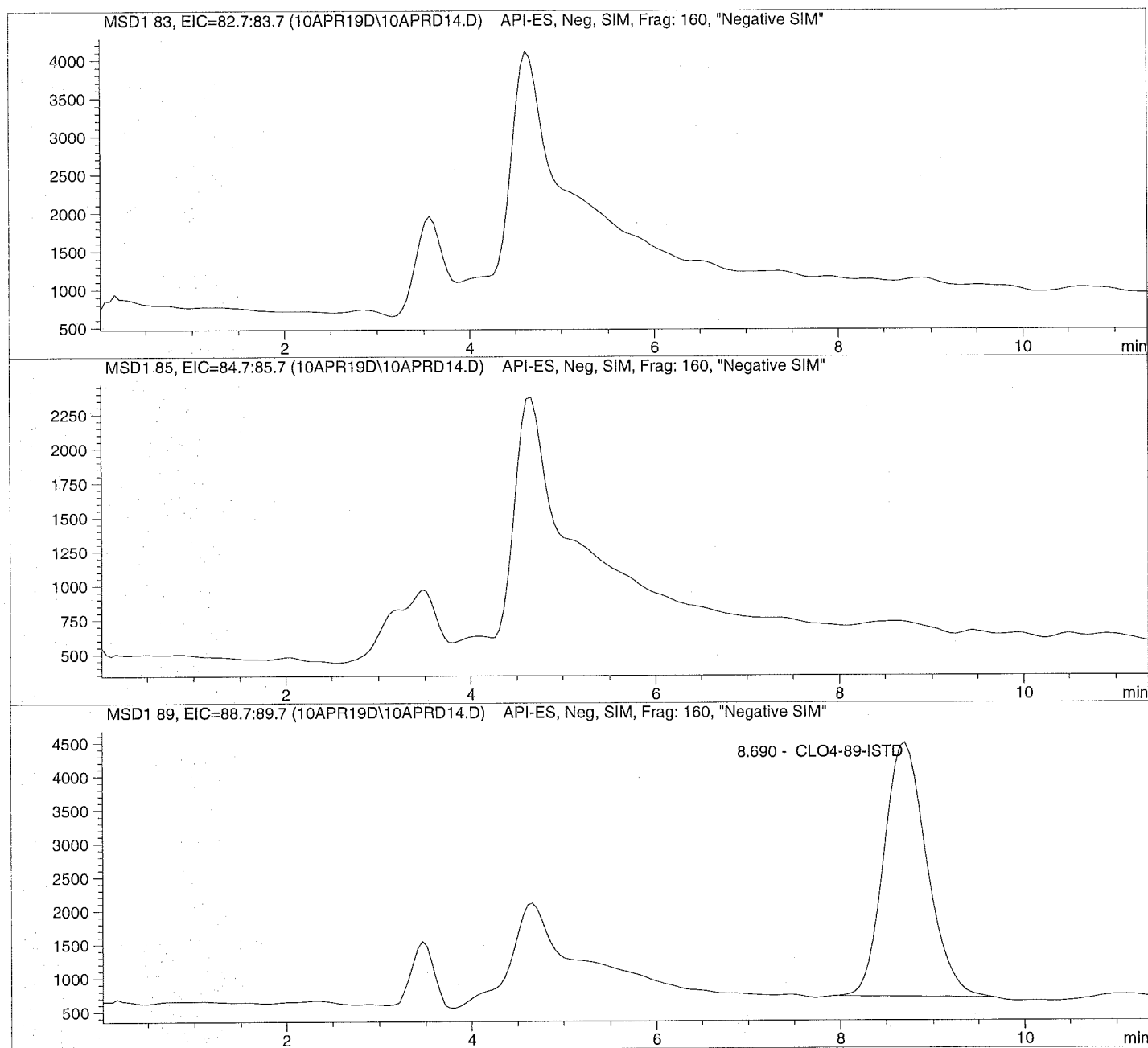
Sample Name: 1909949004

Injection Date: 4/10/2019 12:55:15
Sample Name: 1909949004
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD14.D

Sample Name: 1909949004

```

=====
Injection Date:  4/10/2019  12:55:15      Seq Line:      14
Sample Name:    1909949004      Location:      Vial 84
Acq Operator:   TNB              Inj. No.:      1
                                      Inj. Vol.:     30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.690	BBA	120241.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

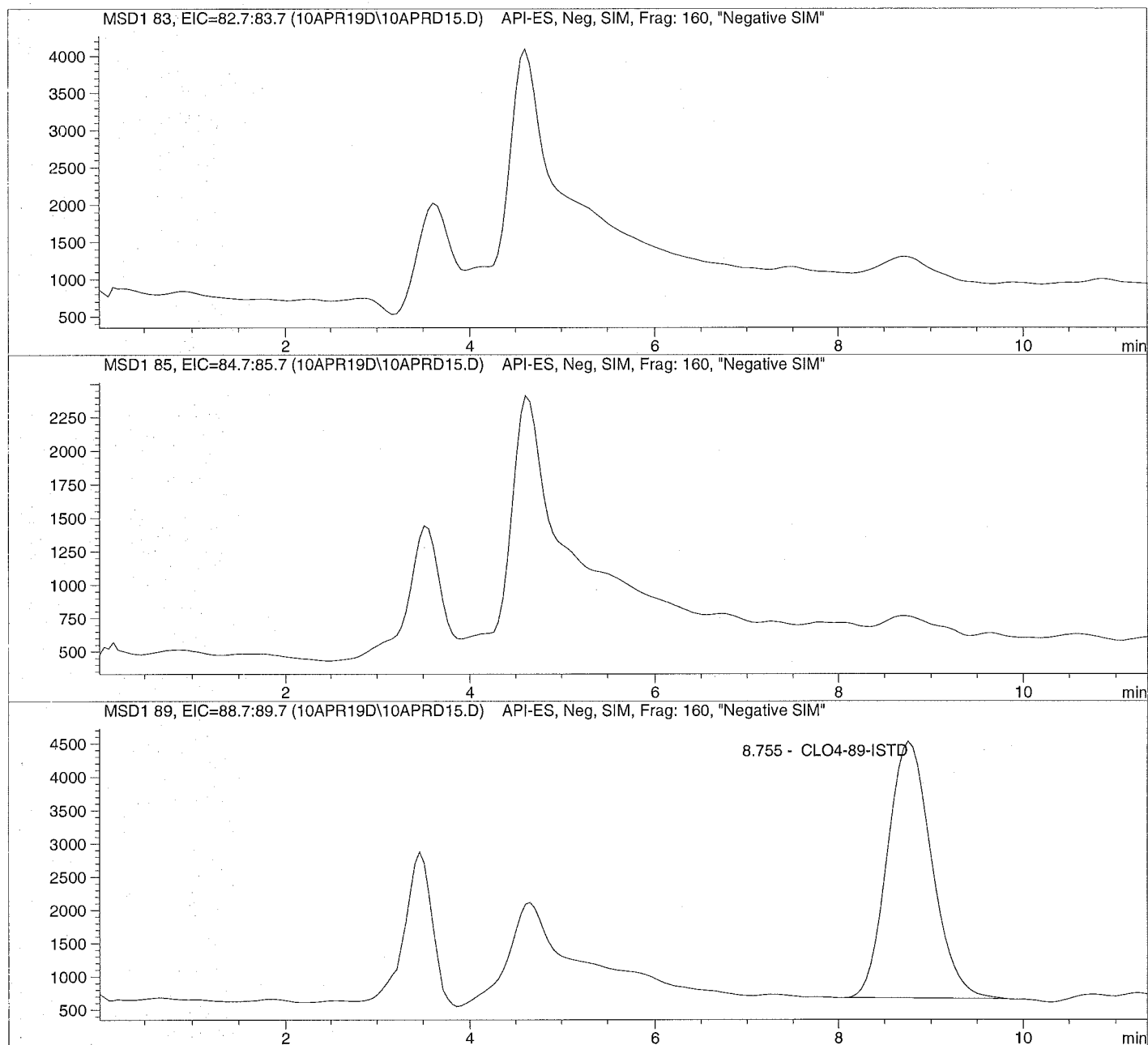

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD15.D

Sample Name: 1909949005

```
=====
Injection Date: 4/10/2019 13:08:30      Seq Line: 15
Sample Name: 1909949005                 Location: Vial 85
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD15.D Sample Name: 1909949005

```

=====
Injection Date: 4/10/2019 13:08:30      Seq Line: 15
Sample Name: 1909949005                Location: Vial 85
Acq Operator: TNB                      Inj. No.: 1
                                         Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	PBA	127767.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD16.D

Sample Name: 1909949006

Injection Date: 4/10/2019 13:21:46

Seq Line: 16

Sample Name: 1909949006

Location: Vial 86

Acq Operator: TNB

Inj. No.: 1

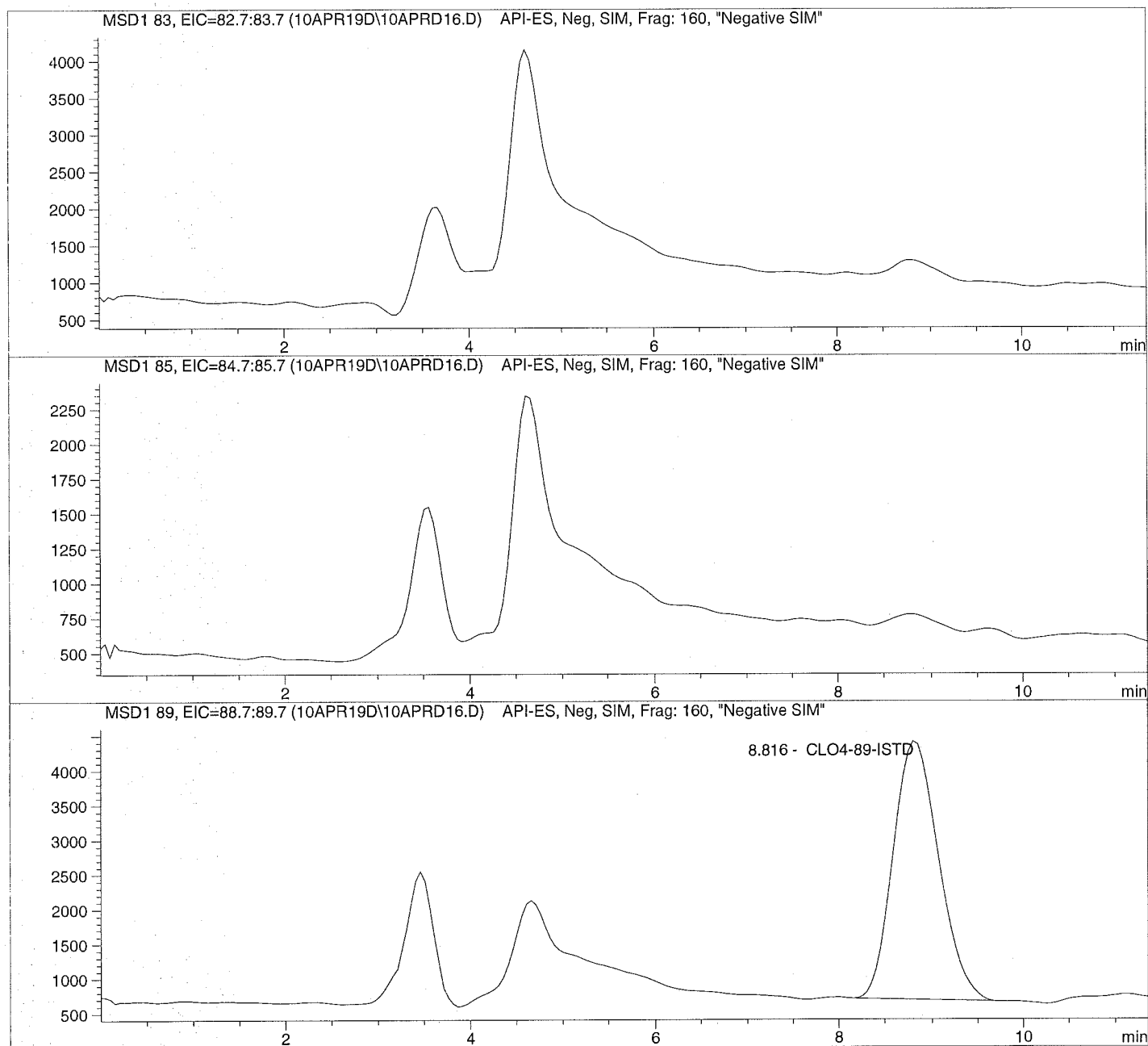
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD16.D Sample Name: 1909949006

```

=====
Injection Date: 4/10/2019 13:21:46      Seq Line: 16
Sample Name: 1909949006                Location: Vial 86
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.816	BBA	123964.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD17.D

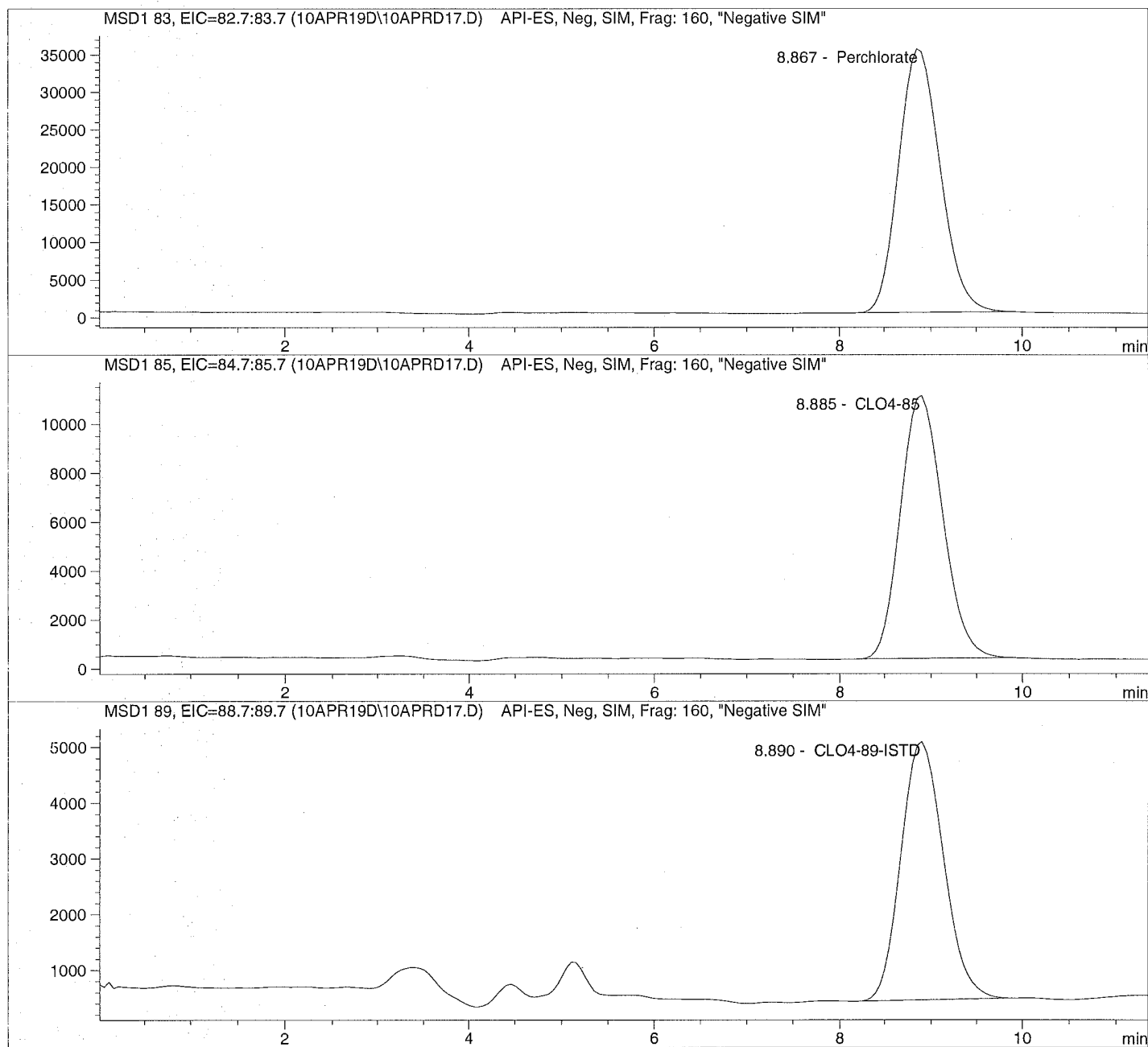
Sample Name: 647202 CCV@25

=====
Injection Date: 4/10/2019 13:35:04
Sample Name: 647202 CCV@25
Acq Operator: TNB

Seq Line: 17
Location: Vial 71
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD17.D Sample Name: 647202 CCV@25

```

=====
Injection Date:  4/10/2019  13:35:04                    Seq Line:                    17
Sample Name:     647202    CCV@25                      Location:                    Vial 71
Acq Operator:    TNB                                    Inj. No.:                    1
                                                          Inj. Vol.:                   30 µl
=====

```

```

Acq. Method:     CLO4-AQN.M
Analysis Method:  C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:    3/19/2019  14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:                    Signal
Calib. Data Modified:        Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                  1.000000
Dilution:                    1.000000
Sample Amount:                25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.867	PBA	1080071.3	22.5952	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.885	PBA	336319.0	23.6329	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.890	PBA	146280.5	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
Calibration Table
=====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
Based on : Peak Area

Rel. Reference Window : 20.000 %
Abs. Reference Window : 0.000 min
Rel. Non-ref. Window : 20.000 %
Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
Uncalibrated Peaks : not reported
Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
Origin : Ignored (some peaks differ, see below)
Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
Average Response : Average all calibrations
Average Retention Time: Floating Average New 75%

Calibration Report Options :
Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
Signal 2: MSD1 85, EIC=84.7:85.7
Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl	Amount	Area	Amt/Area	Ref	Grp	Name
8.744	1	1.00000	7.76074e4	1.28854e-5	1		Perchlorate
	2	2.00000	1.35273e5	1.47849e-5			
	3	5.00000	3.37764e5	1.48033e-5			
	4	10.00000	6.83454e5	1.46316e-5			
	5	25.00000	2.08433e6	1.19943e-5			
	6	50.00000	4.13334e6	1.20968e-5			
	7	75.00000	5.99313e6	1.25143e-5			
8.755	2	1.00000	2.36780e4	4.22333e-5	1		CLO4-85
	2	2.00000	4.69486e4	4.25998e-5			
	3	5.00000	1.06124e5	4.71147e-5			
	4	10.00000	2.13523e5	4.68335e-5			
	5	25.00000	6.14295e5	4.06971e-5			
	6	50.00000	1.19814e6	4.17315e-5			
	7	75.00000	1.78355e6	4.20509e-5			
8.766	3	5.00000	2.73208e5	1.83011e-5	+I1		CLO4-89-ISTD
	2	5.00000	2.24886e5	2.22335e-5			
	3	5.00000	2.33196e5	2.14412e-5			
	4	5.00000	2.34454e5	2.13262e-5			
	5	5.00000	2.50568e5	1.99547e-5			
	6	5.00000	2.30977e5	2.16472e-5			

RetTime	Lvl	Amount	Area	Amt/Area	Ref Grp Name
[min]	Sig				
----- ---	-----	-----	-----	-----	-----
7		5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-85

Time Window : From 6.650 min To 12.505 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

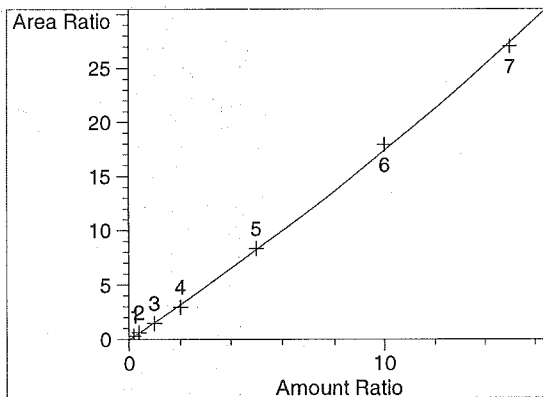
Compound: CLO4-89-ISTD

Time Window : From 6.659 min To 12.466 min
Curve Type : Linear
Origin : Included
Calibration Level Weights:/
Level 1 : 1
Level 2 : 1
Level 3 : 1
Level 4 : 1
Level 5 : 1
Level 6 : 1
Level 7 : 1

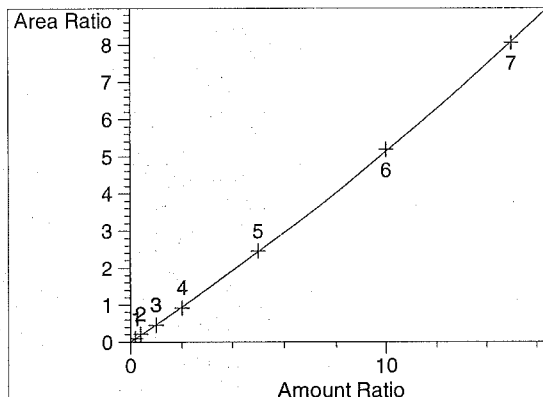
=====
Peak Sum Table
=====

No Entries in table
=====

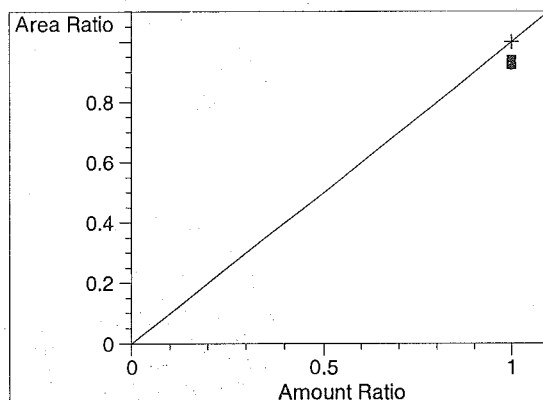
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

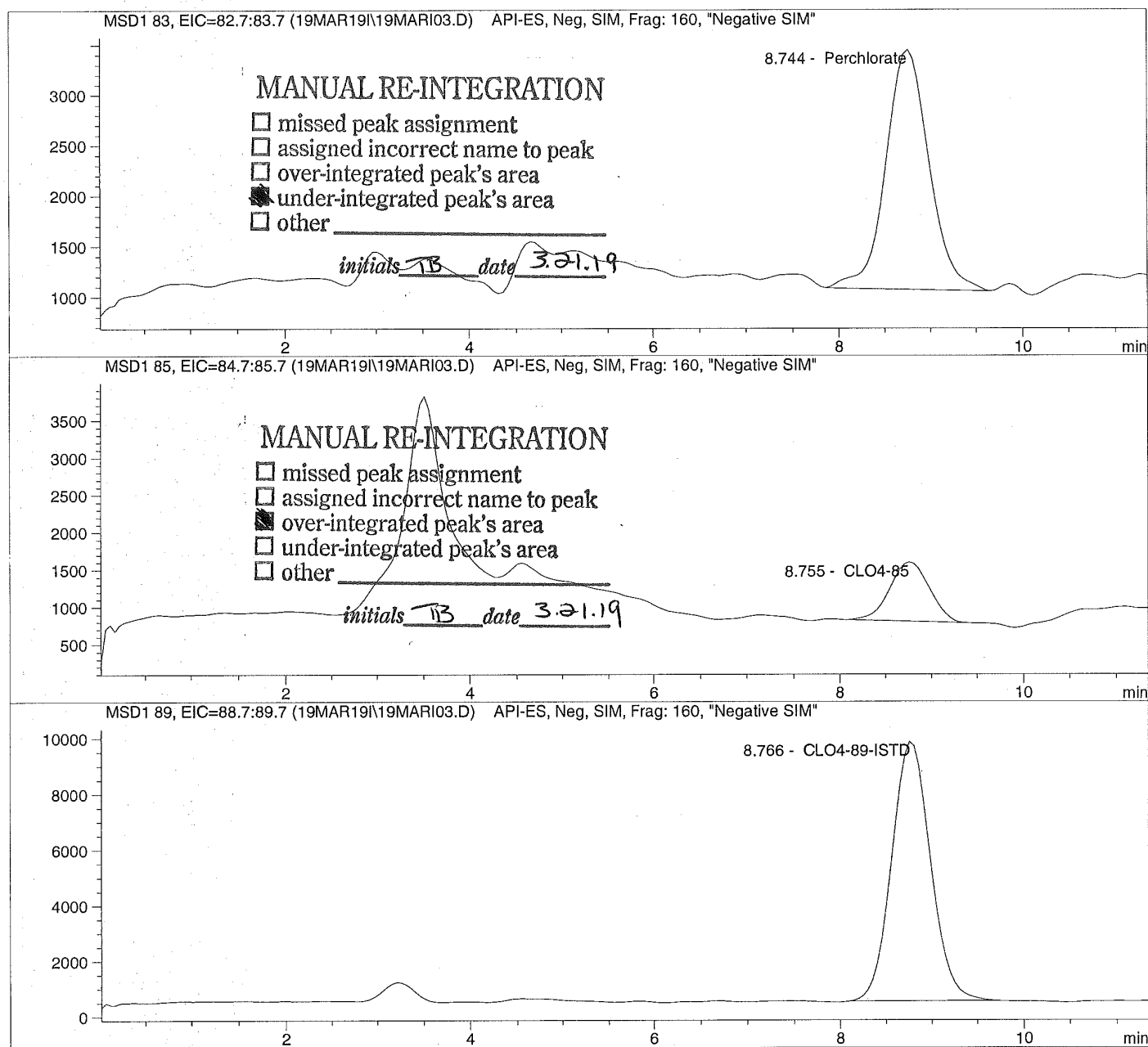
Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40 Seq Line: 3
Sample Name: CLO4@ 1.0ug/L Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl
  
```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22
  
```

Perchlorate analysis

Sample Information

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 1.000
  
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI04.D

Sample Name: CLO4@ 2.0ug/L

Injection Date: 3/19/2019 09:53:00

Seq Line: 4

Sample Name: CLO4@ 2.0ug/L

Location: Vial 74

Acq Operator: TNB

Inj. No.: 1

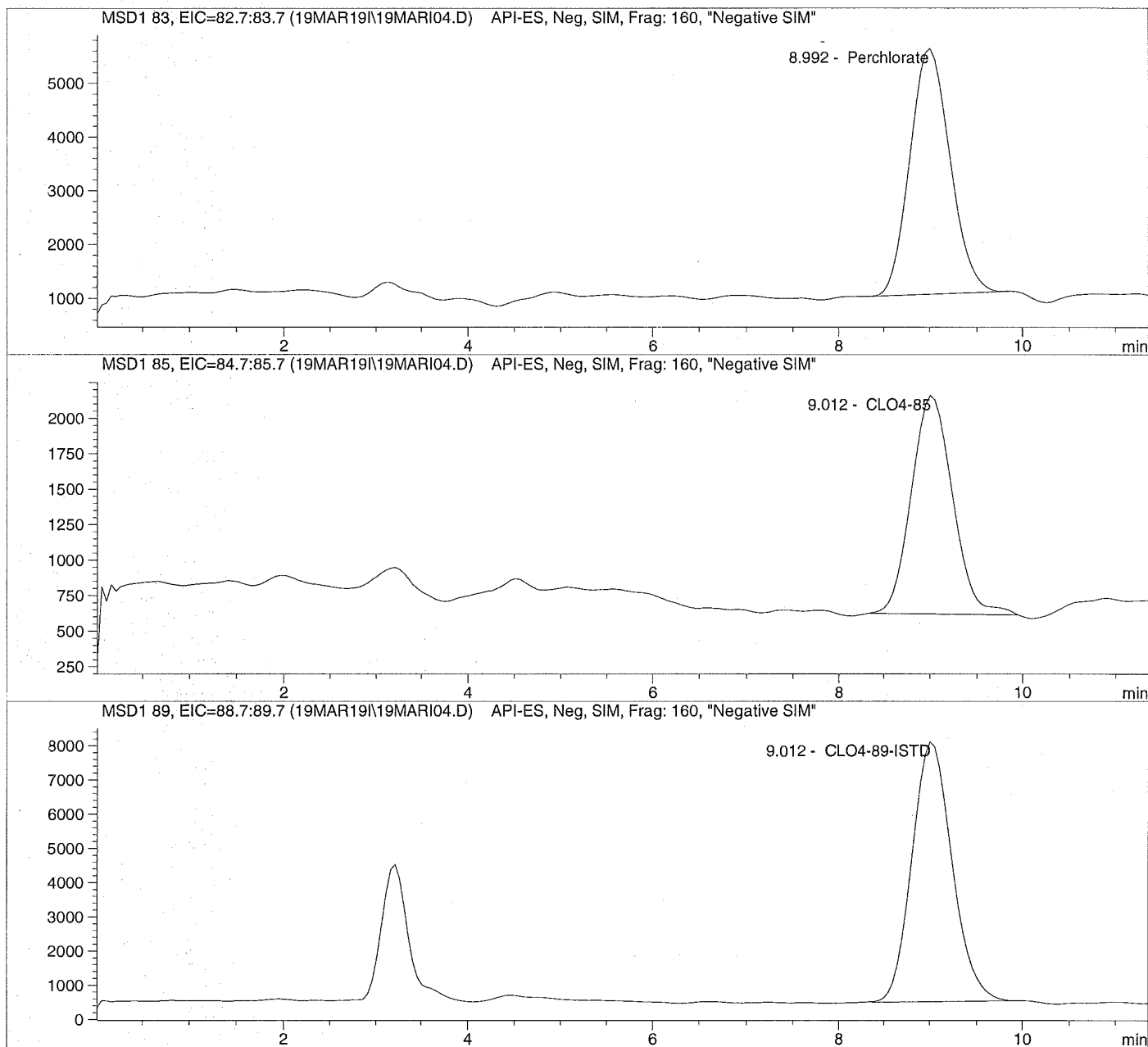
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line: 4
Sample Name:    CLO4@ 2.0ug/L           Location:  Vial 74
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  2.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D

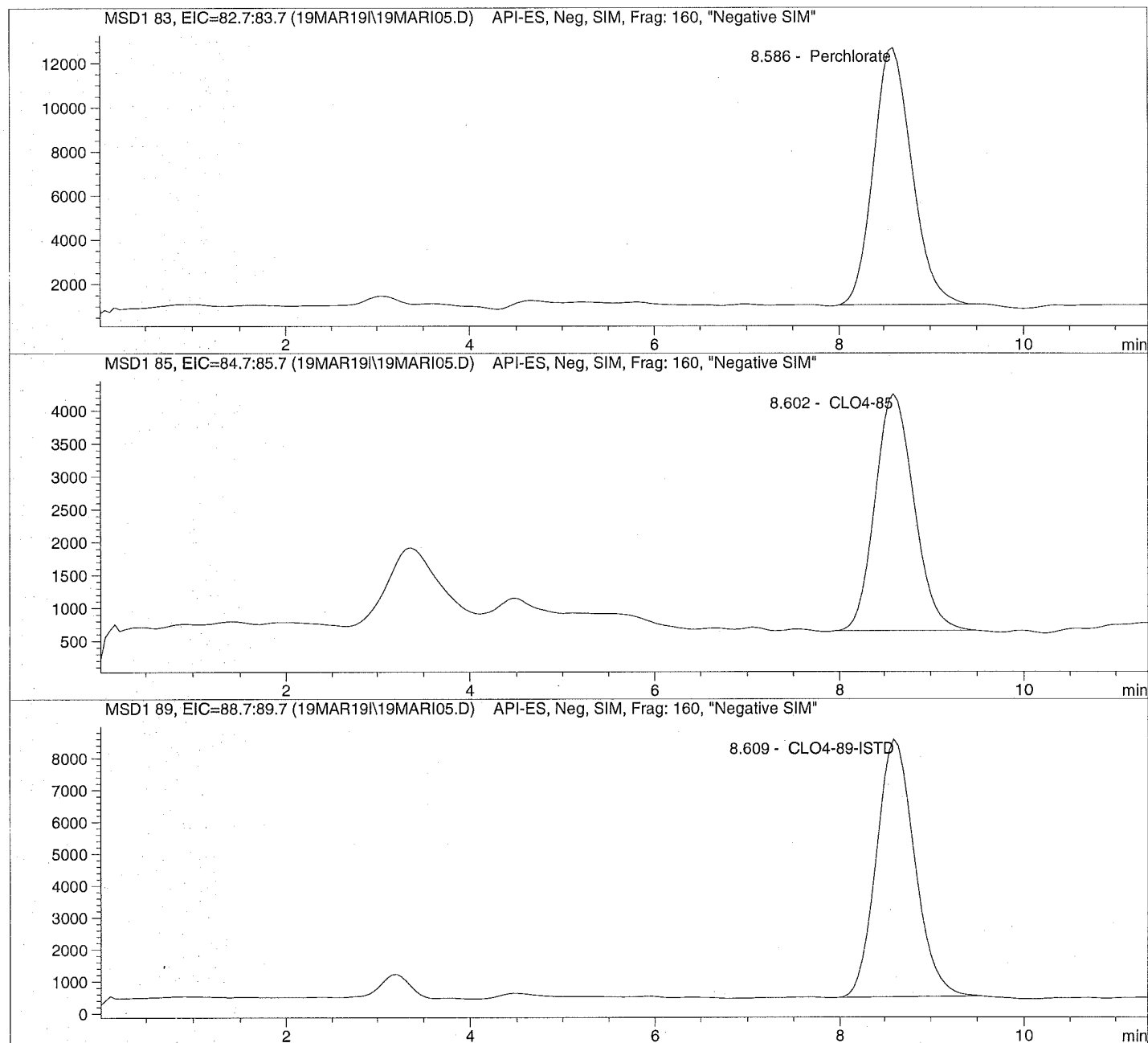
Sample Name: CLO4@ 5.0ug/L

Injection Date: 3/19/2019 10:06:16
Sample Name: CLO4@ 5.0ug/L
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line: 5
Sample Name:    CLO4@ 5.0ug/L          Location:  Vial 75
Acq Operator:  TNB                    Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 5.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D

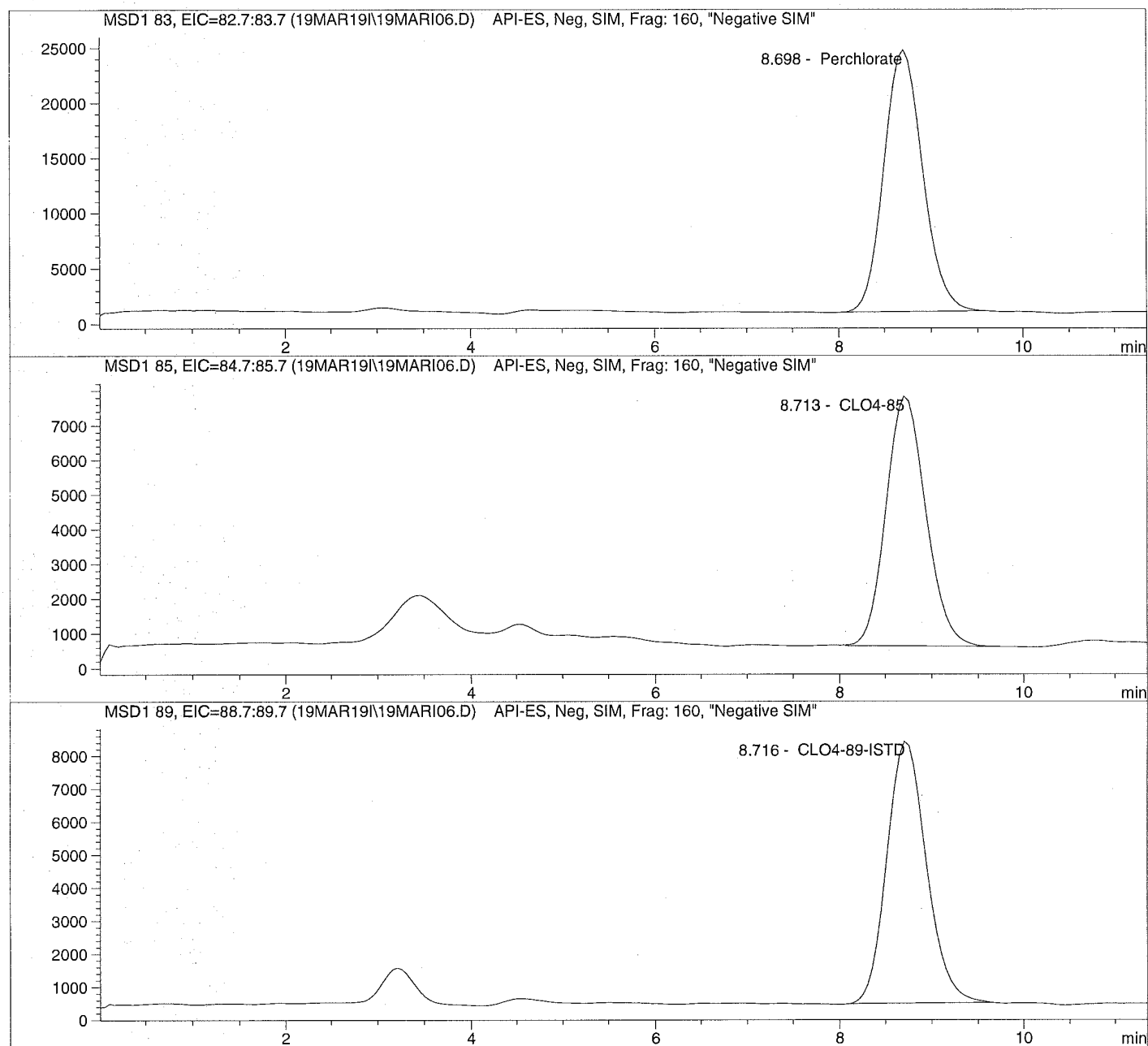
Sample Name: CLO4@ 10.ug/L

Injection Date: 3/19/2019 10:19:32
Sample Name: CLO4@ 10.ug/L
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D

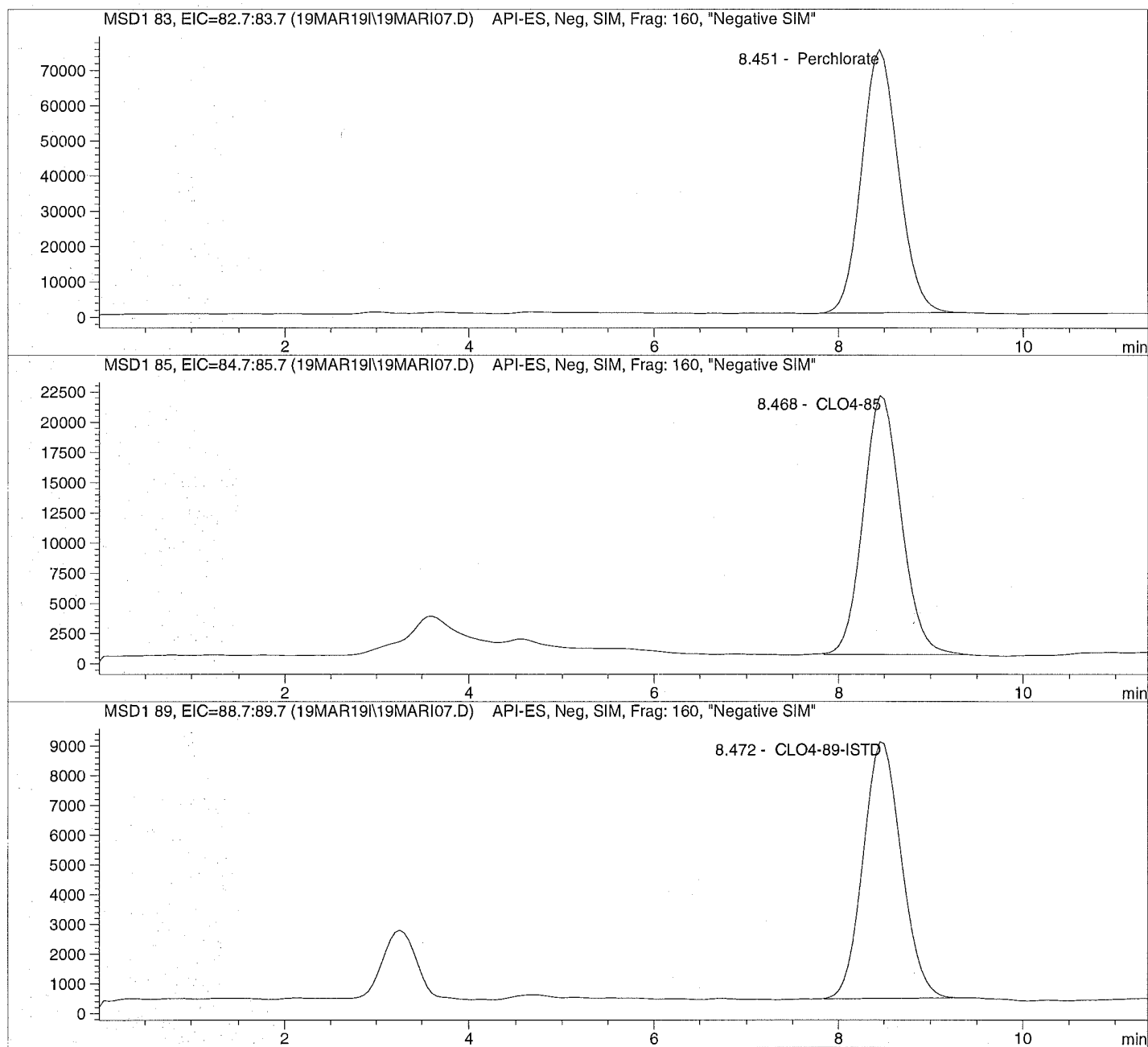
Sample Name: CLO4@ 25.ug/L

Injection Date: 3/19/2019 10:32:49
Sample Name: CLO4@ 25.ug/L
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 3/19/2019 10:32:49      Seq Line: 7
Sample Name:    CLO4@ 25.ug/L          Location:  Vial 77
Acq Operator:   TNB                   Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D

Sample Name: CLO4@ 50.ug/L

Injection Date: 3/19/2019 10:46:05

Seq Line: 8

Sample Name: CLO4@ 50.ug/L

Location: Vial 78

Acq Operator: TNB

Inj. No.: 1

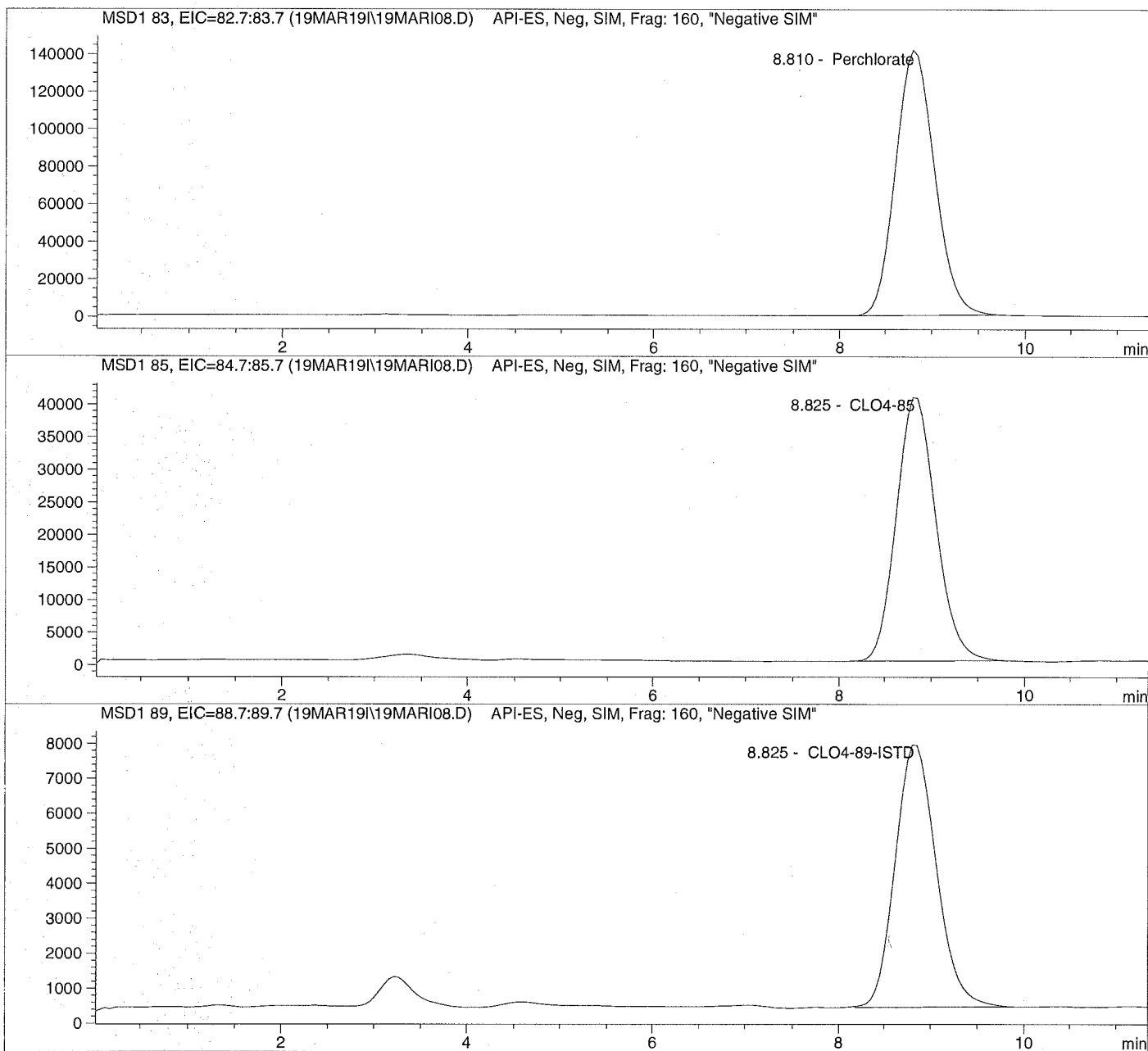
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L          Location:      Vial 78
Acq Operator:   TNB                    Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  50.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```


Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 3/19/2019 10:59:22

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

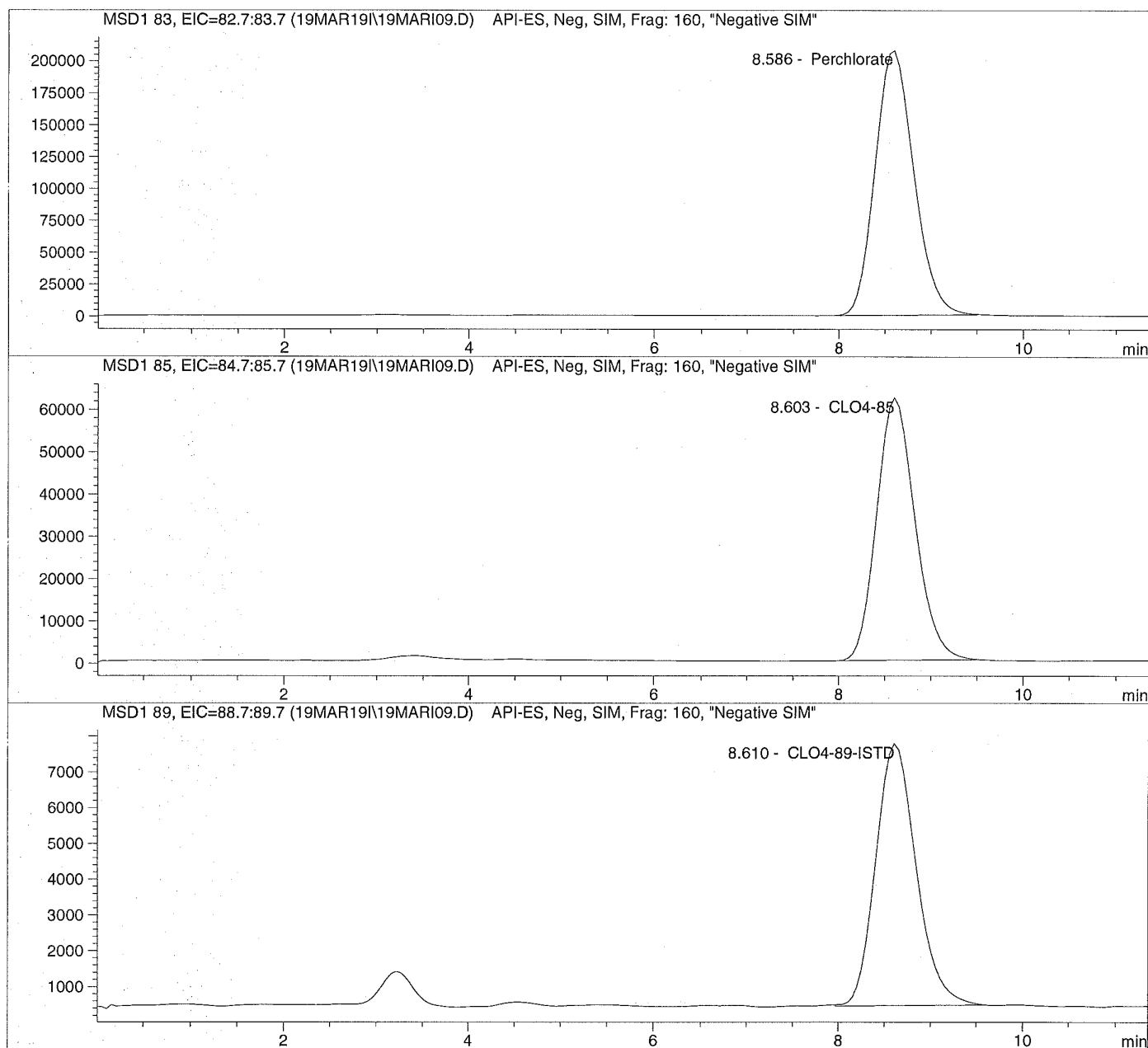
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

=====
 Injection Date: 3/19/2019 10:59:22 Seq Line: 9
 Sample Name: CLO4@ 75.ug/L Location: Vial 79
 Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
 Last Changed: 3/19/2019 14:35:22

Perchlorate analysis

=====
 Sample Information
 =====

Sorted By: Signal
 Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
 Multiplier: 1.000000
 Dilution: 1.000000
 Sample Amount: 75.000

=====
 LCMS Results
 =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

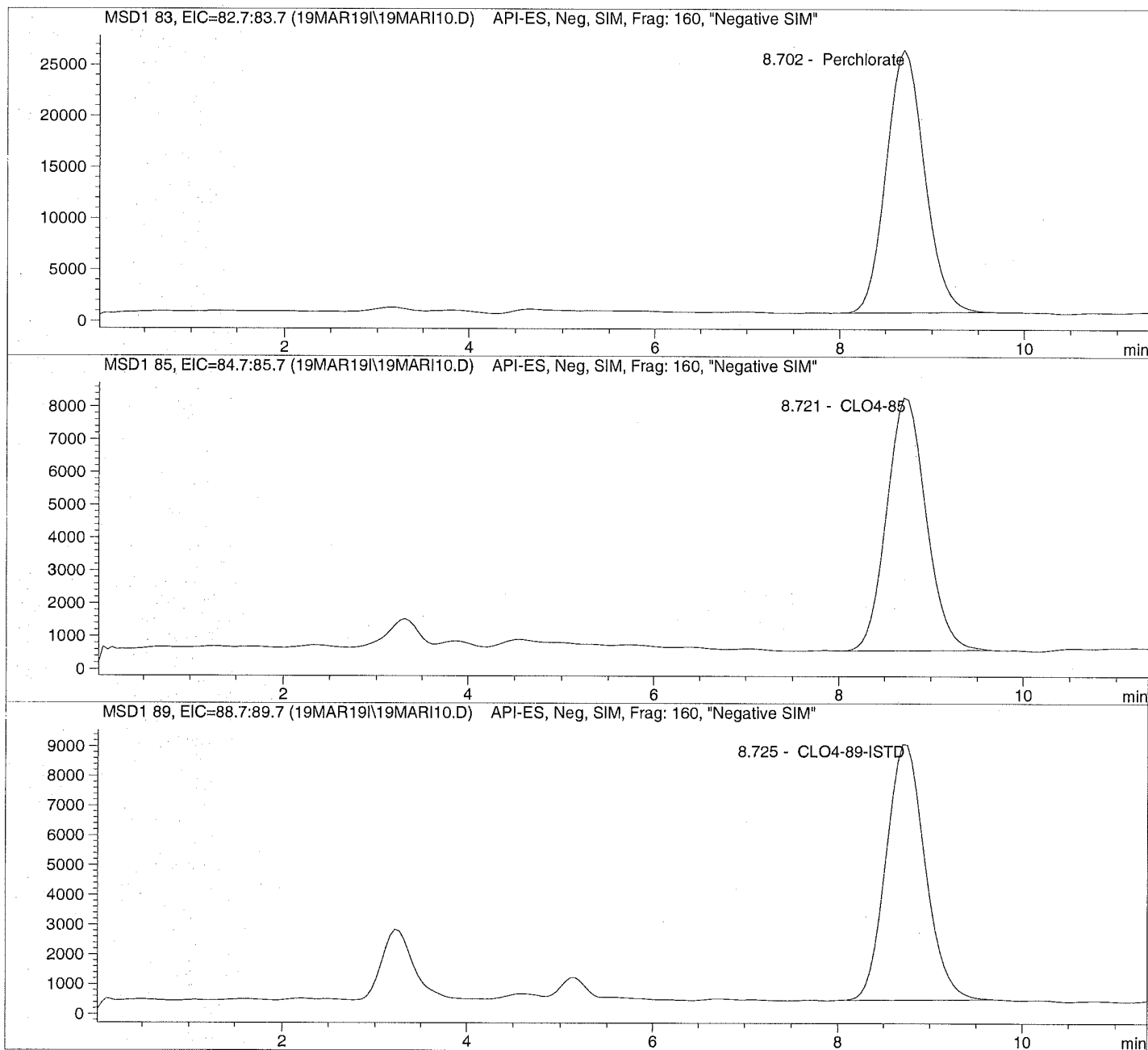
=====
 *** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```
=====
Injection Date: 3/19/2019 11:12:42      Seq Line:      10
Sample Name:    ICAL Verf@10ug/L        Location:      Vial 80
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 3/19/2019 11:12:42      Seq Line: 10
Sample Name: ICAL Verf@10ug/L      Location: Vial 80
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

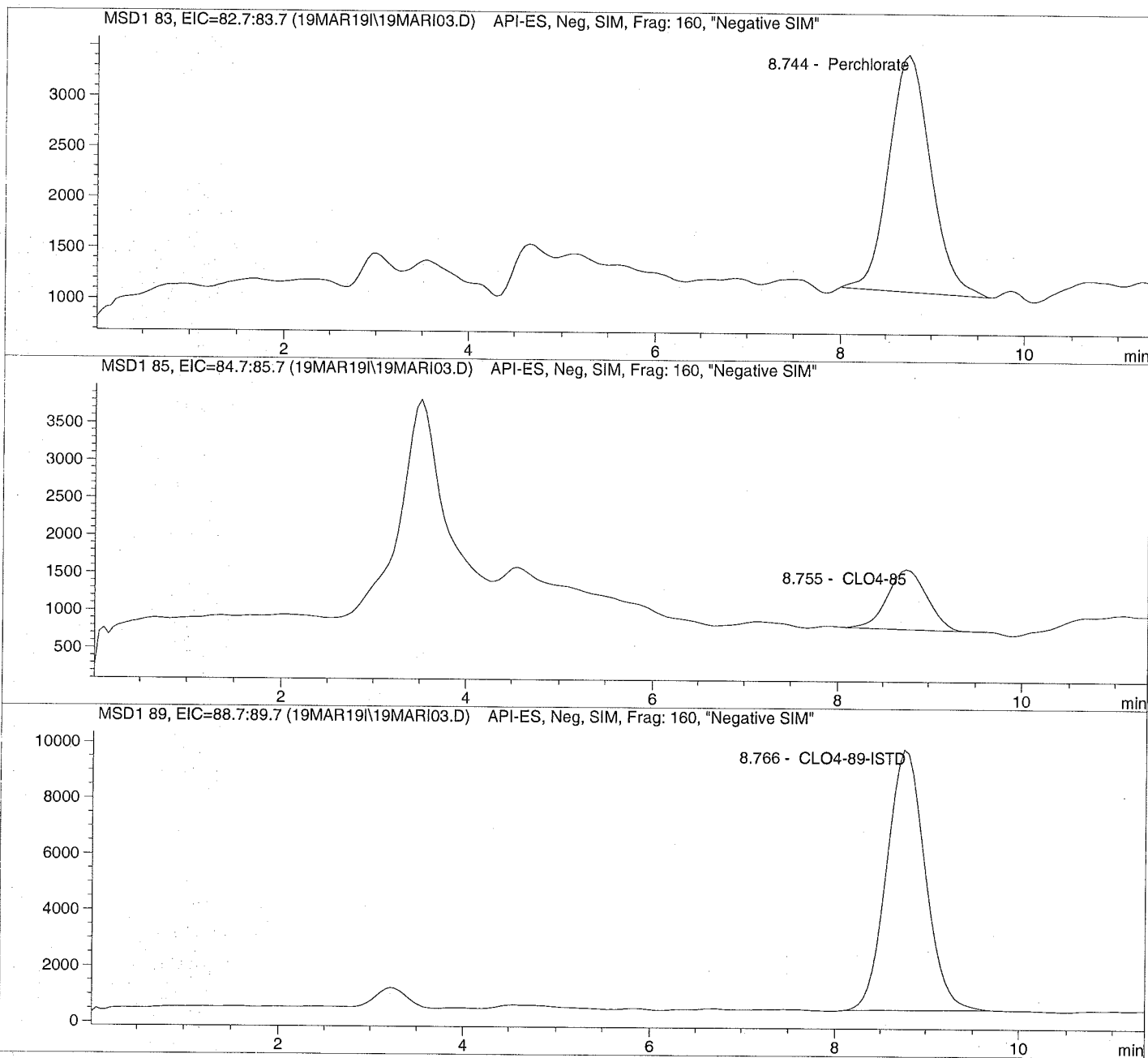
Sample Name: CLO4@ 1.0ug/L

=====
Injection Date: 3/19/2019 09:39:40
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date:  3/19/2019  09:39:40      Seq Line:                    3
Sample Name:    CLO4@ 1.0ug/L            Location:                    Vial 73
Acq Operator:   TNB                        Inj. No.:                    1
                                          Inj. Vol.:                    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019  14:38:25
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:                    Signal
Calib. Data Modified:        Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                  1.000000
Dilution:                    1.000000
Sample Amount:                1.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

*** End of Report ***



10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
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April 12, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19031511**

Laboratory Results for: **LH18/24 Longhorn GW Treatment Plant Weekly Samples**

Dear Marcia,

ALS Environmental received 3 sample(s) on Mar 28, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: JUMOKE.LAWAL
RJ Modashia
Project Manager

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031511

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19031511-01	LH18/24-SP650_032719	Water		27-Mar-2019 14:00	28-Mar-2019 08:30	<input type="checkbox"/>
HS19031511-02	LH18/24-SP650_032719_BIX	Water		27-Mar-2019 14:00	28-Mar-2019 08:30	<input type="checkbox"/>
HS19031511-03	Trip Blank -ALS-020119-75	Water		27-Mar-2019 00:00	28-Mar-2019 08:30	<input type="checkbox"/>

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order:

CASE NARRATIVE

Work Order Comments

- The analysis for Perchlorate was subcontracted to ALS Salt Lake City, UT. Final report attached.

GCMS Semivolatiles by Method SW8270SIM**Batch ID: 139415****Sample ID: LH18/24-SP650_032719 (HS19031511-01)**

- The surrogate recoveries could not be determined due to dilution below the calibration range.

GCMS Volatiles by Method SW8260**Batch ID: R335882****Sample ID: CCV**

- 1,2-Dibromoethane, Bromoform and Dibromochloromethane exceeded %D limits for CCV. Samples are ND for these compounds.

Sample ID: HS19040036-36MS

- MS and MSD are for an unrelated sample

Metals by Method SW6020**Batch ID: 139323****Sample ID: HS19031584-01MS**

- MS/MSD and DUPs are for an unrelated sample

Metals by Method SW7470**Batch ID: 139318**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

WetChemistry by Method E1664A**Batch ID: R336080**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

WetChemistry by Method SW9056**Batch ID: R335875**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.

WetChemistry by Method E410.4**Batch ID: R335804**

- The test results meet requirements of the current NELAP standards, state requirements or programs where applicable.
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ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032719
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031511
 Lab ID:HS19031511-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2-Dichloroethane	0.57	J	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	03-Apr-2019 17:41	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	03-Apr-2019 17:41	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	03-Apr-2019 17:41	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	03-Apr-2019 17:41	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	03-Apr-2019 17:41	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032719
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031511
 Lab ID:HS19031511-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED		
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
cis-1,2-Dichloroethene	2.4		0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	03-Apr-2019 17:41		
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	03-Apr-2019 17:41		
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	03-Apr-2019 17:41		
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Trichloroethene	0.85	J	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:41		
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>89.2</i>			0	<i>81-118</i>	%REC	<i>1</i>	<i>03-Apr-2019 17:41</i>		
<i>Surr: 4-Bromofluorobenzene</i>	<i>99.6</i>			0	<i>85-114</i>	%REC	<i>1</i>	<i>03-Apr-2019 17:41</i>		
<i>Surr: Dibromofluoromethane</i>	<i>91.3</i>			0	<i>80-119</i>	%REC	<i>1</i>	<i>03-Apr-2019 17:41</i>		
<i>Surr: Toluene-d8</i>	<i>107</i>			0	<i>89-112</i>	%REC	<i>1</i>	<i>03-Apr-2019 17:41</i>		
SEMIVOLATILES SIM		Method:SW8270SIM							Prep:SW3510 / 03-Apr-2019	Analyst: QX
1,4-Dioxane	11		1.0	1.0	1.0	ug/L	100	03-Apr-2019 14:53		
<i>Surr: 2-Fluorobiphenyl</i>	<i>0</i>	<i>S</i>		0	<i>40-140</i>	%REC	<i>100</i>	<i>03-Apr-2019 14:53</i>		
<i>Surr: 4-Terphenyl-d14</i>	<i>0</i>	<i>S</i>		0	<i>40-140</i>	%REC	<i>100</i>	<i>03-Apr-2019 14:53</i>		
<i>Surr: Nitrobenzene-d5</i>	<i>0</i>	<i>S</i>		0	<i>40-140</i>	%REC	<i>100</i>	<i>03-Apr-2019 14:53</i>		

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032719
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT
 WorkOrder:HS19031511
 Lab ID:HS19031511-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
ICP-MS METALS BY SW6020A		Method:SW6020				Prep:SW3010A / 01-Apr-2019		Analyst: JHD
Aluminum	0.0311		0.00180	0.00500	0.0100	mg/L	1	03-Apr-2019 13:24
Antimony	0.00101	J	0.000400	0.000500	0.00200	mg/L	1	03-Apr-2019 13:24
Arsenic	0.000860	J	0.000400	0.000500	0.00200	mg/L	1	03-Apr-2019 13:24
Barium	0.132		0.00190	0.00250	0.00400	mg/L	1	03-Apr-2019 13:24
Beryllium	0.00250	U	0.000200	0.00250	0.00200	mg/L	1	03-Apr-2019 13:24
Cadmium	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	03-Apr-2019 13:24
Calcium	14.5		0.0340	0.0500	0.500	mg/L	1	03-Apr-2019 13:24
Chromium	0.00142	J	0.000400	0.000500	0.00400	mg/L	1	03-Apr-2019 13:24
Cobalt	0.00112	J	0.000200	0.000500	0.00500	mg/L	1	03-Apr-2019 13:24
Iron	0.125	J	0.0120	0.0500	0.200	mg/L	1	03-Apr-2019 13:24
Lead	0.00100	U	0.000600	0.00100	0.00200	mg/L	1	03-Apr-2019 13:24
Magnesium	12.9		0.0100	0.0500	0.200	mg/L	1	03-Apr-2019 13:24
Manganese	0.0678		0.000700	0.00100	0.00500	mg/L	1	03-Apr-2019 13:24
Nickel	0.00344		0.000600	0.00100	0.00200	mg/L	1	03-Apr-2019 13:24
Potassium	1.27		0.0180	0.0500	0.200	mg/L	1	03-Apr-2019 13:24
Selenium	0.00250	U	0.00110	0.00250	0.00200	mg/L	1	03-Apr-2019 13:24
Silver	0.000500	U	0.000200	0.000500	0.00200	mg/L	1	03-Apr-2019 13:24
Sodium	339		0.140	0.500	2.00	mg/L	10	02-Apr-2019 16:04
Thallium	0.00100	U	0.000200	0.00100	0.00200	mg/L	1	03-Apr-2019 13:24
Vanadium	0.00382	J	0.000600	0.00100	0.00500	mg/L	1	03-Apr-2019 13:24
Zinc	0.0363		0.00200	0.00250	0.00400	mg/L	1	03-Apr-2019 13:24
MERCURY BY SW7470A		Method:SW7470				Prep:SW7470 / 01-Apr-2019		Analyst: FO
Mercury	0.000100	U	0.0000300	0.000100	0.000200	mg/L	1	01-Apr-2019 17:54
OIL & GREASE (HEM) BY E1664A		Method:E1664A						Analyst: KAH
Oil and Grease	2.08		0.610	1.00	2.00	mg/L	1	05-Apr-2019 14:45
CHEMICAL OXYGEN DEMAND BY E410.4		Method:E410.4						Analyst: AJH
Chemical Oxygen Demand	21.0		5.00	5.00	15.0	mg/L	1	02-Apr-2019 16:45
ANIONS BY SW9056A		Method:SW9056						Analyst: KMU
Chloride	317		2.00	5.00	5.00	mg/L	10	03-Apr-2019 05:57
Sulfate	28.5		2.00	5.00	5.00	mg/L	10	03-Apr-2019 05:57

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: LH18/24-SP650_032719_BIX
 Collection Date: 27-Mar-2019 14:00

ANALYTICAL REPORT

WorkOrder:HS19031511
 Lab ID:HS19031511-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA		Analyst: SUB				
Subcontract Analysis	See Attached		0	0		NA	1	12-Apr-2019 18:22

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: Trip Blank -ALS-020119-75
 Collection Date: 27-Mar-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19031511
 Lab ID:HS19031511-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,1-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	03-Apr-2019 17:17	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	03-Apr-2019 17:17	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	03-Apr-2019 17:17	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	03-Apr-2019 17:17	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	03-Apr-2019 17:17	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
 Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
 Sample ID: Trip Blank -ALS-020119-75
 Collection Date: 27-Mar-2019 00:00

ANALYTICAL REPORT
 WorkOrder:HS19031511
 Lab ID:HS19031511-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD		Method:SW8260							Analyst: PC
8260C									
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
cis-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	03-Apr-2019 17:17	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	03-Apr-2019 17:17	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	03-Apr-2019 17:17	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Trichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	03-Apr-2019 17:17	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>88.6</i>			<i>0</i>	<i>81-118</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 17:17</i>	
<i>Surr: 4-Bromofluorobenzene</i>	<i>96.5</i>			<i>0</i>	<i>85-114</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 17:17</i>	
<i>Surr: Dibromofluoromethane</i>	<i>89.9</i>			<i>0</i>	<i>80-119</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 17:17</i>	
<i>Surr: Toluene-d8</i>	<i>108</i>			<i>0</i>	<i>89-112</i>	<i>%REC</i>	<i>1</i>	<i>03-Apr-2019 17:17</i>	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

WEIGHT LOG

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

Batch ID: 139318 **Method:** MERCURY BY SW7470A **Prep:** HG_WPR

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19031511-01	1	10 (mL)	10 (mL)	1

Batch ID: 139323 **Method:** ICP-MS METALS BY SW6020A **Prep:** 3010A

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19031511-01	1	10	10 (mL)	1

Batch ID: 139415 **Method:** SEMIVOLATILES SIM **Prep:** 3510_B_SIM

SampleID	Container	Sample Wt/Vol	Final Volume	Prep Factor
HS19031511-01	1	1000	1 (mL)	0.001

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID 139318	Test Name : MERCURY BY SW7470A				Matrix: Water	
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00		01 Apr 2019 11:00	01 Apr 2019 17:54	1
Batch ID 139323	Test Name : ICP-MS METALS BY SW6020A				Matrix: Water	
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00		01 Apr 2019 12:30	03 Apr 2019 13:24	1
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00		01 Apr 2019 12:30	02 Apr 2019 16:04	10
Batch ID 139415	Test Name : SEMIVOLATILES SIM				Matrix: Water	
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00		03 Apr 2019 10:00	03 Apr 2019 14:53	100
Batch ID R335804	Test Name : CHEMICAL OXYGEN DEMAND BY E410.4				Matrix: Water	
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00			02 Apr 2019 16:45	1
Batch ID R335875	Test Name : ANIONS BY SW9056A				Matrix: Water	
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00			03 Apr 2019 05:57	10
Batch ID R335882	Test Name : VOLATILES ORGANICS BY METHOD 8260C				Matrix: Water	
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00			03 Apr 2019 17:41	1
HS19031511-03	Trip Blank -ALS-020119-75	27 Mar 2019 00:00			03 Apr 2019 17:17	1
Batch ID R336080	Test Name : OIL & GREASE (HEM) BY E1664A				Matrix: Water	
HS19031511-01	LH18/24-SP650_032719	27 Mar 2019 14:00			05 Apr 2019 14:45	1
Batch ID R336536	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)				Matrix: Water	
HS19031511-02	LH18/24-SP650_032719_BIX	27 Mar 2019 14:00			12 Apr 2019 18:22	1

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139318 (0)		Instrument: HG03		Method: MERCURY BY SW7470A						
MBLK	Sample ID: MBLK-139318	Units: mg/L		Analysis Date: 01-Apr-2019 17:33						
Client ID:	Run ID: HG03_335717	SeqNo: 5016992		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.000100	0.000200							U	
LCS	Sample ID: LCS-139318	Units: mg/L		Analysis Date: 01-Apr-2019 17:35						
Client ID:	Run ID: HG03_335717	SeqNo: 5016993		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.00509	0.000200	0.005	0	102	80 - 120				
MS	Sample ID: HS19031507-01MS	Units: mg/L		Analysis Date: 01-Apr-2019 17:38						
Client ID:	Run ID: HG03_335717	SeqNo: 5016995		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.00514	0.000200	0.005	-0.000029	103	75 - 125				
MSD	Sample ID: HS19031507-01MSD	Units: mg/L		Analysis Date: 01-Apr-2019 17:40						
Client ID:	Run ID: HG03_335717	SeqNo: 5016996		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
Mercury	0.00518	0.000200	0.005	-0.000029	104	75 - 125	0.00514	0.775	20	

The following samples were analyzed in this batch: HS19031511-01

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
MBLK	Sample ID: MBLK-139323	Units: mg/L			Analysis Date: 03-Apr-2019 13:13					
Client ID:	Run ID: ICPMS05_335865	SeqNo: 5020436	PrepDate: 01-Apr-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.00500	0.0100								U
Antimony	0.000500	0.00200								U
Arsenic	0.000500	0.00200								U
Barium	0.00250	0.00400								U
Beryllium	0.00250	0.00200								U
Cadmium	0.000500	0.00200								U
Calcium	0.0500	0.500								U
Chromium	0.000500	0.00400								U
Cobalt	0.000500	0.00500								U
Iron	0.0500	0.200								U
Lead	0.00100	0.00200								U
Magnesium	0.0500	0.200								U
Manganese	0.00100	0.00500								U
Nickel	0.00100	0.00200								U
Potassium	0.0500	0.200								U
Selenium	0.00250	0.00200								U
Silver	0.000500	0.00200								U
Sodium	0.0500	0.200								U
Thallium	0.00100	0.00200								U
Vanadium	0.001347	0.00500								J
Zinc	0.00250	0.00400								U

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
LCS	Sample ID: LCS-139323	Units: mg/L			Analysis Date: 02-Apr-2019 15:47					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018925	PrepDate: 01-Apr-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.1079	0.0100	0.1	0	108	80 - 120				
Antimony	0.05106	0.00200	0.05	0	102	80 - 120				
Arsenic	0.05071	0.00200	0.05	0	101	80 - 120				
Barium	0.05062	0.00400	0.05	0	101	80 - 120				
Beryllium	0.04871	0.00200	0.05	0	97.4	80 - 120				
Cadmium	0.05226	0.00200	0.05	0	105	80 - 120				
Calcium	5.088	0.500	5	0	102	80 - 120				
Chromium	0.05165	0.00400	0.05	0	103	80 - 120				
Cobalt	0.05122	0.00500	0.05	0	102	80 - 120				
Iron	5.174	0.200	5	0	103	80 - 120				
Lead	0.04996	0.00200	0.05	0	99.9	80 - 120				
Magnesium	4.996	0.200	5	0	99.9	80 - 120				
Manganese	0.05141	0.00500	0.05	0	103	80 - 120				
Nickel	0.05127	0.00200	0.05	0	103	80 - 120				
Potassium	4.971	0.200	5	0	99.4	80 - 120				
Selenium	0.05018	0.00200	0.05	0	100	80 - 120				
Silver	0.05235	0.00200	0.05	0	105	80 - 120				
Sodium	5.458	0.200	5	0	109	80 - 120				
Thallium	0.04811	0.00200	0.05	0	96.2	80 - 120				
Vanadium	0.05077	0.00500	0.05	0	102	80 - 120				
Zinc	0.05274	0.00400	0.05	0	105	80 - 120				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
MS	Sample ID: HS19031584-01MS	Units: mg/L			Analysis Date: 02-Apr-2019 15:53					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018928	PrepDate: 01-Apr-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.106	0.0100	0.1	0.004897	101	80 - 120				
Antimony	0.05139	0.00200	0.05	0.000336	102	80 - 120				
Arsenic	0.05303	0.00200	0.05	0.000014	106	80 - 120				
Barium	1.01	0.00400	0.05	0.9881	44.4	80 - 120				SO
Beryllium	0.04853	0.00200	0.05	0.000012	97.0	80 - 120				
Cadmium	0.04927	0.00200	0.05	0.000018	98.5	80 - 120				
Calcium	81.44	0.500	5	74.56	138	80 - 120				SO
Chromium	0.05181	0.00400	0.05	-0.00017	104	80 - 120				
Cobalt	0.0509	0.00500	0.05	0.000005	102	80 - 120				
Iron	5.334	0.200	5	0.2078	103	80 - 120				
Lead	0.04855	0.00200	0.05	0.000157	96.8	80 - 120				
Magnesium	28.82	0.200	5	23.36	109	80 - 120				O
Manganese	0.1757	0.00500	0.05	0.1227	106	80 - 120				
Nickel	0.05074	0.00200	0.05	0.000118	101	80 - 120				
Potassium	7.73	0.200	5	2.581	103	80 - 120				
Selenium	0.0507	0.00200	0.05	0.00072	100.0	80 - 120				
Silver	0.04762	0.00200	0.05	0.000047	95.1	80 - 120				
Sodium	360.5	0.200	5	354.5	119	80 - 120				EO
Thallium	0.04538	0.00200	0.05	0.000243	90.3	80 - 120				
Vanadium	0.05283	0.00500	0.05	0.000556	105	80 - 120				
Zinc	0.1307	0.00400	0.05	0.07703	107	80 - 120				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
MSD	Sample ID: HS19031584-01MSD	Units: mg/L			Analysis Date: 02-Apr-2019 15:56					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018929	PrepDate: 01-Apr-2019	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.1071	0.0100	0.1	0.004897	102	80 - 120	0.106	1.01	20	
Antimony	0.05105	0.00200	0.05	0.000336	101	80 - 120	0.05139	0.658	20	
Arsenic	0.05169	0.00200	0.05	0.000014	103	80 - 120	0.05303	2.56	20	
Barium	1.007	0.00400	0.05	0.9881	37.2	80 - 120	1.01	0.356	20	SO
Beryllium	0.04992	0.00200	0.05	0.000012	99.8	80 - 120	0.04853	2.83	20	
Cadmium	0.05007	0.00200	0.05	0.000018	100	80 - 120	0.04927	1.62	20	
Calcium	78.73	0.500	5	74.56	83.3	80 - 120	81.44	3.38	20	O
Chromium	0.05081	0.00400	0.05	-0.00017	102	80 - 120	0.05181	1.95	20	
Cobalt	0.04989	0.00500	0.05	0.000005	99.8	80 - 120	0.0509	2	20	
Iron	5.239	0.200	5	0.2078	101	80 - 120	5.334	1.78	20	
Lead	0.04939	0.00200	0.05	0.000157	98.5	80 - 120	0.04855	1.72	20	
Magnesium	28.59	0.200	5	23.36	105	80 - 120	28.82	0.802	20	O
Manganese	0.1711	0.00500	0.05	0.1227	96.7	80 - 120	0.1757	2.64	20	
Nickel	0.04944	0.00200	0.05	0.000118	98.6	80 - 120	0.05074	2.59	20	
Potassium	7.883	0.200	5	2.581	106	80 - 120	7.73	1.95	20	
Selenium	0.05281	0.00200	0.05	0.00072	104	80 - 120	0.0507	4.08	20	
Silver	0.04711	0.00200	0.05	0.000047	94.1	80 - 120	0.04762	1.07	20	
Sodium	357.6	0.200	5	354.5	60.2	80 - 120	360.5	0.816	20	SEO
Thallium	0.04656	0.00200	0.05	0.000243	92.6	80 - 120	0.04538	2.57	20	
Vanadium	0.05292	0.00500	0.05	0.000556	105	80 - 120	0.05283	0.161	20	
Zinc	0.1301	0.00400	0.05	0.07703	106	80 - 120	0.1307	0.456	20	

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05			Method: ICP-MS METALS BY SW6020A					
PDS	Sample ID: HS19031584-01PDS	Units: mg/L			Analysis Date: 02-Apr-2019 15:58					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018930		PrepDate: 01-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Aluminum	0.09986	0.0100	0.1	0.004897	95.0	75 - 125				
Antimony	0.08829	0.00200	0.1	0.000336	88.0	75 - 125				
Arsenic	0.0993	0.00200	0.1	0.000014	99.3	75 - 125				
Barium	1.062	0.00400	0.1	0.9881	74.1	75 - 125				SO
Beryllium	0.1001	0.00200	0.1	0.000012	100	75 - 125				
Cadmium	0.09637	0.00200	0.1	0.000018	96.4	75 - 125				
Calcium	80.82	0.500	10	74.56	62.6	75 - 125				SO
Chromium	0.09674	0.00400	0.1	-0.00017	96.9	75 - 125				
Cobalt	0.09603	0.00500	0.1	0.000005	96.0	75 - 125				
Iron	9.848	0.200	10	0.2078	96.4	75 - 125				
Lead	0.0916	0.00200	0.1	0.000157	91.4	75 - 125				
Magnesium	31.73	0.200	10	23.36	83.7	75 - 125				
Manganese	0.2125	0.00500	0.1	0.1227	89.8	75 - 125				
Nickel	0.09352	0.00200	0.1	0.000118	93.4	75 - 125				
Potassium	12.44	0.200	10	2.581	98.6	75 - 125				
Selenium	0.09928	0.00200	0.1	0.00072	98.6	75 - 125				
Silver	0.08785	0.00200	0.1	0.000047	87.8	75 - 125				
Thallium	0.08932	0.00200	0.1	0.000243	89.1	75 - 125				
Vanadium	0.09766	0.00500	0.1	0.000556	97.1	75 - 125				
Zinc	0.1737	0.00400	0.1	0.07703	96.7	75 - 125				
PDS	Sample ID: HS19031584-01PDS	Units: mg/L			Analysis Date: 03-Apr-2019 13:20					
Client ID:	Run ID: ICPMS05_335865	SeqNo: 5020439		PrepDate: 01-Apr-2019		DF: 10				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Sodium	430.6	2.00	100	334.6	96.0	75 - 125				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139323 (0)		Instrument: ICPMS05		Method: ICP-MS METALS BY SW6020A						
SD	Sample ID: HS19031584-01SD	Units: mg/L			Analysis Date: 02-Apr-2019 15:51					
Client ID:	Run ID: ICPMS05_335757	SeqNo: 5018927	PrepDate: 01-Apr-2019	DF: 5						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	%D Limit	Qual
Aluminum	0.009832	0.0500					0.004897	0	10	J
Antimony	0.00250	0.0100					0.000336	0	10	U
Arsenic	0.00250	0.0100					0.000014	0	10	U
Barium	0.9199	0.0200					0.9881	6.9	10	
Beryllium	0.0125	0.0100					0.000012	0	10	U
Cadmium	0.00250	0.0100					0.000018	0	10	U
Calcium	70.09	2.50					74.56	5.99	10	
Chromium	0.00250	0.0200					-0.00017	0	10	U
Cobalt	0.00250	0.0250					0.000005	0	10	U
Iron	0.2043	1.00					0.2078	0	10	J
Lead	0.00500	0.0100					0.000157	0	10	U
Magnesium	21.3	1.00					23.36	8.79	10	
Manganese	0.1188	0.0250					0.1227	3.19	10	
Nickel	0.00500	0.0100					0.000118	0	10	U
Potassium	2.471	1.00					2.581	4.24	10	
Selenium	0.0125	0.0100					0.00072	0	10	U
Silver	0.00250	0.0100					0.000047	0	10	U
Thallium	0.00500	0.0100					0.000243	0	10	U
Vanadium	0.005215	0.0250					0.000556	0	10	J
Zinc	0.07308	0.0200					0.07703	5.12	10	
SD	Sample ID: HS19031584-01SD	Units: mg/L			Analysis Date: 03-Apr-2019 13:18					
Client ID:	Run ID: ICPMS05_335865	SeqNo: 5020438	PrepDate: 01-Apr-2019	DF: 50						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%D	%D Limit	Qual
Sodium	340.7	10.0					334.6	1.83	10	

The following samples were analyzed in this batch: HS19031511-01

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: 139415 (0)		Instrument: SV-5		Method: SEMIVOLATILES SIM						
MBLK	Sample ID: MBLK-139415	Units: ug/L			Analysis Date: 03-Apr-2019 12:47					
Client ID:	Run ID: SV-5_335930	SeqNo: 5021779		PrepDate: 03-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
1,4-Dioxane	0.010	0.010							U	
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.09201</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>115</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.07352</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>91.9</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.08203</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>103</i>	<i>40 - 140</i>				
LCS	Sample ID: LCS-139415	Units: ug/L			Analysis Date: 03-Apr-2019 13:08					
Client ID:	Run ID: SV-5_335930	SeqNo: 5021780		PrepDate: 03-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
1,4-Dioxane	0.07995	0.010	0.08	0	99.9	40 - 140				
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.08927</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>112</i>	<i>40 - 140</i>				
<i>Surr: 4-Terphenyl-d14</i>	<i>0.07072</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>88.4</i>	<i>40 - 140</i>				
<i>Surr: Nitrobenzene-d5</i>	<i>0.07169</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>89.6</i>	<i>40 - 140</i>				
LCSD	Sample ID: LCSD-139415	Units: ug/L			Analysis Date: 03-Apr-2019 13:29					
Client ID:	Run ID: SV-5_335930	SeqNo: 5021781		PrepDate: 03-Apr-2019		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	RPD %RPD	RPD Limit Qual	
1,4-Dioxane	0.08058	0.010	0.08	0	101	40 - 140	0.07995	0.787	20	
<i>Surr: 2-Fluorobiphenyl</i>	<i>0.08169</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>102</i>	<i>40 - 140</i>	<i>0.08927</i>	<i>8.86</i>	<i>20</i>	
<i>Surr: 4-Terphenyl-d14</i>	<i>0.07017</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>87.7</i>	<i>40 - 140</i>	<i>0.07072</i>	<i>0.788</i>	<i>20</i>	
<i>Surr: Nitrobenzene-d5</i>	<i>0.07503</i>	<i>0</i>	<i>0.08</i>	<i>0</i>	<i>93.8</i>	<i>40 - 140</i>	<i>0.07169</i>	<i>4.56</i>	<i>20</i>	
The following samples were analyzed in this batch: HS19031511-01										

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 12:28					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020560	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	0.50	1.0								U
1,1,1-Trichloroethane	0.50	1.0								U
1,1,2,2-Tetrachloroethane	0.50	1.0								U
1,1,2-Trichloroethane	0.50	1.0								U
1,1-Dichloroethane	0.50	1.0								U
1,1-Dichloroethene	0.50	1.0								U
1,1-Dichloropropene	0.50	1.0								U
1,2,3-Trichlorobenzene	0.50	1.0								U
1,2,3-Trichloropropane	0.50	1.0								U
1,2,4-Trichlorobenzene	0.50	1.0								U
1,2,4-Trimethylbenzene	0.50	1.0								U
1,2-Dibromo-3-chloropropane	0.50	1.0								U
1,2-Dibromoethane	0.50	1.0								U
1,2-Dichlorobenzene	0.50	1.0								U
1,2-Dichloroethane	0.50	1.0								U
1,2-Dichloropropane	0.50	1.0								U
1,3,5-Trimethylbenzene	0.50	1.0								U
1,3-Dichlorobenzene	0.50	1.0								U
1,3-Dichloropropane	0.50	1.0								U
1,4-Dichlorobenzene	0.50	1.0								U
2,2-Dichloropropane	0.50	1.0								U
2-Butanone	1.0	2.0								U
2-Chlorotoluene	0.50	1.0								U
2-Hexanone	1.0	2.0								U
4-Chlorotoluene	0.50	1.0								U
4-Isopropyltoluene	0.50	1.0								U
4-Methyl-2-pentanone	1.0	2.0								U
Acetone	2.0	2.0								U
Benzene	0.50	1.0								U
Bromobenzene	0.50	1.0								U
Bromochloromethane	0.50	1.0								U
Bromodichloromethane	0.50	1.0								U
Bromoform	0.50	1.0								U
Bromomethane	0.50	1.0								U

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MBLK	Sample ID: VBLKW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 12:28					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020560	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	43.62	1.0	50	0	87.2	81 - 118				
Surr: 4-Bromofluorobenzene	49.43	1.0	50	0	98.9	85 - 114				
Surr: Dibromofluoromethane	45.1	1.0	50	0	90.2	80 - 119				
Surr: Toluene-d8	53	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 11:40					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020559	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	23.34	1.0	20	0	117	78 - 124				
1,1,1-Trichloroethane	19.93	1.0	20	0	99.7	74 - 131				
1,1,2,2-Tetrachloroethane	20.44	1.0	20	0	102	71 - 121				
1,1,2-Trichloroethane	22.06	1.0	20	0	110	80 - 119				
1,1-Dichloroethane	19.85	1.0	20	0	99.2	77 - 125				
1,1-Dichloroethene	19.49	1.0	20	0	97.4	71 - 131				
1,1-Dichloropropene	19.9	1.0	20	0	99.5	78 - 125				
1,2,3-Trichlorobenzene	21.21	1.0	20	0	106	69 - 129				
1,2,3-Trichloropropane	21.32	1.0	20	0	107	73 - 122				
1,2,4-Trichlorobenzene	21.68	1.0	20	0	108	69 - 130				
1,2,4-Trimethylbenzene	23.62	1.0	20	0	118	76 - 124				
1,2-Dibromo-3-chloropropane	21.63	1.0	20	0	108	62 - 128				
1,2-Dibromoethane	22.93	1.0	20	0	115	77 - 121				
1,2-Dichlorobenzene	21.63	1.0	20	0	108	80 - 119				
1,2-Dichloroethane	21.35	1.0	20	0	107	73 - 128				
1,2-Dichloropropane	20.9	1.0	20	0	104	78 - 122				
1,3,5-Trimethylbenzene	22.82	1.0	20	0	114	75 - 124				
1,3-Dichlorobenzene	21.78	1.0	20	0	109	80 - 119				
1,3-Dichloropropane	22.02	1.0	20	0	110	80 - 119				
1,4-Dichlorobenzene	21.6	1.0	20	0	108	79 - 118				
2,2-Dichloropropane	20.72	1.0	20	0	104	60 - 139				
2-Butanone	39.59	2.0	40	0	99.0	56 - 143				
2-Chlorotoluene	21.51	1.0	20	0	108	79 - 122				
2-Hexanone	42.13	2.0	40	0	105	57 - 139				
4-Chlorotoluene	21.85	1.0	20	0	109	78 - 122				
4-Isopropyltoluene	22.25	1.0	20	0	111	77 - 127				
4-Methyl-2-pentanone	41.02	2.0	40	0	103	67 - 130				
Acetone	36.89	2.0	40	0	92.2	39 - 160				
Benzene	21.72	1.0	20	0	109	79 - 120				
Bromobenzene	22.97	1.0	20	0	115	80 - 120				
Bromochloromethane	20.97	1.0	20	0	105	78 - 123				
Bromodichloromethane	22.16	1.0	20	0	111	79 - 125				
Bromoform	23.37	1.0	20	0	117	66 - 130				
Bromomethane	24.03	1.0	20	0	120	53 - 141				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
LCS	Sample ID: VLCSW-190403	Units: UG/L			Analysis Date: 03-Apr-2019 11:40					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020559	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	37.4	2.0	40	0	93.5	64 - 133				
Carbon tetrachloride	22.65	1.0	20	0	113	72 - 136				
Chlorobenzene	21.88	1.0	20	0	109	82 - 118				
Chloroethane	18.95	1.0	20	0	94.7	60 - 138				
Chloroform	20.52	1.0	20	0	103	79 - 124				
Chloromethane	19.74	1.0	20	0	98.7	50 - 139				
cis-1,2-Dichloroethene	20.43	1.0	20	0	102	78 - 123				
cis-1,3-Dichloropropene	20.84	1.0	20	0	104	75 - 124				
Dibromochloromethane	22.56	1.0	20	0	113	74 - 126				
Dibromomethane	21.6	1.0	20	0	108	79 - 123				
Dichlorodifluoromethane	21.85	1.0	20	0	109	32 - 152				
Ethylbenzene	23.41	1.0	20	0	117	79 - 121				
Hexachlorobutadiene	23.54	1.0	20	0	118	66 - 134				
Isopropylbenzene	22.97	1.0	20	0	115	72 - 131				
m,p-Xylene	45.47	2.0	40	0	114	80 - 121				
Methylene chloride	20.82	2.0	20	0	104	74 - 124				
Naphthalene	21.06	1.0	20	0	105	61 - 128				
n-Butylbenzene	22.89	1.0	20	0	114	75 - 128				
n-Propylbenzene	21.9	1.0	20	0	109	76 - 126				
o-Xylene	22.22	1.0	20	0	111	78 - 122				
sec-Butylbenzene	21.56	1.0	20	0	108	77 - 126				
Styrene	23.58	1.0	20	0	118	78 - 123				
tert-Butylbenzene	21.9	1.0	20	0	109	78 - 124				
Tetrachloroethene	22.75	1.0	20	0	114	74 - 129				
Toluene	22.69	1.0	20	0	113	80 - 121				
trans-1,2-Dichloroethene	20.31	1.0	20	0	102	75 - 124				
trans-1,3-Dichloropropene	22.21	1.0	20	0	111	73 - 127				
Trichloroethene	20.76	1.0	20	0	104	79 - 123				
Trichlorofluoromethane	19.87	1.0	20	0	99.3	65 - 141				
Vinyl chloride	18.34	1.0	20	0	91.7	58 - 137				
Surr: 1,2-Dichloroethane-d4	44.85	1.0	50	0	89.7	81 - 118				
Surr: 4-Bromofluorobenzene	49.44	1.0	50	0	98.9	85 - 114				
Surr: Dibromofluoromethane	46.66	1.0	50	0	93.3	80 - 119				
Surr: Toluene-d8	52.9	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19040036-36MS	Units: UG/L			Analysis Date: 03-Apr-2019 14:52					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020790	PrepDate:	DF: 1000						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	24370	1000	20000	0	122	78 - 124				
1,1,1-Trichloroethane	22280	1000	20000	0	111	74 - 131				
1,1,2,2-Tetrachloroethane	22750	1000	20000	0	114	71 - 121				
1,1,2-Trichloroethane	23590	1000	20000	0	118	80 - 119				
1,1-Dichloroethane	21090	1000	20000	0	105	77 - 125				
1,1-Dichloroethene	22220	1000	20000	0	111	71 - 131				
1,1-Dichloropropene	22920	1000	20000	0	115	78 - 125				
1,2,3-Trichlorobenzene	21690	1000	20000	0	108	69 - 129				
1,2,3-Trichloropropane	22780	1000	20000	0	114	73 - 122				
1,2,4-Trichlorobenzene	22700	1000	20000	0	114	69 - 130				
1,2,4-Trimethylbenzene	24950	1000	20000	0	125	76 - 124				S
1,2-Dibromo-3-chloropropane	23950	1000	20000	0	120	62 - 128				
1,2-Dibromoethane	23610	1000	20000	0	118	77 - 121				
1,2-Dichlorobenzene	23700	1000	20000	0	118	80 - 119				
1,2-Dichloroethane	23200	1000	20000	0	116	73 - 128				
1,2-Dichloropropane	21950	1000	20000	0	110	78 - 122				
1,3,5-Trimethylbenzene	24840	1000	20000	0	124	75 - 124				S
1,3-Dichlorobenzene	24090	1000	20000	0	120	80 - 119				S
1,3-Dichloropropane	23000	1000	20000	0	115	80 - 119				
1,4-Dichlorobenzene	23350	1000	20000	0	117	79 - 118				
2,2-Dichloropropane	22210	1000	20000	0	111	60 - 139				
2-Butanone	41410	2000	40000	0	104	56 - 143				
2-Chlorotoluene	23870	1000	20000	0	119	79 - 122				
2-Hexanone	44400	2000	40000	0	111	57 - 139				
4-Chlorotoluene	23820	1000	20000	0	119	78 - 122				
4-Isopropyltoluene	25150	1000	20000	0	126	77 - 127				
4-Methyl-2-pentanone	44310	2000	40000	0	111	67 - 130				
Acetone	39450	2000	40000	0	98.6	39 - 160				
Benzene	163700	1000	20000	136400	137	79 - 120				SO
Bromobenzene	24010	1000	20000	0	120	80 - 120				S
Bromochloromethane	20610	1000	20000	0	103	78 - 123				
Bromodichloromethane	22820	1000	20000	0	114	79 - 125				
Bromoform	23650	1000	20000	0	118	66 - 130				
Bromomethane	24600	1000	20000	0	123	53 - 141				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MS	Sample ID: HS19040036-36MS	Units: UG/L			Analysis Date: 03-Apr-2019 14:52					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020790	PrepDate:	DF: 1000						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	43460	2000	40000	0	109	64 - 133				
Carbon tetrachloride	25400	1000	20000	0	127	72 - 136				
Chlorobenzene	23430	1000	20000	0	117	82 - 118				
Chloroethane	20830	1000	20000	0	104	60 - 138				
Chloroform	21220	1000	20000	0	106	79 - 124				
Chloromethane	23190	1000	20000	0	116	50 - 139				
cis-1,2-Dichloroethene	21440	1000	20000	0	107	78 - 123				
cis-1,3-Dichloropropene	22680	1000	20000	0	113	75 - 124				
Dibromochloromethane	24100	1000	20000	0	120	74 - 126				
Dibromomethane	22210	1000	20000	0	111	79 - 123				
Dichlorodifluoromethane	22700	1000	20000	0	113	32 - 152				
Ethylbenzene	25690	1000	20000	1292	122	79 - 121				S
Hexachlorobutadiene	26030	1000	20000	0	130	66 - 134				
Isopropylbenzene	24720	1000	20000	0	124	72 - 131				
m,p-Xylene	53610	2000	40000	4701	122	80 - 121				S
Methylene chloride	22110	2000	20000	0	111	74 - 124				
Naphthalene	21950	1000	20000	0	110	61 - 128				
n-Butylbenzene	25170	1000	20000	0	126	75 - 128				
n-Propylbenzene	25030	1000	20000	0	125	76 - 126				
o-Xylene	26700	1000	20000	2345	122	78 - 122				
sec-Butylbenzene	24850	1000	20000	0	124	77 - 126				
Styrene	26250	1000	20000	1811	122	78 - 123				
tert-Butylbenzene	24760	1000	20000	0	124	78 - 124				
Tetrachloroethene	25340	1000	20000	0	127	74 - 129				
Toluene	52760	1000	20000	28020	124	80 - 121				S
trans-1,2-Dichloroethene	22170	1000	20000	0	111	75 - 124				
trans-1,3-Dichloropropene	22600	1000	20000	0	113	73 - 127				
Trichloroethene	24510	1000	20000	0	123	79 - 123				
Trichlorofluoromethane	23470	1000	20000	0	117	65 - 141				
Vinyl chloride	20510	1000	20000	0	103	58 - 137				
Surr: 1,2-Dichloroethane-d4	43910	1000	50000	0	87.8	81 - 118				
Surr: 4-Bromofluorobenzene	49050	1000	50000	0	98.1	85 - 114				
Surr: Dibromofluoromethane	45590	1000	50000	0	91.2	80 - 119				
Surr: Toluene-d8	52160	1000	50000	0	104	89 - 112				

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19040036-36MSD	Units: UG/L			Analysis Date: 03-Apr-2019 15:16					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020791	PrepDate:	DF: 1000						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	24320	1000	20000	0	122	78 - 124	24370	0.228	20	
1,1,1-Trichloroethane	21690	1000	20000	0	108	74 - 131	22280	2.69	20	
1,1,2,2-Tetrachloroethane	22810	1000	20000	0	114	71 - 121	22750	0.276	20	
1,1,2-Trichloroethane	23280	1000	20000	0	116	80 - 119	23590	1.3	20	
1,1-Dichloroethane	20610	1000	20000	0	103	77 - 125	21090	2.28	20	
1,1-Dichloroethene	21230	1000	20000	0	106	71 - 131	22220	4.59	20	
1,1-Dichloropropene	22490	1000	20000	0	112	78 - 125	22920	1.87	20	
1,2,3-Trichlorobenzene	22690	1000	20000	0	113	69 - 129	21690	4.52	20	
1,2,3-Trichloropropane	23960	1000	20000	0	120	73 - 122	22780	5.05	20	
1,2,4-Trichlorobenzene	22750	1000	20000	0	114	69 - 130	22700	0.208	20	
1,2,4-Trimethylbenzene	24690	1000	20000	0	123	76 - 124	24950	1.03	20	
1,2-Dibromo-3-chloropropane	24840	1000	20000	0	124	62 - 128	23950	3.68	20	
1,2-Dibromoethane	23750	1000	20000	0	119	77 - 121	23610	0.569	20	
1,2-Dichlorobenzene	23560	1000	20000	0	118	80 - 119	23700	0.565	20	
1,2-Dichloroethane	23300	1000	20000	0	116	73 - 128	23200	0.421	20	
1,2-Dichloropropane	21950	1000	20000	0	110	78 - 122	21950	0.0278	20	
1,3,5-Trimethylbenzene	24400	1000	20000	0	122	75 - 124	24840	1.78	20	
1,3-Dichlorobenzene	23770	1000	20000	0	119	80 - 119	24090	1.35	20	
1,3-Dichloropropane	23040	1000	20000	0	115	80 - 119	23000	0.159	20	
1,4-Dichlorobenzene	23460	1000	20000	0	117	79 - 118	23350	0.488	20	
2,2-Dichloropropane	21530	1000	20000	0	108	60 - 139	22210	3.12	20	
2-Butanone	42690	2000	40000	0	107	56 - 143	41410	3.06	20	
2-Chlorotoluene	23450	1000	20000	0	117	79 - 122	23870	1.75	20	
2-Hexanone	45910	2000	40000	0	115	57 - 139	44400	3.33	20	
4-Chlorotoluene	23550	1000	20000	0	118	78 - 122	23820	1.15	20	
4-Isopropyltoluene	24580	1000	20000	0	123	77 - 127	25150	2.31	20	
4-Methyl-2-pentanone	45920	2000	40000	0	115	67 - 130	44310	3.58	20	
Acetone	43390	2000	40000	0	108	39 - 160	39450	9.5	20	
Benzene	161100	1000	20000	136400	123	79 - 120	163700	1.64	20	SO
Bromobenzene	23980	1000	20000	0	120	80 - 120	24010	0.139	20	
Bromochloromethane	20980	1000	20000	0	105	78 - 123	20610	1.77	20	
Bromodichloromethane	22510	1000	20000	0	113	79 - 125	22820	1.39	20	
Bromoform	24000	1000	20000	0	120	66 - 130	23650	1.45	20	
Bromomethane	22690	1000	20000	0	113	53 - 141	24600	8.06	20	

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335882 (0)		Instrument: VOA6		Method: VOLATILES ORGANICS BY METHOD 8260C						
MSD	Sample ID: HS19040036-36MSD	Units: UG/L			Analysis Date: 03-Apr-2019 15:16					
Client ID:	Run ID: VOA6_335882	SeqNo: 5020791		PrepDate:		DF: 1000				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	41860	2000	40000	0	105	64 - 133	43460	3.75	20	
Carbon tetrachloride	24790	1000	20000	0	124	72 - 136	25400	2.42	20	
Chlorobenzene	23190	1000	20000	0	116	82 - 118	23430	1.07	20	
Chloroethane	19980	1000	20000	0	99.9	60 - 138	20830	4.15	20	
Chloroform	20990	1000	20000	0	105	79 - 124	21220	1.06	20	
Chloromethane	21200	1000	20000	0	106	50 - 139	23190	8.96	20	
cis-1,2-Dichloroethene	20790	1000	20000	0	104	78 - 123	21440	3.08	20	
cis-1,3-Dichloropropene	22920	1000	20000	0	115	75 - 124	22680	1.02	20	
Dibromochloromethane	24120	1000	20000	0	121	74 - 126	24100	0.0892	20	
Dibromomethane	22680	1000	20000	0	113	79 - 123	22210	2.1	20	
Dichlorodifluoromethane	21730	1000	20000	0	109	32 - 152	22700	4.35	20	
Ethylbenzene	25210	1000	20000	1292	120	79 - 121	25690	1.92	20	
Hexachlorobutadiene	26760	1000	20000	0	134	66 - 134	26030	2.75	20	
Isopropylbenzene	24210	1000	20000	0	121	72 - 131	24720	2.07	20	
m,p-Xylene	52410	2000	40000	4701	119	80 - 121	53610	2.27	20	
Methylene chloride	21630	2000	20000	0	108	74 - 124	22110	2.19	20	
Naphthalene	23380	1000	20000	0	117	61 - 128	21950	6.3	20	
n-Butylbenzene	24790	1000	20000	0	124	75 - 128	25170	1.55	20	
n-Propylbenzene	24480	1000	20000	0	122	76 - 126	25030	2.2	20	
o-Xylene	26170	1000	20000	2345	119	78 - 122	26700	2.01	20	
sec-Butylbenzene	24340	1000	20000	0	122	77 - 126	24850	2.07	20	
Styrene	26210	1000	20000	1811	122	78 - 123	26250	0.147	20	
tert-Butylbenzene	24250	1000	20000	0	121	78 - 124	24760	2.09	20	
Tetrachloroethene	24480	1000	20000	0	122	74 - 129	25340	3.45	20	
Toluene	51510	1000	20000	28020	117	80 - 121	52760	2.38	20	
trans-1,2-Dichloroethene	21410	1000	20000	0	107	75 - 124	22170	3.5	20	
trans-1,3-Dichloropropene	23180	1000	20000	0	116	73 - 127	22600	2.52	20	
Trichloroethene	23240	1000	20000	0	116	79 - 123	24510	5.32	20	
Trichlorofluoromethane	23010	1000	20000	0	115	65 - 141	23470	1.98	20	
Vinyl chloride	19890	1000	20000	0	99.5	58 - 137	20510	3.06	20	
Surr: 1,2-Dichloroethane-d4	43970	1000	50000	0	87.9	81 - 118	43910	0.129	20	
Surr: 4-Bromofluorobenzene	49970	1000	50000	0	99.9	85 - 114	49050	1.86	20	
Surr: Dibromofluoromethane	45550	1000	50000	0	91.1	80 - 119	45590	0.0906	20	
Surr: Toluene-d8	51790	1000	50000	0	104	89 - 112	52160	0.709	20	

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW**Batch ID:** R335882 (0)**Instrument:** VOA6**Method:** VOLATILES ORGANICS BY METHOD
8260C

The following samples were analyzed in this batch:

HS19031511-01	HS19031511-03
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ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID:	R335804 (0)	Instrument:	WetChem_HS	Method:	CHEMICAL OXYGEN DEMAND BY E410.4					
MBLK	Sample ID: MBLK-R335804	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018767	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	5.00	15.0								U
LCS	Sample ID: LCS-R335804	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018766	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	97	15.0	100	0	97.0	85 - 115				
MS	Sample ID: HS19040002-01MS	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5020059	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	60	15.0	50	7	106	80 - 120				
MS	Sample ID: HS19031461-01MS	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018770	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	94	15.0	50	37	114	80 - 120				
DUP	Sample ID: HS19040002-01DUP	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5020060	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	8	15.0					7	0	20	J
DUP	Sample ID: HS19031461-01DUP	Units:	mg/L	Analysis Date:	02-Apr-2019 16:45					
Client ID:	Run ID: WetChem_HS_335804	SeqNo:	5018771	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Chemical Oxygen Demand	37	15.0					37	0	20	

The following samples were analyzed in this batch: HS19031511-01

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID: R335875 (0)		Instrument: ICS2100		Method: ANIONS BY SW9056A						
MBLK	Sample ID: WBLKW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:16					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020414		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	0.500	0.500							U	
Sulfate	0.500	0.500							U	
LCS	Sample ID: WLCSW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:31					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020415		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	20.22	0.500	20	0	101	80 - 120				
Sulfate	19.98	0.500	20	0	99.9	80 - 120				
LCSD	Sample ID: WLCSDW2-040219	Units: mg/L			Analysis Date: 03-Apr-2019 01:46					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020416		PrepDate:			DF: 1			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	18.95	0.500	20	0	94.8	80 - 120	20.22	6.51	20	
Sulfate	18.67	0.500	20	0	93.3	80 - 120	19.98	6.79	20	
MS	Sample ID: HS19031013-07MS	Units: mg/L			Analysis Date: 03-Apr-2019 04:28					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020427		PrepDate:			DF: 50			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	1535	25.0	500	1090	89.1	80 - 120				
Sulfate	1565	25.0	500	1119	89.2	80 - 120				
MSD	Sample ID: HS19031013-07MSD	Units: mg/L			Analysis Date: 03-Apr-2019 04:43					
Client ID:	Run ID: ICS2100_335875	SeqNo: 5020428		PrepDate:			DF: 50			
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Chloride	1605	25.0	500	1090	103	80 - 120	1535	4.46	20	
Sulfate	1638	25.0	500	1119	104	80 - 120	1565	4.57	20	

The following samples were analyzed in this batch: HS19031511-01

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
WorkOrder: HS19031511

QC BATCH REPORT NEW

Batch ID:	R336080 (0)	Instrument:	Balance1	Method:	OIL & GREASE (HEM) BY E1664A					
MBLK	Sample ID: WBLKW-040519	Units:	mg/L	Analysis Date:	05-Apr-2019 14:45					
Client ID:	Run ID: Balance1_336080	SeqNo:	5024526	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	1.00	2.00							U	
LCS	Sample ID: WLCSW-040519	Units:	mg/L	Analysis Date:	05-Apr-2019 14:45					
Client ID:	Run ID: Balance1_336080	SeqNo:	5024528	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	38.8	2.00	40	0	97.0	78 - 114				
LCSD	Sample ID: WLCSDW-040519	Units:	mg/L	Analysis Date:	05-Apr-2019 14:45					
Client ID:	Run ID: Balance1_336080	SeqNo:	5024527	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	38.8	2.00	40	0	97.0	78 - 114	38.8	0	18	
MS	Sample ID: HS19031453-01MS	Units:	mg/L	Analysis Date:	05-Apr-2019 14:45					
Client ID:	Run ID: Balance1_336080	SeqNo:	5024506	PrepDate:	DF: 1					
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit Qual	
Oil and Grease	39.58	2.00	40	1.25	95.8	78 - 114				

The following samples were analyzed in this batch: HS19031511-01

ALS Houston, US

Date: 12-Apr-19

Client:	Bhate Environmental Associates, Inc.	QUALIFIERS, ACRONYMS, UNITS
Project:	LH18/24 Longhorn GW Treatment Plant Weekly Samples	
WorkOrder:	HS19031511	

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

Unit Reported	Description
mg/L	Milligrams per Liter

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019
Arkansas	19-028-0	27-Mar-2020

ALS Houston, US

Date: 12-Apr-19

Client: Bhate Environmental Associates, Inc.
Project: LH18/24 Longhorn GW Treatment Plant Weekly Samples
Work Order: HS19031511

SAMPLE TRACKING

Lab Samp ID	Client Sample ID	Action	Date	Person	New Location
HS19031511-01	LH18/24-SP650_032719	Login	3/29/2019 10:44:05 AM	NDR	Sub
HS19031511-01	LH18/24-SP650_032719	Login	3/29/2019 10:44:05 AM	NDR	WET169
HS19031511-01	LH18/24-SP650_032719	Login	3/29/2019 10:44:05 AM	NDR	EXT125
HS19031511-01	LH18/24-SP650_032719	Login	3/29/2019 10:44:05 AM	NDR	WET169
HS19031511-01	LH18/24-SP650_032719	Login	3/29/2019 10:44:05 AM	NDR	WET169
HS19031511-01	LH18/24-SP650_032719	Login	3/29/2019 10:44:05 AM	NDR	MET012
HS19031511-01	LH18/24-SP650_032719	Login	3/29/2019 10:44:05 AM	NDR	VOA230
HS19031511-03	Trip Blank -ALS-020119-75	Login	3/29/2019 10:44:05 AM	NDR	VOA230

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19031511

Date/Time Received: **28-Mar-2019 08:30**
 Received by: **PMG**

Checklist completed by: Nilesh D. Ranchod 29-Mar-2019
 eSignature | Date

Reviewed by: RJ Modashia 29-Mar-2019
 eSignature | Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

- Shipping container/cooler in good condition? Yes No Not Present
- Custody seals intact on shipping container/cooler? Yes No Not Present
- Custody seals intact on sample bottles? Yes No Not Present
- VOA/TX1005/TX1006 Solids in hermetically sealed vials? Yes No Not Present
- Chain of custody present? Yes No 1 Page(s)
- Chain of custody signed when relinquished and received? Yes No COC IDs:N/A
- Samplers name present on COC? Yes No
- Chain of custody agrees with sample labels? Yes No
- Samples in proper container/bottle? Yes No
- Sample containers intact? Yes No
- Sufficient sample volume for indicated test? Yes No
- All samples received within holding time? Yes No
- Container/Temp Blank temperature in compliance? Yes No

Temperature(s)/Thermometer(s): 1.5c UC/C | IR25
 Cooler(s)/Kit(s): 44581
 Date/Time sample(s) sent to storage: 03/28/2019 18:00

- Water - VOA vials have zero headspace? Yes No No VOA vials submitted
- Water - pH acceptable upon receipt? Yes No N/A
- pH adjusted? Yes No N/A

pH adjusted by:

Login Notes:

Client Contacted: Date Contacted: Person Contacted:
 Contacted By: Regarding:

Comments:

Corrective Action:

HS19031511

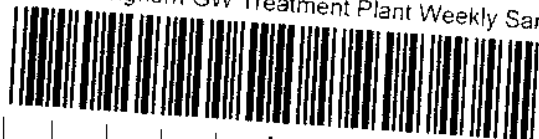
Bhate Environmental Associates, Inc.
LH18/24 Longhorn GW Treatment Plant Weekly Sar

CHAIN OF CUSTODY

Name Of Lab Shipping To: ALS 10450 Stancliff Rd., Suite 210 Houston, Tx. 77099 ATTN: RJ Modashia

Project: BHATE
LONGHORN ARMY AMMN. PLANT (LHAAP)
GROUNDWATER TREATMENT PLANT (GWTP)
KARNACK, TEXAS

Project No.
NWO1312.0150.0
16.0001



Job:
**GROUNDWATER TREATMENT PLANT
QUARTERLY EFFLUENT SAMPLES**

Prepared By:
Scott Beesinger


P. O. Number

Field Sample I.D.	Sample Matrix	Date / Time	MS / MSD	No. OF CONTAINERS	ROD Volatiles	Total Metals	Oil & Grease	Chemical Oxygen Demand	Chloride & Sulfate	1, 4 - DIOXANE	PERCHLORATE	Remarks (Preservatives, etc.)	Lab I.D.#
LH18/24-SP650_032719	Water	03/27/19 / 14:00		4	3		1					HCL	
LH18/24-SP650_032719	Water	03/27/19 / 14:00		1		1						HNO3	
LH18/24-SP650_032719	Water	03/27/19 / 14:00		2					1	1		NONE	
LH18/24-SP650_032719	Water	03/27/19 / 14:00		1				1				H2SO4	
LH18/24-SP650_032719_BIX	Water	03/27/19 / 14:00		1							1	NONE	
Trip Blank	Water	03/27/19		2	2							HCL	

Additional Remarks: STANDARD TURN AROUND TIME

Relinquished By: <i>Scott Beesinger</i>	Date: 03/27/19	Time: 14:30	Received By: <i>[Signature]</i>	Date: 3/28/19	Time: 08:30	Relinquished By:	Date:	Time:	Received By:	Date:	Time:
---	--------------------------	-----------------------	---	-------------------------	-----------------------	-------------------------	--------------	--------------	---------------------	--------------	--------------

For Lab Use Only										
Received At Lab By:	Date	Time	Airbill No.	Opened By:	Date	Time	Temp of Container	Seal No.	Condition	
Remarks:										
g.c. 44581 1-50 #25 C/1000										

 ALS Environmental 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CUSTODY SEAL		Seal Broken By: <i>SM</i>
	Date: 3/27/19	Time: 11:30	Date: 03/28/19
44581	Name: Scott Beehner	Company: DLAATX	

44581 MAR 28 2019



Must Deliver Next Business Day
Time and Temperature Sensitive!

44581

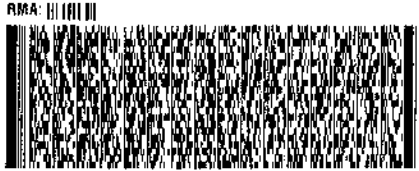
Part 9 15678-24 RITE EXP 1-101

ORIGIN ID: SGRA (303) 597-2450
SCOTT BEEHNER
STATE ENVIRONMENTAL ASSOCIATES
1203-B EAST GRAND AVE. PMB202
MARSHALL, TX 75670
UNITED STATES US

SHIP DATE: 06MAR19
ACTWT: 1.00 LB MAN
CAD: 30013D/CAF3211
DIHS: 28x19x14 IN

TO **CLIENT SERVICES**
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON TX 77099
(281) 530-5656
REF: LHAAP-69-BD 64113-RJ

551C1/4603/8C0A



FedEx
TRK# 4809 7831 3421
0221

THU - 28 MAR 10:30A
PRIORITY OVERNIGHT

AB SGRA

77099
TX-US
IAH



FD 152785 23MAR19 66GA 552C1/4603/8C0A



Case Narrative

Method: 6850
Analysis: Perchlorate
Analysis SOP: LC-MS-CLO4
ALS WO ID(s): 1909152; 1909153; 1909154;
1909947; 1909949

Client: ALS Laboratories (Houston, TX)
Matrix: Water
ELMS Batch (HBN): 2233 (236356)

General Set Information: There were ten field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ¹⁸O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50μL of an ¹⁸O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Method QC data: The method blank (LMB 647198) was less than 1/2 the CRDL. The recovery for the LCS (647199) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on sample 1909152001 (Client ID's: LH18/24-SP650_032719_BIX). 4.0 μ l of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4. μ g/L. The MS/MSD – 647200/01 failed QC acceptance criteria for percent recoveries, biased high. This is due to the fact that the unspiked sample result of 2.574 μ g/L was not subtracted from the MS/MSD results. The Matrix Spike and Matrix Spike duplicate is reported for the clients' information only. The sample matrix may be inappropriate for the method selected. The MS/MSD relative percent difference (RPD) was within the performance limits.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in μ g/L. Results were calculated in μ g/L by the equation (A)x(B),

where: A = Analyte concentration from the standard curve (μ g/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 647196) is reported from the analysis of the Laboratory Control Sample (LCS – 647199) at a level of 4.0 μ g/L. Due to limitations of the Chemstation Software, some of the chromatographic peaks may require manual integrations. A manual integration was performed for one of the Initial Calibration analyses (datafile: 19MARI03).

Thomas Bosch April 11, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: April 11, 2019

RJ Modashia
 ALS Environmental (Houston)
 10450 Stancliff Road
 Suite 210
 Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1909154**

Project ID: HS19031511

Purchase Order: HS19031511

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
LH18/24-SP650_032719_BIX	1909154001	03/27/19	03/30/19	

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40 of 127



ANALYTICAL REPORT

Workorder: 34-1909154

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: LH18/24-SP650_032719_BIX	Sampling Site: NA	Collected: 03/27/2019				
Lab ID: 1909154001	Media: 125 mL Nalgene	Received: 03/30/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2233 (HBN: 236356) Analyzed: 04/10/2019 11:48	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	2.3	1.0	2.0	4.0	1	J

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 236356)

Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 04/11/2019 09:27	/S/ Stephen Brose 04/11/2019 15:00

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1909154

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00935965

Analysis Information

Workorder: 1909154

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2233 (HBN: 236356)
Analyzed By: Thomas Bosch

Blank

LMB: 647198 Analyzed: 04/10/2019 10:42 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 647199 Analyzed: 04/10/2019 10:15 Dilution: 1 Units: ug/L					
Analyte	Result	Target	% Rec	QC Limits	
Perchlorate	3.68	4.00	91.9	78.8	123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1909152001 Analyzed: 04/10/2019 10:55 Dilution: 1 Units: ug/L		MS: 647200 Analyzed: 04/10/2019 11:08 Dilution: 1 Units: ug/L				MSD: 647201 Analyzed: 04/10/2019 11:22 Dilution: 1 Units: ug/L					
Analyte	Result	Result	Target	% Rec	QC Limits		Result	% Rec	RPD	QC Limits	
Perchlorate	2.60	6.18	4	# 154	78.8	123.8	6.36	# 159	2.92	0.0	20.0

Comments

Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 04/11/2019 13:48	/S/ Stephen Brose 04/11/2019 15:00

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



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18698/# 2

Subcontract Chain of Custody

COC ID: 11014

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

1909154

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19031511
TSR: Danielle Winnings

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
1. HS19031511-02	LH18/24-SP650_032719_BIX	Water	27 Mar 2019 14:00
SUB_Perch-6850			05 Apr 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. MAUMY
Received By: [Signature]
Cooler ID(s): 9274

Date/Time: 3/29/19 18:00
Date/Time: 3/20/19 858
Temperature(s): 2

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ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: Ms Houston Project/Task/Site: 1909154
 Date/Time of Receipt: 3/20/19 858 Number of Coolers Received: 1

Condition of Coolers:	Acceptable/Unacceptable	Temperature Control:	Present/Not Included
Cooler Custody Seals:	Present/Absent/NA	Location Temp Taken:	Control/Between Samples
Container Custody Seals:	Present/Absent/NA	Are all temperatures within project specific guidelines?	Yes/No/NA
Ice Present:	Yes/No/NA	VOA Headspace Present?	Yes/No/NA
	Frozen/Melted/NA		

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 <u>9270</u>	2 °C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: Meredith Edrington Signature Meredith Edrington Printed Name 3/20/19 Date

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Must Deliver Next Business Day
Time and Temperature Sensitive!

Part # 159469-434 RITZ EXP 11/19

ORIGIN ID:SGRA (281) 530-5656
CLIENT SERVICES
ALS LABORATORY GROUP
10450 STANCLIFF ROAD
SUITE 210
HOUSTON, TX 77099
UNITED STATES US

SHIP DATE: 29MAR19
ACTWGT: 9.25 LB
CAD: 300130/CAFE3211
DIMS: 14x11x10 IN

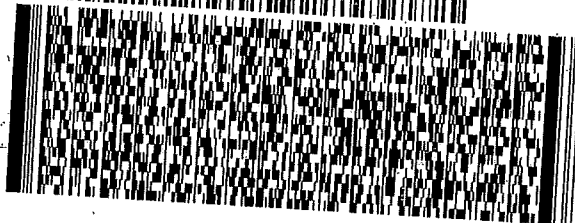
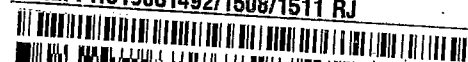
BILL THIRD PARTY

TO **SAMPLE RECEIVING**
ALS ENVIRONMENTAL
960 W. LEVOY DRIVE

SALT LAKE CITY UT 84123

(801) 268-7700

REF: HS19031492/1508/1511 RJ



FedEx
Express



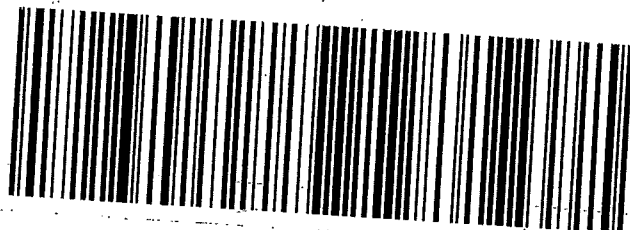
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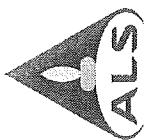
TRK# 4809 7832 2367
0201

SATURDAY 12:00P
PRIORITY OVERNIGHT

XO BTFA

84123
UT-US SLC





Batch Worklist

Batch: ELMS/ 2233 **Created:** 4/9/2019 08:42 **Instrument:** LCMS04 **HBN:** 236356
Rule: EPA 6850, DoD QSM Water **Analyst:** T. Bosch **Status:** WP



- Workorder: 1909152 [ENV_LVL4]
- Workorder: 1909153 [ENV_LVL4]
- Workorder: 1909154 [ENV_LVL4]
- Workorder: 1909947 [ENV_LVL4]
- Workorder: 1909949 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mx	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	647195	CCV for HBN 236356 [ELMS/2233]				CCV	3		E685041C3Q	5311		4/11/2019	4/10/2019
2	647199	LCS for HBN 236356 [ELMS/2233]				LCS	3		E6850Q413Q	5311		4/11/2019	4/10/2019
3	647197	ICS for HBN 236356 [ELMS/2233]				ICS	3		E6850.D3Q	5311		4/11/2019	4/10/2019
4	647198	LMB for HBN 236356 [ELMS/2233]				LMB	3		E6850Q413Q	5311		4/11/2019	4/10/2019
5	1909152001	LH18/24-SP650_032719_BIX				SAMPLE	3	1909152001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
6	647200	LH18/24-SP650...(1909152001MS)				MS	3		E6850Q413Q	5311		4/11/2019	4/10/2019
7	647201	LH18/24-SP65...(1909152001MSD)				MSD	3		E6850Q413Q	5311		4/11/2019	4/10/2019
8	1909153001	LH18-24-SP140_032719				SAMPLE	3	1909153001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
9	1909154001	LH18/24-SP650_032719_BIX				SAMPLE	3	1909154001-A	E6850Q41.3	5480	4/24/2019	4/11/2019	4/10/2019
10	1909947001	LH18/25-SP650_040419_BIX				SAMPLE	3	1909947001-A	E6850Q41.3	5480	5/2/2019	4/18/2019	4/10/2019
11	1909949001	HBW7_040119				SAMPLE	3	1909949001-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
12	1909949002	HBW7_040119-a				SAMPLE	3	1909949002-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
13	1909949003	HBW10_040119				SAMPLE	3	1909949003-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
14	1909949004	HBW1_040119				SAMPLE	3	1909949004-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
15	1909949005	GPW1_040119				SAMPLE	3	1909949005-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
16	1909949006	GPW3_040119				SAMPLE	3	1909949006-A	E6850Q41.3	5480	5/1/2019	4/18/2019	4/10/2019
17	647196	RLVS for HBN 236356 [ELMS/2233]				RLVS	3		E685041C3Q	5311		4/11/2019	4/10/2019
18	647202	CCV for HBN 236356 [ELMS/2233]				CCV	3		E685041C3Q	5311		4/11/2019	4/10/2019



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1909152 (001); 1909153 (001); 1909154 (001); 1909947 (001); 1909949 (001-06) ELMS Batch/HBN ID: 2233 (236356)
 Prep Date: 04/09/2019 Analysis Date: 04/10/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\28MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by TNB. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4) / 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 03/19/2019, sequence 19MAR19D.s Offline Quantitation Method: CLO4-DP2.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 6 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 647199; Target = 4.0µg/L. ASTM type II water was used for LMB 647198.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on sample 1909152001 (Client ID's: LH18/24-SP650_032719_BIX). 4.0µl of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- Results reported in µg/L. Field sample 1909153001 was analyzed and reported from a 1:1,000 dilution. The reporting limit has been adjusted accordingly.
- All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The MS/MSD – 647200/01 failed QC acceptance criteria for percent recoveries, biased high. This is due to the fact that the unspiked sample result of 2.574µg/L was not subtracted from the MS/MSD results. The Matrix Spike and Matrix Spike duplicate is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\APR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- Notebook: \\slstws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\236356-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATA\REVIEW\HBN#
- The Reporting Limit Verification Standard (RLVS – 647196) is reported from the analysis of the Laboratory Control Sample (LCS – 647199) at a level 4.0µg/L.
- Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 19MARI03).

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
Batch(es)/SDG: E LMS: 2233 HBN: 236356 1209947 / 1209949		
Sample Set IDs if Applicable: 1909152/1909153/1909154		
Calibration standards analyzed and meets criteria	TB	SB
Standards traceability checked and meets criteria	TB	SB
Standard curve coefficients evaluated and meet criteria	TB	SB
ICVs analyzed and meet acceptance criteria	TB	SB
CCVs analyzed and meet acceptance criteria	TB	SB
Method Blanks analyzed and meet acceptance criteria	TB	SB
Retention Time Windows checked	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
Surrogate recoveries checked and appropriately addressed	—	—
Method Preparation Blanks analyzed and meet acceptance criteria	TB	SB
MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed	TB	SB
RLVS analyzed	TB	SB
Preparation and analysis hold times met	TB	SB
Preparation deviations and re-preparations noted when performed	TB	SB
Analysis deviations and re-analyses noted when performed	TB	SB
Sample dilution factors noted on reports	TB	SB
Electronic records in HBN transcription accuracy and completeness checked	TB	SB
Preparation and analysis calculations checked	TB	SB
NCRs are completed as necessary NC/CAR#	—	—
Report forms are complete and accurate	TB	SB
Manual integrations checked	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK			Description - 6850 WKG Std 100.ug/L		
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10.ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK		Description - 6850 Stock AccStd 1,000ug/mL	
Standard: 43659	Created By: Thomas Bosch	Amount: 100 mL	
MFG: AccuStandard	Create Date: 09/17/2018 09:09AM	Expires: 07/25/2020	
MFG Lot: 218065075		Usable: No	
Part ID: IC-PER-10X-1		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT		Description - 6850 Intermdt AccStd 10.ug/mL			
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK		Description - 6850 QC WKG STD 100ug/L			
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte:	Name:	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK		Description - 6850 QC Stock STD 1,000ug/mL	
Standard: 36748		Created By: Thomas Bosch	Amount: 100 mL
MFG: Ultra Scientific		Create Date: 05/11/2017 01:05PM	Expires: 03/31/2020
MFG Lot: CP-0860			Usable: Yes
Part ID: ICC-013			Lab Lot: CLO4 QC STOCK
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFF-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:

This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis



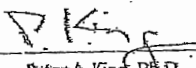
ISO Guide 34 Reference Material

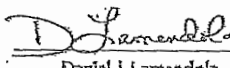
Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:

The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QA/RA



125 Market Street
New Haven, CT 06513
USA



AccuStandard®

Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



S 43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:
ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$
Labeled CAS Number: NA
Unlabeled CAS Number: 7601-89-0
MW*: 130.4
Chemical Formula: NaCl*O4
Storage: Store at room temperature away from light and moisture.
Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 $\mu\text{g/mL}$
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 \pm 2.8 $\mu\text{g/mL}$ ($k=2$)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method

['*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	647195	CCV@25	Vial 71	1	Control	1	1.06336e6	21.31933
#*	647199	QC@4.0	Vial 72	1	Control	2	1.97600e5	3.67714
#*	647197	ICS@4.0	Vial 73	1	Control	3	1.49399e5	3.21517
#*	647198	LMB	Vial 74	1	Control	4	0.00000	0.00000
#*	1909152001		Vial 75	1	Sample	5	9.15156e4	2.57433
#*	647200	91521MS	Vial 76	1	Sample	6	2.25049e5	6.17769
#*	647201	91521SD	Vial 77	1	Sample	7	2.49579e5	6.36091
#*	1909153001	1K	Vial 78	1	Sample	8	3.70174e5	6532.86203
#*	1909154001		Vial 79	1	Sample	9	7.79123e4	2.34023
#*	1909947001		Vial 80	1	Sample	10	0.00000	0.00000
#*	1909949001		Vial 81	1	Sample	11	0.00000	0.00000
#*	1909949002		Vial 82	1	Sample	12	0.00000	0.00000
#*	1909949003		Vial 83	1	Sample	13	0.00000	0.00000
#*	1909949004		Vial 84	1	Sample	14	0.00000	0.00000
#*	1909949005		Vial 85	1	Sample	15	0.00000	0.00000
#*	1909949006		Vial 86	1	Sample	16	0.00000	0.00000
*	647202	CCV@25	Vial 71	1	Control	17	1.08007e6	22.59518

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	647195	CCV@25	Vial 71	1	Control	1	3.43635e5	23.09461
#*	647199	QC@4.0	Vial 72	1	Control	2	7.01681e4	4.22460
#*	647197	ICS@4.0	Vial 73	1	Control	3	5.45965e4	3.77625
#*	647198	LMB	Vial 74	1	Control	4	0.00000	0.00000
#*	1909152001		Vial 75	1	Sample	5	3.18275e4	2.84107
#*	647200	91521MS	Vial 76	1	Sample	6	7.61114e4	6.88356
#*	647201	91521SD	Vial 77	1	Sample	7	8.42686e4	7.08137
#*	1909153001	1K	Vial 78	1	Sample	8	1.21054e5	7051.77497
#*	1909154001		Vial 79	1	Sample	9	2.76117e4	2.61418
#*	1909947001		Vial 80	1	Sample	10	0.00000	0.00000
#*	1909949001		Vial 81	1	Sample	11	0.00000	0.00000
#*	1909949002		Vial 82	1	Sample	12	0.00000	0.00000
#*	1909949003		Vial 83	1	Sample	13	0.00000	0.00000
#*	1909949004		Vial 84	1	Sample	14	0.00000	0.00000
#*	1909949005		Vial 85	1	Sample	15	0.00000	0.00000
#*	1909949006		Vial 86	1	Sample	16	0.00000	0.00000
*	647202	CCV@25	Vial 71	1	Control	17	3.36319e5	23.63294

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	647195	CCV@25	Vial 71	1	Control	1	1.53120e5	5.00000
#*	647199	QC@4.0	Vial 72	1	Control	2	1.77838e5	5.00000
#*	647197	ICS@4.0	Vial 73	1	Control	3	1.54931e5	5.00000
#*	647198	LMB	Vial 74	1	Control	4	1.76915e5	5.00000
#*	1909152001		Vial 75	1	Sample	5	1.20240e5	5.00000
#*	647200	91521MS	Vial 76	1	Sample	6	1.17757e5	5.00000
#*	647201	91521SD	Vial 77	1	Sample	7	1.26683e5	5.00000
#*	1909153001	1K	Vial 78	1	Sample	8	1.82758e5	5000.00000
#*	1909154001		Vial 79	1	Sample	9	1.13404e5	5.00000
#*	1909947001		Vial 80	1	Sample	10	1.15130e5	5.00000
#*	1909949001		Vial 81	1	Sample	11	1.14791e5	5.00000
#*	1909949002		Vial 82	1	Sample	12	1.24046e5	5.00000
#*	1909949003		Vial 83	1	Sample	13	1.23373e5	5.00000
#*	1909949004		Vial 84	1	Sample	14	1.20241e5	5.00000
#*	1909949005		Vial 85	1	Sample	15	1.27767e5	5.00000
#*	1909949006		Vial 86	1	Sample	16	1.23965e5	5.00000
*	647202	CCV@25	Vial 71	1	Control	17	1.46281e5	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

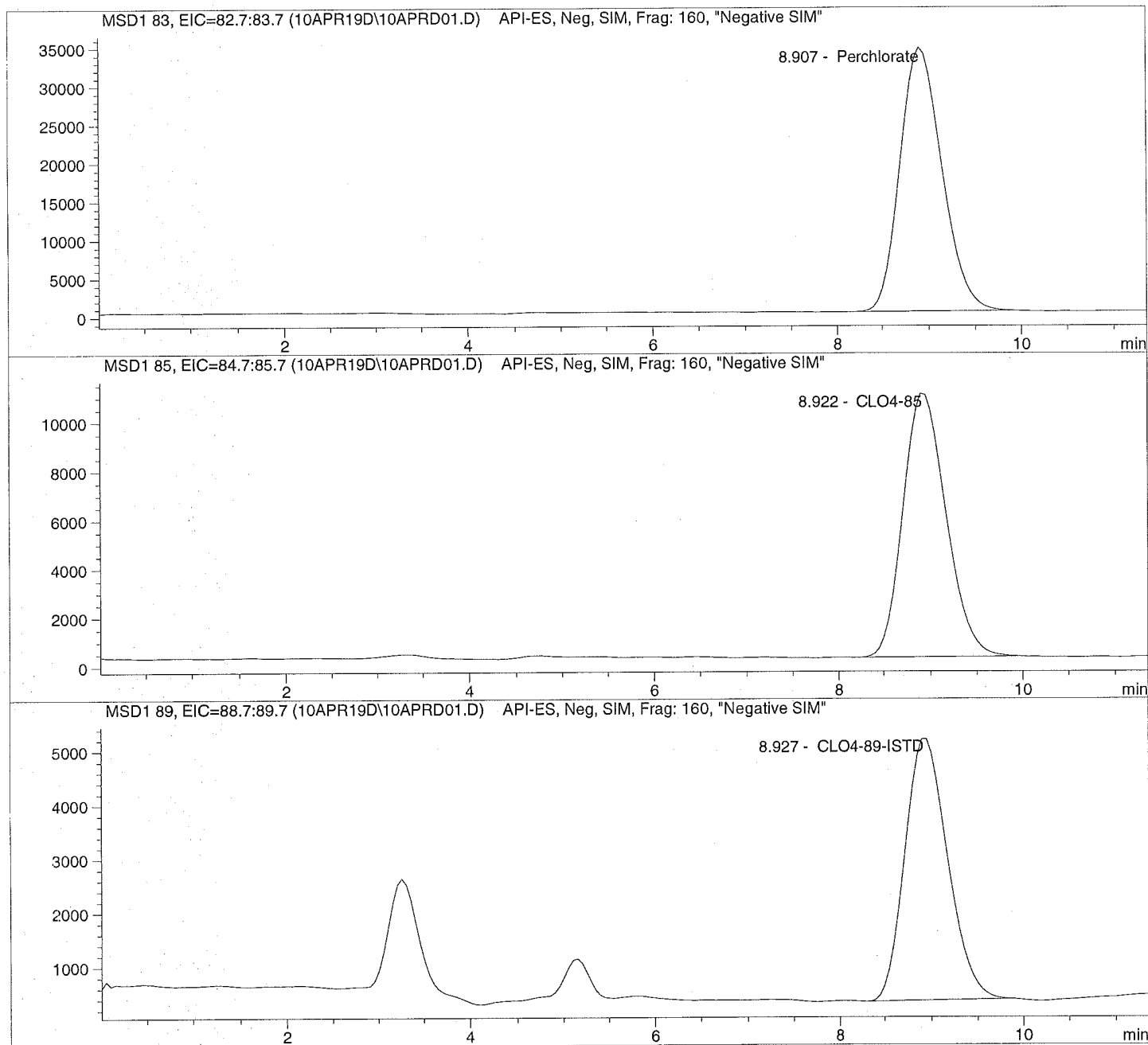
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	647195	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	647199	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	647197	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	647198	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1909152001		CLO4-AQN	1	Sample	
6	Vial 76	647200	91521MS	CLO4-AQN	1	Sample	
7	Vial 77	647201	91521SD	CLO4-AQN	1	Sample	
8	Vial 78	1909153001	1K	CLO4-AQN	1	Sample	
9	Vial 79	1909154001		CLO4-AQN	1	Sample	
10	Vial 80	1909947001		CLO4-AQN	1	Sample	
11	Vial 81	1909949001		CLO4-AQN	1	Sample	
12	Vial 82	1909949002		CLO4-AQN	1	Sample	
13	Vial 83	1909949003		CLO4-AQN	1	Sample	
14	Vial 84	1909949004		CLO4-AQN	1	Sample	
15	Vial 85	1909949005		CLO4-AQN	1	Sample	
16	Vial 86	1909949006		CLO4-AQN	1	Sample	
17	Vial 71	647202	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD01.D Sample Name: 647195 CCV025

=====
Injection Date: 4/10/2019 10:01:38 Seq Line: 1
Sample Name: 647195 CCV025 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis
=====



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD01.D Sample Name: 647195 CCV@25

```

=====
Injection Date: 4/10/2019 10:01:38      Seq Line: 1
Sample Name:    647195   CCV@25         Location:  Vial 71
Acq Operator:  TNB              Inj. No.: 1
                                      Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.907	PBA	1063358.2	21.3193	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.922	PBA	343635.4	23.0946	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.927	PBA	153120.0	5.0000	CLO4-89-ISTD

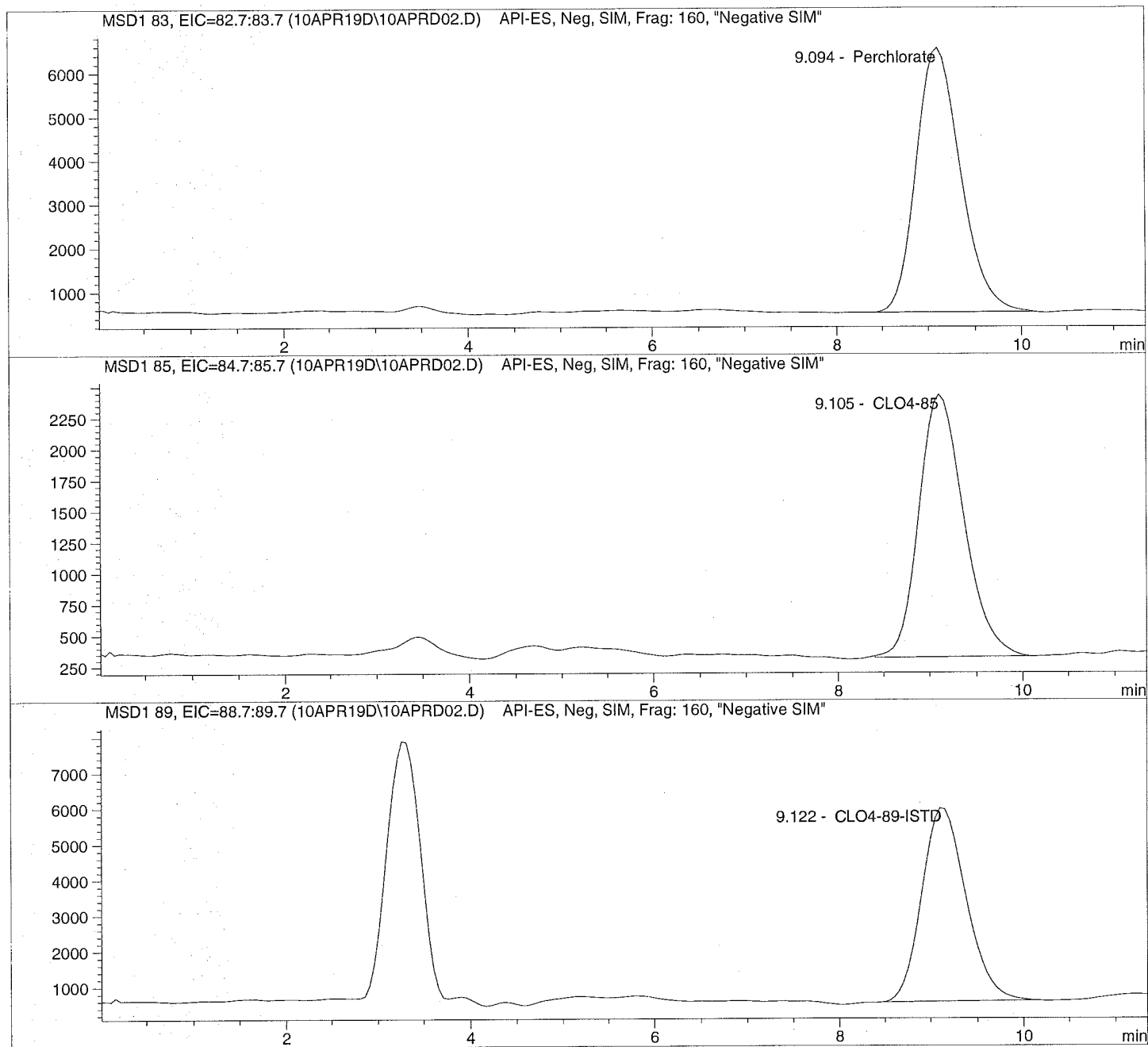
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD02.D Sample Name: 647199 QC@4.0

```
=====
Injection Date: 4/10/2019 10:15:40      Seq Line:      2
Sample Name:    647199 QC@4.0           Location:      Vial 72
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD02.D Sample Name: 647199 QC@4.0

```

=====
Injection Date: 4/10/2019 10:15:40      Seq Line:      2
Sample Name:    647199   QC@4.0         Location:      Vial 72
Acq Operator:   TNB                Inj. No.:     1
                                           Inj. Vol.:    30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 4.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.094	BBA	197600.1	3.6771	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.105	BBA	70168.1	4.2246	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.122	BBA	177837.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

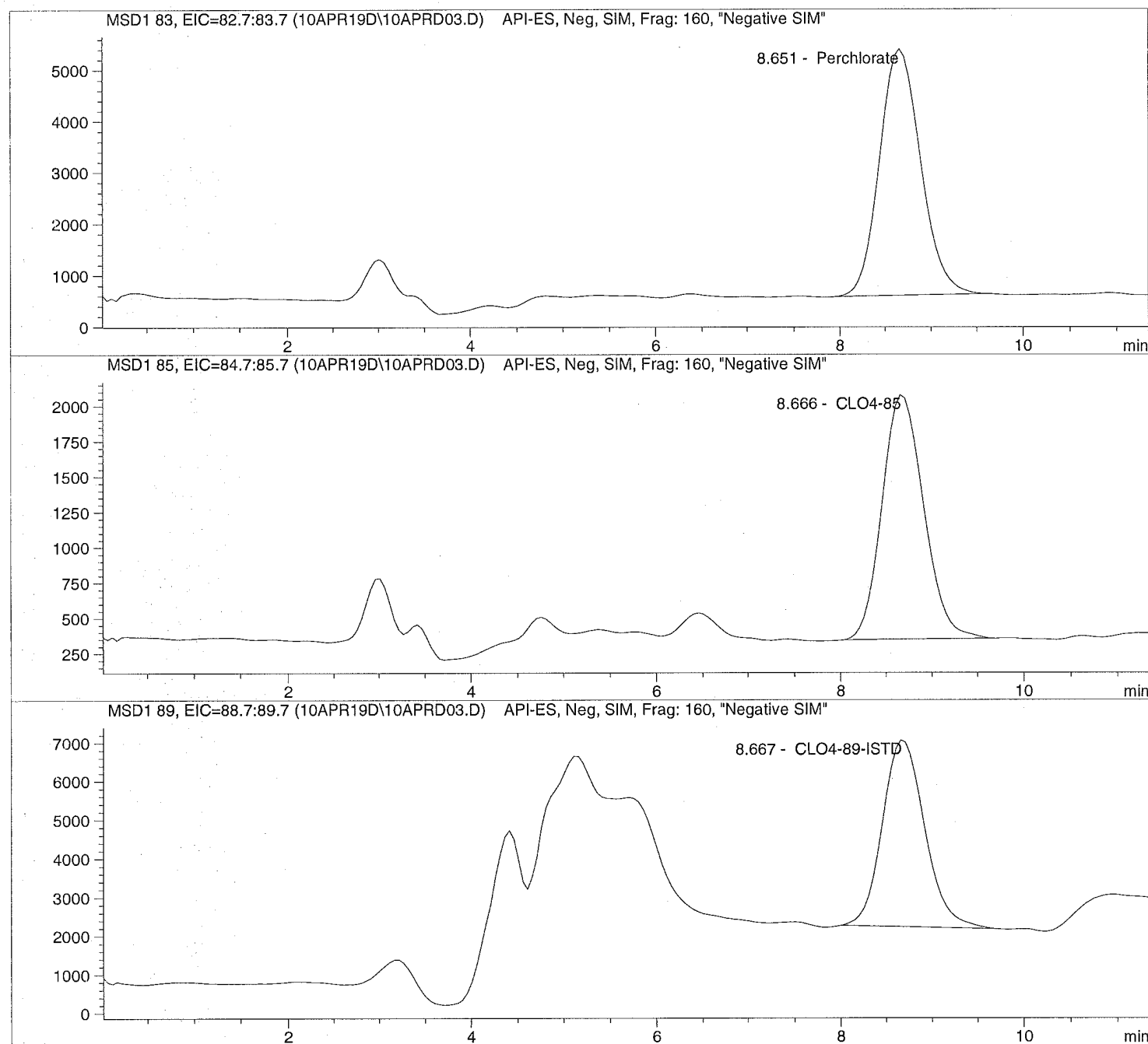
```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD03.D Sample Name: 647197 ICS@4.0

=====
Injection Date: 4/10/2019 10:28:57 Seq Line: 3
Sample Name: 647197 ICS@4.0 Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD03.D Sample Name: 647197 ICS@4.0

```

=====
Injection Date: 4/10/2019 10:28:57      Seq Line:            3
Sample Name:    647197    ICS@4.0            Location:            Vial 73
Acq Operator:   TNB                        Inj. No.:            1
                                             Inj. Vol.:            30 µl
  
```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
  
```

Perchlorate analysis

Sample Information

```

=====
Sorted By:            Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:           1.000000
Dilution:             1.000000
Sample Amount:        4.000
  
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.651	PBA	149398.5	3.2152	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.666	BBA	54596.5	3.7763	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.667	PBA	154930.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD04.D

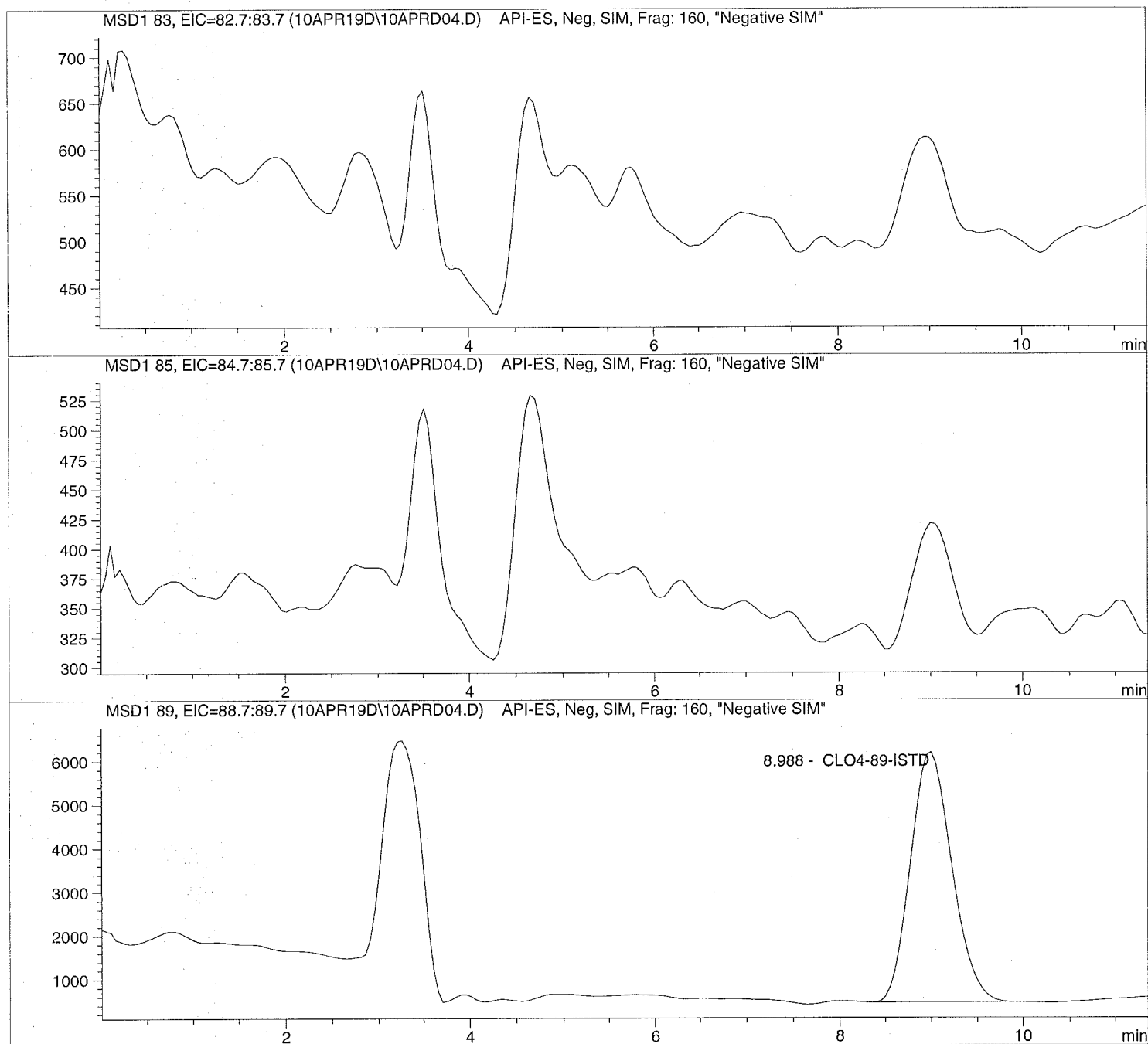
Sample Name: 647198 LMB

Injection Date: 4/10/2019 10:42:16
Sample Name: 647198 LMB
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD04.D Sample Name: 647198 LMB

```

=====
Injection Date: 4/10/2019 10:42:16      Seq Line: 4
Sample Name: 647198 LMB                  Location: Vial 74
Acq Operator: TNB                         Inj. No.: 1
                                           Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.988	PBA	176915.0	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD05.D

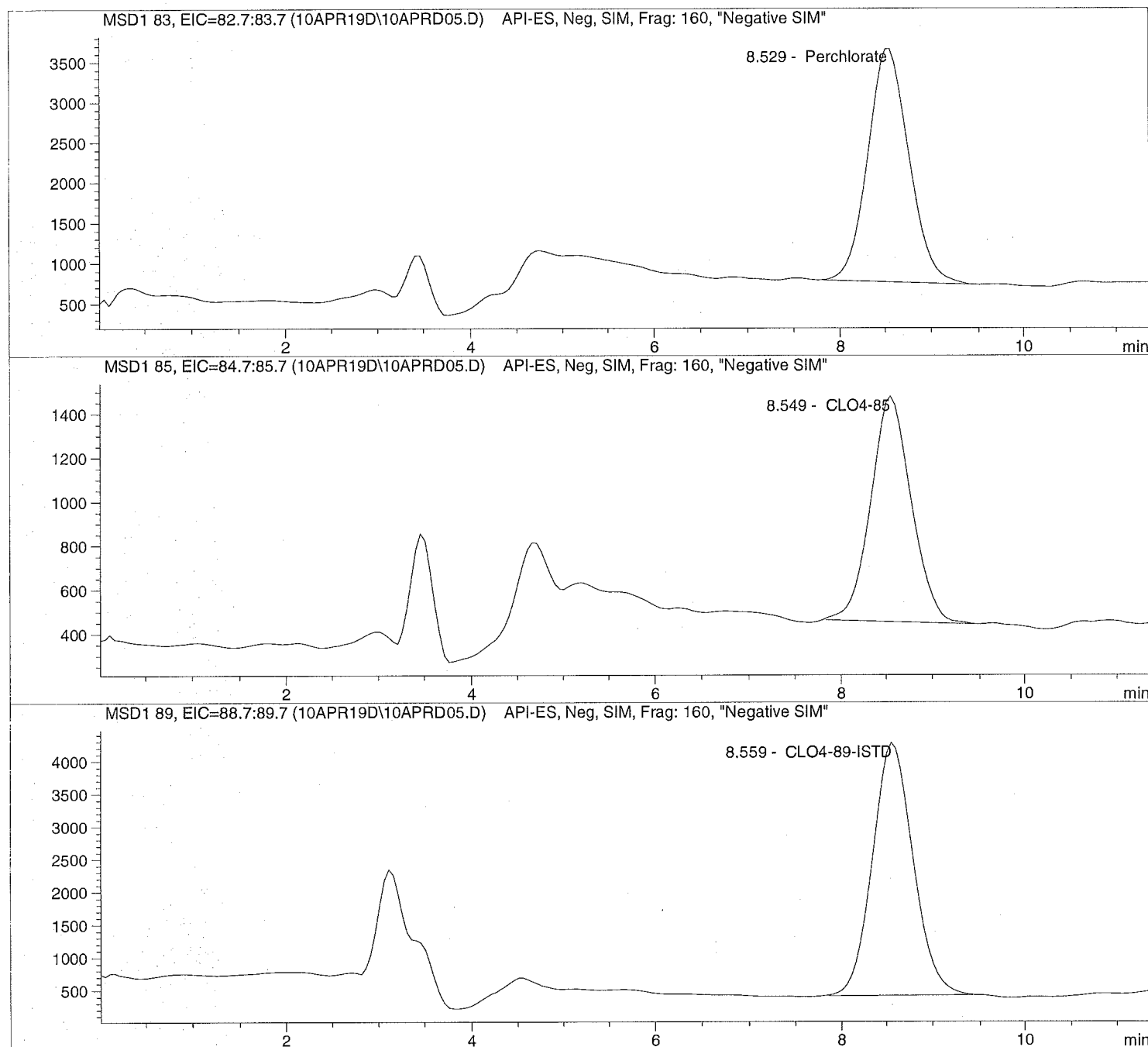
Sample Name: 1909152001

Injection Date: 4/10/2019 10:55:33
Sample Name: 1909152001
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD05.D Sample Name: 1909152001

```

=====
Injection Date: 4/10/2019 10:55:33      Seq Line: 5
Sample Name: 1909152001                Location: Vial 75
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.529	BBA	91515.6	2.5743	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.549	BBA	31827.5	2.8411	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.559	BBA	120239.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD06.D

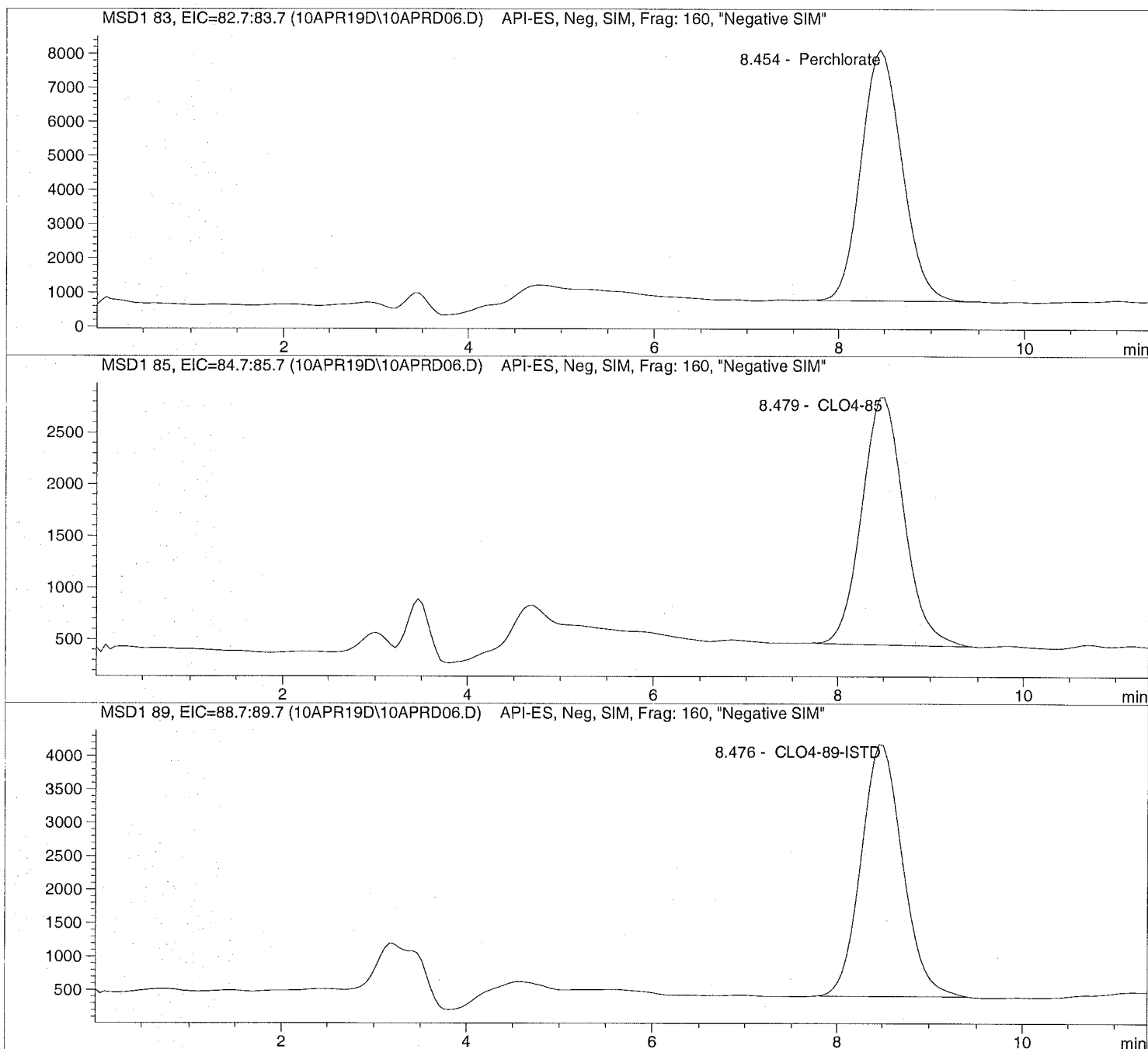
Sample Name: 647200 91521MS

Injection Date: 4/10/2019 11:08:49
Sample Name: 647200 91521MS
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD06.D Sample Name: 647200 91521MS

```

=====
Injection Date: 4/10/2019 11:08:49 Seq Line: 6
Sample Name: 647200 91521MS Location: Vial 76
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.454	BBA	225049.5	6.1777	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.479	BBA	76111.4	6.8836	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.476	BBA	117756.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD07.D

Sample Name: 647201 91521SD

Injection Date: 4/10/2019 11:22:09

Seq Line: 7

Sample Name: 647201 91521SD

Location: Vial 77

Acq Operator: TNB

Inj. No.: 1

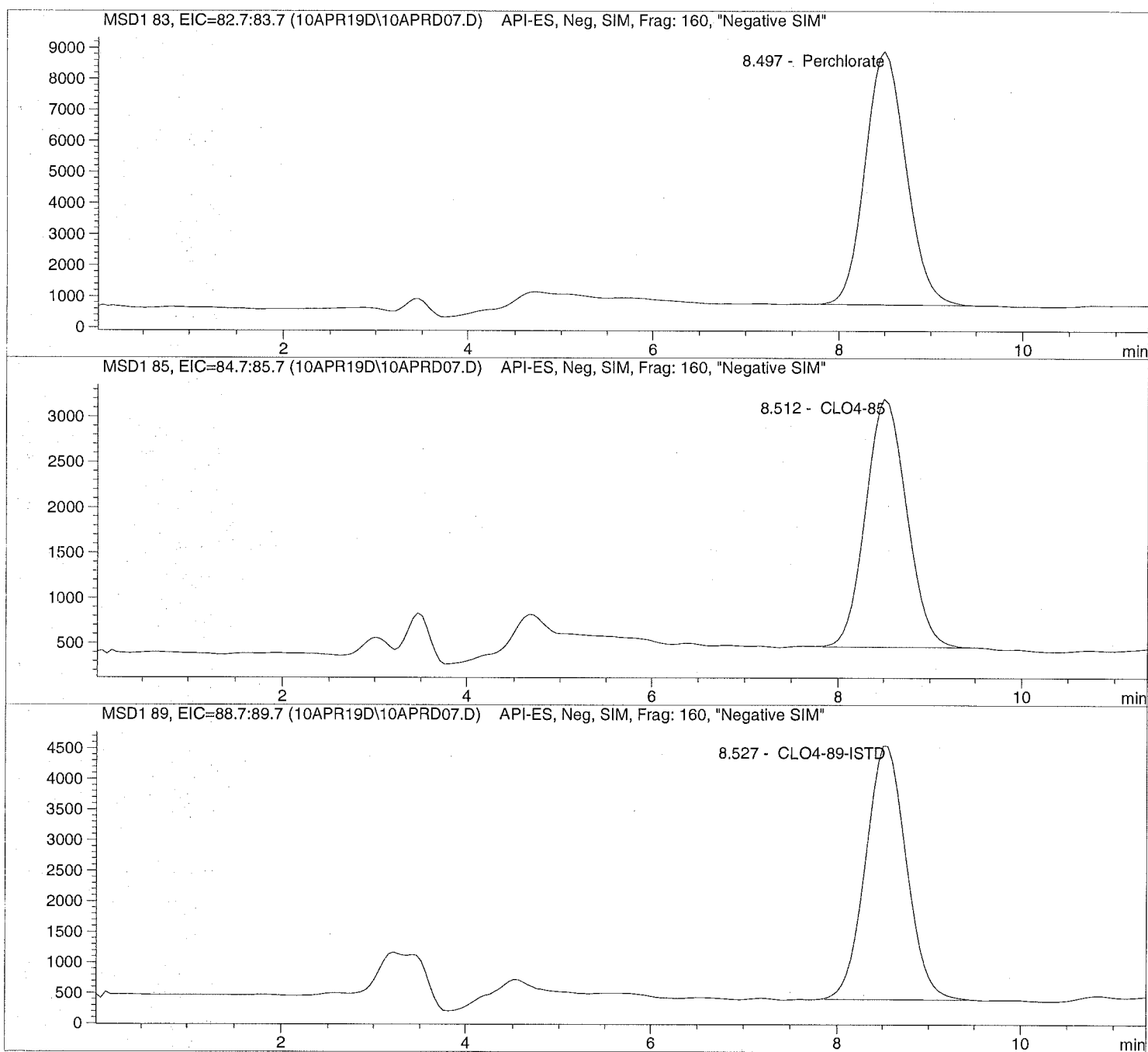
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD07.D Sample Name: 647201 91521SD

```

=====
Injection Date: 4/10/2019 11:22:09      Seq Line: 7
Sample Name: 647201 91521SD      Location: Vial 77
Acq Operator: TNB      Inj. No.: 1
                                         Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.497	BBA	249579.0	6.3609	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.512	BBA	84268.6	7.0814	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.527	BBA	126683.1	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD08.D

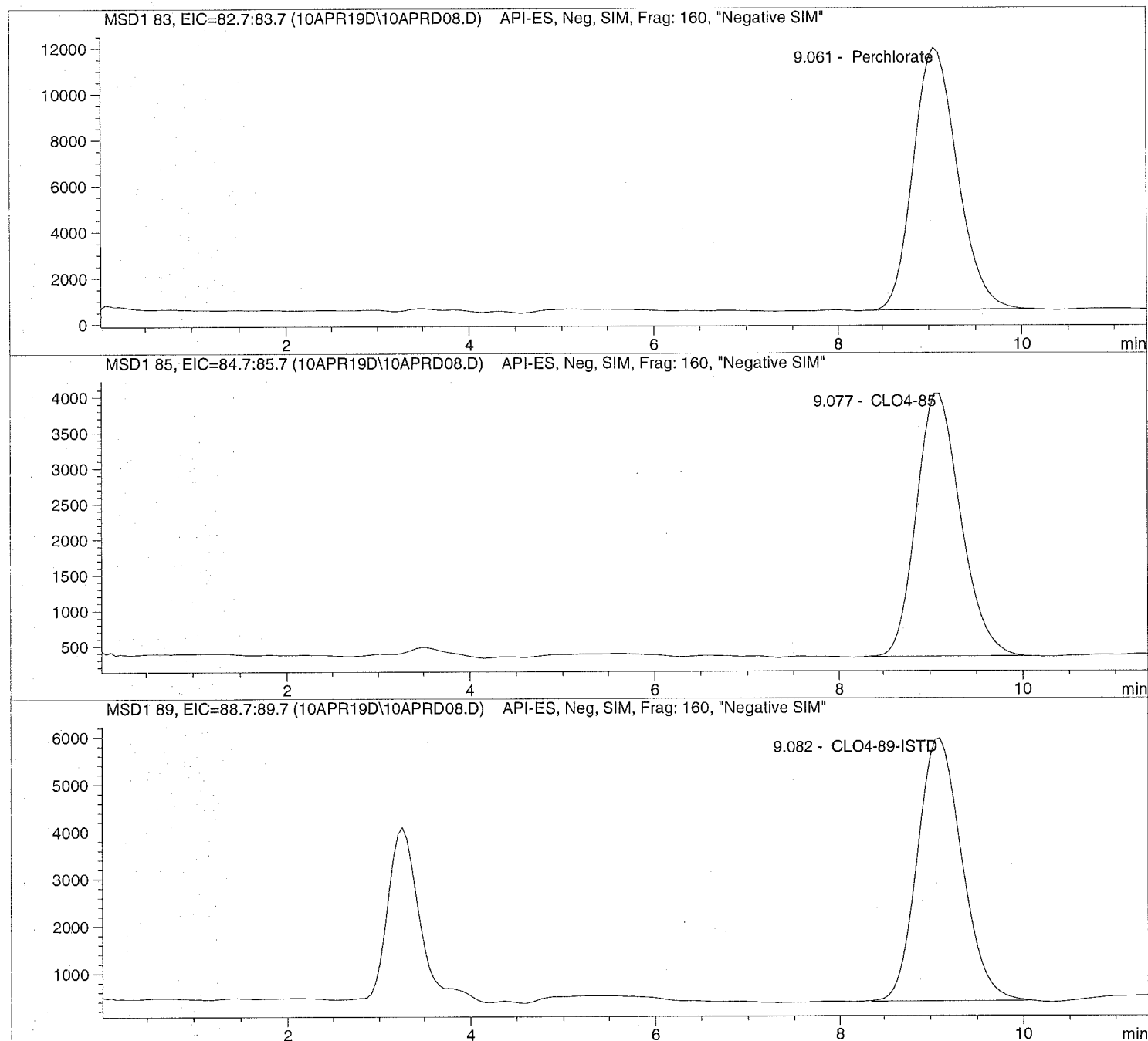
Sample Name: 1909153001 1K

Injection Date: 4/10/2019 11:35:27
Sample Name: 1909153001 1K
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD08.D Sample Name: 1909153001 1K

```
=====
Injection Date: 4/10/2019 11:35:27 Seq Line: 8
Sample Name: 1909153001 1K Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
```

Perchlorate analysis

=====

Sample Information

=====

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1000.000000
Sample Amount: 0.000
```

=====

LCMS Results

=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.061	PBA	370174.3	6532.8620	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.077	BBA	121053.5	7051.7750	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.082	BBA	182757.9	5000.0000	CLO4-89-ISTD

=====

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD09.D

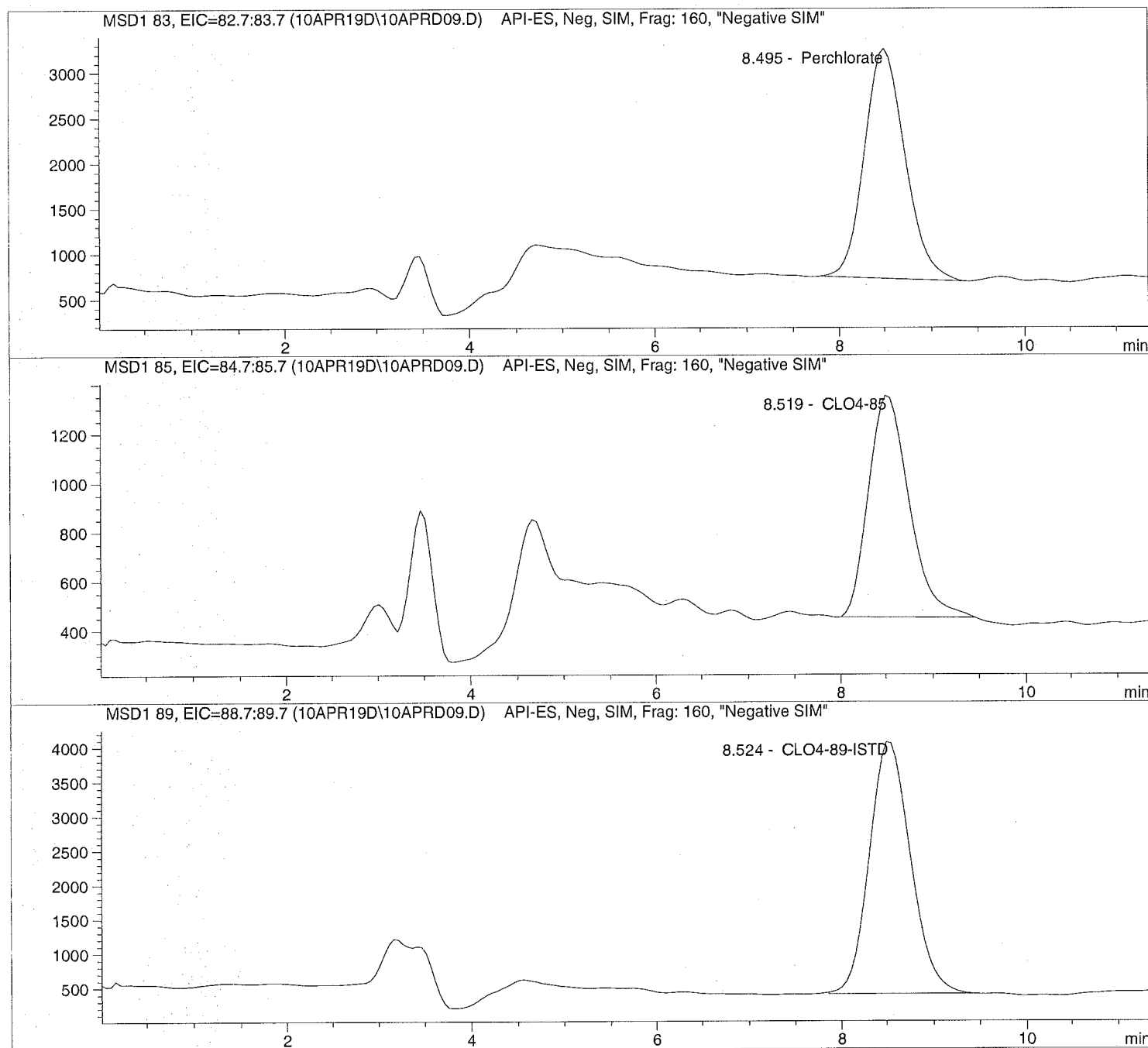
Sample Name: 1909154001

=====
Injection Date: 4/10/2019 11:48:52
Sample Name: 1909154001
Acq Operator: TNB

Seq Line: 9
Location: Vial 79
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD09.D Sample Name: 1909154001

```

=====
Injection Date: 4/10/2019 11:48:52      Seq Line: 9
Sample Name: 1909154001                Location: Vial 79
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.495	PBA	77912.3	2.3402	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.519	PBA	27611.7	2.6142	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.524	BBA	113404.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD10.D

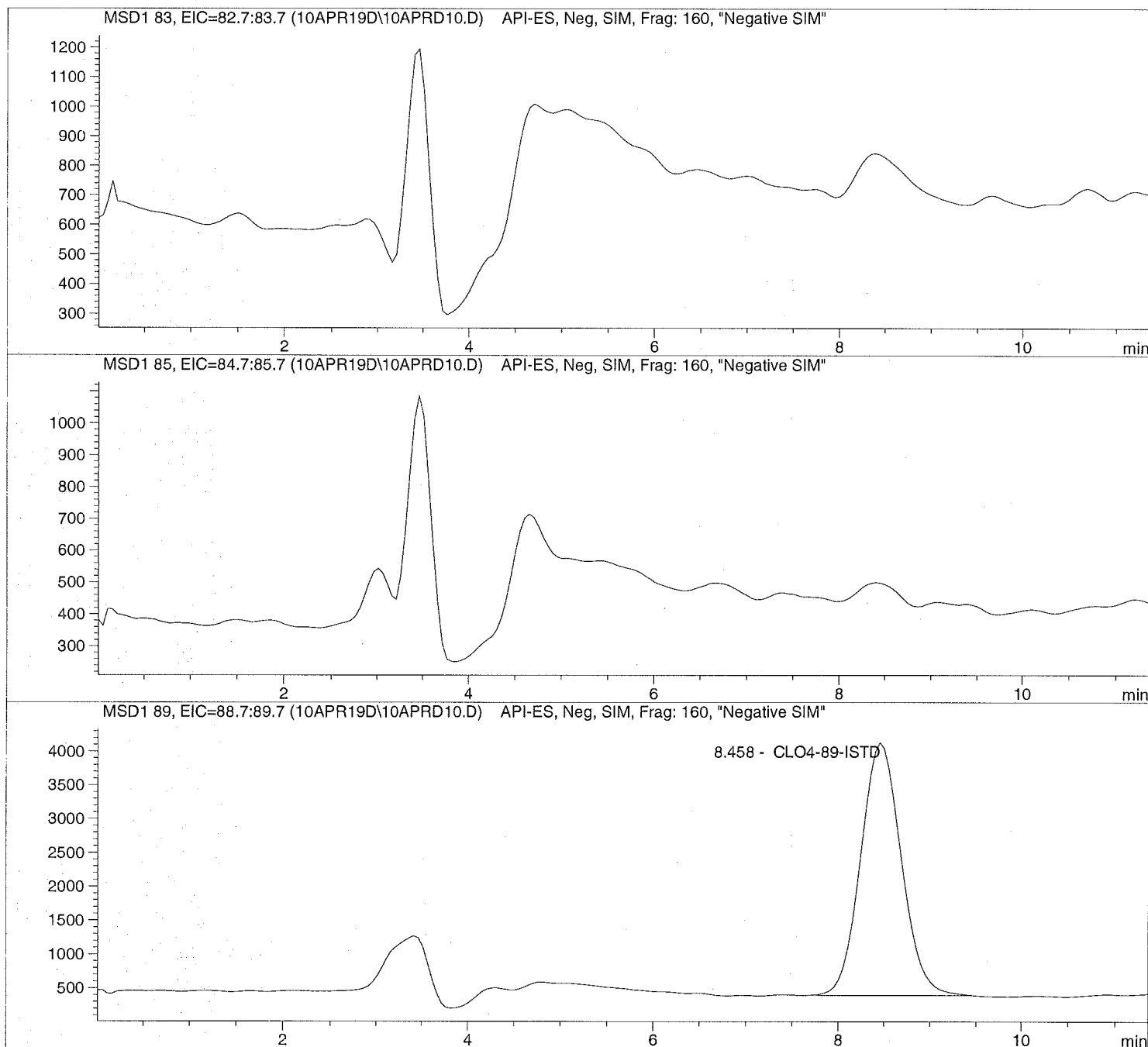
Sample Name: 1909947001

Injection Date: 4/10/2019 12:02:10
Sample Name: 1909947001
Acq Operator: TNB

Seq Line: 10
Location: Vial 80
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD10.D Sample Name: 1909947001

```

=====
Injection Date: 4/10/2019 12:02:10 Seq Line: 10
Sample Name: 1909947001 Location: Vial 80
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.458	BBA	115129.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

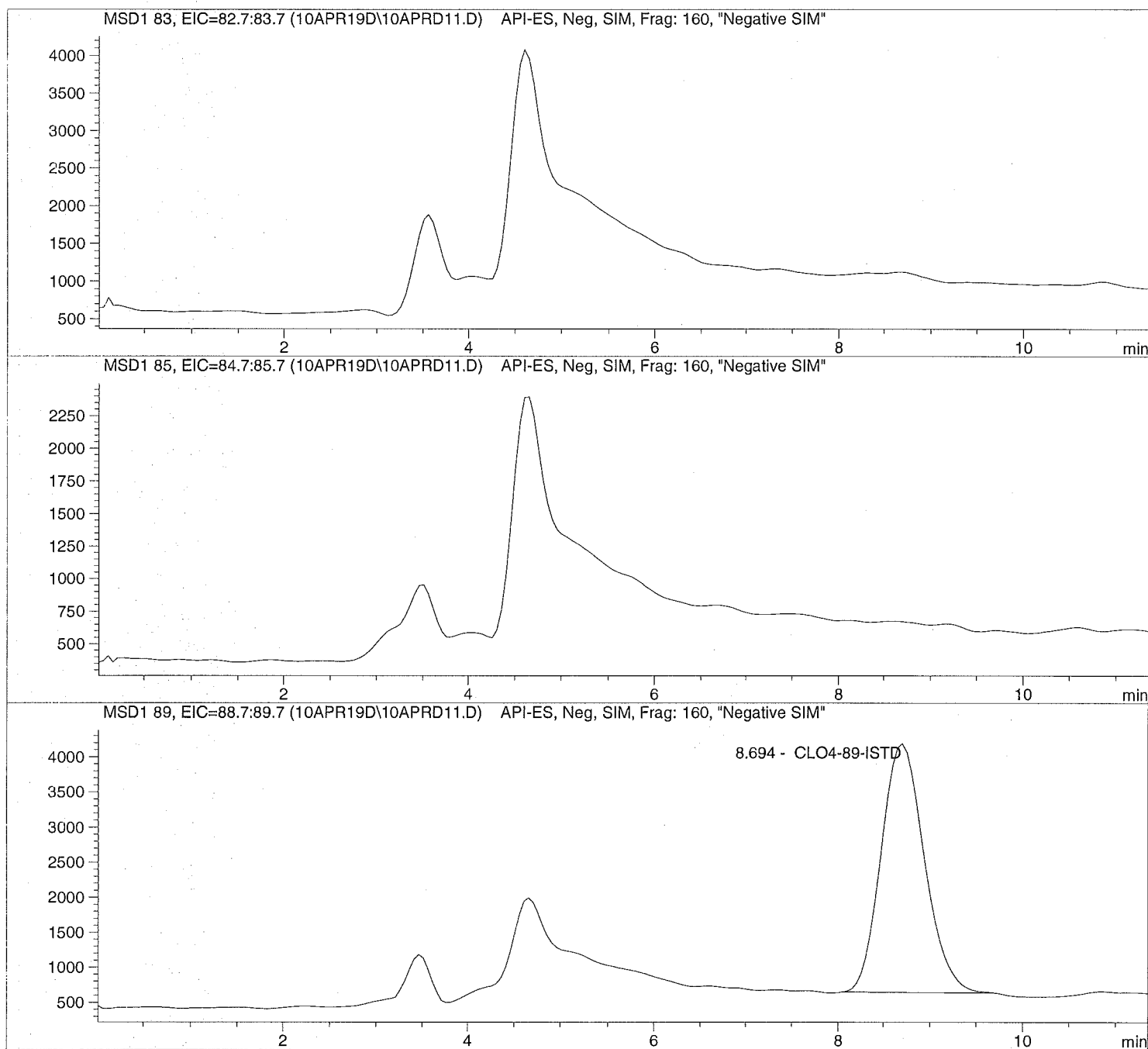
```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD11.D Sample Name: 1909949001

=====
Injection Date: 4/10/2019 12:15:27 Seq Line: 11
Sample Name: 1909949001 Location: Vial 81
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD11.D Sample Name: 1909949001

```

=====
Injection Date: 4/10/2019 12:15:27      Seq Line: 11
Sample Name: 1909949001                Location: Vial 81
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.694	PBA	114791.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD12.D

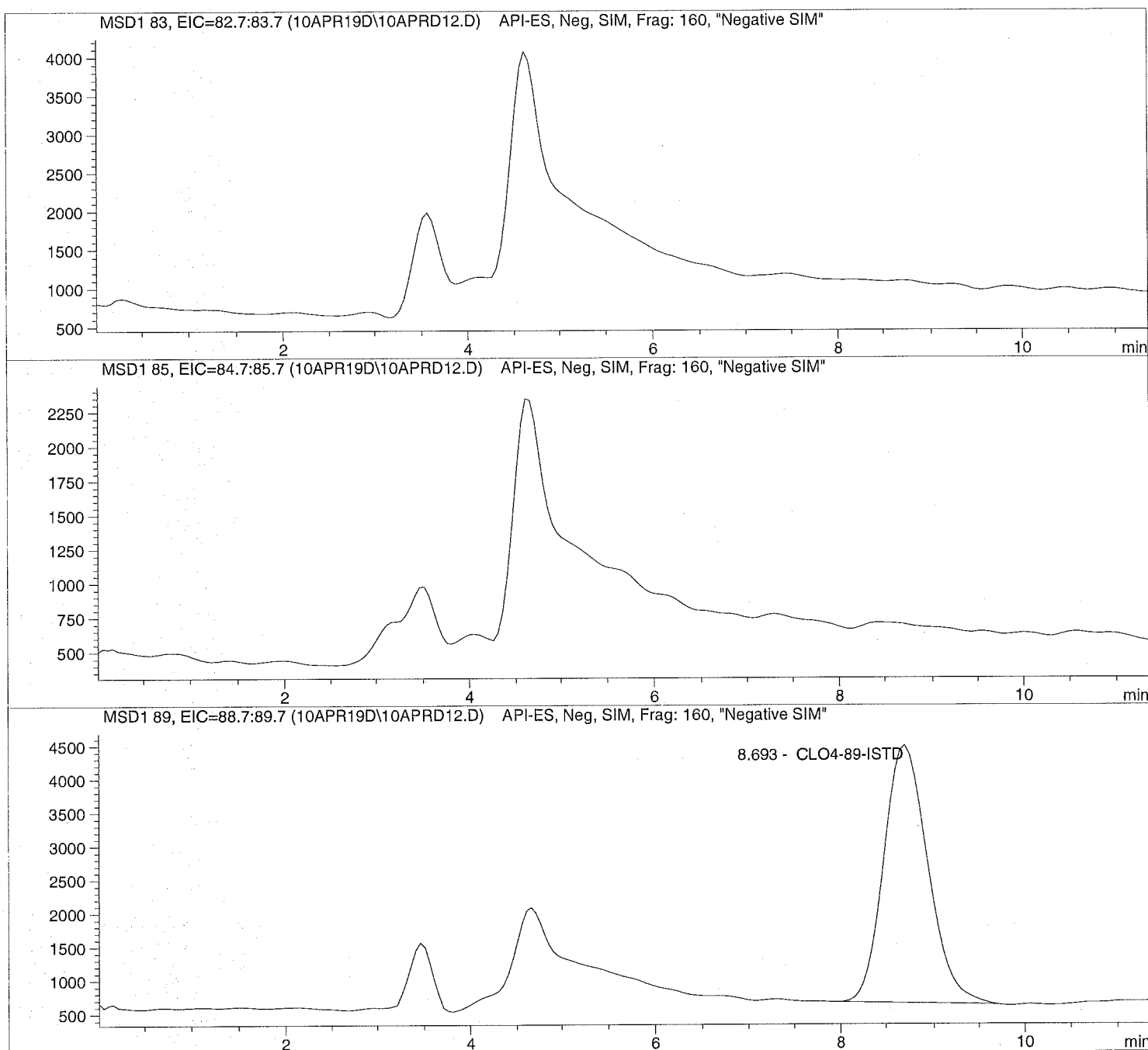
Sample Name: 1909949002

Injection Date: 4/10/2019 12:28:41
Sample Name: 1909949002
Acq Operator: TNB

Seq Line: 12
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD12.D Sample Name: 1909949002

```

=====
Injection Date: 4/10/2019 12:28:41      Seq Line: 12
Sample Name: 1909949002                Location: Vial 82
Acq Operator: TNB                      Inj. No.: 1
                                         Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.693	PBA	124045.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD13.D

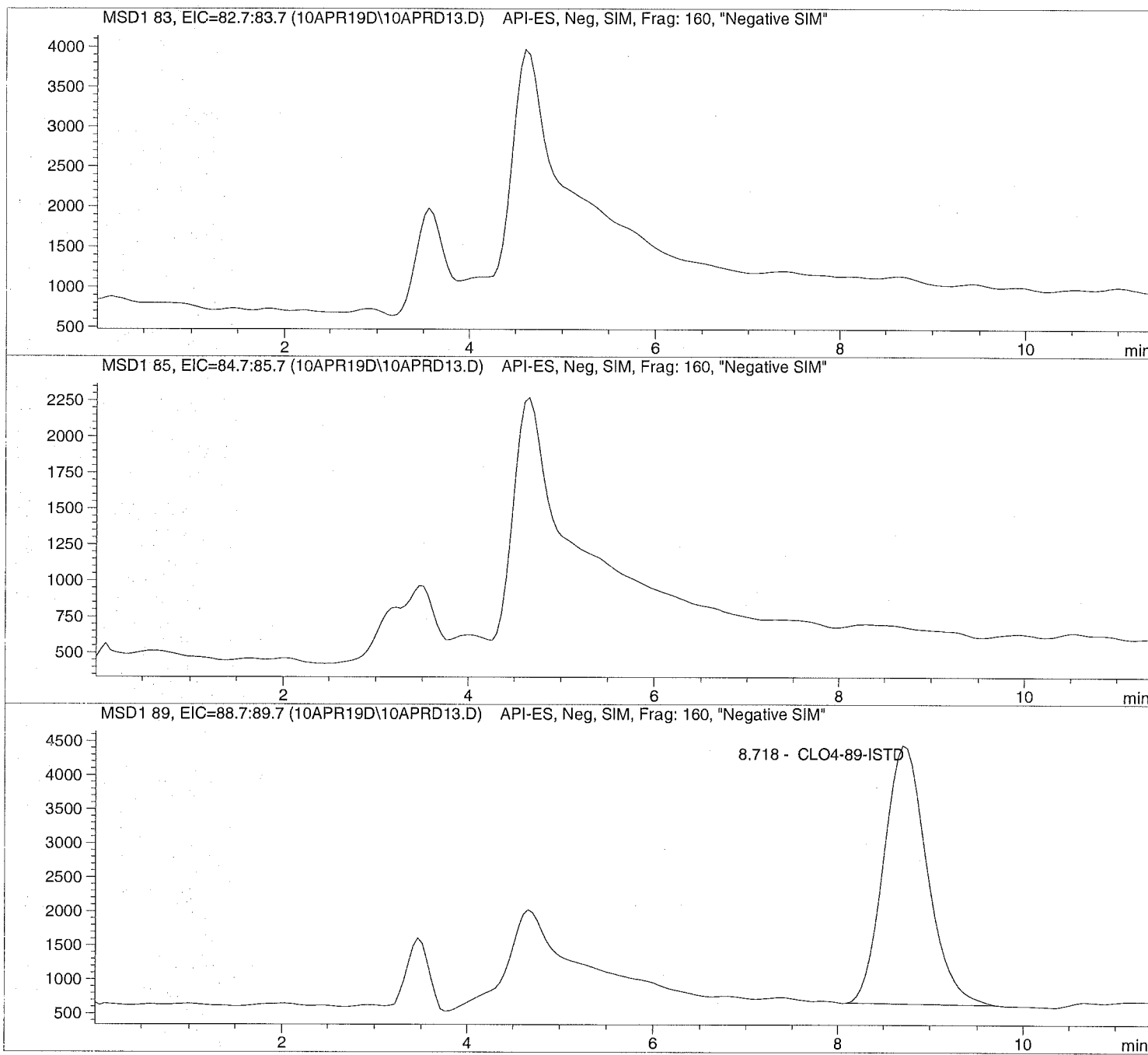
Sample Name: 1909949003

=====
Injection Date: 4/10/2019 12:41:59
Sample Name: 1909949003
Acq Operator: TNB

Seq Line: 13
Location: Vial 83
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD13.D

Sample Name: 1909949003

```

=====
Injection Date: 4/10/2019 12:41:59      Seq Line:          13
Sample Name:   1909949003              Location:         Vial 83
Acq Operator:  TNB                     Inj. No.:        1
                                           Inj. Vol.:       30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.718	PBA	123373.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD14.D

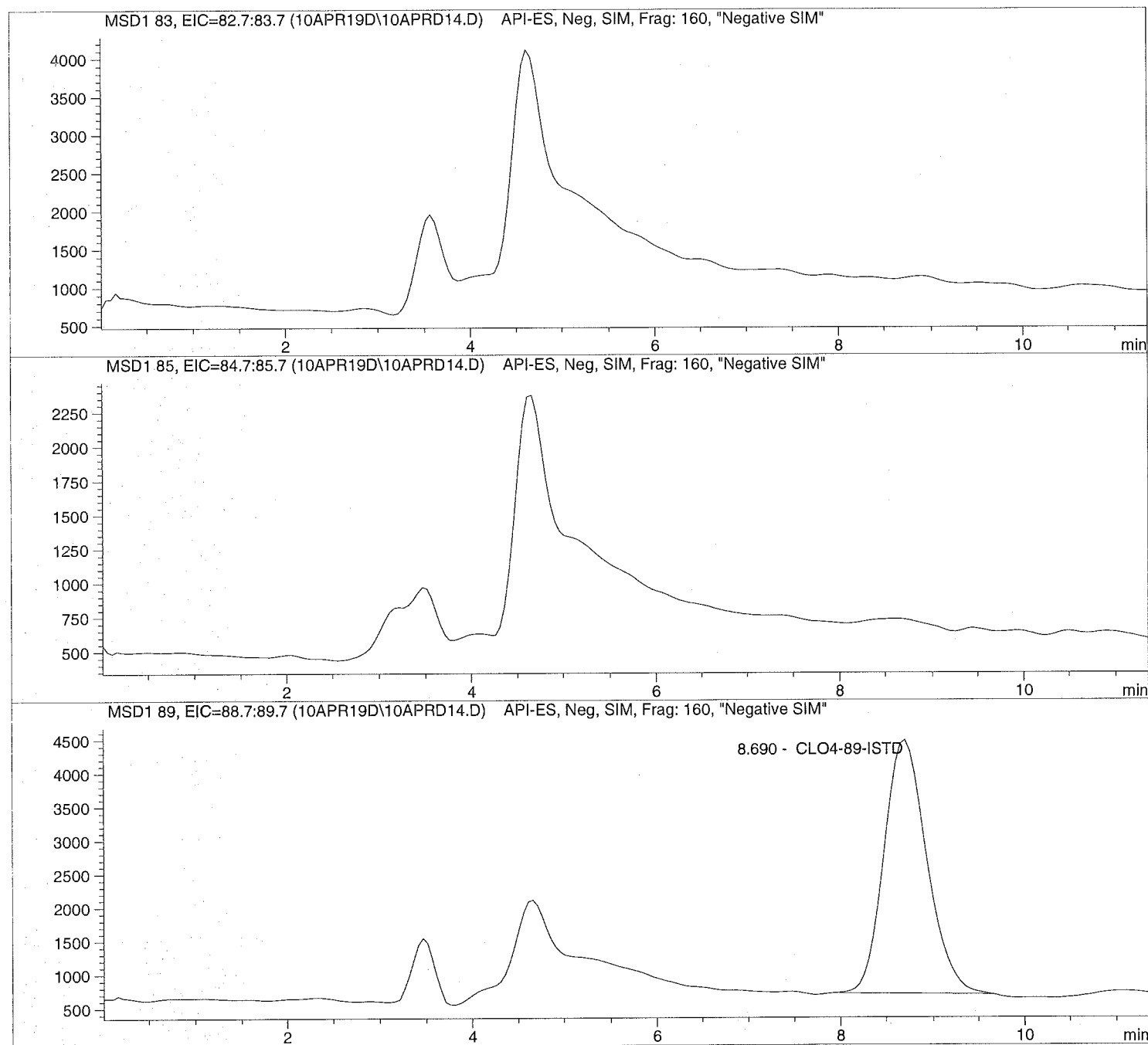
Sample Name: 1909949004

=====
Injection Date: 4/10/2019 12:55:15
Sample Name: 1909949004
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD14.D

Sample Name: 1909949004

```

=====
Injection Date: 4/10/2019 12:55:15      Seq Line:      14
Sample Name:    1909949004              Location:      Vial 84
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.690	BBA	120241.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD15.D

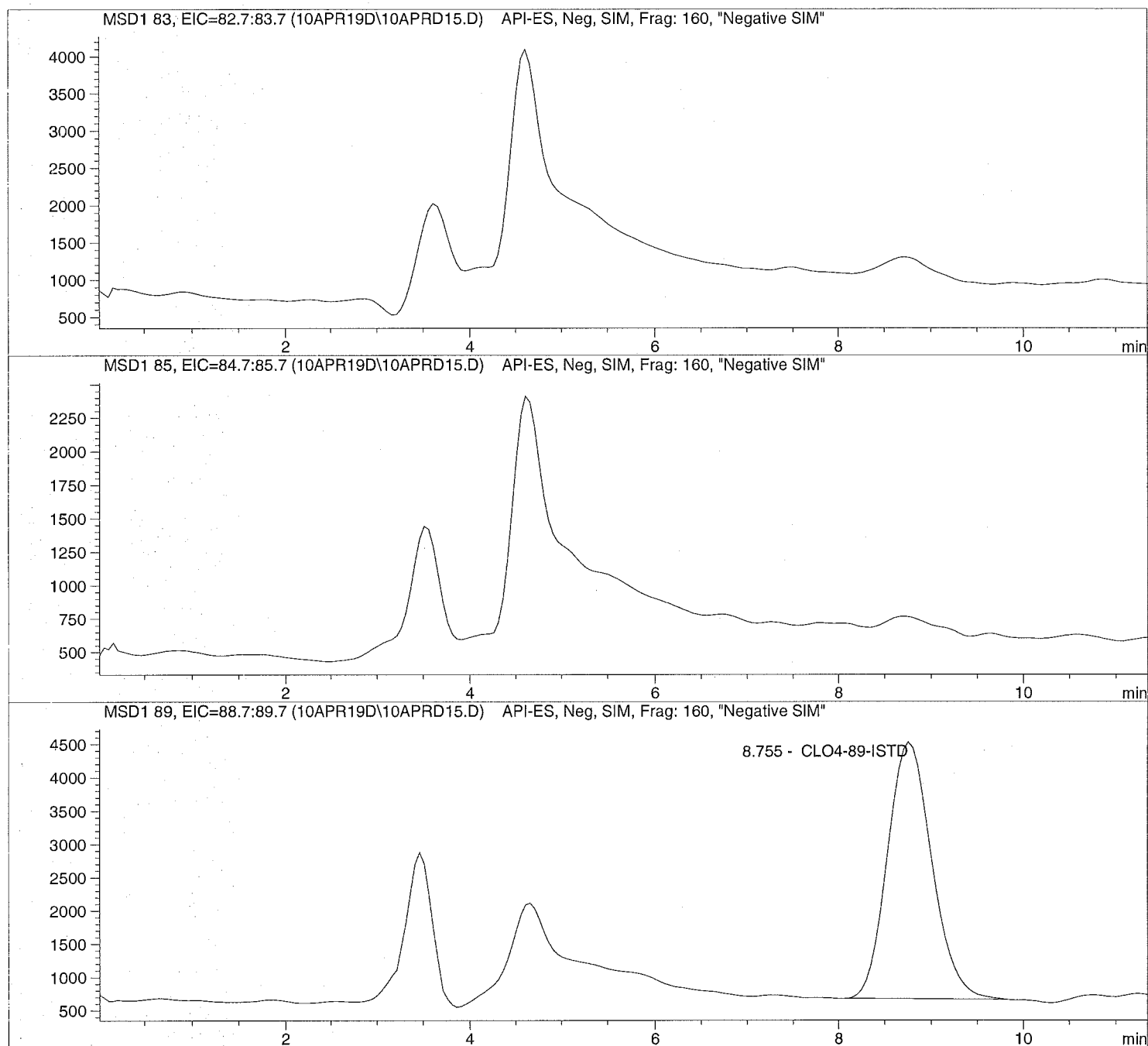
Sample Name: 1909949005

Injection Date: 4/10/2019 13:08:30
Sample Name: 1909949005
Acq Operator: TNB

Seq Line: 15
Location: Vial 85
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD15.D

Sample Name: 1909949005

```

=====
Injection Date: 4/10/2019 13:08:30      Seq Line:      15
Sample Name:   1909949005                Location:      Vial 85
Acq Operator:  TNB                       Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	PBA	127767.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD16.D

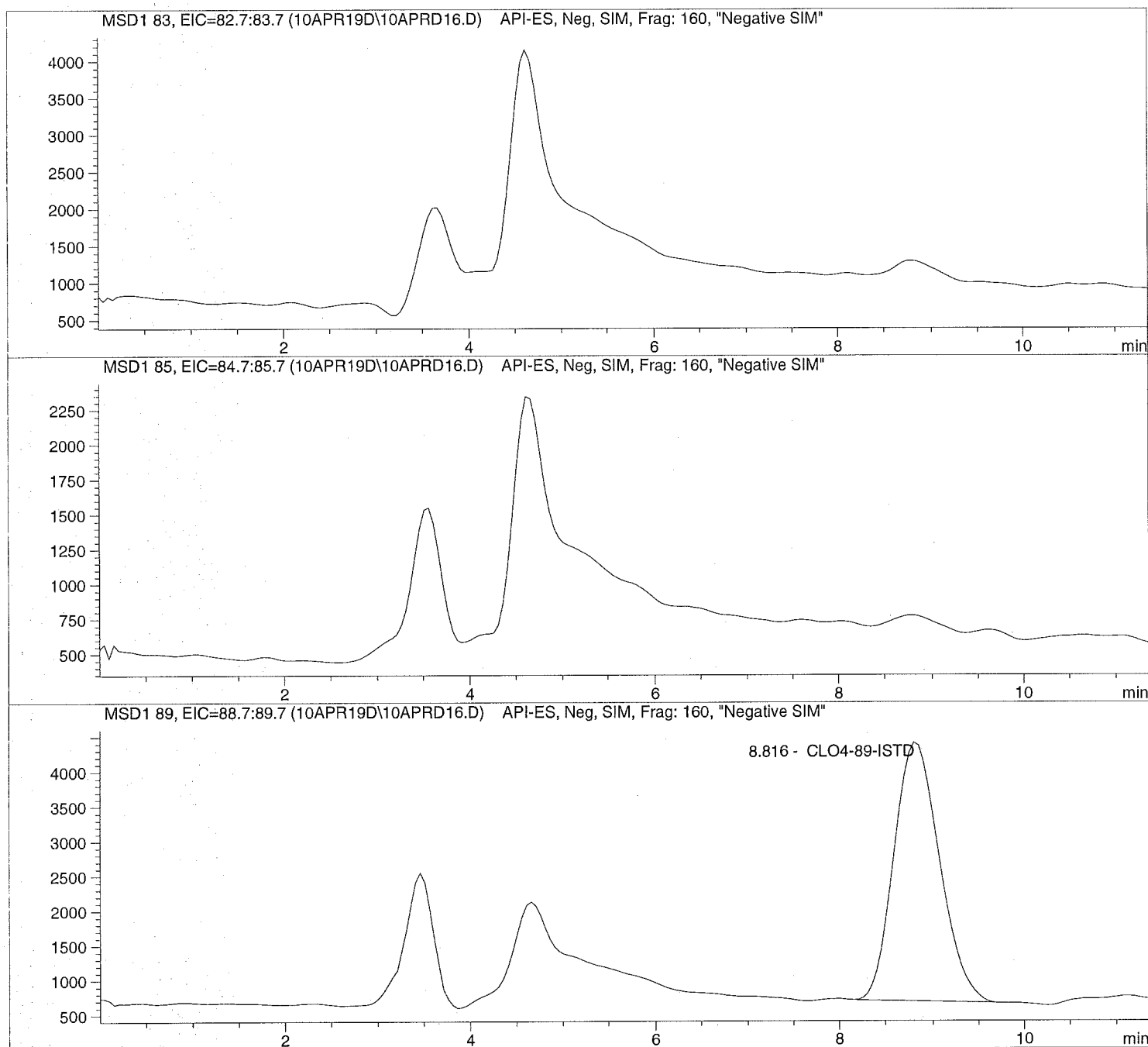
Sample Name: 1909949006

Injection Date: 4/10/2019 13:21:46
Sample Name: 1909949006
Acq Operator: TNB

Seq Line: 16
Location: Vial 86
Inj. No.: 1
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD16.D

Sample Name: 1909949006

```

=====
Injection Date: 4/10/2019 13:21:46      Seq Line:          16
Sample Name:   1909949006              Location:          Vial 86
Acq Operator:  TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:  3/19/2019 14:48:09
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:    1.000000
Dilution:     1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.816	BBA	123964.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD17.D

Sample Name: 647202 CCV@25

Injection Date: 4/10/2019 13:35:04

Seq Line: 17

Sample Name: 647202 CCV@25

Location: Vial 71

Acq Operator: TNB

Inj. No.: 1

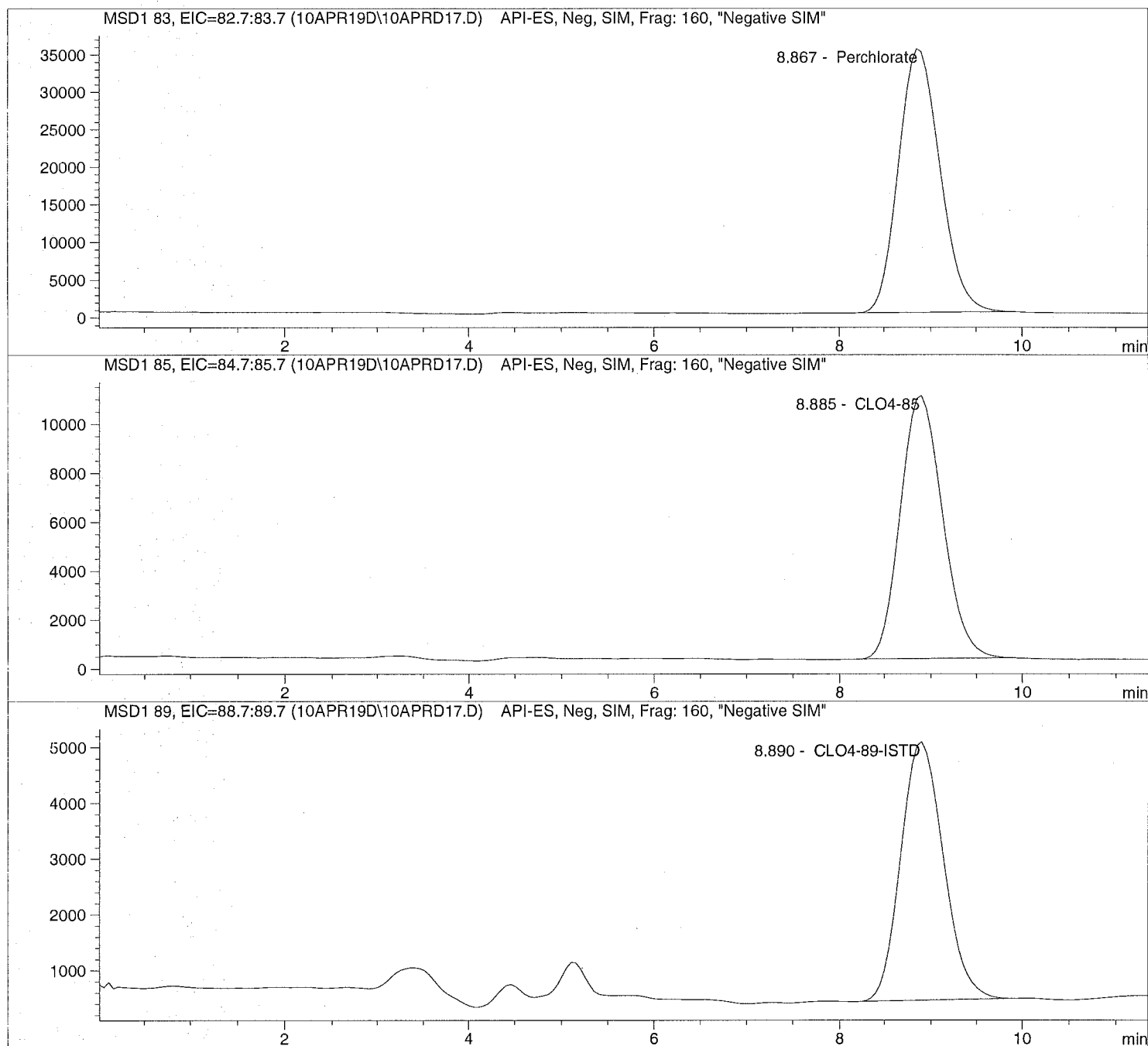
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:48:09

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\10APR19D\10APRD17.D Sample Name: 647202 CCV@25

```

=====
Injection Date: 4/10/2019 13:35:04      Seq Line:            17
Sample Name:    647202    CCV@25            Location:            Vial 71
Acq Operator:   TNB                        Inj. No.:            1
                                             Inj. Vol.:            30 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:48:09

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:            Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:           1.000000
Dilution:             1.000000
Sample Amount:        25.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.867	PBA	1080071.3	22.5952	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.885	PBA	336319.0	23.6329	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.890	PBA	146280.5	5.0000	CLO4-89-ISTD

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Initial Calibration

=====
Calibration Table
=====

Perchlorate

Calib. Data Modified : 3/19/2019 2:35:19 PM

Calculate : Internal Standard
Based on : Peak Area

Rel. Reference Window : 20.000 %
Abs. Reference Window : 0.000 min
Rel. Non-ref. Window : 20.000 %
Abs. Non-ref. Window : 0.000 min

Use Multiplier & Dilution Factor with ISTDs
Uncalibrated Peaks : not reported
Partial Calibration : No recalibration if peaks missing

Curve Type : Quadratic (some peaks differ, see below)
Origin : Ignored (some peaks differ, see below)
Weight : Linear (Amnt) (some peaks differ, see below)

Recalibration Settings:
Average Response : Average all calibrations
Average Retention Time: Floating Average New 75%

Calibration Report Options :
Printout of recalibrations within a sequence:
 Calibration Table after Recalibration
 Normal Report after Recalibration
If the sequence is done with bracketing:
 Results of first cycle (ending previous bracket)

Default Sample ISTD Information (if not set in sample table):

ISTD #	ISTD Amount	Name
1	5.00000	CLO4-89-ISTD

Signal 1: MSD1 83, EIC=82.7:83.7
Signal 2: MSD1 85, EIC=84.7:85.7
Signal 3: MSD1 89, EIC=88.7:89.7

RetTime [min]	Lvl	Amount	Area	Amt/Area	Ref	Grp	Name
8.744	1	1.00000	7.76074e4	1.28854e-5	1		Perchlorate
	2	2.00000	1.35273e5	1.47849e-5			
	3	5.00000	3.37764e5	1.48033e-5			
	4	10.00000	6.83454e5	1.46316e-5			
	5	25.00000	2.08433e6	1.19943e-5			
	6	50.00000	4.13334e6	1.20968e-5			
	7	75.00000	5.99313e6	1.25143e-5			
8.755	2	1.00000	2.36780e4	4.22333e-5	1		CLO4-85
	2	2.00000	4.69486e4	4.25998e-5			
	3	5.00000	1.06124e5	4.71147e-5			
	4	10.00000	2.13523e5	4.68335e-5			
	5	25.00000	6.14295e5	4.06971e-5			
	6	50.00000	1.19814e6	4.17315e-5			
	7	75.00000	1.78355e6	4.20509e-5			
8.766	3	5.00000	2.73208e5	1.83011e-5	+I1		CLO4-89-ISTD
	2	5.00000	2.24886e5	2.22335e-5			
	3	5.00000	2.33196e5	2.14412e-5			
	4	5.00000	2.34454e5	2.13262e-5			
	5	5.00000	2.50568e5	1.99547e-5			
	6	5.00000	2.30977e5	2.16472e-5			

RetTime [min]	Lvl Sig	Amount	Area	Amt/Area	Ref Grp Name
	7	5.00000	2.21504e5	2.25729e-5	

More compound-specific settings:

Compound: Perchlorate

Time Window : From 6.654 min To 12.544 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

Compound: CLO4-85

Time Window : From 6.650 min To 12.505 min
Curve Type : Quadratic
Origin : Ignored
Calibration Level Weights:/
Level 1 : 1
Level 2 : 0.5
Level 3 : 0.2
Level 4 : 0.1
Level 5 : 0.04
Level 6 : 0.02
Level 7 : 0.013333

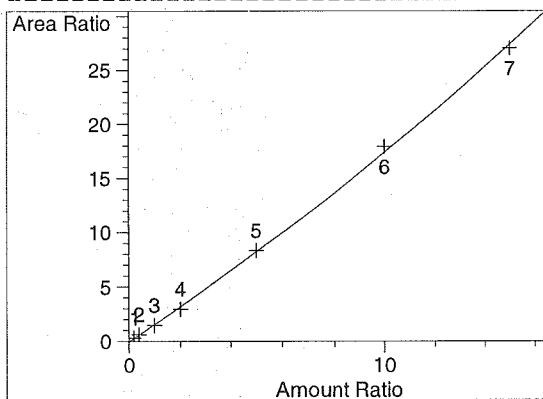
Compound: CLO4-89-ISTD

Time Window : From 6.659 min To 12.466 min
Curve Type : Linear
Origin : Included
Calibration Level Weights:/
Level 1 : 1
Level 2 : 1
Level 3 : 1
Level 4 : 1
Level 5 : 1
Level 6 : 1
Level 7 : 1

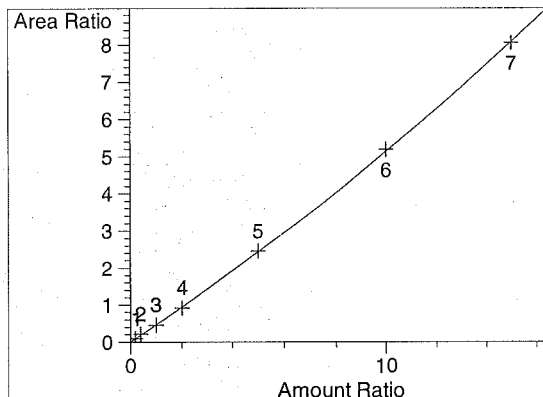
=====
Peak Sum Table
=====

No Entries in table
=====

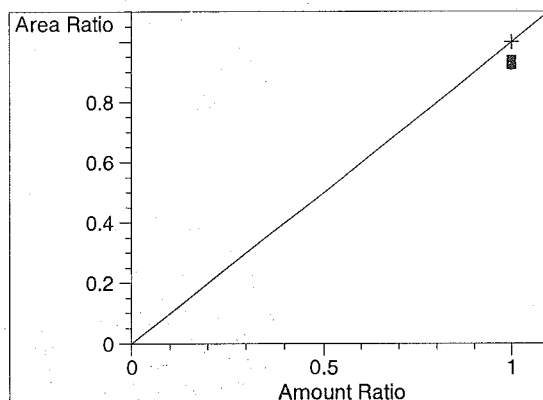
=====
 Calibration Curves
 =====



Perchlorate at exp. RT: 8.744
 MSD1 83, EIC=82.7:83.7
 Correlation: 0.99957
 Residual Std. Dev.: 0.30744
 Formula: $y = ax^2 + bx + c$
 a: 1.76988e-2
 b: 1.56480
 c: -4.92430e-2
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-85 at exp. RT: 8.755
 MSD1 85, EIC=84.7:85.7
 Correlation: 0.99983
 Residual Std. Dev.: 0.03473
 Formula: $y = ax^2 + bx + c$
 a: 5.13396e-3
 b: 4.62055e-1
 c: 4.97209e-4
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 0.5
 Level 3 : 0.2
 Level 4 : 0.1
 Level 5 : 0.04
 Level 6 : 0.02
 Level 7 : 0.013333



CLO4-89-ISTD at exp. RT: 8.766
 MSD1 89, EIC=88.7:89.7
 Correlation: 1.00000
 Residual Std. Dev.: 0.00000
 Formula: $y = mx + b$
 m: 1.00000
 b: 0.00000
 x: Amount Ratio
 y: Area Ratio
 Calibration Level Weights:
 Level 1 : 1
 Level 2 : 1
 Level 3 : 1
 Level 4 : 1
 Level 5 : 1
 Level 6 : 1
 Level 7 : 1

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP2.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	7.76074e4	8.744	1.06245
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.35273e5	8.992	2.06969
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.37764e5	8.586	4.73474
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	6.83454e5	8.698	9.27727
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.08433e6	8.451	25.29036
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	4.13334e6	8.810	51.36844
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	5.99313e6	8.586	74.16754
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	7.34719e5	8.702	9.25940

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.36780e4	8.755	9.30535e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	4.69486e4	9.012	2.24255
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.06124e5	8.602	4.86656
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.13523e5	8.713	9.64312
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	6.14295e5	8.468	25.12159
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.19814e6	8.825	50.46721
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	1.78355e6	8.603	74.72019
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.27495e5	8.721	9.54024

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	2.73208e5	8.766	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.24886e5	9.012	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	2.33196e5	8.609	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.34454e5	8.716	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.50568e5	8.472	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	2.30977e5	8.825	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.21504e5	8.610	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.52544e5	8.725	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ 0.2ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI03.D

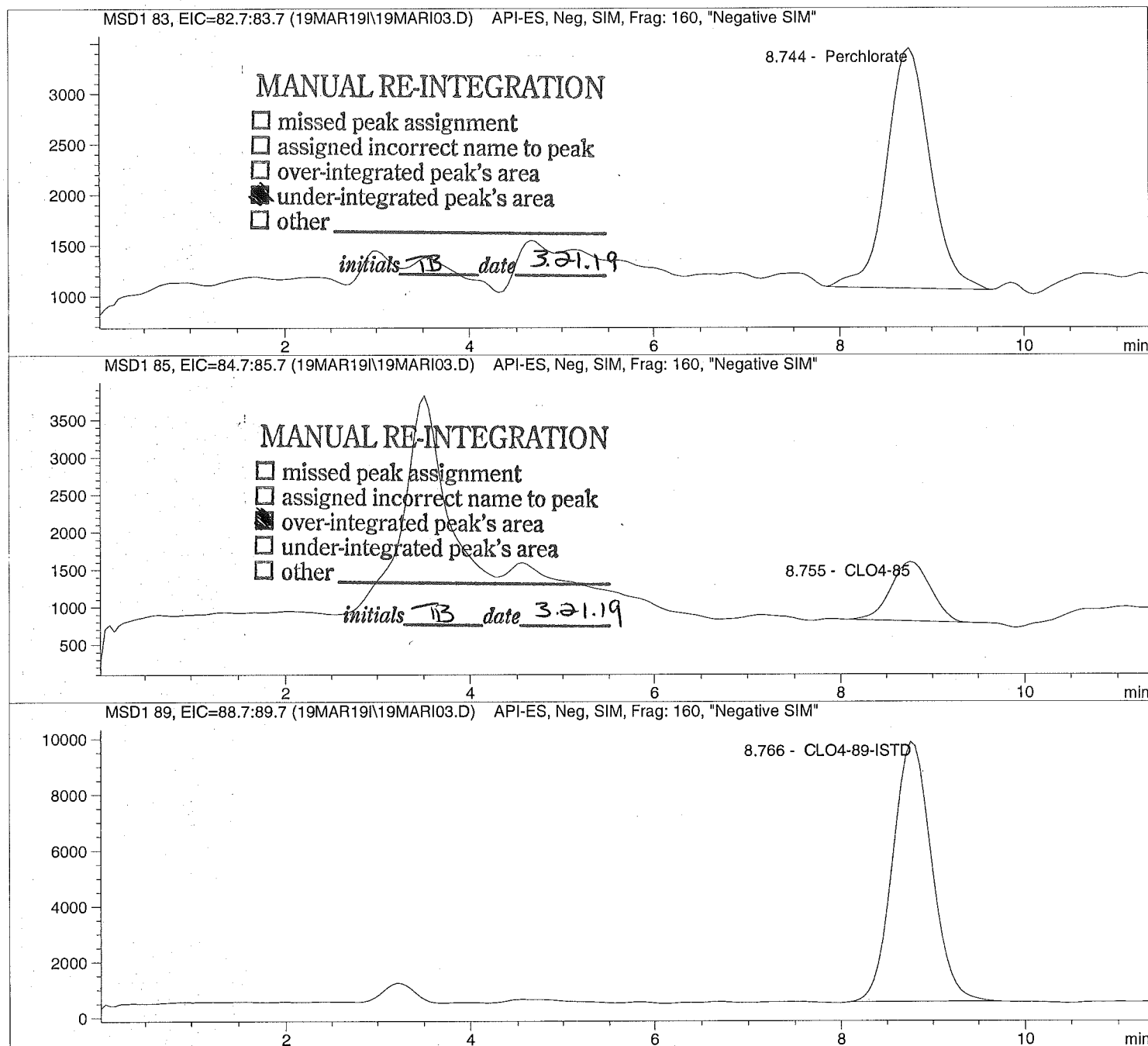
Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L           Location:  Vial 73
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  1.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	MM	77607.4	1.0625	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	MM	23678.0	0.9305	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

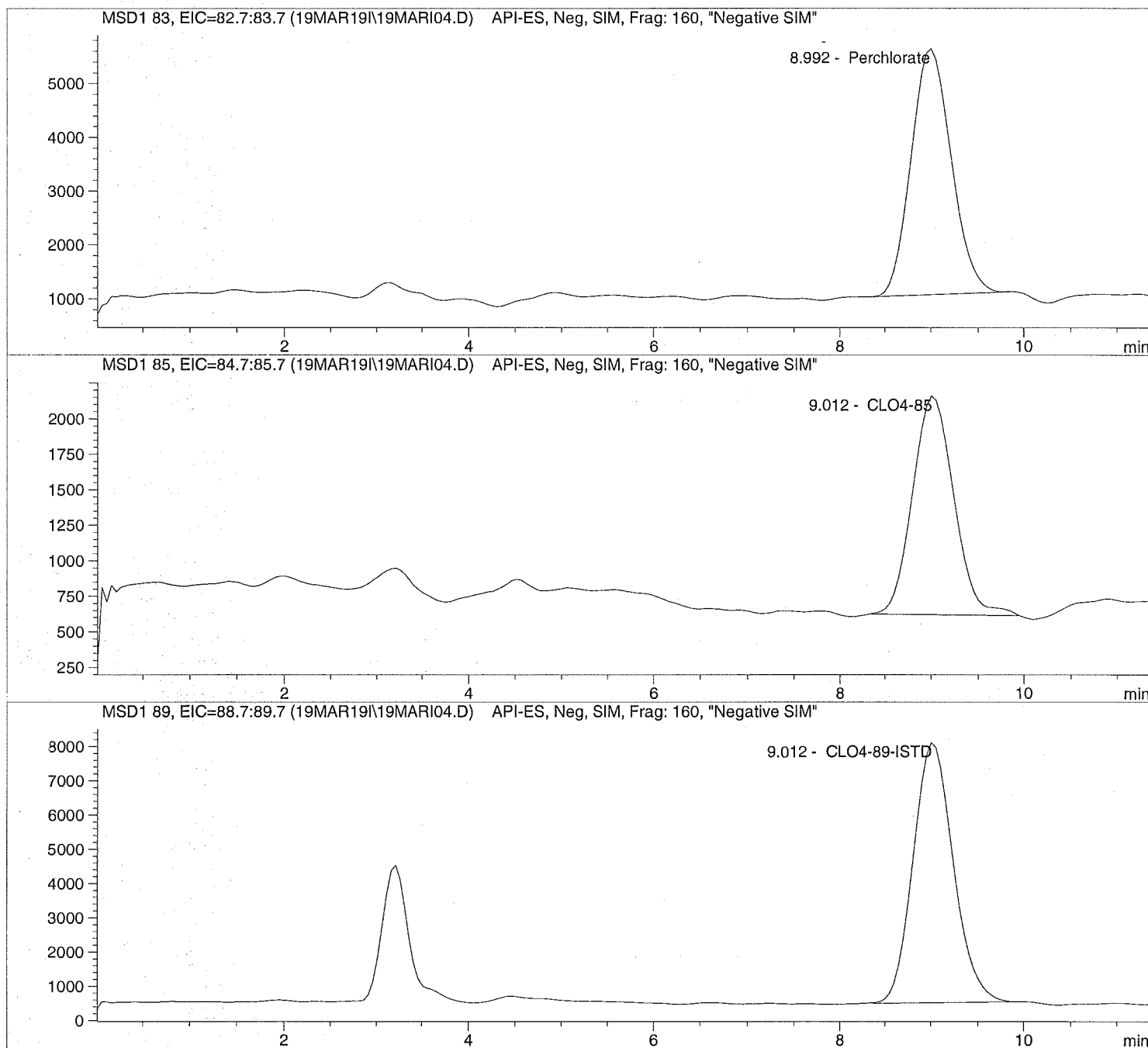
```

Data file: C:\HPCHEM\1\DATA\19MAR19\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```
=====
Injection Date: 3/19/2019 09:53:00      Seq Line: 4
Sample Name:    CLO4@ 2.0ug/L           Location: Vial 74
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 3/19/2019 09:53:00      Seq Line:                    4
Sample Name:    CLO4@ 2.0ug/L            Location:                Vial 74
Acq Operator:   TNB                      Inj. No.:                1
                                          Inj. Vol.:               30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:                    Signal
Calib. Data Modified:      Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:                1.000000
Dilution:                  1.000000
Sample Amount:             2.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.992	BBA	135272.8	2.0697	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	46948.6	2.2425	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
9.012	BBA	224885.9	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D

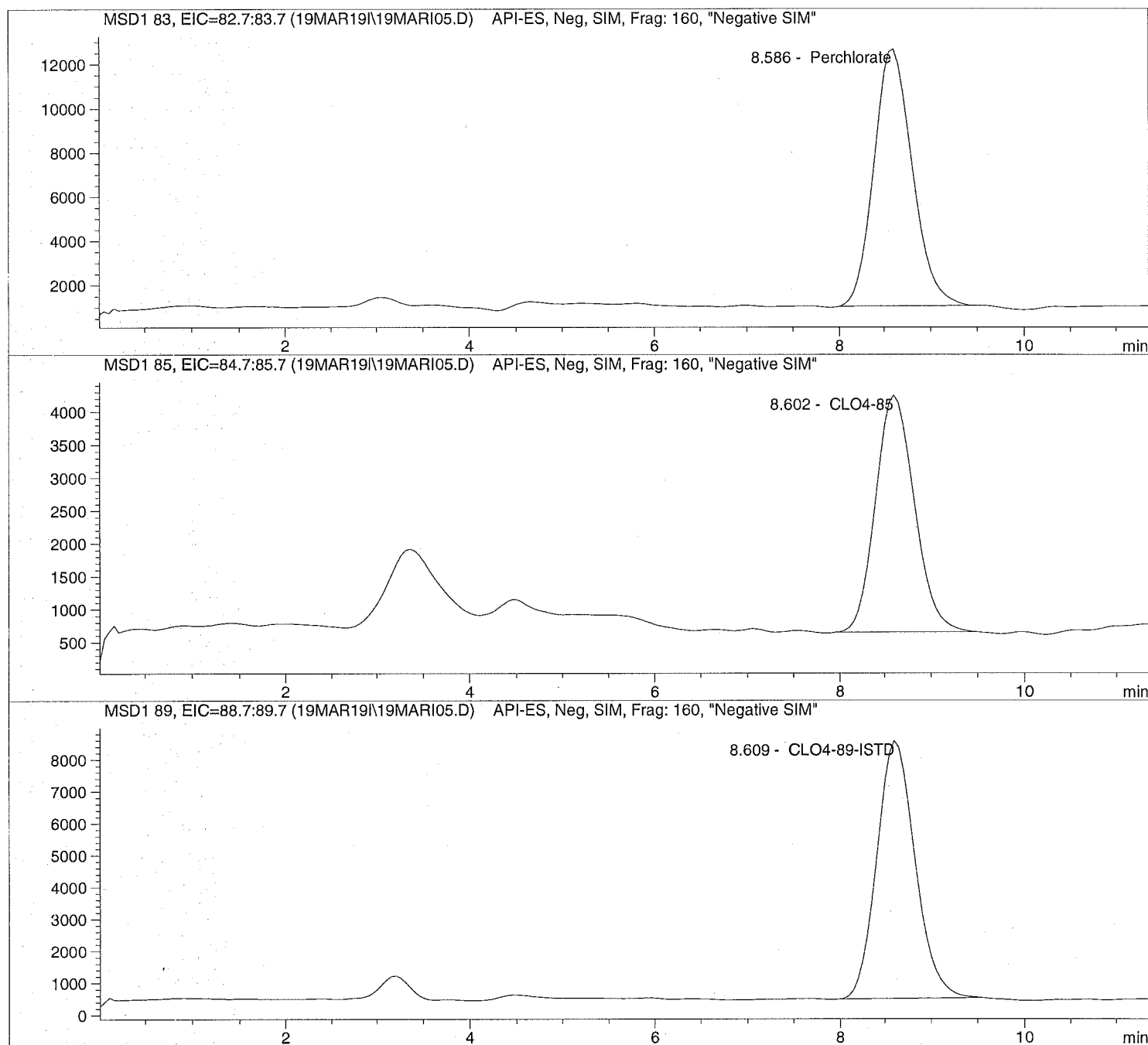
Sample Name: CLO4@ 5.0ug/L

Injection Date: 3/19/2019 10:06:16
Sample Name: CLO4@ 5.0ug/L
Acq Operator: TNB

Seq Line: 5
Location: Vial 75
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 3/19/2019 10:06:16      Seq Line:          5
Sample Name:    CLO4@ 5.0ug/L           Location:          Vial 75
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  5.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	337763.6	4.7347	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.602	PBA	106124.0	4.8666	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.609	PBA	233196.3	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D

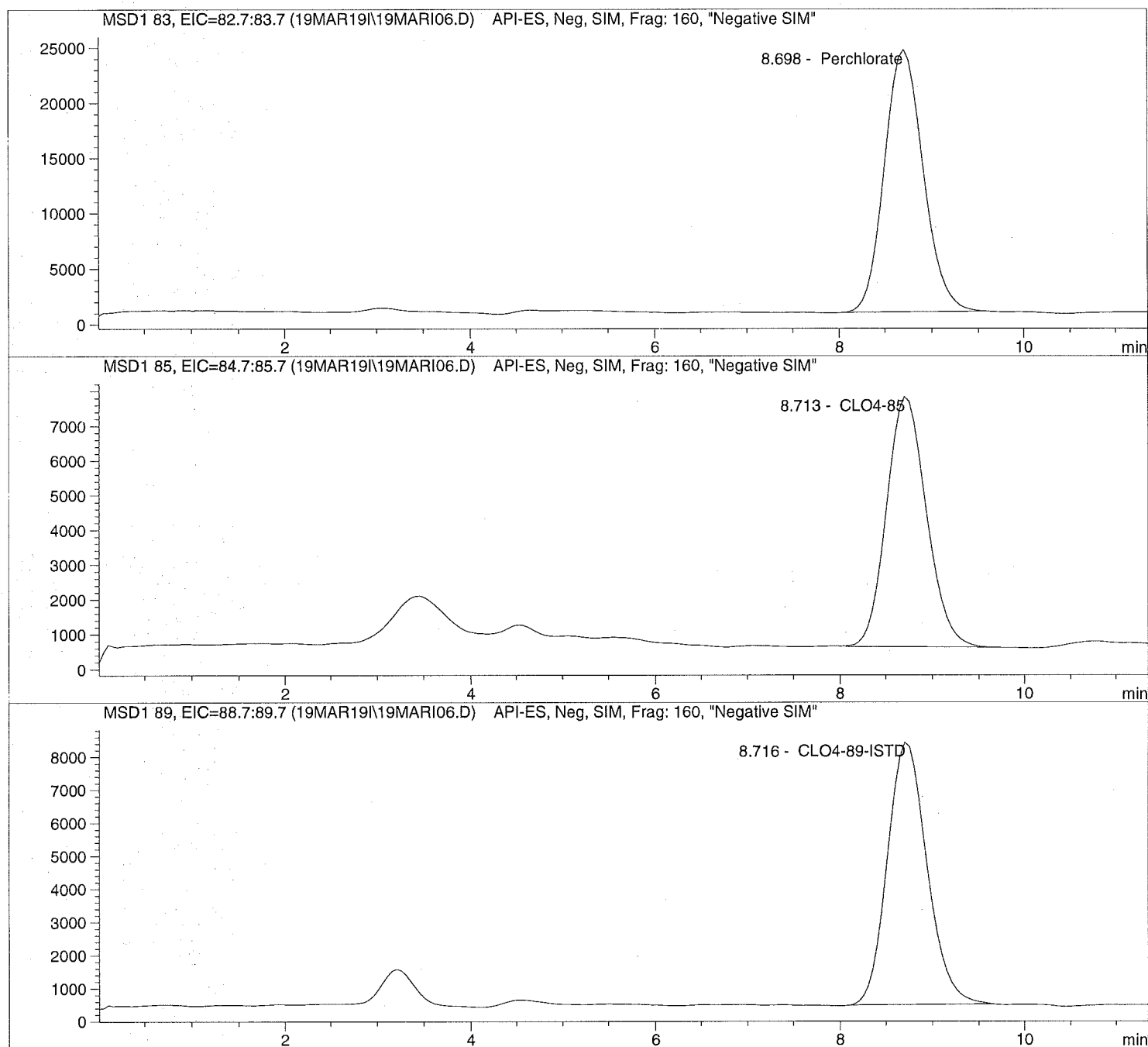
Sample Name: CLO4@ 10.ug/L

=====
Injection Date: 3/19/2019 10:19:32
Sample Name: CLO4@ 10.ug/L
Acq Operator: TNB

Seq Line: 6
Location: Vial 76
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI06.D Sample Name: CLO4@ 10.ug/L

```

=====
Injection Date: 3/19/2019 10:19:32      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.698	PBA	683454.4	9.2773	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.713	BBA	213522.6	9.6431	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.716	PBA	234453.6	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D

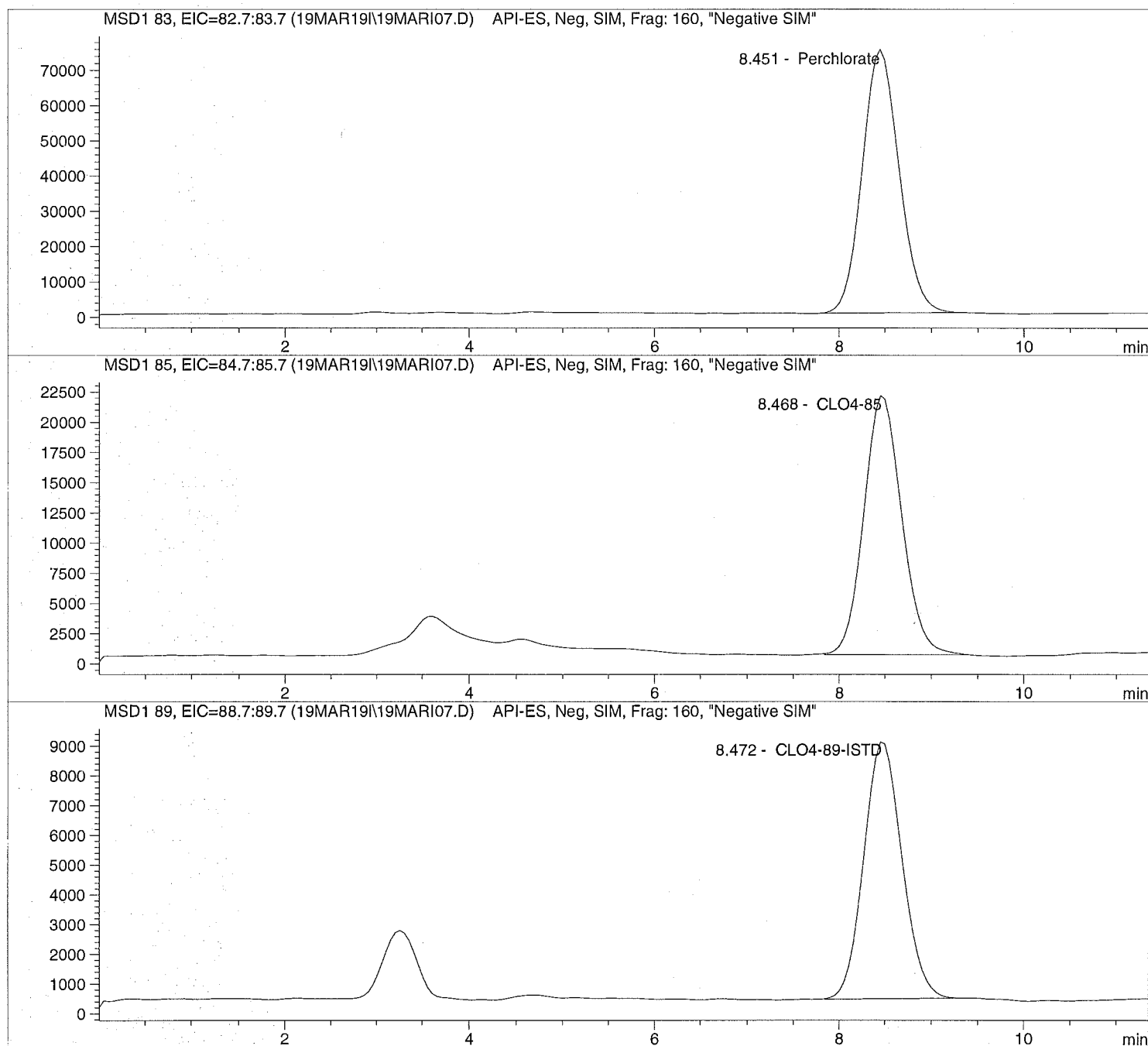
Sample Name: CLO4@ 25.ug/L

Injection Date: 3/19/2019 10:32:49
Sample Name: CLO4@ 25.ug/L
Acq Operator: TNB

Seq Line: 7
Location: Vial 77
Inj. No.: 1
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 3/19/2019 10:32:49      Seq Line:          7
Sample Name:    CLO4@ 25.ug/L           Location:          Vial 77
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019, 02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.451	PBA	2084327.4	25.2904	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.468	BBA	614294.8	25.1216	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.472	BBA	250568.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

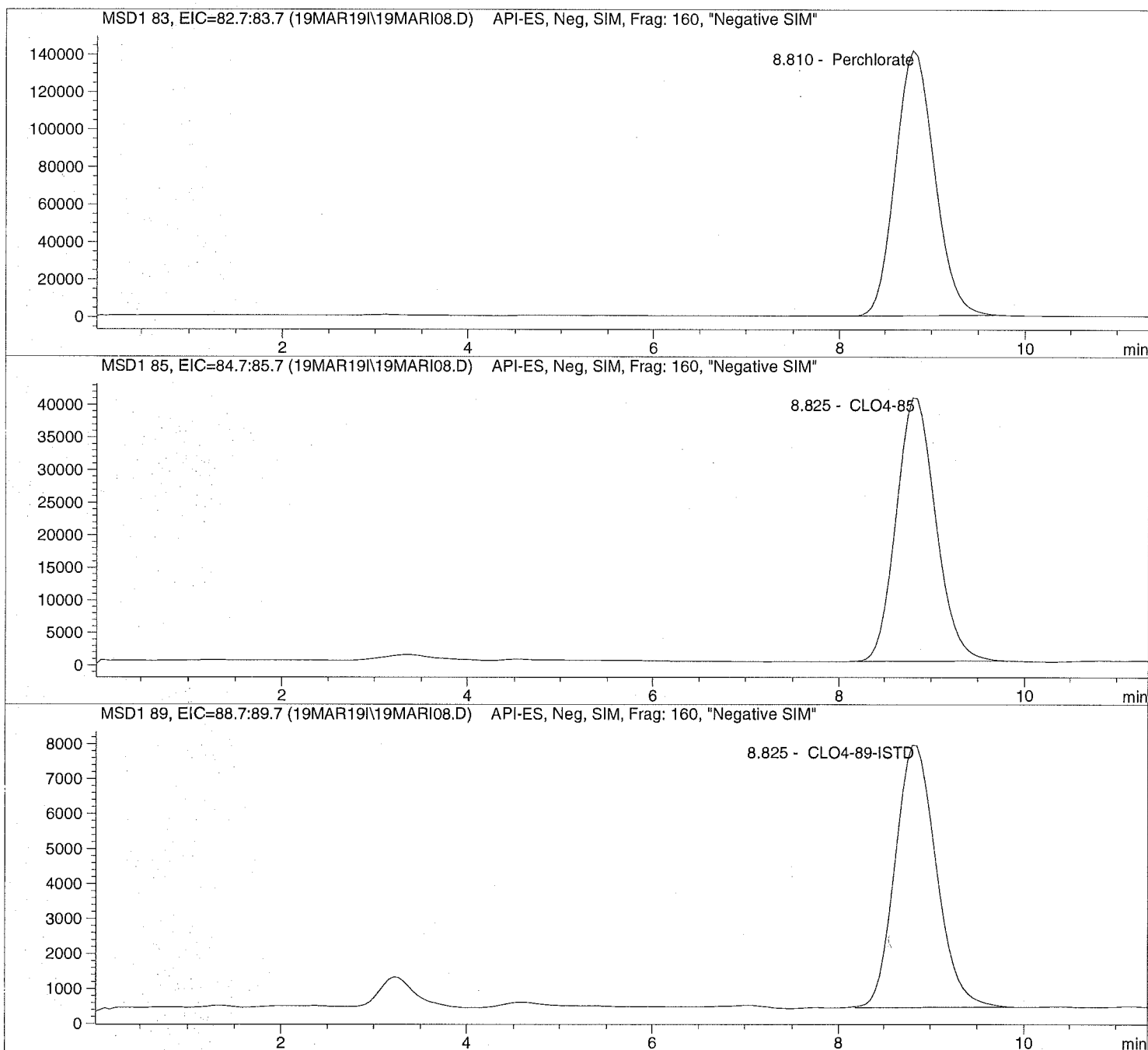
```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```
=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 3/19/2019 10:46:05      Seq Line:      8
Sample Name:    CLO4@ 50.ug/L           Location:      Vial 78
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  50.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.810	PBA	4133340.5	51.3684	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	1198135.6	50.4672	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.825	BBA	230976.7	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

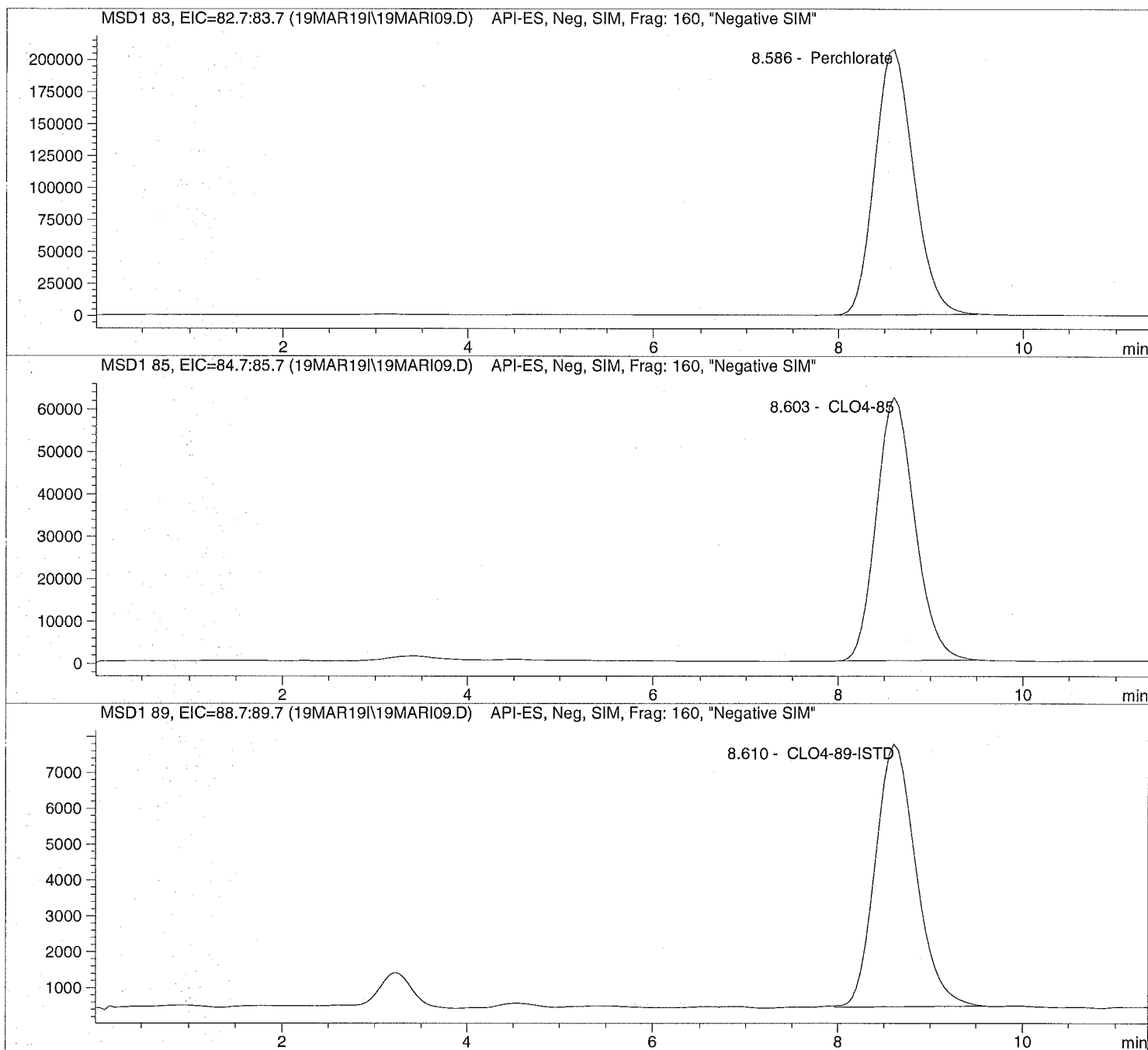
```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```
=====
Injection Date: 3/19/2019 10:59:22      Seq Line:          9
Sample Name:    CLO4@ 75.ug/L           Location:          Vial 79
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date: 3/19/2019 10:59:22      Seq Line:          9
Sample Name:    CLO4@ 75.ug/L           Location:          Vial 79
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:       30 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  75.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.586	PBA	5993128.0	74.1675	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.603	PBA	1783554.4	74.7202	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.610	BBA	221504.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D

Sample Name: ICAL Verf@10ug/L

Injection Date: 3/19/2019 11:12:42

Seq Line: 10

Sample Name: ICAL Verf@10ug/L

Location: Vial 80

Acq Operator: TNB

Inj. No.: 1

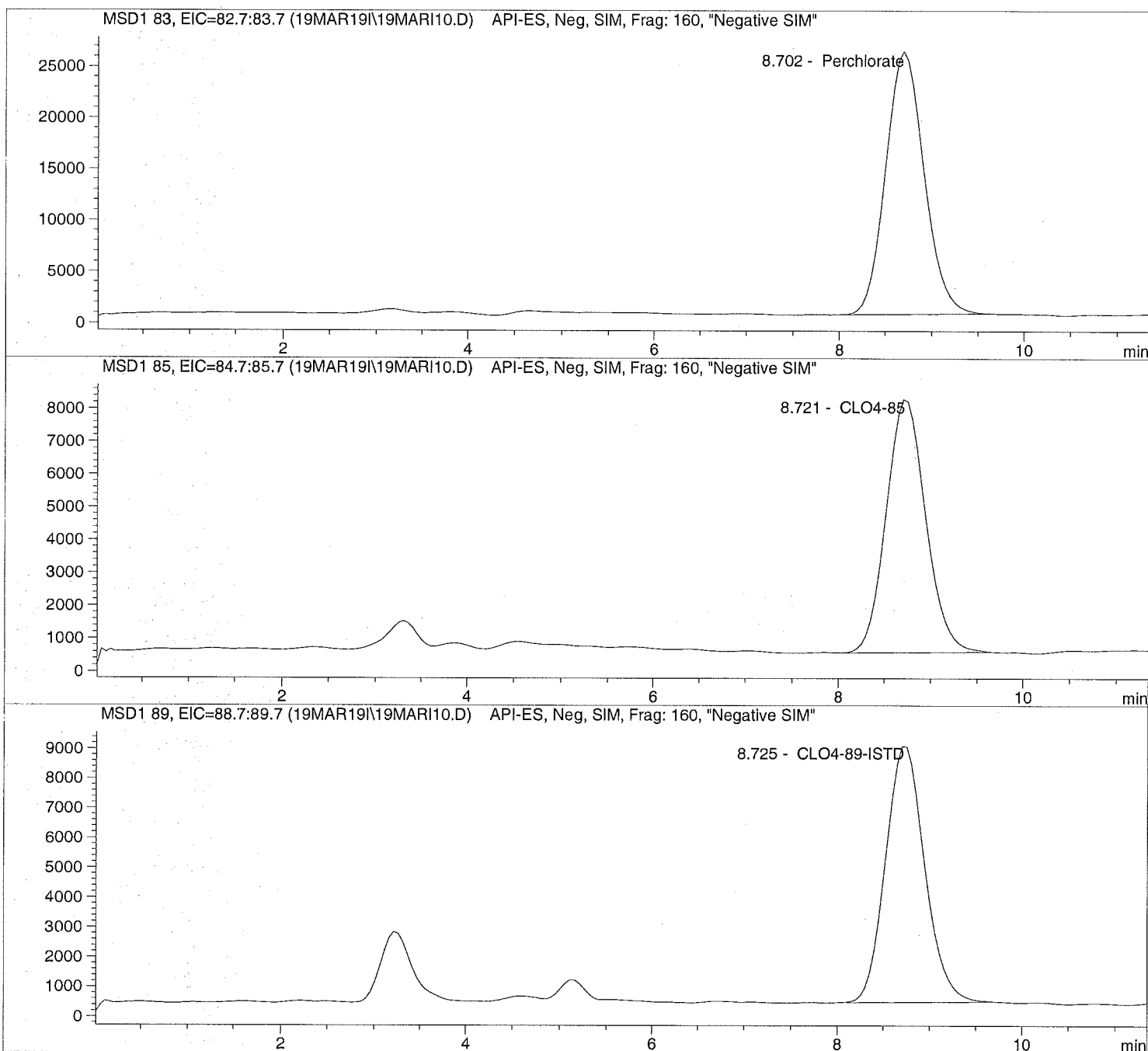
Inj. Vol.: 30 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:35:22

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI10.D Sample Name: ICAL Verf@10ug/L

```
=====
Injection Date: 3/19/2019 11:12:42      Seq Line:      10
Sample Name:    ICAL Verf@10ug/L        Location:      Vial 80
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    30 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed:   3/19/2019 14:35:22
```

Perchlorate analysis

=====

Sample Information

=====

```
Sorted By:      Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  10.000
```

=====

LCMS Results

=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.702	PBA	734718.7	9.2594	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.721	PBA	227494.7	9.5402	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.725	BBA	252544.4	5.0000	CLO4-89-ISTD

=====

*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D

Sample Name: CLO4@ 1.0ug/L

Injection Date: 3/19/2019 09:39:40

Seq Line: 3

Sample Name: CLO4@ 1.0ug/L

Location: Vial 73

Acq Operator: TNB

Inj. No.: 1

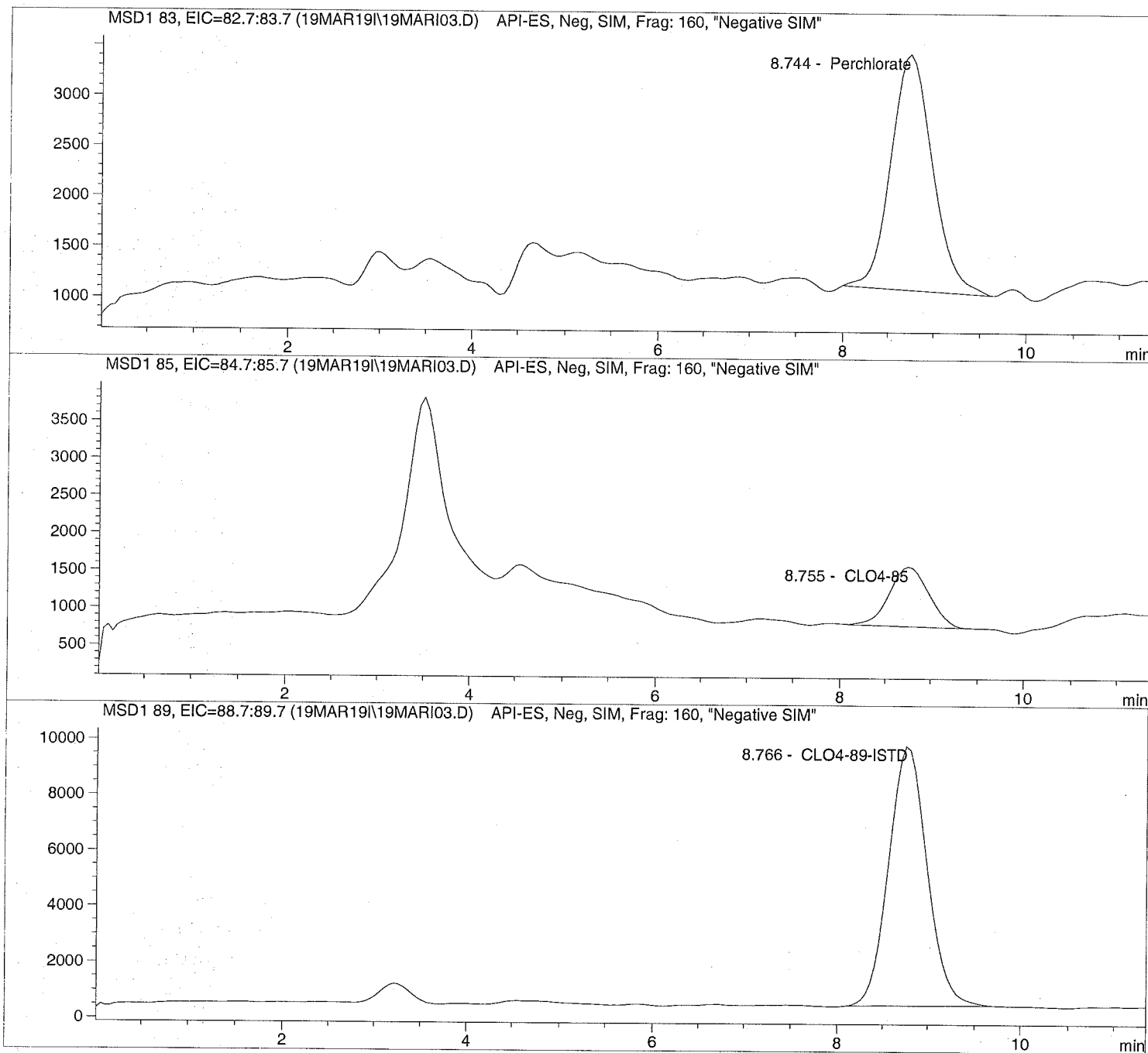
Inj. Vol.: 30 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M

Last Changed: 3/19/2019 14:38:25

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\19MAR19I\19MARI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 3/19/2019 09:39:40 Seq Line: 3
Sample Name: CLO4@ 1.0ug/L Location: Vial 73
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 30 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP2.M
Last Changed: 3/19/2019 14:38:25
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Mar. 2019,02:35:19 pm
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 1.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.744	BBA	74166.3	1.0224	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.755	BBA	24138.1	0.9487	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.766	BBA	273207.6	5.0000	CLO4-89-ISTD

*** End of Report ***

GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

APPENDIX D
LABORATORY ANALYTICAL RESULTS FOR LHAAP-16 (PROVIDED ON CD
ONLY)

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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10450 Stancliff Rd. Suite 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887

March 08, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Ste 129
Lakewood, CO 80228

Work Order: **HS19021428**

Laboratory Results for: **LHAAP Site 16**

Dear Marcia,

ALS Environmental received 9 sample(s) on Feb 27, 2019 for the analysis presented in the following report.

The analytical data provided relates directly to the samples received by ALS Environmental and for only the analyses requested. Results are expressed as "as received" unless otherwise noted.

QC sample results for this data met EPA or laboratory specifications except as noted in the Case Narrative or as noted with qualifiers in the QC batch information. Should this laboratory report need to be reproduced, it should be reproduced in full unless written approval has been obtained by ALS Environmental. Samples will be disposed in 30 days unless storage arrangements are made.

If you have any questions regarding this report, please feel free to call me.

Sincerely,

A handwritten signature in black ink, appearing to read 'Raj. P. Modashia', enclosed in a simple oval scribble.

Generated By: DAYNA.FISHER
RJ Modashia
Project Manager

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
Work Order: HS19021428

SAMPLE SUMMARY

Lab Samp ID	Client Sample ID	Matrix	TagNo	Collection Date	Date Received	Hold
HS19021428-01	16EW01_022619	Water		26-Feb-2019 08:30	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-02	16EW05_022619	Water		26-Feb-2019 08:50	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-03	16EW05_022619_a	Water		26-Feb-2019 08:50	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-04	16EW02_022619	Water		26-Feb-2019 09:00	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-05	16EW06_022619	Water		26-Feb-2019 09:10	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-06	16EW03_022619	Water		26-Feb-2019 09:20	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-07	16EW07_022619	Water		26-Feb-2019 09:30	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-08	16EW04_022619	Water		26-Feb-2019 09:40	27-Feb-2019 09:10	<input type="checkbox"/>
HS19021428-09	16EW08_022619	Water		26-Feb-2019 09:50	27-Feb-2019 09:10	<input type="checkbox"/>

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
Work Order: HS19021428

CASE NARRATIVE**Work Order Comments**

- The analysis for Perchlorate was subcontracted to ALS in Salt Lake City, Utah. Final report attached.
-

GCMS Volatiles by Method SW8260**Batch ID: R334128****Sample ID: 16EW01_022619 (HS19021428-01MS/MSD)**

- MS and/or MSD recovered outside control limits for multiple compounds

Batch ID: R334221**Sample ID: 16EW08_022619 (HS19021428-09MSD)**

- MSD recovered outside control limits for multiple compounds
-

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW01_022619
 Collection Date: 26-Feb-2019 08:30

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,1,1-Trichloroethane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,1,2,2-Tetrachloroethane	2.5	U	2.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,1,2-Trichloroethane	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,1-Dichloroethane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,1-Dichloroethene	14		1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,1-Dichloropropene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2,3-Trichlorobenzene	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2,3-Trichloropropane	2.5	U	2.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2,4-Trichlorobenzene	2.5	U	2.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2,4-Trimethylbenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2-Dibromo-3-chloropropane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2-Dibromoethane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2-Dichlorobenzene	2.5	U	2.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2-Dichloroethane	12		1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,2-Dichloropropane	2.5	U	2.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,3,5-Trimethylbenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,3-Dichlorobenzene	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,3-Dichloropropane	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
1,4-Dichlorobenzene	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
2,2-Dichloropropane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
2-Butanone	5.0	U	2.5	5.0	10	UG/L	5	06-Mar-2019 13:27	
2-Chlorotoluene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
2-Hexanone	5.0	U	5.0	5.0	10	UG/L	5	06-Mar-2019 13:27	
4-Chlorotoluene	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
4-Isopropyltoluene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
4-Methyl-2-pentanone	5.0	U	3.5	5.0	10	UG/L	5	06-Mar-2019 13:27	
Acetone	10	U	2.0	10	10	UG/L	5	06-Mar-2019 13:27	
Benzene	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Bromobenzene	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Bromochloromethane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Bromodichloromethane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Bromoform	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Bromomethane	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Carbon disulfide	5.0	U	3.0	5.0	10	UG/L	5	06-Mar-2019 13:27	
Carbon tetrachloride	2.5	U	2.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Chlorobenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Chloroethane	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Chloroform	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW01_022619
 Collection Date: 26-Feb-2019 08:30

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-01
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
cis-1,2-Dichloroethene	3,700		10	25	50	UG/L	50	06-Mar-2019 13:03	
cis-1,3-Dichloropropene	2.5	U	0.50	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Dibromochloromethane	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Dibromomethane	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Dichlorodifluoromethane	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Ethylbenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Hexachlorobutadiene	5.0	U	5.0	5.0	5.0	UG/L	5	06-Mar-2019 13:27	
Isopropylbenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
m,p-Xylene	5.0	U	2.5	5.0	10	UG/L	5	06-Mar-2019 13:27	
Methylene chloride	2.5	U	2.0	2.5	10	UG/L	5	06-Mar-2019 13:27	
n-Butylbenzene	2.5	U	2.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
n-Propylbenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Naphthalene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
o-Xylene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
sec-Butylbenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Styrene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
tert-Butylbenzene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Tetrachloroethene	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Toluene	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
trans-1,2-Dichloroethene	8.7		1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
trans-1,3-Dichloropropene	2.5	U	1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Trichloroethene	3,700		10	25	50	UG/L	50	06-Mar-2019 13:03	
Trichlorofluoromethane	2.5	U	1.5	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Vinyl chloride	45		1.0	2.5	5.0	UG/L	5	06-Mar-2019 13:27	
Surr: 1,2-Dichloroethane-d4	81.5			0	81-118	%REC	50	06-Mar-2019 13:03	
Surr: 1,2-Dichloroethane-d4	82.0			0	81-118	%REC	5	06-Mar-2019 13:27	
Surr: 4-Bromofluorobenzene	99.9			0	85-114	%REC	50	06-Mar-2019 13:03	
Surr: 4-Bromofluorobenzene	102			0	85-114	%REC	5	06-Mar-2019 13:27	
Surr: Dibromofluoromethane	85.1			0	80-119	%REC	50	06-Mar-2019 13:03	
Surr: Dibromofluoromethane	84.5			0	80-119	%REC	5	06-Mar-2019 13:27	
Surr: Toluene-d8	106			0	89-112	%REC	50	06-Mar-2019 13:03	
Surr: Toluene-d8	105			0	89-112	%REC	5	06-Mar-2019 13:27	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB	
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW05_022619
 Collection Date: 26-Feb-2019 08:50

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,1-Dichloroethane	1.4		0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,1-Dichloroethene	6.0		0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2-Dichloroethane	6.0		0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 15:52	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	06-Mar-2019 15:52	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	06-Mar-2019 15:52	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	06-Mar-2019 15:52	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	06-Mar-2019 15:52	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW05_022619
 Collection Date: 26-Feb-2019 08:50

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-02
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
cis-1,2-Dichloroethene	1,000		5.0	12	25	UG/L	25	06-Mar-2019 16:43	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	06-Mar-2019 15:52	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 15:52	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	06-Mar-2019 15:52	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
trans-1,2-Dichloroethene	3.7		0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Trichloroethene	1,700		5.0	12	25	UG/L	25	06-Mar-2019 16:43	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 15:52	
Vinyl chloride	260		5.0	12	25	UG/L	25	06-Mar-2019 16:43	
Surr: 1,2-Dichloroethane-d4	101			0	81-118	%REC	25	06-Mar-2019 16:43	
Surr: 1,2-Dichloroethane-d4	81.6			0	81-118	%REC	1	06-Mar-2019 15:52	
Surr: 4-Bromofluorobenzene	99.0			0	85-114	%REC	1	06-Mar-2019 15:52	
Surr: 4-Bromofluorobenzene	104			0	85-114	%REC	25	06-Mar-2019 16:43	
Surr: Dibromofluoromethane	85.8			0	80-119	%REC	1	06-Mar-2019 15:52	
Surr: Dibromofluoromethane	102			0	80-119	%REC	25	06-Mar-2019 16:43	
Surr: Toluene-d8	106			0	89-112	%REC	1	06-Mar-2019 15:52	
Surr: Toluene-d8	95.6			0	89-112	%REC	25	06-Mar-2019 16:43	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB	
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW05_022619_a
 Collection Date: 26-Feb-2019 08:50

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,1-Dichloroethane	1.4		0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,1-Dichloroethene	6.5		0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2-Dichloroethane	6.4		0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 17:31	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	06-Mar-2019 17:31	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	06-Mar-2019 17:31	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	06-Mar-2019 17:31	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	06-Mar-2019 17:31	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW05_022619_a
 Collection Date: 26-Feb-2019 08:50

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-03
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
cis-1,2-Dichloroethene	1,200		5.0	12	25	UG/L	25	06-Mar-2019 18:21	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	06-Mar-2019 17:31	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 17:31	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	06-Mar-2019 17:31	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
trans-1,2-Dichloroethene	3.9		0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Trichloroethene	1,900		5.0	12	25	UG/L	25	06-Mar-2019 18:21	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 17:31	
Vinyl chloride	420		5.0	12	25	UG/L	25	06-Mar-2019 18:21	
Surr: 1,2-Dichloroethane-d4	86.2			0	81-118	%REC	1	06-Mar-2019 17:31	
Surr: 1,2-Dichloroethane-d4	112			0	81-118	%REC	25	06-Mar-2019 18:21	
Surr: 4-Bromofluorobenzene	98.8			0	85-114	%REC	1	06-Mar-2019 17:31	
Surr: 4-Bromofluorobenzene	103			0	85-114	%REC	25	06-Mar-2019 18:21	
Surr: Dibromofluoromethane	84.9			0	80-119	%REC	1	06-Mar-2019 17:31	
Surr: Dibromofluoromethane	111			0	80-119	%REC	25	06-Mar-2019 18:21	
Surr: Toluene-d8	95.8			0	89-112	%REC	25	06-Mar-2019 18:21	
Surr: Toluene-d8	106			0	89-112	%REC	1	06-Mar-2019 17:31	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB	
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW02_022619
 Collection Date: 26-Feb-2019 09:00

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-04
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						
								Analyst: PC
1,1,1,2-Tetrachloroethane	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
1,1,1-Trichloroethane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
1,1,2,2-Tetrachloroethane	25	U	25	25	50	UG/L	50	07-Mar-2019 15:56
1,1,2-Trichloroethane	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
1,1-Dichloroethane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
1,1-Dichloroethene	86		10	25	50	UG/L	50	07-Mar-2019 15:56
1,1-Dichloropropene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
1,2,3-Trichlorobenzene	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56
1,2,3-Trichloropropane	25	U	25	25	50	UG/L	50	07-Mar-2019 15:56
1,2,4-Trichlorobenzene	25	U	25	25	50	UG/L	50	07-Mar-2019 15:56
1,2,4-Trimethylbenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
1,2-Dibromo-3-chloropropane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
1,2-Dibromoethane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
1,2-Dichlorobenzene	25	U	25	25	50	UG/L	50	07-Mar-2019 15:56
1,2-Dichloroethane	34	J	10	25	50	UG/L	50	07-Mar-2019 15:56
1,2-Dichloropropane	25	U	25	25	50	UG/L	50	07-Mar-2019 15:56
1,3,5-Trimethylbenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
1,3-Dichlorobenzene	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56
1,3-Dichloropropane	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
1,4-Dichlorobenzene	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56
2,2-Dichloropropane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
2-Butanone	50	U	25	50	100	UG/L	50	07-Mar-2019 15:56
2-Chlorotoluene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
2-Hexanone	50	U	50	50	100	UG/L	50	07-Mar-2019 15:56
4-Chlorotoluene	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56
4-Isopropyltoluene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
4-Methyl-2-pentanone	50	U	35	50	100	UG/L	50	07-Mar-2019 15:56
Acetone	100	U	20	100	100	UG/L	50	07-Mar-2019 15:56
Benzene	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
Bromobenzene	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56
Bromochloromethane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
Bromodichloromethane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56
Bromoform	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56
Bromomethane	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56
Carbon disulfide	50	U	30	50	100	UG/L	50	07-Mar-2019 15:56
Carbon tetrachloride	25	U	25	25	50	UG/L	50	07-Mar-2019 15:56
Chlorobenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
Chloroethane	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56
Chloroform	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW02_022619
 Collection Date: 26-Feb-2019 09:00

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-04
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56	
cis-1,2-Dichloroethene	17,000		50	120	250	UG/L	250	07-Mar-2019 19:57	
cis-1,3-Dichloropropene	25	U	5.0	25	50	UG/L	50	07-Mar-2019 15:56	
Dibromochloromethane	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Dibromomethane	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56	
Dichlorodifluoromethane	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Ethylbenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Hexachlorobutadiene	50	U	50	50	50	UG/L	50	07-Mar-2019 15:56	
Isopropylbenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
m,p-Xylene	50	U	25	50	100	UG/L	50	07-Mar-2019 15:56	
Methylene chloride	25	U	20	25	100	UG/L	50	07-Mar-2019 15:56	
n-Butylbenzene	25	U	20	25	50	UG/L	50	07-Mar-2019 15:56	
n-Propylbenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Naphthalene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
o-Xylene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
sec-Butylbenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Styrene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
tert-Butylbenzene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Tetrachloroethene	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Toluene	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56	
trans-1,2-Dichloroethene	24	J	10	25	50	UG/L	50	07-Mar-2019 15:56	
trans-1,3-Dichloropropene	25	U	10	25	50	UG/L	50	07-Mar-2019 15:56	
Trichloroethene	34,000		50	120	250	UG/L	250	07-Mar-2019 19:57	
Trichlorofluoromethane	25	U	15	25	50	UG/L	50	07-Mar-2019 15:56	
Vinyl chloride	51		10	25	50	UG/L	50	07-Mar-2019 15:56	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>84.7</i>			0	<i>81-118</i>	%REC	50	07-Mar-2019 15:56	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>82.7</i>			0	<i>81-118</i>	%REC	250	07-Mar-2019 19:57	
<i>Surr: 4-Bromofluorobenzene</i>	<i>99.7</i>			0	<i>85-114</i>	%REC	50	07-Mar-2019 15:56	
<i>Surr: 4-Bromofluorobenzene</i>	<i>96.8</i>			0	<i>85-114</i>	%REC	250	07-Mar-2019 19:57	
<i>Surr: Dibromofluoromethane</i>	<i>82.6</i>			0	<i>80-119</i>	%REC	50	07-Mar-2019 15:56	
<i>Surr: Dibromofluoromethane</i>	<i>83.0</i>			0	<i>80-119</i>	%REC	250	07-Mar-2019 19:57	
<i>Surr: Toluene-d8</i>	<i>107</i>			0	<i>89-112</i>	%REC	50	07-Mar-2019 15:56	
<i>Surr: Toluene-d8</i>	<i>107</i>			0	<i>89-112</i>	%REC	250	07-Mar-2019 19:57	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB	
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW06_022619
 Collection Date: 26-Feb-2019 09:10

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-05
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,1-Dichloroethene	0.95	J	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2-Dichloroethane	0.56	J	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 20:45	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	06-Mar-2019 20:45	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	06-Mar-2019 20:45	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	06-Mar-2019 20:45	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	06-Mar-2019 20:45	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW06_022619
 Collection Date: 26-Feb-2019 09:10

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-05
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
cis-1,2-Dichloroethene	180		0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	06-Mar-2019 20:45	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	06-Mar-2019 20:45	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	06-Mar-2019 20:45	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
trans-1,2-Dichloroethene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Trichloroethene	250		1.0	2.5	5.0	UG/L	5	07-Mar-2019 17:57	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
Vinyl chloride	0.50	U	0.20	0.50	1.0	UG/L	1	06-Mar-2019 20:45	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>82.9</i>			0	<i>81-118</i>	%REC	1	06-Mar-2019 20:45	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>82.1</i>			0	<i>81-118</i>	%REC	5	07-Mar-2019 17:57	
<i>Surr: 4-Bromofluorobenzene</i>	<i>99.0</i>			0	<i>85-114</i>	%REC	1	06-Mar-2019 20:45	
<i>Surr: 4-Bromofluorobenzene</i>	<i>100.0</i>			0	<i>85-114</i>	%REC	5	07-Mar-2019 17:57	
<i>Surr: Dibromofluoromethane</i>	<i>84.6</i>			0	<i>80-119</i>	%REC	1	06-Mar-2019 20:45	
<i>Surr: Dibromofluoromethane</i>	<i>82.9</i>			0	<i>80-119</i>	%REC	5	07-Mar-2019 17:57	
<i>Surr: Toluene-d8</i>	<i>106</i>			0	<i>89-112</i>	%REC	1	06-Mar-2019 20:45	
<i>Surr: Toluene-d8</i>	<i>107</i>			0	<i>89-112</i>	%REC	5	07-Mar-2019 17:57	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB	
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW03_022619
 Collection Date: 26-Feb-2019 09:20

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-06
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,1,1-Trichloroethane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,1,2,2-Tetrachloroethane	5.0	U	5.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,1,2-Trichloroethane	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,1-Dichloroethane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,1-Dichloroethene	12		2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,1-Dichloropropene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2,3-Trichlorobenzene	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2,3-Trichloropropane	5.0	U	5.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2,4-Trichlorobenzene	5.0	U	5.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2,4-Trimethylbenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2-Dibromo-3-chloropropane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2-Dibromoethane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2-Dichlorobenzene	5.0	U	5.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2-Dichloroethane	6.2	J	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,2-Dichloropropane	5.0	U	5.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,3,5-Trimethylbenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,3-Dichlorobenzene	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,3-Dichloropropane	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
1,4-Dichlorobenzene	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
2,2-Dichloropropane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
2-Butanone	10	U	5.0	10	20	UG/L	10	07-Mar-2019 15:08	
2-Chlorotoluene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
2-Hexanone	10	U	10	10	20	UG/L	10	07-Mar-2019 15:08	
4-Chlorotoluene	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
4-Isopropyltoluene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
4-Methyl-2-pentanone	10	U	7.0	10	20	UG/L	10	07-Mar-2019 15:08	
Acetone	20	U	4.0	20	20	UG/L	10	07-Mar-2019 15:08	
Benzene	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Bromobenzene	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Bromochloromethane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Bromodichloromethane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Bromoform	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Bromomethane	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Carbon disulfide	10	U	6.0	10	20	UG/L	10	07-Mar-2019 15:08	
Carbon tetrachloride	5.0	U	5.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Chlorobenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Chloroethane	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Chloroform	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW03_022619
 Collection Date: 26-Feb-2019 09:20

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-06
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
cis-1,2-Dichloroethene	2,000		20	50	100	UG/L	100	07-Mar-2019 19:09	
cis-1,3-Dichloropropene	5.0	U	1.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Dibromochloromethane	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Dibromomethane	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Dichlorodifluoromethane	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Ethylbenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Hexachlorobutadiene	10	U	10	10	10	UG/L	10	07-Mar-2019 15:08	
Isopropylbenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
m,p-Xylene	10	U	5.0	10	20	UG/L	10	07-Mar-2019 15:08	
Methylene chloride	5.0	U	4.0	5.0	20	UG/L	10	07-Mar-2019 15:08	
n-Butylbenzene	5.0	U	4.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
n-Propylbenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Naphthalene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
o-Xylene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
sec-Butylbenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Styrene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
tert-Butylbenzene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Tetrachloroethene	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Toluene	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
trans-1,2-Dichloroethene	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
trans-1,3-Dichloropropene	5.0	U	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Trichloroethene	7,400		20	50	100	UG/L	100	07-Mar-2019 19:09	
Trichlorofluoromethane	5.0	U	3.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Vinyl chloride	7.5	J	2.0	5.0	10	UG/L	10	07-Mar-2019 15:08	
Surr: 1,2-Dichloroethane-d4	85.6			0	81-118	%REC	10	07-Mar-2019 15:08	
Surr: 1,2-Dichloroethane-d4	82.1			0	81-118	%REC	100	07-Mar-2019 19:09	
Surr: 4-Bromofluorobenzene	100			0	85-114	%REC	100	07-Mar-2019 19:09	
Surr: 4-Bromofluorobenzene	101			0	85-114	%REC	10	07-Mar-2019 15:08	
Surr: Dibromofluoromethane	84.3			0	80-119	%REC	10	07-Mar-2019 15:08	
Surr: Dibromofluoromethane	84.1			0	80-119	%REC	100	07-Mar-2019 19:09	
Surr: Toluene-d8	106			0	89-112	%REC	10	07-Mar-2019 15:08	
Surr: Toluene-d8	108			0	89-112	%REC	100	07-Mar-2019 19:09	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB	
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW07_022619
 Collection Date: 26-Feb-2019 09:30

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-07
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,1-Dichloroethene	1.7		0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	07-Mar-2019 13:32	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	07-Mar-2019 13:32	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	07-Mar-2019 13:32	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	07-Mar-2019 13:32	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	07-Mar-2019 13:32	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW07_022619
 Collection Date: 26-Feb-2019 09:30

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-07
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
cis-1,2-Dichloroethene	340		1.0	2.5	5.0	UG/L	5	07-Mar-2019 18:21	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	07-Mar-2019 13:32	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	07-Mar-2019 13:32	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	07-Mar-2019 13:32	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
trans-1,2-Dichloroethene	0.84	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Trichloroethene	81		0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Vinyl chloride	25		0.20	0.50	1.0	UG/L	1	07-Mar-2019 13:32	
Surr: 1,2-Dichloroethane-d4	82.3			0	81-118	%REC	1	07-Mar-2019 13:32	
Surr: 1,2-Dichloroethane-d4	82.7			0	81-118	%REC	5	07-Mar-2019 18:21	
Surr: 4-Bromofluorobenzene	99.3			0	85-114	%REC	1	07-Mar-2019 13:32	
Surr: 4-Bromofluorobenzene	101			0	85-114	%REC	5	07-Mar-2019 18:21	
Surr: Dibromofluoromethane	82.6			0	80-119	%REC	5	07-Mar-2019 18:21	
Surr: Dibromofluoromethane	84.3			0	80-119	%REC	1	07-Mar-2019 13:32	
Surr: Toluene-d8	107			0	89-112	%REC	1	07-Mar-2019 13:32	
Surr: Toluene-d8	107			0	89-112	%REC	5	07-Mar-2019 18:21	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA							Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW04_022619
 Collection Date: 26-Feb-2019 09:40

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-08
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,1-Dichloroethene	0.88	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2-Dichloroethane	0.55	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	07-Mar-2019 14:20	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	07-Mar-2019 14:20	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	07-Mar-2019 14:20	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	07-Mar-2019 14:20	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	07-Mar-2019 14:20	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW04_022619
 Collection Date: 26-Feb-2019 09:40

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-08
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
cis-1,2-Dichloroethene	260		5.0	12	25	UG/L	25	07-Mar-2019 18:45	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	07-Mar-2019 14:20	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	07-Mar-2019 14:20	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	07-Mar-2019 14:20	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
trans-1,2-Dichloroethene	0.72	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Trichloroethene	900		5.0	12	25	UG/L	25	07-Mar-2019 18:45	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
Vinyl chloride	0.47	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 14:20	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>81.3</i>			0	<i>81-118</i>	%REC	1	07-Mar-2019 14:20	
<i>Surr: 1,2-Dichloroethane-d4</i>	<i>81.9</i>			0	<i>81-118</i>	%REC	25	07-Mar-2019 18:45	
<i>Surr: 4-Bromofluorobenzene</i>	<i>99.5</i>			0	<i>85-114</i>	%REC	25	07-Mar-2019 18:45	
<i>Surr: 4-Bromofluorobenzene</i>	<i>100</i>			0	<i>85-114</i>	%REC	1	07-Mar-2019 14:20	
<i>Surr: Dibromofluoromethane</i>	<i>82.6</i>			0	<i>80-119</i>	%REC	1	07-Mar-2019 14:20	
<i>Surr: Dibromofluoromethane</i>	<i>82.9</i>			0	<i>80-119</i>	%REC	25	07-Mar-2019 18:45	
<i>Surr: Toluene-d8</i>	<i>108</i>			0	<i>89-112</i>	%REC	25	07-Mar-2019 18:45	
<i>Surr: Toluene-d8</i>	<i>106</i>			0	<i>89-112</i>	%REC	1	07-Mar-2019 14:20	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA							Analyst: SUB
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW08_022619
 Collection Date: 26-Feb-2019 09:50

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-09
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260							Analyst: PC
1,1,1,2-Tetrachloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,1,1-Trichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,1,2,2-Tetrachloroethane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,1,2-Trichloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,1-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,1-Dichloroethene	0.76	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,1-Dichloropropene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2,3-Trichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2,3-Trichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2,4-Trichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2,4-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2-Dibromo-3-chloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2-Dibromoethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2-Dichlorobenzene	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2-Dichloroethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,2-Dichloropropane	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,3,5-Trimethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,3-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,3-Dichloropropane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
1,4-Dichlorobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
2,2-Dichloropropane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
2-Butanone	1.0	U	0.50	1.0	2.0	UG/L	1	07-Mar-2019 12:44	
2-Chlorotoluene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
2-Hexanone	1.0	U	1.0	1.0	2.0	UG/L	1	07-Mar-2019 12:44	
4-Chlorotoluene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
4-Isopropyltoluene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
4-Methyl-2-pentanone	1.0	U	0.70	1.0	2.0	UG/L	1	07-Mar-2019 12:44	
Acetone	2.0	U	0.40	2.0	2.0	UG/L	1	07-Mar-2019 12:44	
Benzene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Bromobenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Bromochloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Bromodichloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Bromoform	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Bromomethane	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Carbon disulfide	1.0	U	0.60	1.0	2.0	UG/L	1	07-Mar-2019 12:44	
Carbon tetrachloride	0.50	U	0.50	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Chlorobenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Chloroethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Chloroform	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP Site 16
 Sample ID: 16EW08_022619
 Collection Date: 26-Feb-2019 09:50

ANALYTICAL REPORT
 WorkOrder:HS19021428
 Lab ID:HS19021428-09
 Matrix:Water

ANALYSES	RESULT	QUAL	DL	LOD	LOQ	UNITS	DILUTION FACTOR	DATE ANALYZED	
VOLATILES ORGANICS BY METHOD 8260C		Method:SW8260						Analyst: PC	
Chloromethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
cis-1,2-Dichloroethene	170		1.0	2.5	5.0	UG/L	5	07-Mar-2019 16:44	
cis-1,3-Dichloropropene	0.50	U	0.10	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Dibromochloromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Dibromomethane	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Dichlorodifluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Ethylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Hexachlorobutadiene	1.0	U	1.0	1.0	1.0	UG/L	1	07-Mar-2019 12:44	
Isopropylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
m,p-Xylene	1.0	U	0.50	1.0	2.0	UG/L	1	07-Mar-2019 12:44	
Methylene chloride	0.50	U	0.40	0.50	2.0	UG/L	1	07-Mar-2019 12:44	
n-Butylbenzene	0.50	U	0.40	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
n-Propylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Naphthalene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
o-Xylene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
sec-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Styrene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
tert-Butylbenzene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Tetrachloroethene	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Toluene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
trans-1,2-Dichloroethene	0.64	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
trans-1,3-Dichloropropene	0.50	U	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Trichloroethene	170		1.0	2.5	5.0	UG/L	5	07-Mar-2019 16:44	
Trichlorofluoromethane	0.50	U	0.30	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Vinyl chloride	0.86	J	0.20	0.50	1.0	UG/L	1	07-Mar-2019 12:44	
Surr: 1,2-Dichloroethane-d4	82.8			0	81-118	%REC	1	07-Mar-2019 12:44	
Surr: 1,2-Dichloroethane-d4	81.7			0	81-118	%REC	5	07-Mar-2019 16:44	
Surr: 4-Bromofluorobenzene	100			0	85-114	%REC	1	07-Mar-2019 12:44	
Surr: 4-Bromofluorobenzene	97.1			0	85-114	%REC	5	07-Mar-2019 16:44	
Surr: Dibromofluoromethane	83.3			0	80-119	%REC	5	07-Mar-2019 16:44	
Surr: Dibromofluoromethane	84.2			0	80-119	%REC	1	07-Mar-2019 12:44	
Surr: Toluene-d8	109			0	89-112	%REC	5	07-Mar-2019 16:44	
Surr: Toluene-d8	108			0	89-112	%REC	1	07-Mar-2019 12:44	
SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)		Method:NA						Analyst: SUB	
Subcontract Analysis	See Attached		0	0		NA	1	07-Mar-2019 17:47	

Note: See Qualifiers Page for a list of qualifiers and their explanation.

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

DATES REPORT

Sample ID	Client Samp ID	Collection Date	TCLP Date	Prep Date	Analysis Date	DF
Batch ID R334128	Test Name : VOLATILES ORGANICS BY METHOD 8260C			Matrix: Water		
HS19021428-01	16EW01_022619	26 Feb 2019 08:30			06 Mar 2019 13:27	5
HS19021428-01	16EW01_022619	26 Feb 2019 08:30			06 Mar 2019 13:03	50
HS19021428-02	16EW05_022619	26 Feb 2019 08:50			06 Mar 2019 16:43	25
HS19021428-02	16EW05_022619	26 Feb 2019 08:50			06 Mar 2019 15:52	1
HS19021428-03	16EW05_022619_a	26 Feb 2019 08:50			06 Mar 2019 18:21	25
HS19021428-03	16EW05_022619_a	26 Feb 2019 08:50			06 Mar 2019 17:31	1
HS19021428-05	16EW06_022619	26 Feb 2019 09:10			06 Mar 2019 20:45	1
Batch ID R334176	Test Name : SUBCONTRACT ANALYSIS - PERCHLORATE (EPA 6850)			Matrix: Water		
HS19021428-01	16EW01_022619	26 Feb 2019 08:30			07 Mar 2019 17:47	1
HS19021428-02	16EW05_022619	26 Feb 2019 08:50			07 Mar 2019 17:47	1
HS19021428-03	16EW05_022619_a	26 Feb 2019 08:50			07 Mar 2019 17:47	1
HS19021428-04	16EW02_022619	26 Feb 2019 09:00			07 Mar 2019 17:47	1
HS19021428-05	16EW06_022619	26 Feb 2019 09:10			07 Mar 2019 17:47	1
HS19021428-06	16EW03_022619	26 Feb 2019 09:20			07 Mar 2019 17:47	1
HS19021428-07	16EW07_022619	26 Feb 2019 09:30			07 Mar 2019 17:47	1
HS19021428-08	16EW04_022619	26 Feb 2019 09:40			07 Mar 2019 17:47	1
HS19021428-09	16EW08_022619	26 Feb 2019 09:50			07 Mar 2019 17:47	1
Batch ID R334221	Test Name : VOLATILES ORGANICS BY METHOD 8260C			Matrix: Water		
HS19021428-04	16EW02_022619	26 Feb 2019 09:00			07 Mar 2019 19:57	250
HS19021428-04	16EW02_022619	26 Feb 2019 09:00			07 Mar 2019 15:56	50
HS19021428-05	16EW06_022619	26 Feb 2019 09:10			07 Mar 2019 17:57	5
HS19021428-06	16EW03_022619	26 Feb 2019 09:20			07 Mar 2019 19:09	100
HS19021428-06	16EW03_022619	26 Feb 2019 09:20			07 Mar 2019 15:08	10
HS19021428-07	16EW07_022619	26 Feb 2019 09:30			07 Mar 2019 18:21	5
HS19021428-07	16EW07_022619	26 Feb 2019 09:30			07 Mar 2019 13:32	1
HS19021428-08	16EW04_022619	26 Feb 2019 09:40			07 Mar 2019 18:45	25
HS19021428-08	16EW04_022619	26 Feb 2019 09:40			07 Mar 2019 14:20	1
HS19021428-09	16EW08_022619	26 Feb 2019 09:50			07 Mar 2019 16:44	5
HS19021428-09	16EW08_022619	26 Feb 2019 09:50			07 Mar 2019 12:44	1

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334128		Instrument: VOA6		Method: SW8260						
MBLK	Sample ID: VBLKW-190306	Units: UG/L			Analysis Date: 06-Mar-2019 11:51					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977685		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	41.51	1.0	50	0	83.0	81 - 118				
Surr: 4-Bromofluorobenzene	49.32	1.0	50	0	98.6	85 - 114				
Surr: Dibromofluoromethane	43.47	1.0	50	0	86.9	80 - 119				
Surr: Toluene-d8	53.24	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334128		Instrument: VOA6		Method: SW8260						
LCS	Sample ID: VLCSW-190306	Units: UG/L			Analysis Date: 06-Mar-2019 11:03					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977684	PrepDate:	DF: 1						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	20.28	1.0	20	0	101	78 - 124				
1,1,1-Trichloroethane	20.23	1.0	20	0	101	74 - 131				
1,1,2,2-Tetrachloroethane	19.2	1.0	20	0	96.0	71 - 121				
1,1,2-Trichloroethane	19.93	1.0	20	0	99.6	80 - 119				
1,1-Dichloroethane	19.67	1.0	20	0	98.3	77 - 125				
1,1-Dichloroethene	19.73	1.0	20	0	98.6	71 - 131				
1,1-Dichloropropene	19.78	1.0	20	0	98.9	78 - 125				
1,2,3-Trichlorobenzene	20.06	1.0	20	0	100	69 - 129				
1,2,3-Trichloropropane	18.96	1.0	20	0	94.8	73 - 122				
1,2,4-Trichlorobenzene	19.89	1.0	20	0	99.4	69 - 130				
1,2,4-Trimethylbenzene	19.37	1.0	20	0	96.8	76 - 124				
1,2-Dibromo-3-chloropropane	18.77	1.0	20	0	93.9	62 - 128				
1,2-Dibromoethane	20.52	1.0	20	0	103	77 - 121				
1,2-Dichlorobenzene	19.02	1.0	20	0	95.1	80 - 119				
1,2-Dichloroethane	20.54	1.0	20	0	103	73 - 128				
1,2-Dichloropropane	20.31	1.0	20	0	102	78 - 122				
1,3,5-Trimethylbenzene	19.43	1.0	20	0	97.1	75 - 124				
1,3-Dichlorobenzene	19.44	1.0	20	0	97.2	80 - 119				
1,3-Dichloropropane	20.12	1.0	20	0	101	80 - 119				
1,4-Dichlorobenzene	19.16	1.0	20	0	95.8	79 - 118				
2,2-Dichloropropane	19.96	1.0	20	0	99.8	60 - 139				
2-Butanone	42.82	2.0	40	0	107	56 - 143				
2-Chlorotoluene	19.13	1.0	20	0	95.7	79 - 122				
2-Hexanone	40.07	2.0	40	0	100	57 - 139				
4-Chlorotoluene	19.38	1.0	20	0	96.9	78 - 122				
4-Isopropyltoluene	19.42	1.0	20	0	97.1	77 - 127				
4-Methyl-2-pentanone	39.73	2.0	40	0	99.3	67 - 130				
Acetone	42.44	2.0	40	0	106	39 - 160				
Benzene	19.99	1.0	20	0	99.9	79 - 120				
Bromobenzene	19.33	1.0	20	0	96.6	80 - 120				
Bromochloromethane	19.66	1.0	20	0	98.3	78 - 123				
Bromodichloromethane	19.92	1.0	20	0	99.6	79 - 125				
Bromoform	20.37	1.0	20	0	102	66 - 130				
Bromomethane	24.28	1.0	20	0	121	53 - 141				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334128		Instrument: VOA6		Method: SW8260						
LCS	Sample ID: VLCSW-190306	Units: UG/L			Analysis Date: 06-Mar-2019 11:03					
Client ID:	Run ID: VOA6_334128	SeqNo: 4977684		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	40.1	2.0	40	0	100	64 - 133				
Carbon tetrachloride	20.59	1.0	20	0	103	72 - 136				
Chlorobenzene	19.9	1.0	20	0	99.5	82 - 118				
Chloroethane	19.71	1.0	20	0	98.5	60 - 138				
Chloroform	19.55	1.0	20	0	97.7	79 - 124				
Chloromethane	20.54	1.0	20	0	103	50 - 139				
cis-1,2-Dichloroethene	19.71	1.0	20	0	98.6	78 - 123				
cis-1,3-Dichloropropene	20.44	1.0	20	0	102	75 - 124				
Dibromochloromethane	20.25	1.0	20	0	101	74 - 126				
Dibromomethane	20.09	1.0	20	0	100	79 - 123				
Dichlorodifluoromethane	20.24	1.0	20	0	101	32 - 152				
Ethylbenzene	20.01	1.0	20	0	100	79 - 121				
Hexachlorobutadiene	21.69	1.0	20	0	108	66 - 134				
Isopropylbenzene	19.97	1.0	20	0	99.8	72 - 131				
m,p-Xylene	40.08	2.0	40	0	100	80 - 121				
Methylene chloride	20.41	2.0	20	0	102	74 - 124				
Naphthalene	18.14	1.0	20	0	90.7	61 - 128				
n-Butylbenzene	19.01	1.0	20	0	95.0	75 - 128				
n-Propylbenzene	19.22	1.0	20	0	96.1	76 - 126				
o-Xylene	20.19	1.0	20	0	101	78 - 122				
sec-Butylbenzene	19	1.0	20	0	95.0	77 - 126				
Styrene	20.48	1.0	20	0	102	78 - 123				
tert-Butylbenzene	18.99	1.0	20	0	95.0	78 - 124				
Tetrachloroethene	19.96	1.0	20	0	99.8	74 - 129				
Toluene	19.89	1.0	20	0	99.5	80 - 121				
trans-1,2-Dichloroethene	20.32	1.0	20	0	102	75 - 124				
trans-1,3-Dichloropropene	20.39	1.0	20	0	102	73 - 127				
Trichloroethene	20.14	1.0	20	0	101	79 - 123				
Trichlorofluoromethane	19.72	1.0	20	0	98.6	65 - 141				
Vinyl chloride	19.12	1.0	20	0	95.6	58 - 137				
Surr: 1,2-Dichloroethane-d4	50.81	1.0	50	0	102	81 - 118				
Surr: 4-Bromofluorobenzene	51.81	1.0	50	0	104	85 - 114				
Surr: Dibromofluoromethane	51.53	1.0	50	0	103	80 - 119				
Surr: Toluene-d8	48.19	1.0	50	0	96.4	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334128		Instrument: VOA6		Method: SW8260						
MS	Sample ID: HS19021428-01MS	Units: UG/L			Analysis Date: 06-Mar-2019 14:39					
Client ID: 16EW01_022619	Run ID: VOA6_334128	SeqNo: 4977690	PrepDate:	DF: 50						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	1131	50	1000	0	113	78 - 124				
1,1,1-Trichloroethane	987.3	50	1000	0	98.7	74 - 131				
1,1,2,2-Tetrachloroethane	1076	50	1000	0	108	71 - 121				
1,1,2-Trichloroethane	1111	50	1000	0	111	80 - 119				
1,1-Dichloroethane	940.2	50	1000	0	94.0	77 - 125				
1,1-Dichloroethene	1011	50	1000	0	101	71 - 131				
1,1-Dichloropropene	1112	50	1000	0	111	78 - 125				
1,2,3-Trichlorobenzene	1082	50	1000	0	108	69 - 129				
1,2,3-Trichloropropane	1033	50	1000	0	103	73 - 122				
1,2,4-Trichlorobenzene	1136	50	1000	0	114	69 - 130				
1,2,4-Trimethylbenzene	1203	50	1000	0	120	76 - 124				
1,2-Dibromo-3-chloropropane	1044	50	1000	0	104	62 - 128				
1,2-Dibromoethane	1119	50	1000	0	112	77 - 121				
1,2-Dichlorobenzene	1128	50	1000	0	113	80 - 119				
1,2-Dichloroethane	1026	50	1000	0	103	73 - 128				
1,2-Dichloropropane	1034	50	1000	0	103	78 - 122				
1,3,5-Trimethylbenzene	1243	50	1000	0	124	75 - 124				S
1,3-Dichlorobenzene	1180	50	1000	0	118	80 - 119				
1,3-Dichloropropane	1110	50	1000	0	111	80 - 119				
1,4-Dichlorobenzene	1153	50	1000	0	115	79 - 118				
2,2-Dichloropropane	1001	50	1000	0	100	60 - 139				
2-Butanone	1842	100	2000	0	92.1	56 - 143				
2-Chlorotoluene	1181	50	1000	0	118	79 - 122				
2-Hexanone	2034	100	2000	0	102	57 - 139				
4-Chlorotoluene	1190	50	1000	0	119	78 - 122				
4-Isopropyltoluene	1296	50	1000	0	130	77 - 127				S
4-Methyl-2-pentanone	2036	100	2000	0	102	67 - 130				
Acetone	1745	100	2000	0	87.3	39 - 160				
Benzene	1045	50	1000	0	104	79 - 120				
Bromobenzene	1145	50	1000	0	114	80 - 120				
Bromochloromethane	903.7	50	1000	0	90.4	78 - 123				
Bromodichloromethane	1027	50	1000	0	103	79 - 125				
Bromoform	1094	50	1000	0	109	66 - 130				
Bromomethane	1198	50	1000	0	120	53 - 141				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334128		Instrument: VOA6		Method: SW8260						
MS		Sample ID: HS19021428-01MS		Units: UG/L		Analysis Date: 06-Mar-2019 14:39				
Client ID: 16EW01_022619		Run ID: VOA6_334128		SeqNo: 4977690		PrepDate:		DF: 50		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1925	100	2000	0	96.2	64 - 133				
Carbon tetrachloride	1169	50	1000	0	117	72 - 136				
Chlorobenzene	1159	50	1000	0	116	82 - 118				
Chloroethane	922.9	50	1000	0	92.3	60 - 138				
Chloroform	938.1	50	1000	0	93.8	79 - 124				
Chloromethane	804.4	50	1000	0	80.4	50 - 139				
cis-1,2-Dichloroethene	4602	50	1000	3749	85.3	78 - 123				
cis-1,3-Dichloropropene	1038	50	1000	0	104	75 - 124				
Dibromochloromethane	1148	50	1000	0	115	74 - 126				
Dibromomethane	1016	50	1000	0	102	79 - 123				
Dichlorodifluoromethane	668.3	50	1000	0	66.8	32 - 152				
Ethylbenzene	1202	50	1000	0	120	79 - 121				
Hexachlorobutadiene	1396	50	1000	0	140	66 - 134				S
Isopropylbenzene	1253	50	1000	0	125	72 - 131				
m,p-Xylene	2393	100	2000	0	120	80 - 121				
Methylene chloride	955.4	100	1000	0	95.5	74 - 124				
Naphthalene	970.7	50	1000	0	97.1	61 - 128				
n-Butylbenzene	1323	50	1000	0	132	75 - 128				S
n-Propylbenzene	1256	50	1000	0	126	76 - 126				
o-Xylene	1200	50	1000	0	120	78 - 122				
sec-Butylbenzene	1306	50	1000	0	131	77 - 126				S
Styrene	1164	50	1000	0	116	78 - 123				
tert-Butylbenzene	1270	50	1000	0	127	78 - 124				S
Tetrachloroethene	1293	50	1000	0	129	74 - 129				S
Toluene	1184	50	1000	0	118	80 - 121				
trans-1,2-Dichloroethene	997.2	50	1000	0	99.7	75 - 124				
trans-1,3-Dichloropropene	1039	50	1000	0	104	73 - 127				
Trichloroethene	4573	50	1000	3686	88.7	79 - 123				
Trichlorofluoromethane	1014	50	1000	0	101	65 - 141				
Vinyl chloride	865.2	50	1000	0	86.5	58 - 137				
Surr: 1,2-Dichloroethane-d4	2074	50	2500	0	83.0	81 - 118				
Surr: 4-Bromofluorobenzene	2472	50	2500	0	98.9	85 - 114				
Surr: Dibromofluoromethane	2130	50	2500	0	85.2	80 - 119				
Surr: Toluene-d8	2648	50	2500	0	106	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334128		Instrument: VOA6		Method: SW8260						
MSD		Sample ID: HS19021428-01MSD		Units: UG/L		Analysis Date: 06-Mar-2019 15:03				
Client ID: 16EW01_022619		Run ID: VOA6_334128		SeqNo: 4977691		PrepDate:		DF: 50		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	1074	50	1000	0	107	78 - 124	1131	5.23	20	
1,1,1-Trichloroethane	1004	50	1000	0	100	74 - 131	987.3	1.64	20	
1,1,2,2-Tetrachloroethane	977.6	50	1000	0	97.8	71 - 121	1076	9.59	20	
1,1,2-Trichloroethane	1026	50	1000	0	103	80 - 119	1111	7.97	20	
1,1-Dichloroethane	918.7	50	1000	0	91.9	77 - 125	940.2	2.32	20	
1,1-Dichloroethene	1003	50	1000	0	100	71 - 131	1011	0.851	20	
1,1-Dichloropropene	1155	50	1000	0	116	78 - 125	1112	3.85	20	
1,2,3-Trichlorobenzene	1116	50	1000	0	112	69 - 129	1082	3.06	20	
1,2,3-Trichloropropane	947	50	1000	0	94.7	73 - 122	1033	8.7	20	
1,2,4-Trichlorobenzene	1120	50	1000	0	112	69 - 130	1136	1.43	20	
1,2,4-Trimethylbenzene	1139	50	1000	0	114	76 - 124	1203	5.47	20	
1,2-Dibromo-3-chloropropane	993.9	50	1000	0	99.4	62 - 128	1044	4.94	20	
1,2-Dibromoethane	1034	50	1000	0	103	77 - 121	1119	7.94	20	
1,2-Dichlorobenzene	1051	50	1000	0	105	80 - 119	1128	7.08	20	
1,2-Dichloroethane	1012	50	1000	0	101	73 - 128	1026	1.39	20	
1,2-Dichloropropane	1017	50	1000	0	102	78 - 122	1034	1.63	20	
1,3,5-Trimethylbenzene	1168	50	1000	0	117	75 - 124	1243	6.24	20	
1,3-Dichlorobenzene	1093	50	1000	0	109	80 - 119	1180	7.6	20	
1,3-Dichloropropane	1025	50	1000	0	102	80 - 119	1110	7.93	20	
1,4-Dichlorobenzene	1063	50	1000	0	106	79 - 118	1153	8.16	20	
2,2-Dichloropropane	982.5	50	1000	0	98.2	60 - 139	1001	1.83	20	
2-Butanone	1713	100	2000	0	85.7	56 - 143	1842	7.25	20	
2-Chlorotoluene	1102	50	1000	0	110	79 - 122	1181	6.9	20	
2-Hexanone	1939	100	2000	0	96.9	57 - 139	2034	4.78	20	
4-Chlorotoluene	1105	50	1000	0	110	78 - 122	1190	7.4	20	
4-Isopropyltoluene	1246	50	1000	0	125	77 - 127	1296	3.91	20	
4-Methyl-2-pentanone	1953	100	2000	0	97.6	67 - 130	2036	4.18	20	
Acetone	1701	100	2000	0	85.0	39 - 160	1745	2.58	20	
Benzene	1032	50	1000	0	103	79 - 120	1045	1.18	20	
Bromobenzene	1050	50	1000	0	105	80 - 120	1145	8.62	20	
Bromochloromethane	886.4	50	1000	0	88.6	78 - 123	903.7	1.92	20	
Bromodichloromethane	1012	50	1000	0	101	79 - 125	1027	1.47	20	
Bromoform	1026	50	1000	0	103	66 - 130	1094	6.41	20	
Bromomethane	1111	50	1000	0	111	53 - 141	1198	7.53	20	

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334128		Instrument: VOA6		Method: SW8260						
MSD	Sample ID: HS19021428-01MSD	Units: UG/L			Analysis Date: 06-Mar-2019 15:03					
Client ID: 16EW01_022619	Run ID: VOA6_334128	SeqNo: 4977691	PrepDate:	DF: 50						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1914	100	2000	0	95.7	64 - 133	1925	0.566	20	
Carbon tetrachloride	1209	50	1000	0	121	72 - 136	1169	3.36	20	
Chlorobenzene	1092	50	1000	0	109	82 - 118	1159	5.98	20	
Chloroethane	929.6	50	1000	0	93.0	60 - 138	922.9	0.728	20	
Chloroform	908	50	1000	0	90.8	79 - 124	938.1	3.27	20	
Chloromethane	783	50	1000	0	78.3	50 - 139	804.4	2.7	20	
cis-1,2-Dichloroethene	4457	50	1000	3749	70.8	78 - 123	4602	3.18	20	S
cis-1,3-Dichloropropene	1021	50	1000	0	102	75 - 124	1038	1.73	20	
Dibromochloromethane	1059	50	1000	0	106	74 - 126	1148	8.06	20	
Dibromomethane	990.7	50	1000	0	99.1	79 - 123	1016	2.54	20	
Dichlorodifluoromethane	688.8	50	1000	0	68.9	32 - 152	668.3	3.02	20	
Ethylbenzene	1160	50	1000	0	116	79 - 121	1202	3.53	20	
Hexachlorobutadiene	1434	50	1000	0	143	66 - 134	1396	2.69	20	S
Isopropylbenzene	1231	50	1000	0	123	72 - 131	1253	1.82	20	
m,p-Xylene	2294	100	2000	0	115	80 - 121	2393	4.21	20	
Methylene chloride	914.4	100	1000	0	91.4	74 - 124	955.4	4.38	20	
Naphthalene	939.7	50	1000	0	94.0	61 - 128	970.7	3.25	20	
n-Butylbenzene	1292	50	1000	0	129	75 - 128	1323	2.38	20	S
n-Propylbenzene	1191	50	1000	0	119	76 - 126	1256	5.27	20	
o-Xylene	1141	50	1000	0	114	78 - 122	1200	5.07	20	
sec-Butylbenzene	1272	50	1000	0	127	77 - 126	1306	2.68	20	S
Styrene	1098	50	1000	0	110	78 - 123	1164	5.82	20	
tert-Butylbenzene	1210	50	1000	0	121	78 - 124	1270	4.84	20	
Tetrachloroethene	1247	50	1000	0	125	74 - 129	1293	3.64	20	
Toluene	1121	50	1000	0	112	80 - 121	1184	5.5	20	
trans-1,2-Dichloroethene	971.6	50	1000	0	97.2	75 - 124	997.2	2.59	20	
trans-1,3-Dichloropropene	1012	50	1000	0	101	73 - 127	1039	2.6	20	
Trichloroethene	4546	50	1000	3686	86.1	79 - 123	4573	0.581	20	
Trichlorofluoromethane	1013	50	1000	0	101	65 - 141	1014	0.0243	20	
Vinyl chloride	860.2	50	1000	0	86.0	58 - 137	865.2	0.586	20	
Surr: 1,2-Dichloroethane-d4	2035	50	2500	0	81.4	81 - 118	2074	1.92	20	
Surr: 4-Bromofluorobenzene	2550	50	2500	0	102	85 - 114	2472	3.09	20	
Surr: Dibromofluoromethane	2117	50	2500	0	84.7	80 - 119	2130	0.579	20	
Surr: Toluene-d8	2586	50	2500	0	103	89 - 112	2648	2.39	20	

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT**Batch ID:** R334128**Instrument:** VOA6**Method:** SW8260

The following samples were analyzed in this batch:

HS19021428-01	HS19021428-02	HS19021428-03	HS19021428-05
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ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221		Instrument: VOA6		Method: SW8260						
MBLK	Sample ID: VBLKW-190307	Units: UG/L			Analysis Date: 07-Mar-2019 12:20					
Client ID:	Run ID: VOA6_334221	SeqNo: 4979743		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	1.0	2.0								U
Carbon tetrachloride	0.50	1.0								U
Chlorobenzene	0.50	1.0								U
Chloroethane	0.50	1.0								U
Chloroform	0.50	1.0								U
Chloromethane	0.50	1.0								U
cis-1,2-Dichloroethene	0.50	1.0								U
cis-1,3-Dichloropropene	0.50	1.0								U
Dibromochloromethane	0.50	1.0								U
Dibromomethane	0.50	1.0								U
Dichlorodifluoromethane	0.50	1.0								U
Ethylbenzene	0.50	1.0								U
Hexachlorobutadiene	1.0	1.0								U
Isopropylbenzene	0.50	1.0								U
m,p-Xylene	1.0	2.0								U
Methylene chloride	0.50	2.0								U
Naphthalene	0.50	1.0								U
n-Butylbenzene	0.50	1.0								U
n-Propylbenzene	0.50	1.0								U
o-Xylene	0.50	1.0								U
sec-Butylbenzene	0.50	1.0								U
Styrene	0.50	1.0								U
tert-Butylbenzene	0.50	1.0								U
Tetrachloroethene	0.50	1.0								U
Toluene	0.50	1.0								U
trans-1,2-Dichloroethene	0.50	1.0								U
trans-1,3-Dichloropropene	0.50	1.0								U
Trichloroethene	0.50	1.0								U
Trichlorofluoromethane	0.50	1.0								U
Vinyl chloride	0.50	1.0								U
Surr: 1,2-Dichloroethane-d4	41.41	1.0	50	0	82.8	81 - 118				
Surr: 4-Bromofluorobenzene	49.94	1.0	50	0	99.9	85 - 114				
Surr: Dibromofluoromethane	41.97	1.0	50	0	83.9	80 - 119				
Surr: Toluene-d8	53.12	1.0	50	0	106	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221		Instrument: VOA6			Method: SW8260					
LCS	Sample ID: VLCSW-190307	Units: UG/L			Analysis Date: 07-Mar-2019 11:32					
Client ID:	Run ID: VOA6_334221	SeqNo: 4979742		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	20.51	1.0	20	0	103	78 - 124				
1,1,1-Trichloroethane	20.24	1.0	20	0	101	74 - 131				
1,1,2,2-Tetrachloroethane	19.6	1.0	20	0	98.0	71 - 121				
1,1,2-Trichloroethane	20.01	1.0	20	0	100	80 - 119				
1,1-Dichloroethane	20.03	1.0	20	0	100	77 - 125				
1,1-Dichloroethene	20.46	1.0	20	0	102	71 - 131				
1,1-Dichloropropene	19.22	1.0	20	0	96.1	78 - 125				
1,2,3-Trichlorobenzene	22.11	1.0	20	0	111	69 - 129				
1,2,3-Trichloropropane	19.17	1.0	20	0	95.8	73 - 122				
1,2,4-Trichlorobenzene	21.49	1.0	20	0	107	69 - 130				
1,2,4-Trimethylbenzene	19.28	1.0	20	0	96.4	76 - 124				
1,2-Dibromo-3-chloropropane	21.13	1.0	20	0	106	62 - 128				
1,2-Dibromoethane	20.86	1.0	20	0	104	77 - 121				
1,2-Dichlorobenzene	19.47	1.0	20	0	97.4	80 - 119				
1,2-Dichloroethane	20.22	1.0	20	0	101	73 - 128				
1,2-Dichloropropane	20.51	1.0	20	0	103	78 - 122				
1,3,5-Trimethylbenzene	19.33	1.0	20	0	96.7	75 - 124				
1,3-Dichlorobenzene	19.5	1.0	20	0	97.5	80 - 119				
1,3-Dichloropropane	20.2	1.0	20	0	101	80 - 119				
1,4-Dichlorobenzene	19.35	1.0	20	0	96.7	79 - 118				
2,2-Dichloropropane	20.6	1.0	20	0	103	60 - 139				
2-Butanone	44.19	2.0	40	0	110	56 - 143				
2-Chlorotoluene	19.2	1.0	20	0	96.0	79 - 122				
2-Hexanone	39.94	2.0	40	0	99.9	57 - 139				
4-Chlorotoluene	19.4	1.0	20	0	97.0	78 - 122				
4-Isopropyltoluene	19.03	1.0	20	0	95.1	77 - 127				
4-Methyl-2-pentanone	39.79	2.0	40	0	99.5	67 - 130				
Acetone	43.92	2.0	40	0	110	39 - 160				
Benzene	20.1	1.0	20	0	100	79 - 120				
Bromobenzene	19.87	1.0	20	0	99.3	80 - 120				
Bromochloromethane	20.57	1.0	20	0	103	78 - 123				
Bromodichloromethane	20.81	1.0	20	0	104	79 - 125				
Bromoform	20.8	1.0	20	0	104	66 - 130				
Bromomethane	24.89	1.0	20	0	124	53 - 141				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221		Instrument: VOA6		Method: SW8260						
LCS	Sample ID: VLCSW-190307	Units: UG/L			Analysis Date: 07-Mar-2019 11:32					
Client ID:	Run ID: VOA6_334221	SeqNo: 4979742		PrepDate:		DF: 1				
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	41.71	2.0	40	0	104	64 - 133				
Carbon tetrachloride	20.37	1.0	20	0	102	72 - 136				
Chlorobenzene	20.04	1.0	20	0	100	82 - 118				
Chloroethane	20.16	1.0	20	0	101	60 - 138				
Chloroform	20.66	1.0	20	0	103	79 - 124				
Chloromethane	20.84	1.0	20	0	104	50 - 139				
cis-1,2-Dichloroethene	20.24	1.0	20	0	101	78 - 123				
cis-1,3-Dichloropropene	21.28	1.0	20	0	106	75 - 124				
Dibromochloromethane	21.15	1.0	20	0	106	74 - 126				
Dibromomethane	20.71	1.0	20	0	104	79 - 123				
Dichlorodifluoromethane	20.85	1.0	20	0	104	32 - 152				
Ethylbenzene	19.75	1.0	20	0	98.8	79 - 121				
Hexachlorobutadiene	21.73	1.0	20	0	109	66 - 134				
Isopropylbenzene	19.51	1.0	20	0	97.5	72 - 131				
m,p-Xylene	39.04	2.0	40	0	97.6	80 - 121				
Methylene chloride	21.08	2.0	20	0	105	74 - 124				
Naphthalene	20.19	1.0	20	0	101	61 - 128				
n-Butylbenzene	18.73	1.0	20	0	93.7	75 - 128				
n-Propylbenzene	18.86	1.0	20	0	94.3	76 - 126				
o-Xylene	20.35	1.0	20	0	102	78 - 122				
sec-Butylbenzene	18.7	1.0	20	0	93.5	77 - 126				
Styrene	20.57	1.0	20	0	103	78 - 123				
tert-Butylbenzene	18.73	1.0	20	0	93.7	78 - 124				
Tetrachloroethene	19.69	1.0	20	0	98.5	74 - 129				
Toluene	20.02	1.0	20	0	100	80 - 121				
trans-1,2-Dichloroethene	20.66	1.0	20	0	103	75 - 124				
trans-1,3-Dichloropropene	21.3	1.0	20	0	106	73 - 127				
Trichloroethene	20.61	1.0	20	0	103	79 - 123				
Trichlorofluoromethane	20.26	1.0	20	0	101	65 - 141				
Vinyl chloride	19.61	1.0	20	0	98.1	58 - 137				
Surr: 1,2-Dichloroethane-d4	51.83	1.0	50	0	104	81 - 118				
Surr: 4-Bromofluorobenzene	52.4	1.0	50	0	105	85 - 114				
Surr: Dibromofluoromethane	51.54	1.0	50	0	103	80 - 119				
Surr: Toluene-d8	47.67	1.0	50	0	95.3	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221		Instrument: VOA6		Method: SW8260						
MS		Sample ID: HS19021428-09MS		Units: UG/L		Analysis Date: 07-Mar-2019 17:08				
Client ID: 16EW08_022619		Run ID: VOA6_334221		SeqNo: 4979750		PrepDate:		DF: 5		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	106.7	5.0	100	0	107	78 - 124				
1,1,1-Trichloroethane	92.75	5.0	100	0	92.8	74 - 131				
1,1,2,2-Tetrachloroethane	101.9	5.0	100	0	102	71 - 121				
1,1,2-Trichloroethane	106.8	5.0	100	0	107	80 - 119				
1,1-Dichloroethane	84.87	5.0	100	0	84.9	77 - 125				
1,1-Dichloroethene	89.03	5.0	100	0	89.0	71 - 131				
1,1-Dichloropropene	103.2	5.0	100	0	103	78 - 125				
1,2,3-Trichlorobenzene	119.4	5.0	100	0	119	69 - 129				
1,2,3-Trichloropropane	98.65	5.0	100	0	98.7	73 - 122				
1,2,4-Trichlorobenzene	117.3	5.0	100	0	117	69 - 130				
1,2,4-Trimethylbenzene	109.1	5.0	100	0	109	76 - 124				
1,2-Dibromo-3-chloropropane	105.8	5.0	100	0	106	62 - 128				
1,2-Dibromoethane	109.1	5.0	100	0	109	77 - 121				
1,2-Dichlorobenzene	105.2	5.0	100	0	105	80 - 119				
1,2-Dichloroethane	97.19	5.0	100	0	97.2	73 - 128				
1,2-Dichloropropane	98.98	5.0	100	0	99.0	78 - 122				
1,3,5-Trimethylbenzene	111	5.0	100	0	111	75 - 124				
1,3-Dichlorobenzene	105.6	5.0	100	0	106	80 - 119				
1,3-Dichloropropane	107.2	5.0	100	0	107	80 - 119				
1,4-Dichlorobenzene	103.7	5.0	100	0	104	79 - 118				
2,2-Dichloropropane	82.15	5.0	100	0	82.1	60 - 139				
2-Butanone	169.4	10	200	0	84.7	56 - 143				
2-Chlorotoluene	106.1	5.0	100	0	106	79 - 122				
2-Hexanone	211.6	10	200	0	106	57 - 139				
4-Chlorotoluene	106	5.0	100	0	106	78 - 122				
4-Isopropyltoluene	115.7	5.0	100	0	116	77 - 127				
4-Methyl-2-pentanone	212.7	10	200	0	106	67 - 130				
Acetone	169.1	10	200	0	84.6	39 - 160				
Benzene	100.6	5.0	100	0	101	79 - 120				
Bromobenzene	103.8	5.0	100	0	104	80 - 120				
Bromochloromethane	81.39	5.0	100	0	81.4	78 - 123				
Bromodichloromethane	99.73	5.0	100	0	99.7	79 - 125				
Bromoform	107.1	5.0	100	0	107	66 - 130				
Bromomethane	95.96	5.0	100	0	96.0	53 - 141				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221		Instrument: VOA6		Method: SW8260						
MS		Sample ID: HS19021428-09MS		Units: UG/L		Analysis Date: 07-Mar-2019 17:08				
Client ID: 16EW08_022619		Run ID: VOA6_334221		SeqNo: 4979750		PrepDate:		DF: 5		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	167.7	10	200	0	83.8	64 - 133				
Carbon tetrachloride	111	5.0	100	0	111	72 - 136				
Chlorobenzene	109	5.0	100	0	109	82 - 118				
Chloroethane	81.3	5.0	100	0	81.3	60 - 138				
Chloroform	85.71	5.0	100	0	85.7	79 - 124				
Chloromethane	79.48	5.0	100	0	79.5	50 - 139				
cis-1,2-Dichloroethene	246.8	5.0	100	168.2	78.7	78 - 123				
cis-1,3-Dichloropropene	99.04	5.0	100	0	99.0	75 - 124				
Dibromochloromethane	111.3	5.0	100	0	111	74 - 126				
Dibromomethane	100.6	5.0	100	0	101	79 - 123				
Dichlorodifluoromethane	56.25	5.0	100	0	56.3	32 - 152				
Ethylbenzene	113.9	5.0	100	0	114	79 - 121				
Hexachlorobutadiene	126	5.0	100	0	126	66 - 134				
Isopropylbenzene	117.8	5.0	100	0	118	72 - 131				
m,p-Xylene	223	10	200	0	112	80 - 121				
Methylene chloride	85.97	10	100	0	86.0	74 - 124				
Naphthalene	107.2	5.0	100	0	107	61 - 128				
n-Butylbenzene	115.1	5.0	100	0	115	75 - 128				
n-Propylbenzene	112.3	5.0	100	0	112	76 - 126				
o-Xylene	112.7	5.0	100	0	113	78 - 122				
sec-Butylbenzene	115.5	5.0	100	0	116	77 - 126				
Styrene	111.5	5.0	100	0	112	78 - 123				
tert-Butylbenzene	112.9	5.0	100	0	113	78 - 124				
Tetrachloroethene	119.9	5.0	100	0	120	74 - 129				
Toluene	111.6	5.0	100	0	112	80 - 121				
trans-1,2-Dichloroethene	86.72	5.0	100	0	86.7	75 - 124				
trans-1,3-Dichloropropene	99.31	5.0	100	0	99.3	73 - 127				
Trichloroethene	276.9	5.0	100	173	104	79 - 123				
Trichlorofluoromethane	88.99	5.0	100	0	89.0	65 - 141				
Vinyl chloride	75.65	5.0	100	0	75.6	58 - 137				
Surr: 1,2-Dichloroethane-d4	206	5.0	250	0	82.4	81 - 118				
Surr: 4-Bromofluorobenzene	254	5.0	250	0	102	85 - 114				
Surr: Dibromofluoromethane	209	5.0	250	0	83.6	80 - 119				
Surr: Toluene-d8	263.5	5.0	250	0	105	89 - 112				

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221		Instrument: VOA6		Method: SW8260						
MSD		Sample ID: HS19021428-09MSD		Units: UG/L		Analysis Date: 07-Mar-2019 17:33				
Client ID: 16EW08_022619		Run ID: VOA6_334221		SeqNo: 4979751		PrepDate:		DF: 5		
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
1,1,1,2-Tetrachloroethane	110.9	5.0	100	0	111	78 - 124	106.7	3.79	20	
1,1,1-Trichloroethane	100.6	5.0	100	0	101	74 - 131	92.75	8.16	20	
1,1,2,2-Tetrachloroethane	110.7	5.0	100	0	111	71 - 121	101.9	8.3	20	
1,1,2-Trichloroethane	109.3	5.0	100	0	109	80 - 119	106.8	2.28	20	
1,1-Dichloroethane	82.49	5.0	100	0	82.5	77 - 125	84.87	2.84	20	
1,1-Dichloroethene	94.43	5.0	100	0	94.4	71 - 131	89.03	5.89	20	
1,1-Dichloropropene	112.7	5.0	100	0	113	78 - 125	103.2	8.86	20	
1,2,3-Trichlorobenzene	138.7	5.0	100	0	139	69 - 129	119.4	15	20	S
1,2,3-Trichloropropane	104.7	5.0	100	0	105	73 - 122	98.65	5.92	20	
1,2,4-Trichlorobenzene	132.7	5.0	100	0	133	69 - 130	117.3	12.3	20	S
1,2,4-Trimethylbenzene	122.7	5.0	100	0	123	76 - 124	109.1	11.8	20	
1,2-Dibromo-3-chloropropane	119.4	5.0	100	0	119	62 - 128	105.8	12	20	
1,2-Dibromoethane	109.7	5.0	100	0	110	77 - 121	109.1	0.562	20	
1,2-Dichlorobenzene	116.2	5.0	100	0	116	80 - 119	105.2	9.86	20	
1,2-Dichloroethane	101.7	5.0	100	0	102	73 - 128	97.19	4.55	20	
1,2-Dichloropropane	101.4	5.0	100	0	101	78 - 122	98.98	2.38	20	
1,3,5-Trimethylbenzene	127.3	5.0	100	0	127	75 - 124	111	13.7	20	S
1,3-Dichlorobenzene	117.3	5.0	100	0	117	80 - 119	105.6	10.5	20	
1,3-Dichloropropane	110.2	5.0	100	0	110	80 - 119	107.2	2.79	20	
1,4-Dichlorobenzene	115.7	5.0	100	0	116	79 - 118	103.7	10.9	20	
2,2-Dichloropropane	91.55	5.0	100	0	91.6	60 - 139	82.15	10.8	20	
2-Butanone	179.3	10	200	0	89.7	56 - 143	169.4	5.7	20	
2-Chlorotoluene	118.3	5.0	100	0	118	79 - 122	106.1	10.9	20	
2-Hexanone	214.2	10	200	0	107	57 - 139	211.6	1.23	20	
4-Chlorotoluene	119.2	5.0	100	0	119	78 - 122	106	11.7	20	
4-Isopropyltoluene	133.6	5.0	100	0	134	77 - 127	115.7	14.4	20	S
4-Methyl-2-pentanone	214.5	10	200	0	107	67 - 130	212.7	0.815	20	
Acetone	173.4	10	200	0	86.7	39 - 160	169.1	2.47	20	
Benzene	104	5.0	100	0	104	79 - 120	100.6	3.38	20	
Bromobenzene	113.4	5.0	100	0	113	80 - 120	103.8	8.83	20	
Bromochloromethane	88.16	5.0	100	0	88.2	78 - 123	81.39	7.98	20	
Bromodichloromethane	101.3	5.0	100	0	101	79 - 125	99.73	1.53	20	
Bromoform	110.3	5.0	100	0	110	66 - 130	107.1	2.9	20	
Bromomethane	91.51	5.0	100	0	91.5	53 - 141	95.96	4.74	20	

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221		Instrument: VOA6		Method: SW8260						
MSD	Sample ID: HS19021428-09MSD	Units: UG/L			Analysis Date: 07-Mar-2019 17:33					
Client ID: 16EW08_022619	Run ID: VOA6_334221	SeqNo: 4979751	PrepDate:	DF: 5						
Analyte	Result	PQL	SPK Val	SPK Ref Value	%REC	Control Limit	RPD Ref Value	%RPD	RPD Limit	Qual
Carbon disulfide	177.8	10	200	0	88.9	64 - 133	167.7	5.87	20	
Carbon tetrachloride	119.3	5.0	100	0	119	72 - 136	111	7.25	20	
Chlorobenzene	113.8	5.0	100	0	114	82 - 118	109	4.35	20	
Chloroethane	86.74	5.0	100	0	86.7	60 - 138	81.3	6.47	20	
Chloroform	89.46	5.0	100	0	89.5	79 - 124	85.71	4.28	20	
Chloromethane	78.81	5.0	100	0	78.8	50 - 139	79.48	0.851	20	
cis-1,2-Dichloroethene	265.7	5.0	100	168.2	97.5	78 - 123	246.8	7.37	20	
cis-1,3-Dichloropropene	100.7	5.0	100	0	101	75 - 124	99.04	1.67	20	
Dibromochloromethane	112.6	5.0	100	0	113	74 - 126	111.3	1.23	20	
Dibromomethane	100.7	5.0	100	0	101	79 - 123	100.6	0.0743	20	
Dichlorodifluoromethane	61.01	5.0	100	0	61.0	32 - 152	56.25	8.12	20	
Ethylbenzene	119.4	5.0	100	0	119	79 - 121	113.9	4.78	20	
Hexachlorobutadiene	155.1	5.0	100	0	155	66 - 134	126	20.7	20	SR
Isopropylbenzene	127	5.0	100	0	127	72 - 131	117.8	7.49	20	
m,p-Xylene	235.5	10	200	0	118	80 - 121	223	5.46	20	
Methylene chloride	85.24	10	100	0	85.2	74 - 124	85.97	0.853	20	
Naphthalene	119.8	5.0	100	0	120	61 - 128	107.2	11.1	20	
n-Butylbenzene	135.9	5.0	100	0	136	75 - 128	115.1	16.6	20	S
n-Propylbenzene	128.3	5.0	100	0	128	76 - 126	112.3	13.2	20	S
o-Xylene	117.5	5.0	100	0	117	78 - 122	112.7	4.16	20	
sec-Butylbenzene	135.3	5.0	100	0	135	77 - 126	115.5	15.8	20	S
Styrene	115.8	5.0	100	0	116	78 - 123	111.5	3.81	20	
tert-Butylbenzene	131.7	5.0	100	0	132	78 - 124	112.9	15.4	20	S
Tetrachloroethene	131.6	5.0	100	0	132	74 - 129	119.9	9.32	20	S
Toluene	118.5	5.0	100	0	118	80 - 121	111.6	5.95	20	
trans-1,2-Dichloroethene	90.99	5.0	100	0	91.0	75 - 124	86.72	4.8	20	
trans-1,3-Dichloropropene	98.7	5.0	100	0	98.7	73 - 127	99.31	0.62	20	
Trichloroethene	289.3	5.0	100	173	116	79 - 123	276.9	4.38	20	
Trichlorofluoromethane	98.03	5.0	100	0	98.0	65 - 141	88.99	9.66	20	
Vinyl chloride	79.57	5.0	100	0	79.6	58 - 137	75.65	5.06	20	
Surr: 1,2-Dichloroethane-d4	207.5	5.0	250	0	83.0	81 - 118	206	0.759	20	
Surr: 4-Bromofluorobenzene	245	5.0	250	0	98.0	85 - 114	254	3.62	20	
Surr: Dibromofluoromethane	212.9	5.0	250	0	85.1	80 - 119	209	1.85	20	
Surr: Toluene-d8	264.6	5.0	250	0	106	89 - 112	263.5	0.388	20	

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

QC BATCH REPORT

Batch ID: R334221	Instrument: VOA6	Method: SW8260
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The following samples were analyzed in this batch:

HS19021428-04	HS19021428-05	HS19021428-06	HS19021428-07
HS19021428-08	HS19021428-09		

ALS Houston, US

Date: 08-Mar-19

Client: Bhate Environmental Associates, Inc.
Project: LHAAP Site 16
WorkOrder: HS19021428

**QUALIFIERS,
ACRONYMS, UNITS**

Qualifier	Description
*	Value exceeds Regulatory Limit
a	Not accredited
B	Analyte detected in the associated Method Blank above the Reporting Limit
E	Value above quantitation range
H	Analyzed outside of Holding Time
J	Analyte detected below quantitation limit
M	Manually integrated, see raw data for justification
n	Not offered for accreditation
ND	Not Detected at the Reporting Limit
O	Sample amount is > 4 times amount spiked
P	Dual Column results percent difference > 40%
R	RPD above laboratory control limit
S	Spike Recovery outside laboratory control limits
U	Analyzed but not detected above the MDL/SDL

Acronym	Description
DCS	Detectability Check Study
DUP	Method Duplicate
LCS	Laboratory Control Sample
LCSD	Laboratory Control Sample Duplicate
MBLK	Method Blank
MDL	Method Detection Limit
MQL	Method Quantitation Limit
MS	Matrix Spike
MSD	Matrix Spike Duplicate
PDS	Post Digestion Spike
PQL	Practical Quantitation Limit
SD	Serial Dilution
SDL	Sample Detection Limit
TRRP	Texas Risk Reduction Program

CERTIFICATIONS,ACCREDITATIONS & LICENSES

Agency	Number	Expire Date
Arkansas	88-0356	27-Mar-2019
Texas	T10470231-18-21	30-Apr-2019
North Dakota	R193 2018-2019	30-Apr-2019
Illinois	004438	29-Jun-2019
Louisiana	03087	30-Jun-2019
Dept of Defense	ANAB L2231	20-Dec-2021
Kentucky	123043 - 2018	30-Apr-2019
Kansas	E-10352 2018-2019	31-Jul-2019
Oklahoma	2018-156	31-Aug-2019
North Carolina	624-2019	31-Dec-2019
California	2919, 2018-2019	30-Apr-2019
Maryland	343, 2018-2019	30-Jun-2019

Sample Receipt Checklist

Client Name: Bhate Environmental
 Work Order: HS19021428

Date/Time Received: **27-Feb-2019 09:10**
 Received by: **RPG**

Checklist completed by: Jared R. Makan 27-Feb-2019
 eSignature Date

Reviewed by: RJ Modashia 28-Feb-2019
 eSignature Date

Matrices: **Water**

Carrier name: **FedEx Priority Overnight**

Shipping container/cooler in good condition?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	Not Present <input type="checkbox"/>
Custody seals intact on shipping container/cooler?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	Not Present <input type="checkbox"/>
Custody seals intact on sample bottles?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	Not Present <input checked="" type="checkbox"/>
VOA/TX1005/TX1006 Solids in hermetically sealed vials?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	Not Present <input checked="" type="checkbox"/>
Chain of custody present?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	1 Page(s)
Chain of custody signed when relinquished and received?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	COC IDs:N/A
Samplers name present on COC?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Chain of custody agrees with sample labels?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Samples in proper container/bottle?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Sample containers intact?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Sufficient sample volume for indicated test?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
All samples received within holding time?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Container/Temp Blank temperature in compliance?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	
Temperature(s)/Thermometer(s):	0.6c/0.6c UC/C IR11		
Cooler(s)/Kit(s):	44309		
Date/Time sample(s) sent to storage:	02/27/2019 15:20		
Water - VOA vials have zero headspace?	Yes <input checked="" type="checkbox"/>	No <input type="checkbox"/>	No VOA vials submitted <input type="checkbox"/>
Water - pH acceptable upon receipt?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	N/A <input checked="" type="checkbox"/>
pH adjusted?	Yes <input type="checkbox"/>	No <input type="checkbox"/>	N/A <input checked="" type="checkbox"/>
pH adjusted by:	<input type="text"/>		

Login Notes:

Client Contacted: _____ Date Contacted: _____ Person Contacted: _____
 Contacted By: _____ Regarding: _____

Comments:

Corrective Action:



2608 13th Avenue South, Suite 300
Birmingham Alabama 35205
Tel: 205-918-4000
Fax: 205-918-4050

Chain of Custody and Analytical Request

Page: _____ of _____
Project/Phase No: NW01312.0150
COC Number(1): _____
LIMS Number: _____

Facility/Base ID: **LHAAP**

Project/Site Name: **LHAAP / Site 16**

Client Name: _____

Collected by: **Scott Beesinger**

Sample Analyte requested⁽¹⁾

Quality Assay of Samples⁽²⁾

Field Sample ID (to Character Mean)	EPHMS/LC/CO (to Character Mean)	Date Collected (dd-mm-YY)	Time Collected (hh:mm)	Sample Depth (beginning - ending)	SA Code (#)	Sample Number (#)	Sample Matrix ⁽³⁾	Number of Containers	Sample Analyte requested ⁽¹⁾	Quality Assay of Samples ⁽²⁾
EW01_022619		26 FEB 2019	0830	-	N		WG	4 X	VOC PERCHLORATE	
EW01_022619_NS		26 FEB 2019	0830	-	MS		WG	4 X		
EW01_022619_MSD		26 FEB 2019	0830	-	SD		WG	4 X		
EW05_022619		26 FEB 2019	0850	-	N		WG	4 X		
EW05_022619_FD		26 FEB 2019	0850	-	FD		WG	4 X		
EW02_022619		26 FEB 2019	0900	-	N		WG	4 X		
EW06_022619		26 FEB 2019	0910	-	N		WG	4 X		
EW03_022619		26 FEB 2019	0920	-	N		WG	4 X		
EW07_022619		26 FEB 2019	0930	-	N		WG	4 X		
EW04_022619		26 FEB 2019	0940	-	N		WG	4 X		
EW08_022619		26 FEB 2019	0950	-	N		WG	4 X		

COMMENTS:



Bhate Environmental Associates, Inc.
LHAAP Site 16

HS19021428

Requested by (Name): Scott Beesinger Date: 2/26/19 Time: 1400

Requested by (Name): RC Date: 2/27/19 Time: 09:10

Custom/Third Party or Receiving Laboratory

Delivered Directly to Lab: _____

Method of Shipment: _____

Sample Delivery Details / Laboratory Receipt

Shipper: _____

Carrier: _____

Number: _____

ANLN - SONIA WEST

Lab Receptor: _____

Delivery Date/Time: _____


ANLN - SONIA WEST

Lab Receptor: _____

Delivery Date/Time: _____

- Chain of Custody Number a date collected + collector number (e.g. 03-02-1989-01)
- Sample Type (SA) Codes: N = Normal Sample, TB = Trip Blank (+) Sample, FD = Field Duplicate (+) Sample, FB = Field replicate (+) Sample, MS = Matrix Spike, SD = Matrix Spike Duplicate, AB = Ambient Blank (+)
- Sample Number: Unique sample number collected from a particular location per day. (e.g. Groundwater sample collected from MW-1 on 10/10/99 = 02, etc.)
- Matrix Codes: GS = Solid Gas, WG = Groundwater, WS = Surface Water, SD = Soil, SF = Sediment, SA = Sludge, SS = Surface Soil Samples, WA = Aqueous Blank Samples (+), EQ = Equipment, AMB = Ambient, etc. SA = Soil Blank
- Sample Analyte Requested: Analytical method requested and number of containers provided for each.
- Quality assurance samples are assigned by data (id number) and the sample number associated with the sample (01, 02, etc.) (e.g. Equipment Blank collected in association with MW-1 on 10/10/99 will be designated 1010901. In this Equipment Blank List Control

44309
4111
CFC

 ALS Environmental 10450 Stancliff Rd., Suite 210 Houston, Texas 77099 Tel. +1 281 530 5656 Fax. +1 281 530 5887	CUSTODY SEAL		Seal Broken By:
	Date: 2/26/19	Time: 1400	FM
	Name: Scott Beesinger		Date: 02/27/19
	Company: BHATG		

44309 FEB 27 2019

Must Deliver Next Business Day
Time and Temperature Sensitive!



44309

ORIGIN ID: SGRA (903) 930-6199
 SCOTT BEESINGER
 BHATE ENVIRONMENTAL ASSOCIATES
 1209-B EAST GRAND AVE. PHB202

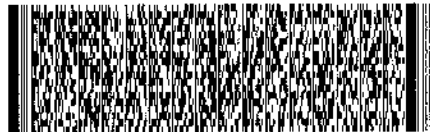
SHIP DATE: 07AUG18
 ACTWT: 1.00 LB MAN
 CAD: 300130/CAFE3111
 DIMS: 26x14x14 IN

MARSHALL, TX 75670
 UNITED STATES US

TO CLIENT SERVICES
 ALS LABORATORY GROUP
 10450 STANCLIFF ROAD
 SUITE 210
 HOUSTON TX 77099

(281) 530-5656
 REF: LHAAP-58-RJ

RMA: ||| ||| |||



FedEx
 TRK# 4380 9531 5094

WED - 27 FEB 10:30A
 PRIORITY OVERNIGHT

AB SGRA

77099
 TX-US
 IAH



FID 162786 26FFB19 GGGA 653C2/RF3D/OCBA



Case Narrative

Method: 6850

Analysis: Perchlorate

Analysis SOP: LC-MS-CLO4

ALS WO ID(s): 1905651; 1906112; 1906330;
1906332; 1906334

Client: ALS Laboratories (Houston, TX)

Matrix: Water

ELMS Batch (HBN): 2223 (233911)

General Set Information: There were thirteen field samples in these Work Orders. The samples were analyzed for perchlorate.

Method Summary: Each sample was prepared as noted below and analyzed using an Agilent 1100 LC/MSD system in select ion monitoring (SIM) mode at m/z 83 and 85, which corresponds to the loss of one oxygen atom from the perchlorate molecule. ChemStation software was used for instrument control and data analysis. The ion ratio of m/z 83 to 85 was used to positively identify the response peak as perchlorate. Quantitation was performed using the m/z 83 peak area. An internal standard (ISTD) of ^{18}O labeled perchlorate was added to each sample to establish the perchlorate peak retention time and used in quantitation.

Sample Preparation: A 10.0mL aliquot of each sample was transferred into a 15-mL centrifuge tube. 50 μL of an ^{18}O labeled perchlorate solution was added to each sample as an internal standard. The samples were then capped, vortexed, and filtered into autosampler vial using Phenex PES membrane 0.45 μm Syringe filters.

Holding Times: Holding times were met for all analyses.

Dilutions: Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Method QC data: The method blank (LMB 642099) was less than 1/2 the CRDL. The recovery for the LCS (642100) was within acceptable parameters.



MS/MSD Analysis: MS/MSD was performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0 μ l of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. The spike target was 4.0 μ g/L. The MS/MSD percent recoveries and relative percent difference (RPD) were within the performance limits, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.

Instrument QC: Instrument initial and continuing calibrations were performed in accordance with published procedures.

NC/CAR(s): NA

Sample Calculation: Samples were reported in μ g/L. Results were calculated in μ g/L by the equation $(A) \times (B)$,

where: A = Analyte concentration from the standard curve (μ g/L)
B = Dilution performed at time of analysis

Miscellaneous Comments: These samples were analyzed in accordance with the requirements found in the DOD QSM Version 5.1.1. The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level of 4.0 μ g/L. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported. Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEBI04) along with datafiles 05MARD07-10.

Thomas Bosch March 06, 2019
Analyst Date



ANALYTICAL REPORT

Report Date: March 07, 2019

RJ Modashia
ALS Environmental (Houston)
10450 Stancliff Road
Suite 210
Houston, TX 77099

Phone: 281 530-5656

E-mail: RJ.Modashia@ALSGlobal.com

Workorder: **34-1906112**

Project ID: HS19021428

Purchase Order: HS19021428

Project Manager Kevin W. Griffiths

Client Sample ID	Lab ID	Collect Date	Receive Date	Sampling Site
EW01_022619	1906112001	02/26/19	02/28/19	
EW05_022619	1906112004	02/26/19	02/28/19	
EW05_022619_FD	1906112005	02/26/19	02/28/19	
EW02_022619	1906112006	02/26/19	02/28/19	
EW06_022619	1906112007	02/26/19	02/28/19	
EW03_022619	1906112008	02/26/19	02/28/19	
EW07_022619	1906112009	02/26/19	02/28/19	
EW04_022619	1906112010	02/26/19	02/28/19	
EW08_022619	1906112011	02/26/19	02/28/19	

Client QC ID *	Lab ID	Collect Date	Receive Date	Sampling Site
EW01_022619MS	1906112002	02/26/19	02/28/19	
EW01_022619MSD	1906112003	02/26/19	02/28/19	

*Client QC is reported as part of the Quality Control results report, if requested.



ANALYTICAL REPORT

Workorder: **34-1906112**Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: EW01_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112001	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 09:54	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	51	1.0	2.0	4.0	1	

Sample ID: EW05_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112004	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 13:15	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	230	10	20	40	10	

Sample ID: EW05_022619_FD	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112005	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 13:28	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	240	10	20	40	10	

Sample ID: EW02_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112006	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 10:59	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U



ANALYTICAL REPORT

Workorder: **34-1906112**Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: EW06_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112007	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 13:41	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Sample ID: EW03_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112008	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 11:26	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Sample ID: EW07_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112009	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 11:39	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Sample ID: EW04_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112010	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 11:52	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U



ANALYTICAL REPORT

Workorder: **34-1906112**Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

Analytical Results

Sample ID: EW08_022619	Sampling Site: NA	Collected: 02/26/2019				
Lab ID: 1906112011	Media: 125 mL Nalgene	Received: 02/28/2019				
Matrix: Water	Sampling Parameter: NA					
Analysis Method - EPA 6850, DoD QSM						
Preparation: Not Applicable	Analysis: EPA 6850, DoD QSM Water Batch: ELMS/2223 (HBN: 233911) Analyzed: 03/05/2019 12:05	Instrument ID: LCMS04 Percent Solid: NA Report Basis: Wet				
Analyte	Result (ug/L)	DL (ug/L)	LOD (ug/L)	LOQ (ug/L)	Dilution	Qual
Perchlorate	ND	1.0	2.0	4.0	1	U

Comments

Quality Control: EPA 6850, DoD QSM - (HBN: 233911)

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.

Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Method	Analyst	Peer Review
EPA 6850, DoD QSM	/S/ Thomas Bosch 03/06/2019 09:17	/S/ Stephen Brose 03/07/2019 11:04

Laboratory Contact Information

ALS Environmental
960 W Levoy Drive
Salt Lake City, Utah 84123

Phone: (801) 266-7700
Email: als@alst.com
Web: www.alst.com



ANALYTICAL REPORT

Workorder: 34-1906112

Client: ALS Environmental
(Houston)

Project Manager: Kevin W. Griffiths

General Lab Comments

The results provided in this report relate only to the items tested.
 Samples were received in acceptable condition unless otherwise noted.
 Samples have not been blank corrected unless otherwise noted.
 This test report shall not be reproduced, except in full, without written approval of ALS.

ALS provides professional analytical services for all samples submitted. ALS is not in a position to interpret the data and assumes no responsibility for the quality of the samples submitted.

All quality control samples processed with the samples in this report yielded acceptable results unless otherwise noted.

ALS is accredited for specific fields of testing (scopes) in the following testing sectors. The quality system implemented at ALS conforms to accreditation requirements and is applied to all analytical testing performed by ALS. The following table lists testing sector, accreditation body, accreditation number and website. Please contact these accrediting bodies or your ALS project manager for the current scope of accreditation that applies to your analytical testing.

Testing Sector	Accreditation Body (Standard)	Certificate Number	Website
Environmental	PJLA (DoD ELAP)	L17-506	http://www.pjlabs.com
	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com
	Utah (TNI)	UT00953	http://lams.nelac-institute.org/search
	Nevada (TNI)	UT00953201-1	https://ndep.nv.gov/water/lab-certification
	Iowa (TNI)	IA# 376	http://www.shl.uiowa.edu/labcert/idnr/
	Kansas	E-10416	http://www.kdheks.gov/envlab/disclaimer.html
	Oklahoma (TNI)	IJ# 9980	http://www.deq.state.ok.us/CSDnew/labcert.htm
Texas (TNI)	T104704456-18-9	https://www.tceq.texas.gov/assets/public/compliance/compliance_support/qa/txnelap_lab_list.pdf	
Industrial Hygiene	AIHA (ISO 17025 & AIHA IHLAP)	101574	http://www.aihaaccreditedlabs.org
	DOECAP-AP	L18-606	http://www.pjlabs.com
	Washington	C596	https://ecology.wa.gov/Regulations-Permits/Permits-certifications/Laboratory-Accreditation
Dietary Supplements	PJLA (ISO 17025)	L17-507-R1	http://www.pjlabs.com

Result Symbol Definitions

MDL = Method Detection Limit, a statistical estimate of method/media/instrument sensitivity.

RL = Reporting Limit, a verified value of method/media/instrument sensitivity.

CRDL = Contract Required Detection Limit

Reg. Limit = Regulatory Limit.

ND = Not Detected, testing result not detected above the MDL or RL.

< Means this testing result is less than the numerical value.

** No result could be reported, see sample comments for details.

Qualifier Symbol Definitions

U = Qualifier indicates that the analyte was not detected above the MDL.

J = Qualifier Indicates that the analyte value is between the MDL and the RL. It is also used to indicate an estimated value for tentatively identified compounds in mass spectrometry where a 1:1 response is assumed.

B = Qualifier indicates that the analyte was detected in the blank.

E = Qualifier indicates that the analyte result exceeds calibration range.

P = Qualifier indicates that the RPD between the two columns is greater than 40%.



Quality Control Sample Batch Report

00936104

Analysis Information

Workorder: 1906112

Limits: Client SOW/Contract Specified
Basis: DoD QSM

Preparation: NA
Batch: NA
Prepared By: NA

Analysis: EPA 6850, DoD QSM
Batch: ELMS/2223 (HBN: 233911)
Analyzed By: Thomas Bosch

Blank

LMB: 642099 Analyzed: 03/05/2019 09:26 Units: ug/L			
Analyte	Result	MDL	RL
Perchlorate	ND	1	2.00

Laboratory Control Sample

LCS: 642100 Analyzed: 03/05/2019 09:00 Dilution: 1 Units: ug/L				
Analyte	Result	Target	% Rec	QC Limits
Perchlorate	4.11	4.00	103	78.8 123.8

Matrix Spike - Matrix Spike Duplicate

Sample: 1906112001 Analyzed: 03/05/2019 09:54 Dilution: 1 Units: ug/L		MS: 1906112002 Analyzed: 03/05/2019 10:07 Dilution: 1 Units: ug/L				MSD: 1906112003 Analyzed: 03/05/2019 10:20 Dilution: 1 Units: ug/L			
Analyte	Result	Result	Target	% Rec	QC Limits	Result	% Rec	RPD	QC Limits
Perchlorate	51.0	53.9	4	▲ 65.7	78.8 123.8	53.7	▲ 61.2	0.337	0.0 20.0

Continuing Calibration Verification

CCV: 642096 Analyzed: 03/05/2019 08:44 Units: ug/L Criteria: ± 15%			CCV: 642101 Analyzed: 03/05/2019 12:18 Units: ug/L Criteria: ± 15%			CCV: 642102 Analyzed: 03/05/2019 14:07 Units: ug/L Criteria: ± 15%			
Analyte	Result	Target	% Rec.	Result	Target	% Rec.	Result	Target	% Rec.
Perchlorate	25.0	25.0	100	25.0	25.0	100	24.5	25.0	97.8

Interference Check Sample

ICSA: 642098 Analyzed: 03/05/2019 09:13 Units: ug/L Criteria: ± 30%			
Analyte	Result	Target	% Rec.
Perchlorate	3.93	4.00	98.2

Comments

Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly.



Quality Control Sample Batch Report

00936105

Analysis Information

Workorder: 1906112

Limits: Client SOW/Contract Specified

Preparation: NA

Analysis: EPA 6850, DoD QSM

Basis: DoD QSM

Batch: NA

Batch: ELMS/2223 (HBN: 233911)

Prepared By: NA

Analyzed By: Thomas Bosch

QC Report Authorization (/S/ is an electronic signature that complies with 21 CFR Part 11)

Analyst	Peer Review
/S/ Thomas Bosch 03/06/2019 13:45	/S/ Stephen Brose 03/07/2019 11:04

Symbols and Definitions

- * - Analyte above reporting limit or outside of control limits
- ▲ - Sample result is greater than 4 times the spike added
- - Sample and Matrix Duplicate less than 5 times the reporting limit
- - Result is above the calibration range
- # - The Matrix Spike, Matrix Spike duplicate or Matrix Duplicate is reported for your information only. The sample matrix may be inappropriate for the method selected.

- RPD - Relative % Difference (Spike / Spike Duplicate)
- ND - Not Detected (U - Qualifier also flags analyte as not detected)
- NA - Not Applicable
- QC results are not adjusted for moisture correction, where applicable



W 1906112

18698/# 2

10450 Stancliff Rd, Ste 210
Houston, TX 77099
T: +1 281 530 5656
F: +1 281 530 5887
www.alsglobal.com

Subcontract Chain of Custody

COC ID: 10844

SUBCONTRACT TO:

ALS Laboratory Group
960 LeVoy Dr
Salt Lake City, UT 84123

Phone: +1 801 266 7700

1906112

CUSTOMER INFORMATION:

Company: ALS Houston
Contact: RJ Modashia
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Email: RJ.Modashia@alsglobal.com
Alternate Contact: Jumoke M. Lawal
Email: jumoke.lawal@alsglobal.com

INVOICE INFORMATION:

Company: ALS Houston
Contact: Accounts Payable
Address: 10450 Stancliff Rd, Ste 210
Phone: +1 281 530 5656
Reference: HS19021428
TSR: Danielle Winnings

	LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
	ANALYSIS REQUESTED			DUE DATE
1.	HS19021428-01	EW01_022619	Water	26 Feb 2019 08:30
	SUB_Perch-6850			13 Mar 2019
2.	HS19021428-02	EW05_022619	Water	26 Feb 2019 08:50
	SUB_Perch-6850			13 Mar 2019
3.	HS19021428-03	EW05_022619_FD	Water	26 Feb 2019 08:50
	SUB_Perch-6850			13 Mar 2019
4.	HS19021428-04	EW02_022619	Water	26 Feb 2019 09:00
	SUB_Perch-6850			13 Mar 2019
5.	HS19021428-05	EW06_022619	Water	26 Feb 2019 09:10
	SUB_Perch-6850			13 Mar 2019
6.	HS19021428-06	EW03_022619	Water	26 Feb 2019 09:20
	SUB_Perch-6850			13 Mar 2019
7.	HS19021428-07	EW07_022619	Water	26 Feb 2019 09:30
	SUB_Perch-6850			13 Mar 2019
8.	HS19021428-08	EW04_022619	Water	26 Feb 2019 09:40
	SUB_Perch-6850			13 Mar 2019
9.	HS19021428-09	EW08_022619	Water	26 Feb 2019 09:50

ms/msD



Subcontract Chain of Custody

COC ID: 10844

LAB SAMPLE ID	CLIENT SAMPLE ID	MATRIX	COLLECT DATE
ANALYSIS REQUESTED			DUE DATE
SUB_Perch-6850			13 Mar 2019

Comments: Please analyze for the analysis listed above.
Send report to the emails shown above.

HS19021428-01 MS/MSD

QC Level: DOD IV (DoD Data Package)

Relinquished By: J. Wainwright
 Received By: [Signature]
 Cooler ID(s): 9190

Date/Time: 2/27/19 18:00
 Date/Time: 2/28/19 0925
 Temperature(s): 1.5°C

cooler custody seal intact



ALS Environmental CHAIN-OF-CUSTODY

Project / Job / Task: HS19021428		Split:		Workorder ID: 1906112		Level: ENV_LVL4		Requested Analysis															
Client: ALS Environmental (Houston)				Account: 8101				Type: 125Poly															
Comments:																							

ALS-SALT LAKE CITY-RELATED INFORMATION REPORT (CRIR)

1906112

COOLER OR CONTAINER INFORMATION CHECKLIST (Fill In or Circle)

Client Name: ALS Environmental Project/Task/Site: POD Perchlorate
 Date/Time of Receipt: 2/28/19 0925 Number of Coolers Received: 1

Condition of Coolers: Acceptable/Unacceptable
 Cooler Custody Seals: Present/Absent/NA
 Container Custody Seals: Intact/Broken/NA
 Present/Absent/NA
 Intact/Broken/NA
 Ice Present: Yes/No/NA
 Frozen/Melted/NA

Temperature Control: Present/Not Included # 2/28
 Location Temp Taken: Control/Between Samples
 Are all temperatures within project specific guidelines? Yes/No/NA
 VOA Headspace Present? Yes/No/NA

pH Check Performed:	Metals	Yes/No/NA	Total Phenolics	Yes/No/NA	NO3/NO2	Yes/No/NA
	Cyanide	Yes/No/NA	TPH - 418.1	Yes/No/NA	Oil & Grease	Yes/No/NA
	Sulfide	Yes/No/NA	COD	Yes/No/NA	Total Phosphorous	Yes/No/NA
	Ammonia	Yes/No/NA	TKN	Yes/No/NA	Gross A.B, Gamma Spec	Yes/No/NA

Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.	Cooler Received	DCL Cooler No.	Temp.
1	C19 9190	1.5°C	4	C19	°C	7	C19	°C
2	C19	°C	5	C19	°C	8	C19	°C
3	C19	°C	6	C19	°C	9	C19	°C

Taken By: [Signature] Signature Jessica Holland Printed Name 2/28/19 Date

CLIENT-RELATED INFORMATION

- | | | | |
|--|---|--|---|
| <input type="checkbox"/> Missing Cooler | <input type="checkbox"/> Missing Samples/Bottles | <input type="checkbox"/> Incorrect Preservation | <input type="checkbox"/> Insufficient Sample Volume |
| <input type="checkbox"/> Cooler Conditions | <input type="checkbox"/> Broken/Leaking Samples | <input type="checkbox"/> pH Criteria Not Met | <input type="checkbox"/> Chain of Custody Problems |
| <input type="checkbox"/> Missing Paperwork | <input type="checkbox"/> Incorrect Bottle Type | <input type="checkbox"/> Residual Chlorine Present | <input type="checkbox"/> Other: |
| <input type="checkbox"/> Missing/Incorrect Bottle Labels | <input type="checkbox"/> Cooler Temperatures Out of Range | <input type="checkbox"/> Head Space in Bottles | |

BRIEFLY DESCRIBE THE PROBLEM AND THE ACTION TAKEN:

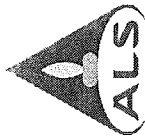
Client Notified? YES NO

Response Required Within 24 Hours

PROJECT MANAGEMENT

PROJECT MANAGER COMMENTS:

ALS Project Manager: _____ Returned to Sample Receipt by: _____ Date: _____
Printed Name Signature



Batch Worklist

HBN: 233911

Instrument: WP
Status: WP

Created: 3/5/2019 08:21
Analyst: T. Bosch

Batch: ELMS/ 2223
Rule: EPA 6850, DoD QSM Water

- Workorder: 1905651 [ENV_LVL4]
- Workorder: 1906112 [ENV_LVL4]
- Workorder: 1906330 [ENV_LVL4]
- Workorder: 1906332 [ENV_LVL4]
- Workorder: 1906334 [ENV_LVL4]

Pos	Lab ID	Sample ID	Prep Initial	Prep Final	Dust Weight	Type	Mix	Container	Procedure	Mgr	Expire Date	Due Date	Run Date
1	642096	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
2	642097	RLVS for HBN 233911 [ELMS/2223]				RLVS	3		E685041C3Q	5311		3/7/2019	
3	642098	ICS for HBN 233911 [ELMS/2223]				ICS	3		E6850.D3Q	5311		3/7/2019	
4	642099	LMB for HBN 233911 [ELMS/2223]				LMB	3		E6850Q413Q	5311		3/7/2019	
5	642100	LCS for HBN 233911 [ELMS/2223]				LCS	3		E6850Q413Q	5311		3/7/2019	
6	1905651001	HS19021158-02/LH18/24-SP650_02				SAMPLE	3	1905651001-A	E6850Q41.3	5480	3/21/2019	3/7/2019	
7	1906112001	EW01_022619				SAMPLE	3	1906112001-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
8	1906112002	EW01_022619MS				MS	3	1906112002-A	E6850Q413Q	5480		3/7/2019	
9	1906112003	EW01_022619MSD				MSD	3	1906112003-A	E6850Q413Q	5480		3/7/2019	
10	1906112004	EW05_022619				SAMPLE	3	1906112004-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
11	1906112005	EW05_022619_FD				FLDDUP	3	1906112005-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
12	1906112006	EW02_022619				SAMPLE	3	1906112006-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
13	1906112007	EW06_022619				SAMPLE	3	1906112007-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
14	1906112008	EW03_022619				SAMPLE	3	1906112008-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
15	1906112009	EW07_022619				SAMPLE	3	1906112009-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
16	642101	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	
17	1906112010	EW04_022619				SAMPLE	3	1906112010-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
18	1906112011	EW08_022619				SAMPLE	3	1906112011-A	E6850Q41.3	5480	3/26/2019	3/13/2019	
19	1906330001	LH18/24-SP140_022819				SAMPLE	3	1906330001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
20	1906332001	LH18/24-SP650_022819_BIX				SAMPLE	3	1906332001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
21	1906334001	LH18-24-SP650_022819_BIX				SAMPLE	3	1906334001-A	E6850Q41.3	5480	3/28/2019	3/14/2019	
22	642102	CCV for HBN 233911 [ELMS/2223]				CCV	3		E685041C3Q	5311		3/7/2019	



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Analytical Documentation

ALS Work Order #'s & Sample #()'s: 1905651 (001); 1906112 (001-11); 1906330 (001); 1906332 (001); 1906334 (001)
 ELMS Batch/HBN ID: 2223 (233911)
 Prep Date: 03/04/2019 Analysis Date: 03/05/2019 Analyst: T. Bosch
 Analyte: **Perchlorate** Matrix: **Water** Method: **6850**
 Sequence: \\HPCHEM\1\SEQUENCE\CLO4\2019\MAR\05MAR19D.s
 Reported DL: **1.0µg/L** Reported LOD: **2.0µg/L** Reported LOQ: **4.0µg/L**

SAMPLE PREPARATION/ANALYSIS:

Water: Samples were prepared by TNB. 10.0mL of each sample was pipetted into a 15-mL centrifuge tube, and 50µL of an oxygen-18 labeled perchlorate solution was added as an internal standard. The samples were capped, vortexed, and filtered with Phenex PES membrane 0.45µm Syringe filters prior to analysis.

REAGENTS: Eluent A1: 95% ASTM Type II water (ALS)/ 5% ACN (B&J Lot AH015-4)/0.1% glacial acetic acid (JT-Baker Lot 04802).
 Eluent B1: 95% ACN (B&J Lot AH015-4)/ 5% ASTM Type II water (ALS)/0.1% glacial acetic acid (JT-Baker Lot 04802).

STANDARDS: Internal Standard Spiking Solution Horizon# 43730. Dilutions of Working Standard Solution ID 43702 used for CCV's, LODV's, RLVS and IPC.

CALIBRATION CURVE: Used curve from 02/15/2019, sequence 15FEB19D.s Offline Quantitation Method: CLO4-DP1.M

INSTRUMENT CONDITIONS: Samples were analyzed with an Agilent 1100 LC/MSD system, in negative SIM mode, monitoring m/z 83, 85, and 89.

Instrument ID: LCMS04 Online Acquisition Method: CLO4-AQN.M Fragmentor: 160 Output Gain: 3 Injection Volume: 30µL
 Column: KP-RPPX C8 separator, 250mm Mobile Phase: 70% Eluent A1; 30% Eluent B1

FLOW GRADIENT:

Time (min.)	Flow (mL/min)
0	0.65
5.8	0.65
5.9	0.25
10.3	0.25
10.5	0.65
12.0	0.65

QC DATA: 4.0µL of QC Solution Horizon ID 41830 was used for LCS 642100; Target = 4.0µg/L. ASTM type II water was used for LMB 642099.

MS/MSD: The Matrix Spike and duplicate (MS/MSD) were performed on samples 1906112002/03 (Client ID's: EW01_022619). 4.0µL of Working Standard Solution Horizon ID 43701 was added to 10.0mL of sample preparation. Spike target = 4.0µg/L.

COMMENTS:

- 1) Results reported in µg/L. Field samples 1906112004/05 were re-analyzed and reported from 1:10 dilutions. Field sample 1906330001 was analyzed and reported from a 1:100 dilution. The reporting limits have been adjusted accordingly. Sample 1906112007 failed the 50-150% method requirement for ISTD recovery. This sample was re-prepped, re-analyzed and reported.
- 2) All QC, Blank, CCV, and MS/MSD results were within method parameters, except for the following. The Matrix Spike and duplicate (MS/MSD) failed QC acceptance criteria for percent recoveries. The MS/MSD is reported for the clients' information only. The sample matrix may be inappropriate for the method selected.
- 3) Sample data can be viewed at two directories within the ALS system: \\ALSLTWS013\LCMS\LCMS04\2019\MAR\HBN# or through NuGenesis\Tree\PrintData\LCMS\DefaultView.
- 4) Notebook: \\alsltws013\ORGANIC\BOSCH\LCMS\Perchlorates\Waters\2019\23911-DoD-ALS-Hstn LCMS4 or through \\ALSLTWS013\DATAREVIEW\HBN#
- 5) The Reporting Limit Verification Standard (RLVS – 642097) is reported from the analysis of the Laboratory Control Sample (LCS – 642100) at a level 4.0µg/L.
- 6) Due to limitations of the Chemstation Software, many of the chromatographic peaks require manual integration. Manual Integrations were performed for one of the Initial Calibration analyses (datafile: 15FEB104) along with datafiles 05MARD07-10.

5.5 Chromatography (GC, HPLC and LC/MS) Technical Review

Note: It is the peer reviewer's responsibility to ensure that appropriate criteria are used as defined in the HORIZON PROFILE. The evaluation criteria are prioritized as per Section 2.2 of this SOP. These items must be checked for all projects. The following checklist will be completed by both the analyst and the peer reviewer and scanned into the HBN folder with the raw data.

<u>Chromatography (GC, HPLC, LC/MS) Technical Review Criteria</u>	<u>Analyst Initials</u>	<u>Reviewer Initials</u>
Batch(es)/SDG: <u>E LMS: 2223 HBN: 233911</u>		
Sample Set IDs if Applicable: <u>1905651 / 190912 / 1906330</u> <u>1906332 / 1906334</u>		
<u>Calibration standards analyzed and meets criteria</u>	TB	SB
<u>Standards traceability checked and meets criteria</u>	TB	SB
<u>Standard curve coefficients evaluated and meet criteria</u>	TB	SB
<u>ICVs analyzed and meet acceptance criteria</u>	TB	SB
<u>CCVs analyzed and meet acceptance criteria</u>	TB	SB
<u>Method Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>Retention Time Windows checked</u>	TB	SB
<u>For method 8081A, Endrin/DDT Breakdown is checked for compliance</u>	—	—
<u>Surrogate recoveries checked and appropriately addressed</u>	—	—
<u>Method Preparation Blanks analyzed and meet acceptance criteria</u>	TB	SB
<u>MSs, MSDs, and/or MDs analyzed and calculations checked; applicable flags applied on QC reports; LCSs analyzed and meet acceptance criteria when performed</u>	TB	SB
<u>RLVS analyzed</u>	TB	SB
<u>Preparation and analysis hold times met</u>	TB	SB
<u>Preparation deviations and re-preparations noted when performed</u>	TB	SB
<u>Analysis deviations and re-analyses noted when performed</u>	TB	SB
<u>Sample dilution factors noted on reports</u>	TB	SB
<u>Electronic records in HBN transcription accuracy and completeness checked</u>	TB	SB
<u>Preparation and analysis calculations checked</u>	TB	SB
<u>NCRs are completed as necessary NC/CAR# _____</u>	—	—
<u>Report forms are complete and accurate</u>	TB	SB
<u>Manual integrations checked</u>	TB	SB



STANDARD REPORT

Working Standard - CLO4 WRK

CLO4 WRK		Description - 6850 WKG Std 100 ug/L			
Standard: 43702		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 WRK	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	0.1 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43701	CLO4 INT	6850 Intermdt AccStd 10 ug/mL	CLO4 INT	0.1 mL	09/18/2019



STANDARD REPORT

Constituent

Stock Standard - CLO4 STOCK

CLO4 STOCK		Description - 6850 Stock AccStd 1,000ug/mL	
Standard: 43659		Created By: Thomas Bosch	
MFG: AccuStandard		Amount: 100 mL	
MFG Lot: 218065075		Create Date: 09/17/2018 09:09AM	
Part ID: IC-PER-10X-1		Expires: 07/25/2020	
		Usable: No	
		Lab Lot: CLO4 STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos.	Analyte	Name	Concentration
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Working Standard - CLO4 INT

CLO4 INT		Description - 6850 Intermdt AccStd 10.ug/mL			
Standard: 43701		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 09/18/2018 02:09PM		Expires: 09/18/2019	
MFG Lot: TNB: 09/18/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 INT	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
43659	CLO4 STOCK	6850 Stock AccStd 1,000ug/mL	CLO4 STOCK	0.1 mL	07/25/2020



STANDARD REPORT

Working Standard - CLO4 QC WRK

CLO4 QC WRK		Description: 6850 QC WKG STD 100ug/L			
Standard: 41831		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC WRK 100.ug/L	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	100 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
41830	CLO4 QC INT	6850 QC Intrmdt Std-QC 10ug/mL	CLO4 QC INT 10.ug/mL	0.1 mL	05/09/2019



STANDARD REPORT

Constituent

Working Standard - CLO4 QC INT

CLO4 QC INT		Description - 6850 QC Intrmdt Std-QC 10ug/mL			
Standard: 41830		Created By: Thomas Bosch		Amount: 10 mL	
MFG: ALS/SLC		Create Date: 05/09/2018 10:05AM		Expires: 05/09/2019	
MFG Lot: TNB: 05/09/2018				Usable: Yes	
Pipette ID: Not Provided				Lab Lot: CLO4 QC INT 10.ug/mL	
Pos.	Analyte	Name	Concentration		
1	14797-73-0	Perchlorate	10 ug/mL		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
109	ASTM H2O	ASTM Type II Water	LAB 109	9.9 mL	11/07/2025
36748	CLO4 QCSTOCK	6850 QC Stock STD 1,000ug/mL	CLO4 QC STOCK	0.1 mL	03/31/2020



STANDARD REPORT

Constituent

Solvent Standard - ASTM H2O

ASTM H2O		Description - ASTM Type II Water	
Standard: 109	Created By: ALS Support (Lims)	Amount: 1000 L	
MFG: DCL In House	Create Date: 10/06/2005 09:10AM	Expires: 11/07/2025	
MFG Lot: Not Provided		Usable: Yes	
Part ID: Not Provided		Lab Lot: LAB 109	
Pos:	Analyte:	Name:	Concentration:
Solvent - Analyte(s) not applicable			



STANDARD REPORT

Constituent

Stock Standard - CLO4 QCSTOCK

CLO4 QCSTOCK		Description: 6850 QC Stock STD 1,000ug/mL	
Standard: 36748	Created By: Thomas Bosch	Amount: 100 mL	
MFG: Ultra Scientific	Create Date: 05/11/2017 01:05PM	Expires: 03/31/2020	
MFG Lot: CP-0860		Usable: Yes	
Part ID: ICC-013		Lab Lot: CLO4 QC STOCK	
Pos.	Analyte	Name	Concentration
1	14797-73-0	Perchlorate	1000 ug/mL



STANDARD REPORT

Working Standard - CLO4ISTDWRK

CLO4ISTDWRK		Description - Perchlorate ISTD Wrk 1,000ug/L			
Standard: 43730	Created By: Thomas Bosch	Amount: 25 mL			
MFG: ALS/SLC	Create Date: 09/20/2018 09:09AM	Expires: 09/20/2019			
MFG Lot: TNB: 05/09/2018	Verified By: Thomas Bosch	Usable: Yes			
Pipette ID: Not Provided	Verify Date:	Lab Lot: CLO4ISTDWRK			
Pos.	Analyte	Name	Concentration		
1	14797-73-0-8385	Perchlorate 83:85 Ratio	1000 ug/L		
2	14797-73-0-89	Perchlorate 89	1000 ug/L		
Composition					
Standard	Standard ID	Description	Lab Lot ID	Volume	Expires
43729	CLO4ISTDSTK	Perchlorate ISTD Stock	CLO4ISTDSTK	0.25 mL	04/28/2026



STANDARD REPORT

Constituent

Stock Standard - CLO4ISTDSTK

CLO4ISTDSTK		Description - Perchlorate ISTD Stock	
Standard: 43729	Created By: Thomas Bosch	Amount: 1 mL	
MFG: Cambridge Isotope	Create Date: 09/20/2018 09:09AM	Expires: 04/28/2026	
MFG Lot: SDFE-012A	Verified By: Thomas Bosch	Usable: Yes	
Part ID: OLM-7310-S	Verify Date:	Lab Lot: CLO4ISTDSTK	
Pos.	Analyte	Name	Concentration
1	14797-73-0-8385	Perchlorate 83:85 Ratio	100 ug/mL
2	14797-73-0-89	Perchlorate 89	100 ug/mL



Certificate of Analysis



ISO Guide 34 Reference Material

Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Product Name: Perchlorate IC Standard

Description:

This Reference Material (RM) was gravimetrically prepared in accordance with ISO Guide 34 and under ULTRA Scientific's ISO 9001 registered quality system. The neat materials used for this product have been verified by ULTRA's ISO 17025 laboratory and under ULTRA's ISO Guide 34 accreditation. The analyte concentrations were verified by ULTRA's ISO 17025 accredited laboratory. For each analyte, the true value, with its uncertainty value calculated at the 95% confidence level, is reported below.

Analyte	Starting Material	Lot Number	Purity (%)	Calculated Value	True Value	Traceability & Method
perchlorate	potassium perchlorate	RM07987	100	1001 ± 5 µg/mL	976 ± 6 µg/mL	NIST SRM 3141A; ICP-OES

Solvent: water (low TOC, < 50 ppb)

Storage: Store at Room Temperature (15° to 30°C).

Traceability:

Traceability has been established through an unbroken chain of comparisons, each having stated uncertainties. Comparisons are based on appropriate physical or chemical measurements, including gravimetric or volumetric dilution, where the mass or volume of a solution before and after dilution is measured. The balances used for these measurements are calibrated with weights traceable to NIST in compliance with ANSI/NCSL Z-540-1, ISO 9001, ISO 17025, and ISO Guide 34. Calibrated Class A glassware is used for volumetric measurements. Thermometers are calibrated against a NIST traceable thermometer in accordance with NIST Special Publication 819.

Estimation of Uncertainties:

The true value is reported, with its uncertainty value calculated at the 95% confidence level.

Homogeneity:

This RM was formulated and unitized according to an in-house procedure and is guaranteed to be homogeneous. There is no minimum sub-sample size required.

Intended Use:

This RM is intended for the preparation of working reference samples for use in routine laboratory analyses, calibration of instruments, validation of analytical methods, assessments of measurement methods and continuing calibration verification.

Instructions for Use:

Sample aliquots for analysis should be withdrawn at 20°C to 25°C immediately after opening and should be processed without delay for the true value to be valid within the stated uncertainties. Do not pipet from the bottle. Do not return any material removed for pipetting to the bottle. Tightly cap the bottle after removing any material and store according to the instructions noted above.

Hazards:

Refer to the Safety Data Sheet for information regarding this RM.

Expiration of Certification:

The certification of this RM is valid, within the measurement uncertainty specified, until the expiration date specified above, provided the RM is handled and stored in accordance with the instructions given in this certificate. This certification is nullified if the RM is damaged, contaminated, or otherwise modified.





Certificate of Analysis



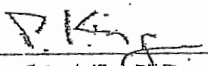
ISO Guide 34 Reference Material

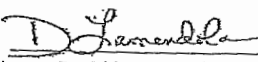
Product Number: ICC-013
Lot Number: CP-0860

Lot Issue Date: 29-Feb 2016
Expiration Date: 31-Mar 2020

Maintenance of Certification:

The real-time, long term stability of the RM may be monitored over the lifetime of the certification. If substantive changes occur that affect the certification before the expiration of this certificate, ULTRA Scientific will notify the purchaser.


Peter A. King, Ph.D.
VP, Technical Operations


Daniel J. Lamendola
Director of QAVRA



125 Market Street
New Haven, CT 06513
USA



Tel (203)786-5290
Fax (203)786-5287
www.AccuStandard.com

CERTIFICATE OF ANALYSIS



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43659

AccuTrace™ Reference Standard

Catalog No: IC-PER-10X-1
Description: Perchlorate Standard
Element: Perchlorate (ClO₄)
SRM: Ind. Std.
Lot: 218065075
Matrix: Water
Hazards: Refer to SDS for complete safety information

Date Certified: Jun 25, 2018
Expiration: Jul 25, 2020
Sample Size: 100 mL
Components: 1
Storage Condition: Ambient (>5 °C)
Included on ISO/IEC 17025 Scope of Accreditation: Yes
Included on ISO 17034 Scope of Accreditation: Yes



Signal Word: None

Component	SRM #	Prepared Concentration (µg/mL)
ClO ₄ Perchlorate	Ind. Std.	1000

The gravimetric uncertainty for this product is ±0.24%.

The final solution was checked against an independent standard to verify its concentration.

We use the highest purity raw materials available to minimize impurity levels in the final solution. Typically 99.999%+ pure starting materials are used as well as ASTM Type I 18 megohm deionized water.

All solutions are filtered through a 0.2 µm filter prior to being bottled.

All glassware used in preparation is Class A and calibrated regularly.

All weights are traceable through NIST, Test No. 822-275872-11

All bottles are triple rinsed with deionized water prior to use.

Shake bottle prior to use and do not pipette directly out of the bottle. Use only cleaned Class A volumetric glassware.

We certify the accuracy of this standard to be ±0.5% of the stated value until its expiration date provided it is kept tightly capped and stored under the conditions stated above.

Certified By:

Meigan O'Leary

Meigan O'Leary, Inorganic QC Manager

Cambridge Isotope Laboratories, Inc.

Certificate of Analysis

Quality Standards:
ISO Guide 34 • ISO/IEC 17025 • ISO 13485 • cGMP



23118

Product Name: PERCHLORIC ACID, SODIUM SALT
(Isotopic Label & Enrichment Specification) (18O4, 90%+) 100 UG/ML IN WATER

Lot Number: SDDG-013

Catalog Number: OLM-7310-S

Product Information

Chemical Purity Specification: $\geq 98\%$
Labeled CAS Number: NA
Unlabeled CAS Number: 7601-89-0
MW*: 130.4
Chemical Formula: NaCl*O4
Storage: Store at room temperature away from light and moisture.
Stability: See storage and expiration date.

Certification

Cambridge Isotope Laboratories, Inc. guarantees that this material meets or exceeds the specifications stated. Absolute identity as well as chemical and isotopic purities are assured by the use of unambiguous synthetic routes and multiple chemical analyses whenever possible. Results are representative of QC testing at time of release from Quality Control unless otherwise stated.

Volumetric measurements were made with Class A glassware. Gravimetry is traceable to the NIST through calibrated balances and certified, calibrated, standard weights. The calibrations are traceable to the NIST under Test No. 822/270236-04. The calibrations also meet specifications outlined in ISO 9001, ISO/IEC 17025, ANSI/NSCL Z540-1-1994, NCR Document 10CFR50 Appendix B, and applicable subdocuments.

This COA references the bulk catalog number before packaging. The COA also applies to the CIL finished good catalog number. Some possible packaging sizes and their corresponding suffix are -1.2, -1, -0.5, -10, or -0.1.

* For isotopically labeled compounds, MW listed is for the fully enriched product.

Approved by: T. J. Eckersley

Timothy J. Eckersley, Ph.D., Quality Assurance

Quality Control Tests and Results

QC Release Date	2/27/2014
Expiration Date	2/27/2024
Concentration Based on Gravimetry	102 $\mu\text{g/mL}$
Chemical Purity of Neat Material(s)	98%
LC/MS for Concentration	109.4 \pm 2.8 $\mu\text{g/mL}$ (k=2)



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount	
*	642096	CCV@25	Vial 71	1	Control	1	2.57589e6	8.017	25.04585
*	642100	QC@4.0	Vial 72	1	Control	2	4.23307e5	8.139	4.10749
*	642098	ICS@4.0	Vial 73	1	Control	3	3.17619e5	7.883	3.92708
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	7.71127e6	7.438	51.25151
*	1906112002	MS	Vial 77	1	Sample	7	8.02696e6	7.464	53.87972
*	1906112003	MSD	Vial 78	1	Sample	8	7.94242e6	7.442	53.69940
*	1906112004		Vial 79	1	Sample	9	2.33017e7	7.227	172.36522
*	1906112005		Vial 80	1	Sample	10	2.30991e7	7.229	180.38568
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.33611e6	8.114	25.01987
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	2.53126e6	7.736	226.48155
*	1906112005	10X	Vial 91	1	Sample	22	2.68237e6	7.739	241.03710
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	3.77363e6	8.090	3708.65666
*	642102	CCV@25	Vial 71	1	Control	25	2.15787e6	8.144	24.45533

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount	
*	642096	CCV@25	Vial 71	1	Control	1	6.79028e5	8.036	25.15984
*	642100	QC@4.0	Vial 72	1	Control	2	1.27412e5	8.156	4.45187
*	642098	ICS@4.0	Vial 73	1	Control	3	9.68721e4	7.906	4.29888
*	642099	LMB	Vial 74	1	Control	4	0.00000	0.000	0.00000
*	1905651001		Vial 75	1	Sample	5	0.00000	0.000	0.00000
*	1906112001		Vial 76	1	Sample	6	2.02786e6	7.452	51.10329
*	1906112002	MS	Vial 77	1	Sample	7	2.14637e6	7.478	54.46797
*	1906112003	MSD	Vial 78	1	Sample	8	2.10991e6	7.456	53.98100
*	1906112004		Vial 79	1	Sample	9	6.79668e6	7.243	180.45962
*	1906112005		Vial 80	1	Sample	10	6.82834e6	7.246	190.34382
*	1906112006		Vial 81	1	Sample	11	0.00000	0.000	0.00000
*	1906112007		Vial 82	1	Sample	12	0.00000	0.000	0.00000
*	1906112008		Vial 83	1	Sample	13	0.00000	0.000	0.00000
*	1906112009		Vial 84	1	Sample	14	0.00000	0.000	0.00000
*	1906112010		Vial 85	1	Sample	15	0.00000	0.000	0.00000
*	1906112011		Vial 86	1	Sample	16	0.00000	0.000	0.00000
*	642101	CCV@25	Vial 71	1	Control	17	6.28961e5	8.128	25.63410
*	1906332001		Vial 88	1	Sample	19	0.00000	0.000	0.00000
*	1906334001		Vial 89	1	Sample	20	0.00000	0.000	0.00000
*	1906112004	10X	Vial 90	1	Sample	21	6.46870e5	7.754	220.91905
*	1906112005	10X	Vial 91	1	Sample	22	6.95643e5	7.756	238.44674
*	1906112007	RE	Vial 82	1	Sample	23	0.00000	0.000	0.00000
*	1906330001	100	Vial 92	1	Sample	24	1.01021e6	8.102	3771.67475
*	642102	CCV@25	Vial 71	1	Control	25	5.83711e5	8.160	25.16652

Batch Report: C:\HPCHEM\1\DATA\05MAR19D\05MAR19S.B

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount	
*	642096	CCV@25	Vial 71	1	Control	1	3.12207e5	8.043	5.00000
*	642100	QC@4.0	Vial 72	1	Control	2	3.41038e5	8.157	5.00000
*	642098	ICS@4.0	Vial 73	1	Control	3	2.68237e5	7.902	5.00000
*	642099	LMB	Vial 74	1	Control	4	3.53313e5	8.102	5.00000
*	1905651001		Vial 75	1	Sample	5	3.26356e5	7.780	5.00000
*	1906112001		Vial 76	1	Sample	6	4.26473e5	7.464	5.00000
*	1906112002	MS	Vial 77	1	Sample	7	4.19549e5	7.491	5.00000
*	1906112003	MSD	Vial 78	1	Sample	8	4.16709e5	7.467	5.00000
*	1906112004		Vial 79	1	Sample	9	2.95705e5	7.253	5.00000
*	1906112005		Vial 80	1	Sample	10	2.75946e5	7.263	5.00000
*	1906112006		Vial 81	1	Sample	11	3.73575e5	7.480	5.00000
*	1906112007		Vial 82	1	Sample	12	5.00533e5	7.256	5.00000
*	1906112008		Vial 83	1	Sample	13	4.65121e5	7.237	5.00000
*	1906112009		Vial 84	1	Sample	14	2.43675e5	7.672	5.00000
*	1906112010		Vial 85	1	Sample	15	2.51865e5	7.710	5.00000
*	1906112011		Vial 86	1	Sample	16	2.80792e5	7.876	5.00000
*	642101	CCV@25	Vial 71	1	Control	17	2.83460e5	8.140	5.00000
*	1906332001		Vial 88	1	Sample	19	2.64674e5	7.827	5.00000
*	1906334001		Vial 89	1	Sample	20	2.65662e5	7.808	5.00000
*	1906112004	10X	Vial 90	1	Sample	21	3.41608e5	7.756	50.00000
*	1906112005	10X	Vial 91	1	Sample	22	3.38724e5	7.764	50.00000
*	1906112007	RE	Vial 82	1	Sample	23	4.31135e5	7.332	5.00000
*	1906330001	100	Vial 92	1	Sample	24	2.98985e5	8.111	500.00000
*	642102	CCV@25	Vial 71	1	Control	25	2.68305e5	8.165	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

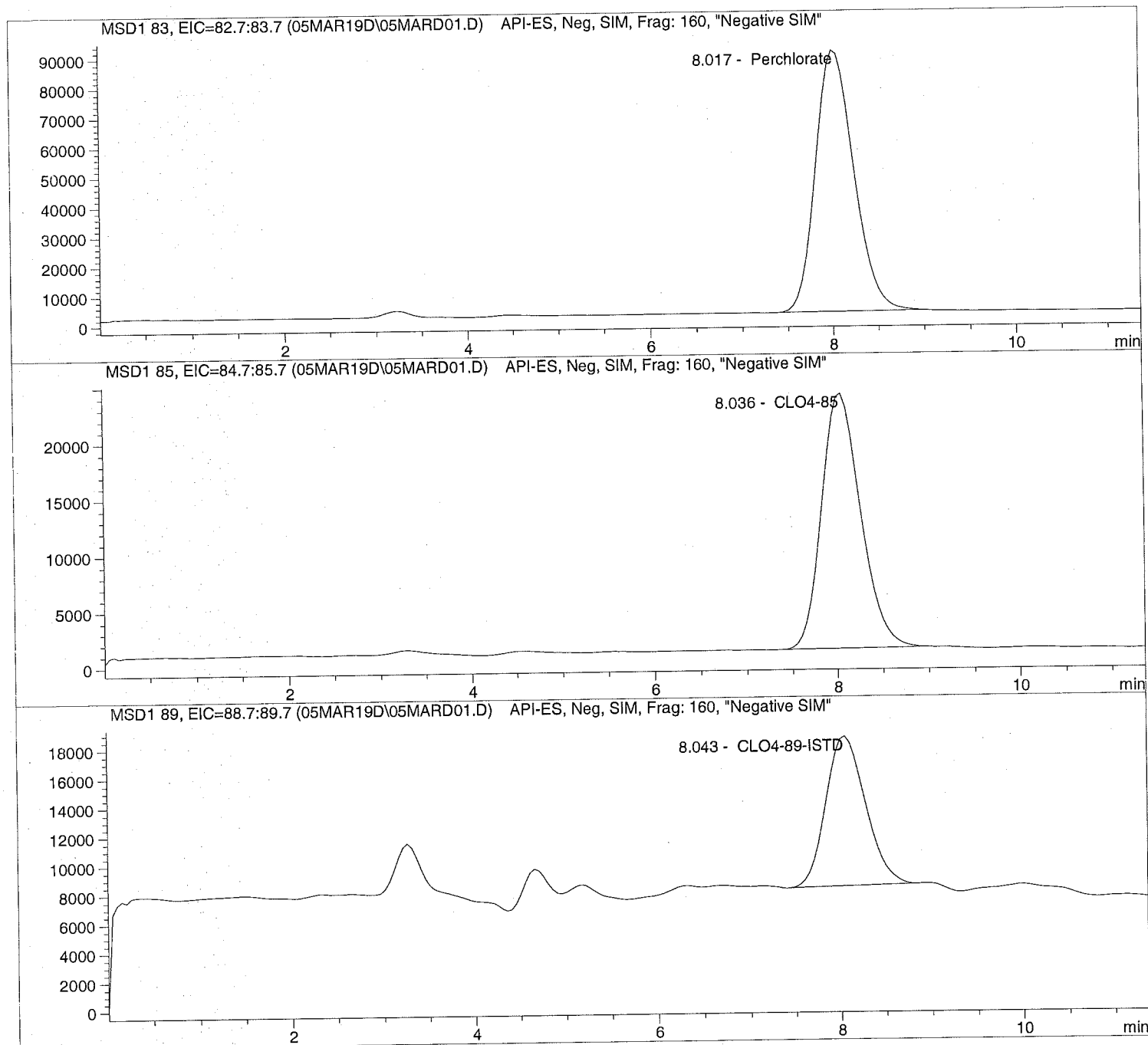
Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	642096	CCV@25	CLO4-AQN	1	Ctrl Samp	
2	Vial 72	642100	QC@4.0	CLO4-AQN	1	Ctrl Samp	
3	Vial 73	642098	ICS@4.0	CLO4-AQN	1	Ctrl Samp	
4	Vial 74	642099	LMB	CLO4-AQN	1	Ctrl Samp	
5	Vial 75	1905651001		CLO4-AQN	1	Sample	
6	Vial 76	1906112001		CLO4-AQN	1	Sample	
7	Vial 77	1906112002	MS	CLO4-AQN	1	Sample	
8	Vial 78	1906112003	MSD	CLO4-AQN	1	Sample	
9	Vial 79	1906112004		CLO4-AQN	1	Sample	
10	Vial 80	1906112005		CLO4-AQN	1	Sample	
11	Vial 81	1906112006		CLO4-AQN	1	Sample	
12	Vial 82	1906112007		CLO4-AQN	1	Sample	
13	Vial 83	1906112008		CLO4-AQN	1	Sample	
14	Vial 84	1906112009		CLO4-AQN	1	Sample	
15	Vial 85	1906112010		CLO4-AQN	1	Sample	
16	Vial 86	1906112011		CLO4-AQN	1	Sample	
17	Vial 71	642101	CCV@25	CLO4-AQN	1	Ctrl Samp	
18	Vial 87	1906330001	1K	CLO4-AQN	1	Sample	
19	Vial 88	1906332001		CLO4-AQN	1	Sample	
20	Vial 89	1906334001		CLO4-AQN	1	Sample	
21	Vial 90	1906112004	10X	CLO4-AQN	1	Sample	
22	Vial 91	1906112005	10X	CLO4-AQN	1	Sample	
23	Vial 82	1906112007	RE	CLO4-AQN	1	Sample	
24	Vial 92	1906330001	100	CLO4-AQN	1	Sample	
25	Vial 71	642102	CCV@25	CLO4-AQN	1	Ctrl Samp	

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD01.D Sample Name: 642096 CCV@25

=====
Injection Date: 3/05/2019 08:44:45 Seq Line: 1
Sample Name: 642096 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis
=====



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD01.D Sample Name: 642096 CCV@25

```

=====
Injection Date:  3/05/2019  08:44:45                    Seq Line:                    1
Sample Name:     642096    CCV@25                      Location:                    Vial 71
Acq Operator:    TNB                                    Inj. No.:                    1
                                                          Inj. Vol.:                    20 µl
=====

```

```

Acq. Method:     CLO4-AQN.M
Analysis Method:  C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:    2/19/2019  12:13:46
=====

```

Perchlorate analysis

Sample Information

```

=====
Sorted By:                    Signal
Calib. Data Modified:        Tue, 19. Feb. 2019,09:07:33 am
Multiplier:                  1.000000
Dilution:                    1.000000
Sample Amount:                25.000
=====

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.017	PBA	2575886.3	25.0459	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.036	PBA	679028.4	25.1598	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.043	PBA	312206.9	5.0000	CLO4-89-ISTD

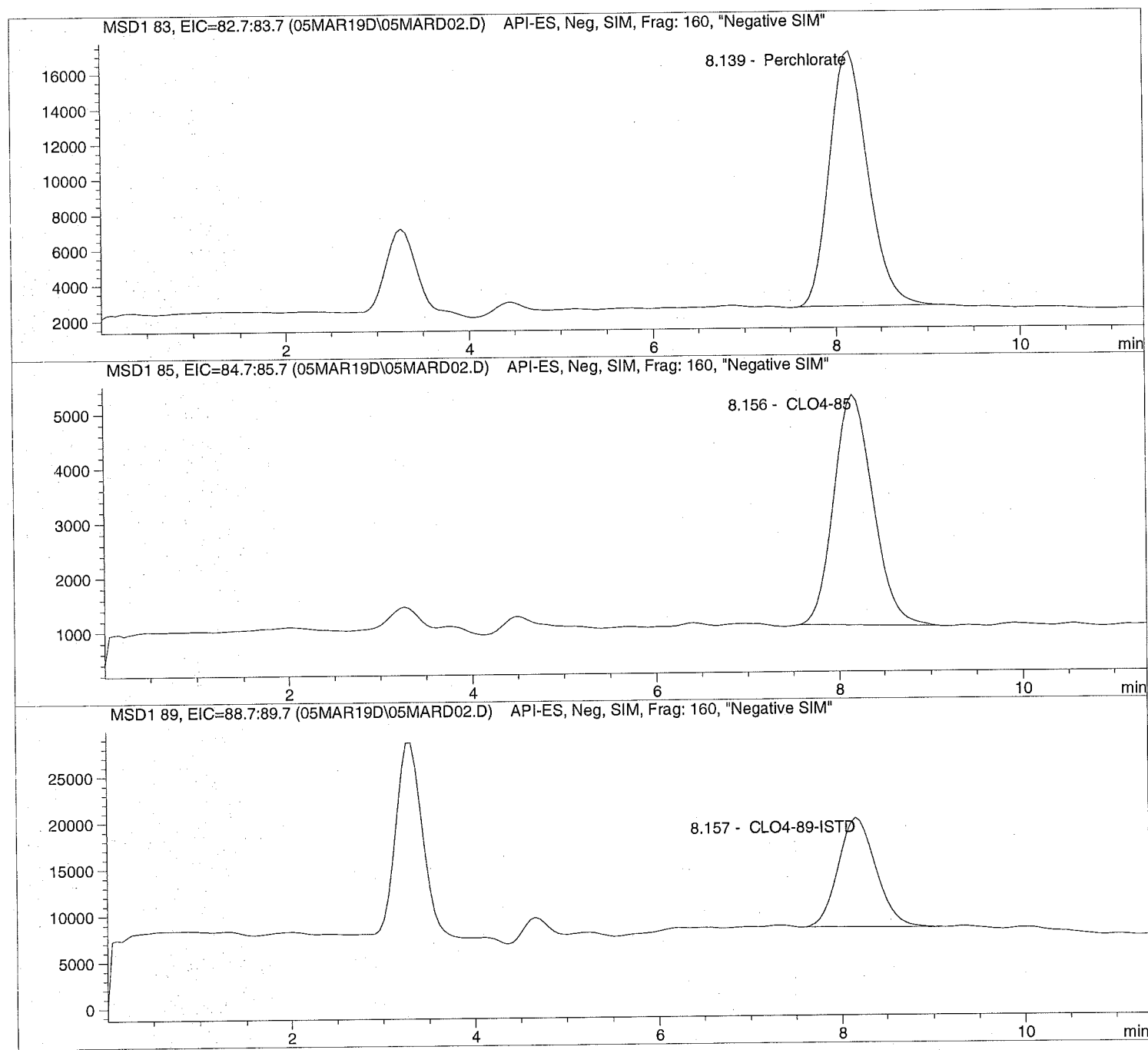
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD02.D Sample Name: 642100 QC@4.0

```
=====
Injection Date: 3/05/2019 09:00:30      Seq Line: 2
Sample Name:    642100 QC@4.0           Location: Vial 72
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD02.D Sample Name: 642100 QC@4.0

=====
 Injection Date: 3/05/2019 09:00:30 Seq Line: 2
 Sample Name: 642100 QC@4.0 Location: Vial 72
 Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
 Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

=====
 Sample Information
 =====

Sorted By: Signal
 Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
 Multiplier: 1.000000
 Dilution: 1.000000
 Sample Amount: 4.000

=====
 LCMS Results
 =====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.139	PBA	423307.1	4.1075	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.156	PBA	127412.0	4.4519	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.157	PBA	341038.3	5.0000	CLO4-89-ISTD

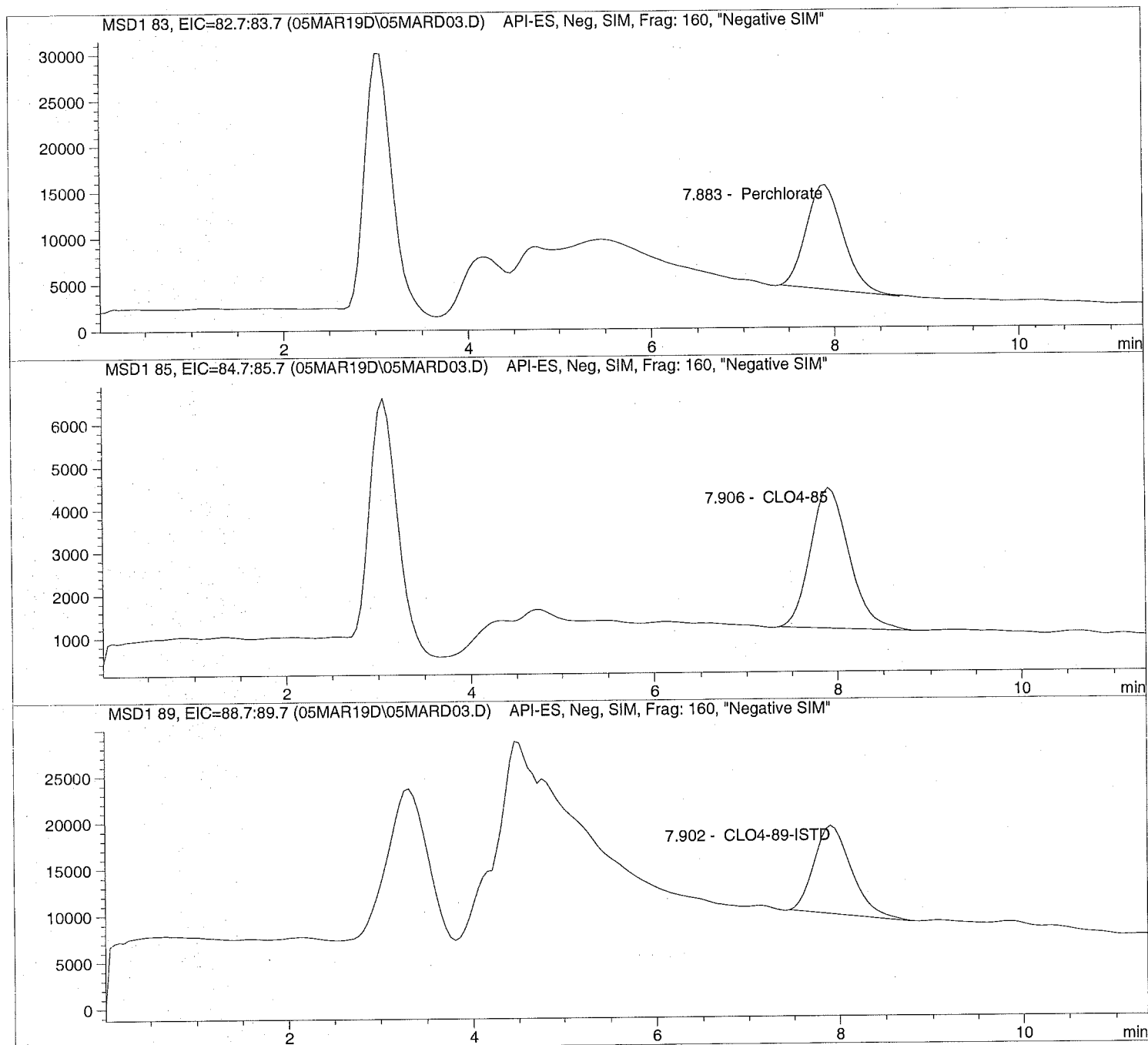
=====
 *** End of Report ***

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD03.D Sample Name: 642098 ICS@4.0

```
=====
Injection Date: 3/05/2019 09:13:34      Seq Line: 3
Sample Name: 642098 ICS@4.0             Location: Vial 73
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD03.D Sample Name: 642098 ICS@4.0

Injection Date: 3/05/2019 09:13:34 Seq Line: 3
 Sample Name: 642098 ICS@4.0 Location: Vial 73
 Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
 Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

Sample Information

Sorted By: Signal
 Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
 Multiplier: 1.000000
 Dilution: 1.000000
 Sample Amount: 4.000

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.883	PBA	317618.9	3.9271	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.906	PBA	96872.1	4.2989	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.902	PBA	268236.7	5.0000	CLO4-89-ISTD

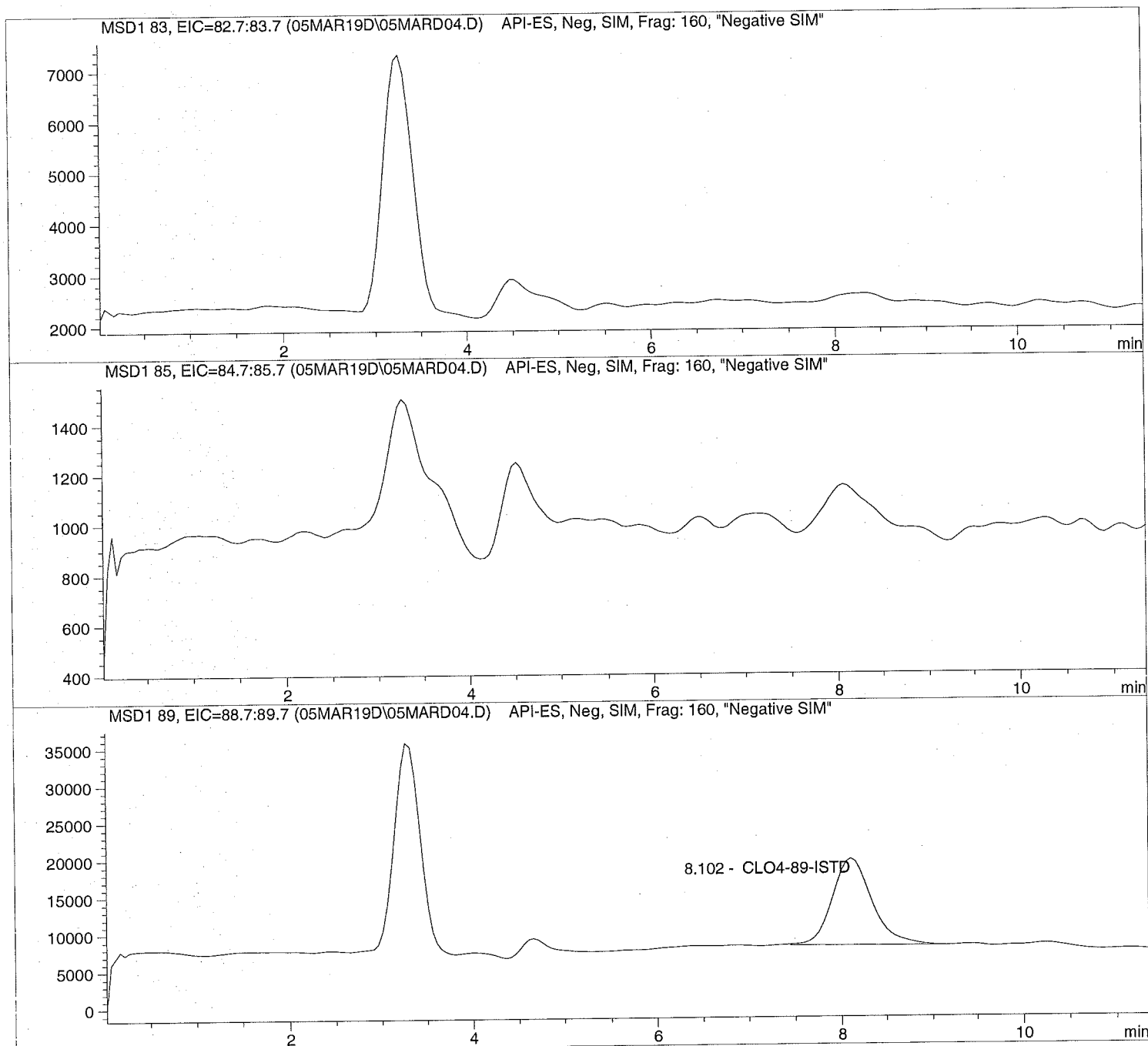
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD04.D Sample Name: 642099 LMB

```
=====
Injection Date: 3/05/2019 09:26:40      Seq Line: 4
Sample Name: 642099 LMB                  Location: Vial 74
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD04.D Sample Name: 642099 LMB

```

=====
Injection Date: 3/05/2019 09:26:40      Seq Line: 4
Sample Name: 642099 LMB                 Location: Vial 74
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	BBA	353313.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

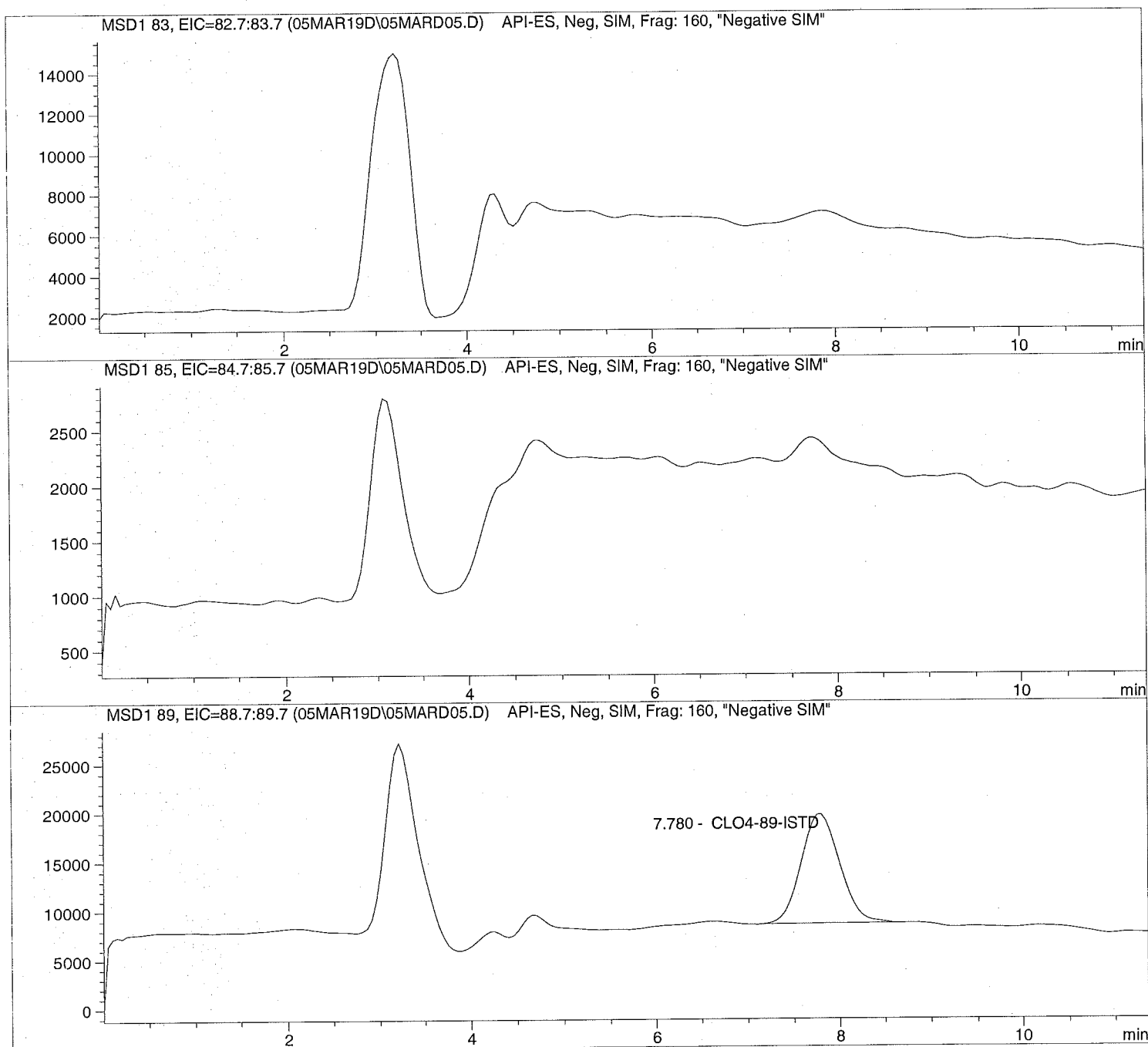
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD05.D Sample Name: 1905651001

```
=====
Injection Date: 3/05/2019 09:40:58      Seq Line:      5
Sample Name:    1905651001              Location:      Vial 75
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD05.D Sample Name: 1905651001

```

=====
Injection Date: 3/05/2019 09:40:58      Seq Line:          5
Sample Name:    1905651001              Location:         Vial 75
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.780	PBA	326356.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

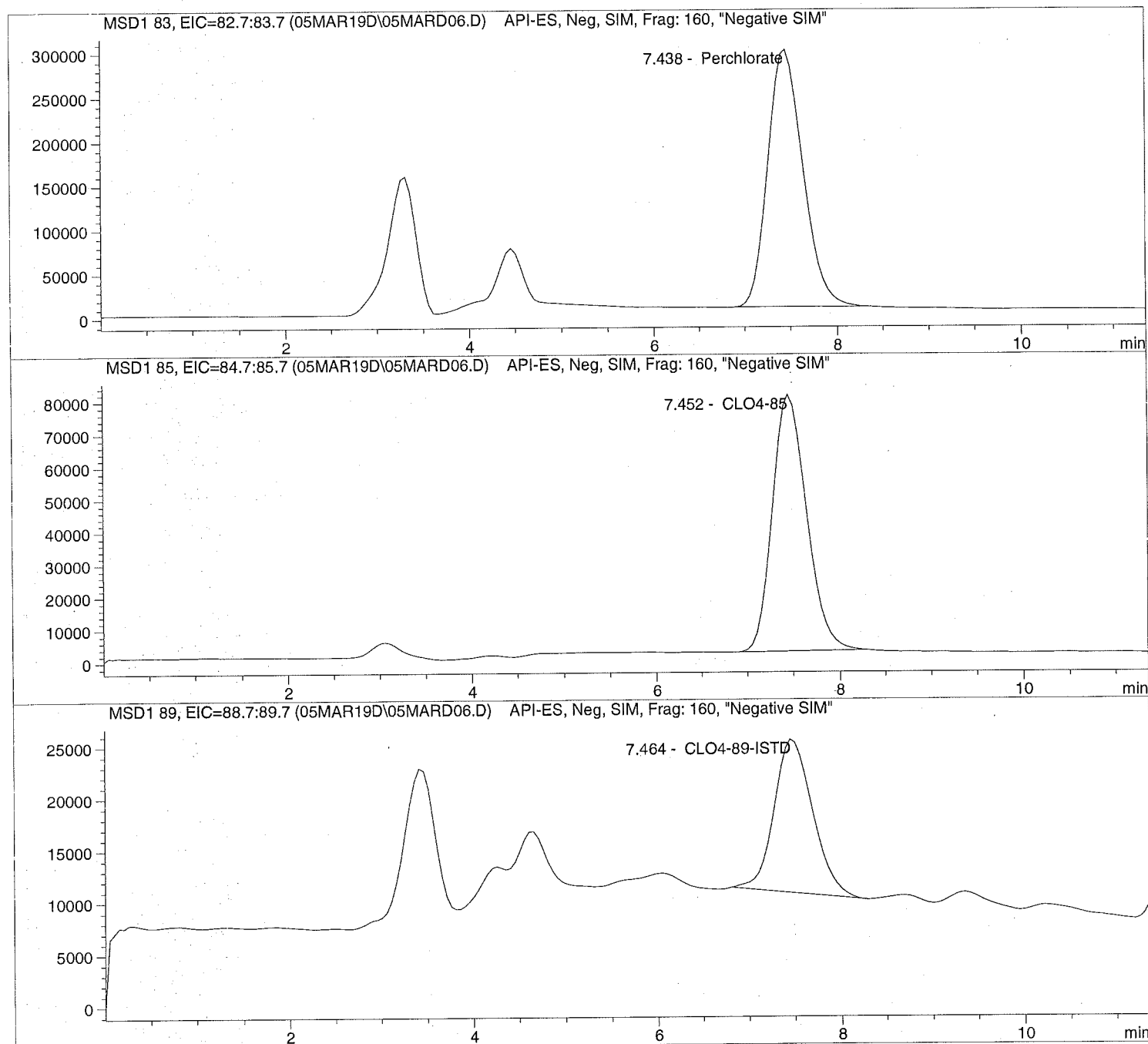
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD06.D Sample Name: 1906112001

=====
Injection Date: 3/05/2019 09:54:04 Seq Line: 6
Sample Name: 1906112001 Location: Vial 76
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD06.D Sample Name: 1906112001

```

=====
Injection Date: 3/05/2019 09:54:04      Seq Line: 6
Sample Name: 1906112001                Location: Vial 76
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.438	PBA	7711270.5	51.2515	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.452	PBA	2027855.1	51.1033	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PB	426473.5	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

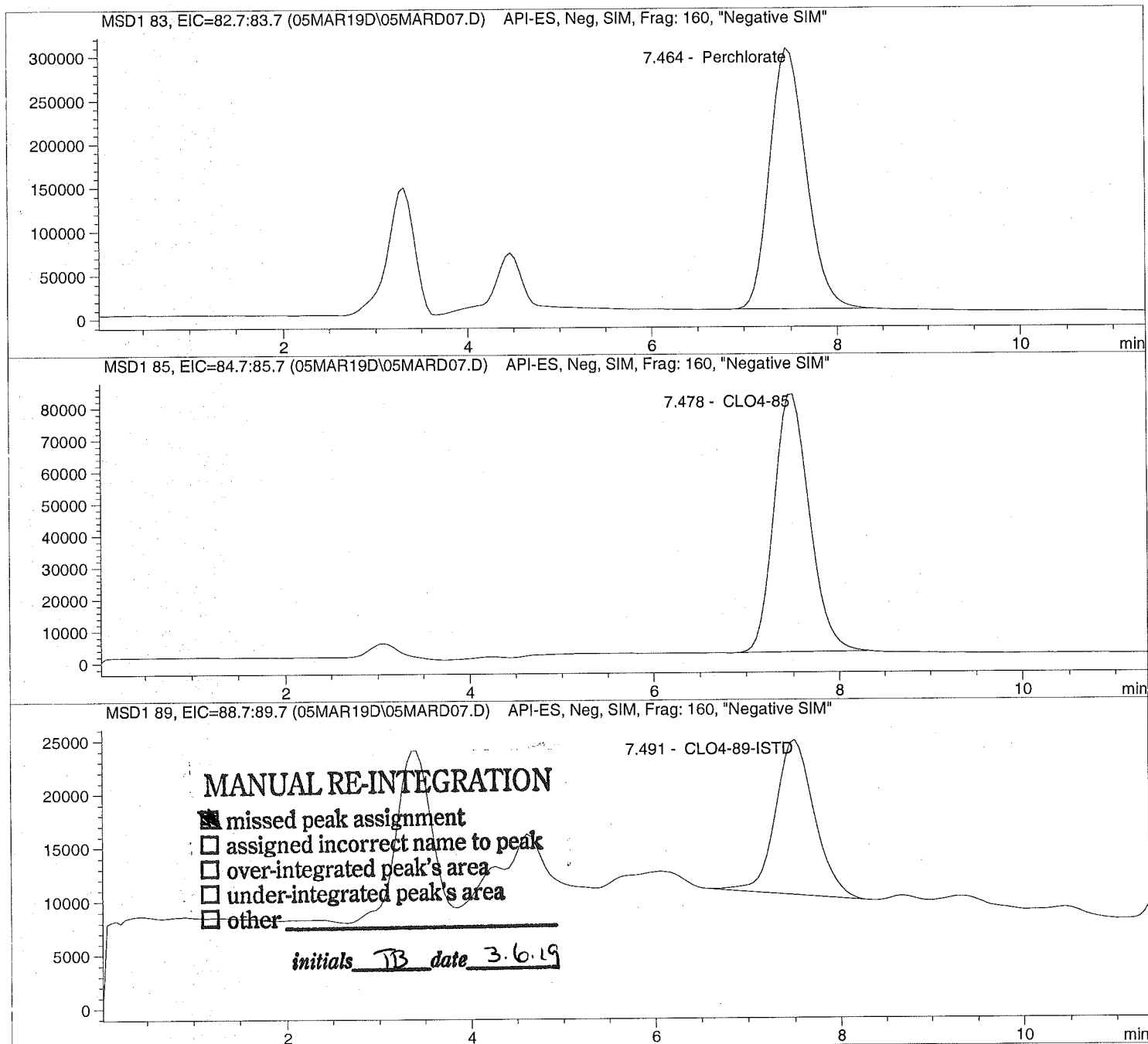
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

Injection Date: 3/05/2019 10:07:11 Seq Line: 7
Sample Name: 1906112002 MS Location: Vial 77
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

```

=====
Injection Date: 3/05/2019 10:07:11      Seq Line:          7
Sample Name:    1906112002 MS           Location:         Vial 77
Acq Operator:   TNB                    Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	53.8797	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	54.4680	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.491	MM	419549.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

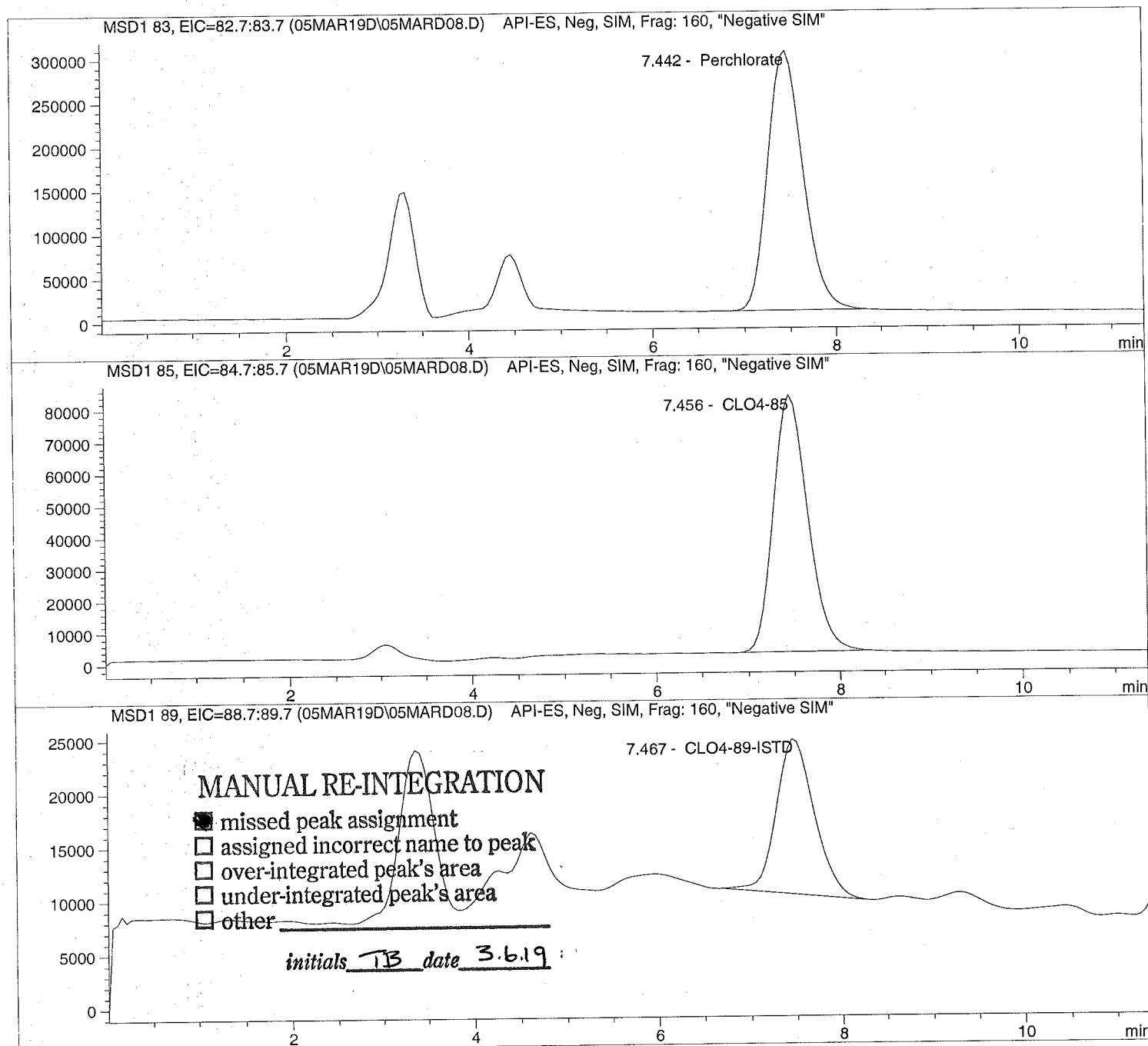
```


Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

```

=====
Injection Date: 3/05/2019 10:20:17      Seq Line:      8
Sample Name:   1906112003   MSD         Location:     Vial 78
Acq Operator:  TNB                Inj. No.:    1
                                           Inj. Vol.:   20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	53.6994	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	53.9810	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.467	MM	416709.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D Sample Name: 1906112004

```

=====
Injection Date: 3/05/2019 10:33:21      Seq Line:          9
Sample Name:    1906112004              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

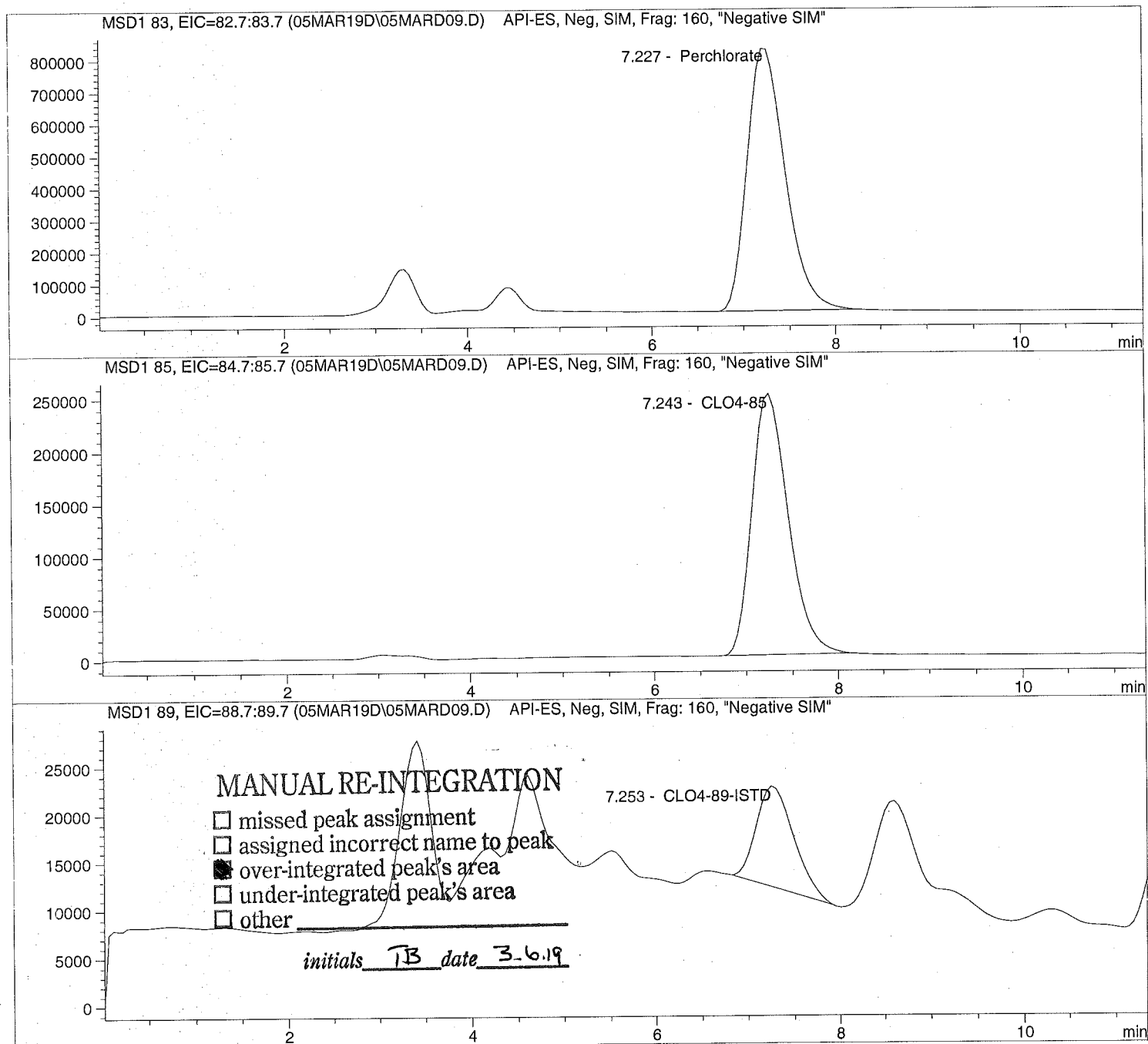
```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D Sample Name: 1906112004

```

=====
Injection Date: 3/05/2019 10:33:21      Seq Line:          9
Sample Name:    1906112004              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	172.3652	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	180.4596	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.253	MM	295705.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D Sample Name: 1906112005

```

=====
Injection Date:  3/05/2019  10:46:26      Seq Line:           10
Sample Name:    1906112005                Location:           Vial 80
Acq Operator:   TNB                       Inj. No.:          1
                                           Inj. Vol.:         20 µl
=====

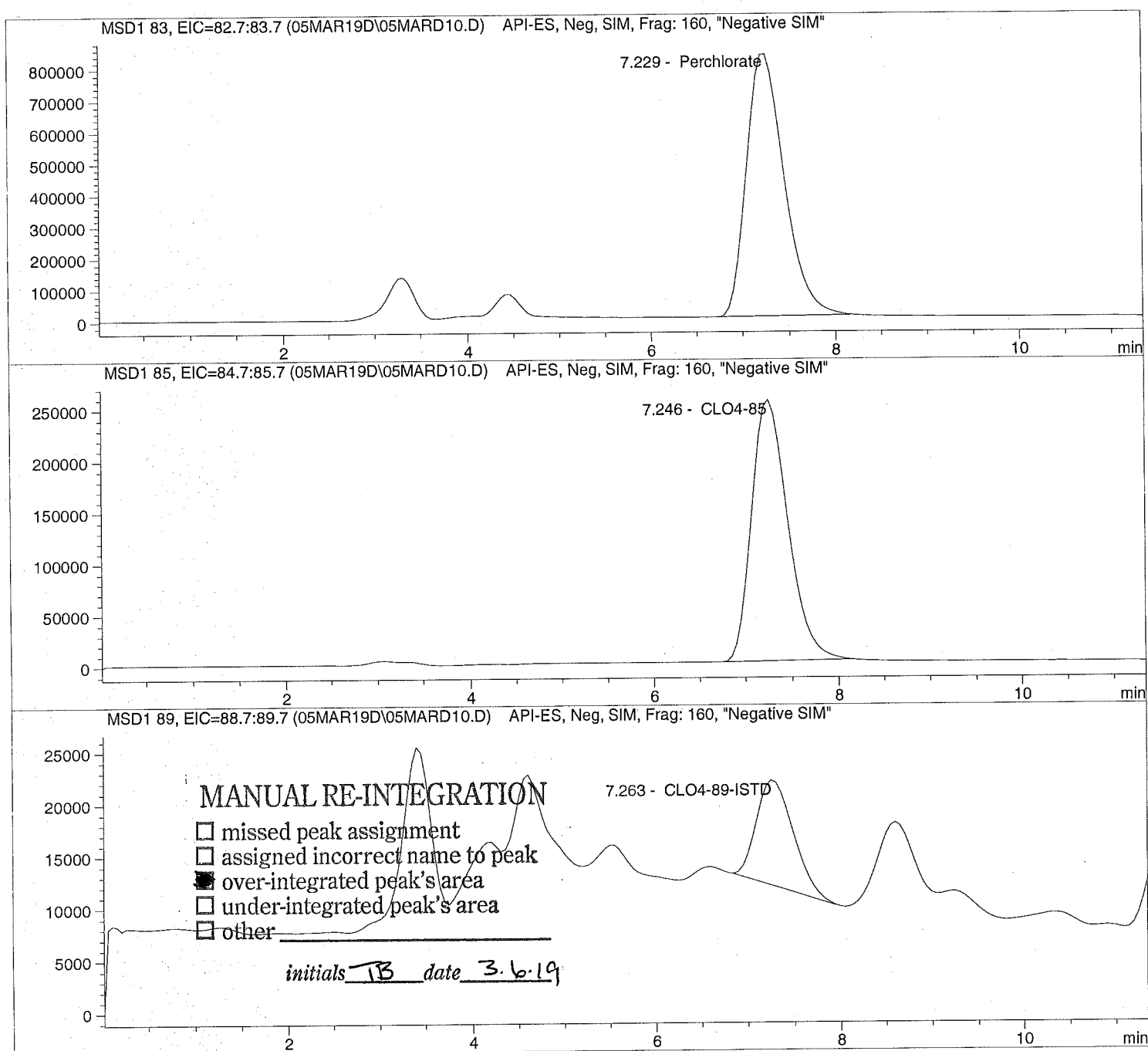
```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019  12:13:46
=====

```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D Sample Name: 1906112005

```

=====
Injection Date: 3/05/2019 10:46:26      Seq Line: 10
Sample Name: 1906112005                Location: Vial 80
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	180.3857	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	190.3438	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	MM	275946.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

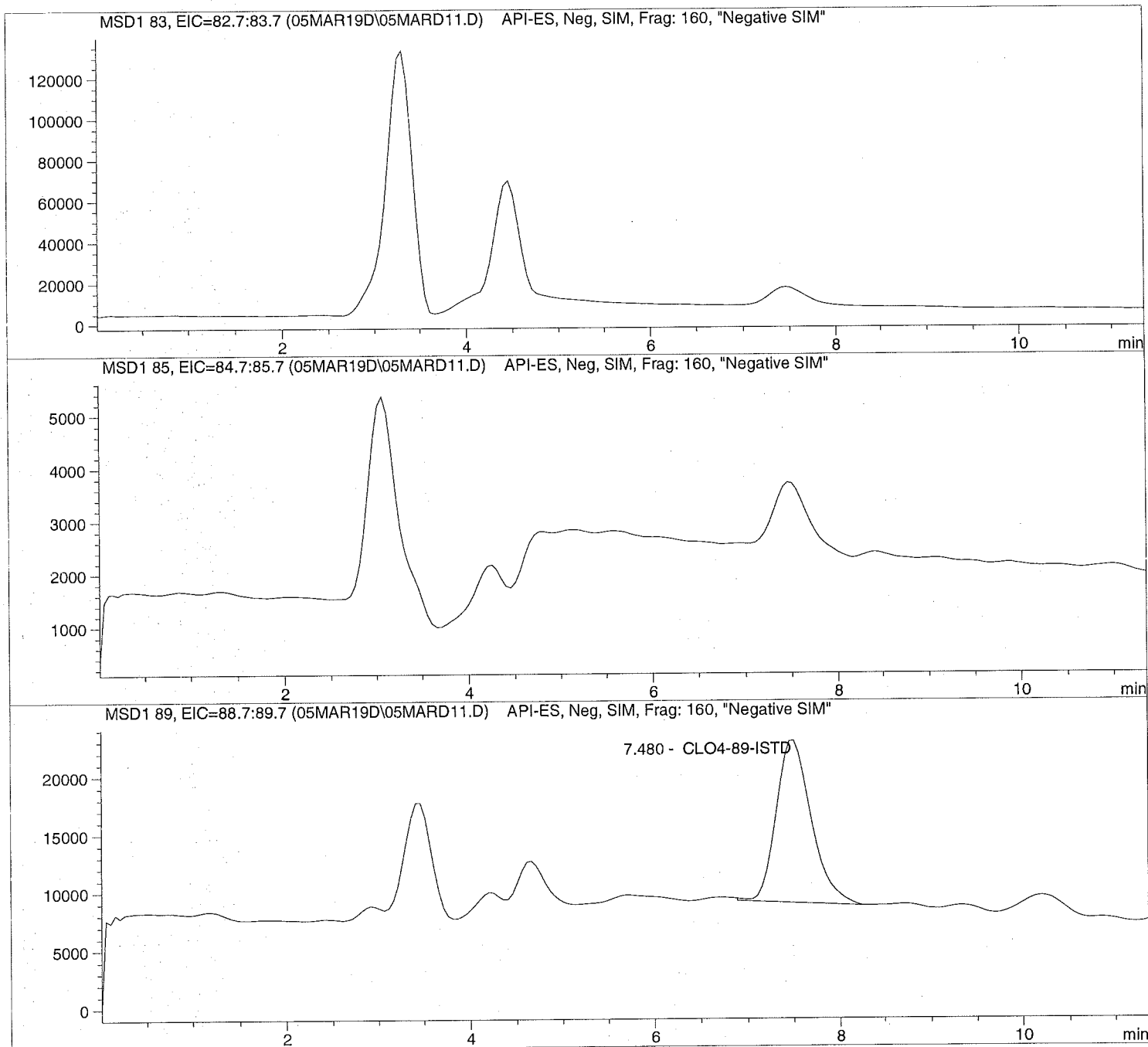
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD11.D Sample Name: 1906112006

```
=====
Injection Date: 3/05/2019 10:59:36      Seq Line:      11
Sample Name:    1906112006              Location:      Vial 81
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD11.D Sample Name: 1906112006

```

=====
Injection Date: 3/05/2019 10:59:36      Seq Line: 11
Sample Name: 1906112006                Location: Vial 81
Acq Operator: TNB                      Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.480	BBA	373575.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD12.D

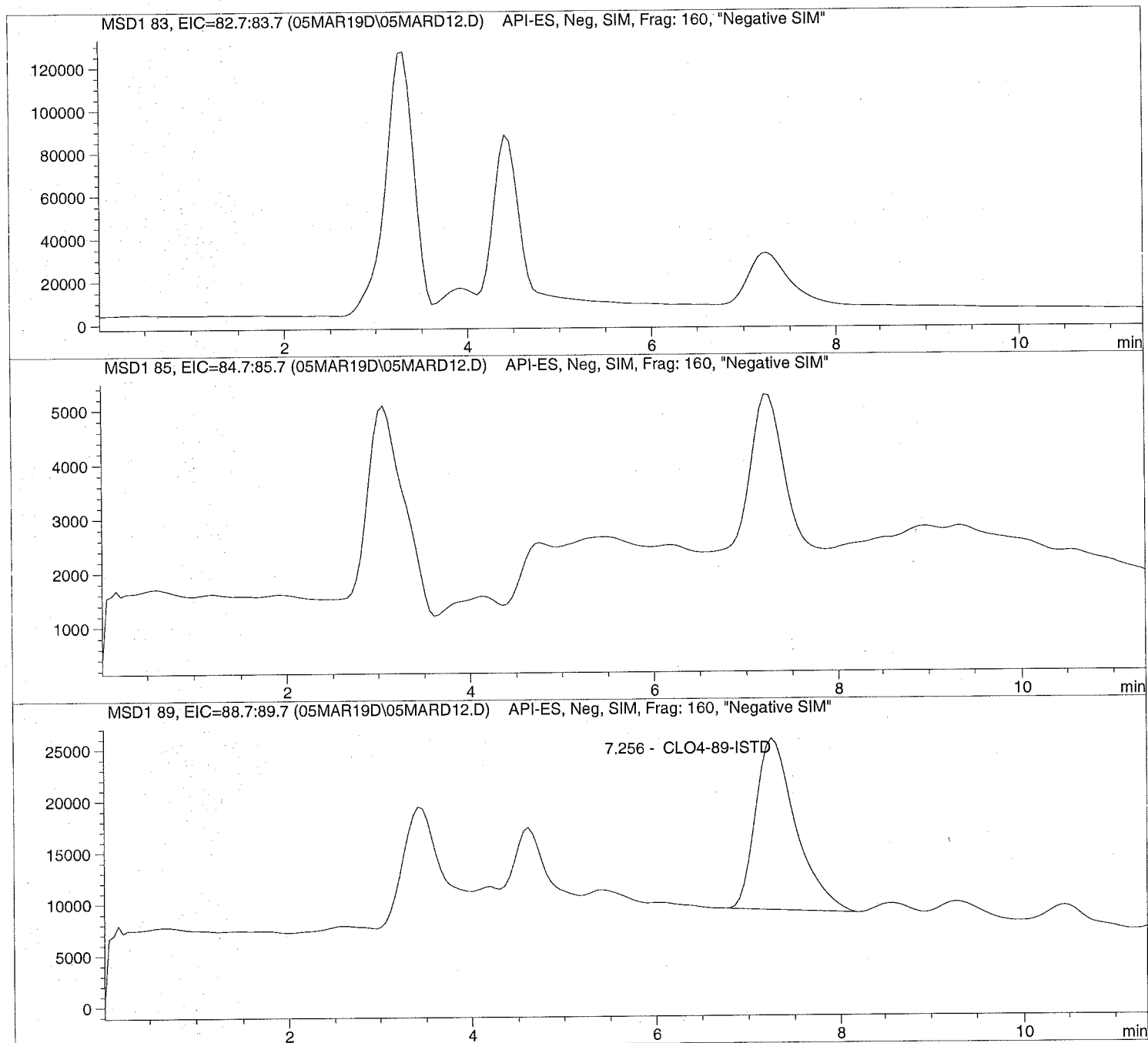
Sample Name: 1906112007

Injection Date: 3/05/2019 11:12:28
Sample Name: 1906112007
Acq Operator: TNB

Seq Line: 12
Location: Vial 82
Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD12.D Sample Name: 1906112007

```

=====
Injection Date: 3/05/2019 11:12:28      Seq Line: 12
Sample Name: 1906112007                 Location: Vial 82
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.256	PB	500532.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

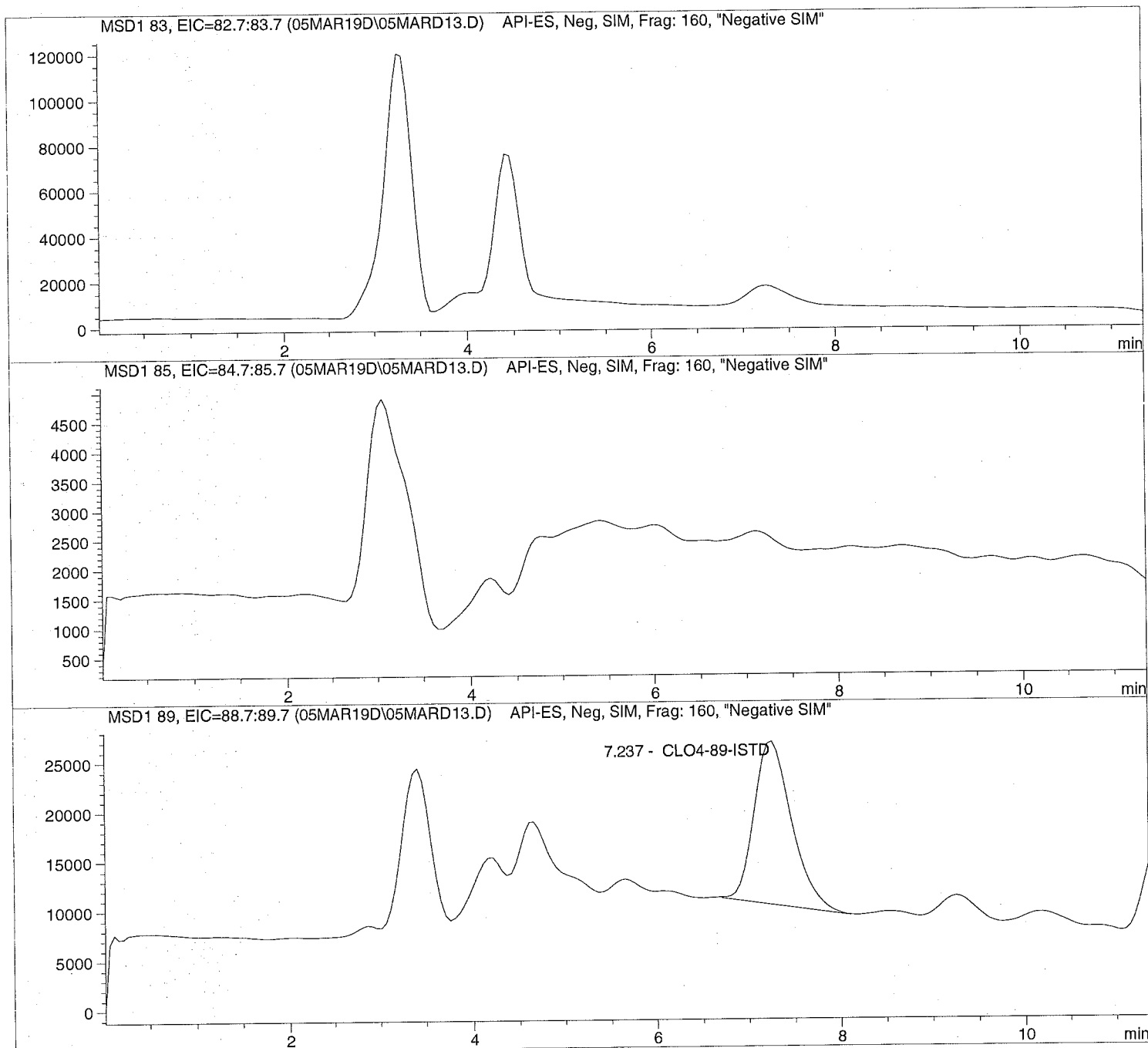
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD13.D Sample Name: 1906112008

```
=====
Injection Date: 3/05/2019 11:26:23      Seq Line:      13
Sample Name:    1906112008              Location:      Vial 83
Acq Operator:   TNB                    Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD13.D Sample Name: 1906112008

```

=====
Injection Date: 3/05/2019 11:26:23      Seq Line: 13
Sample Name: 1906112008                 Location: Vial 83
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.237	BBA	465121.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

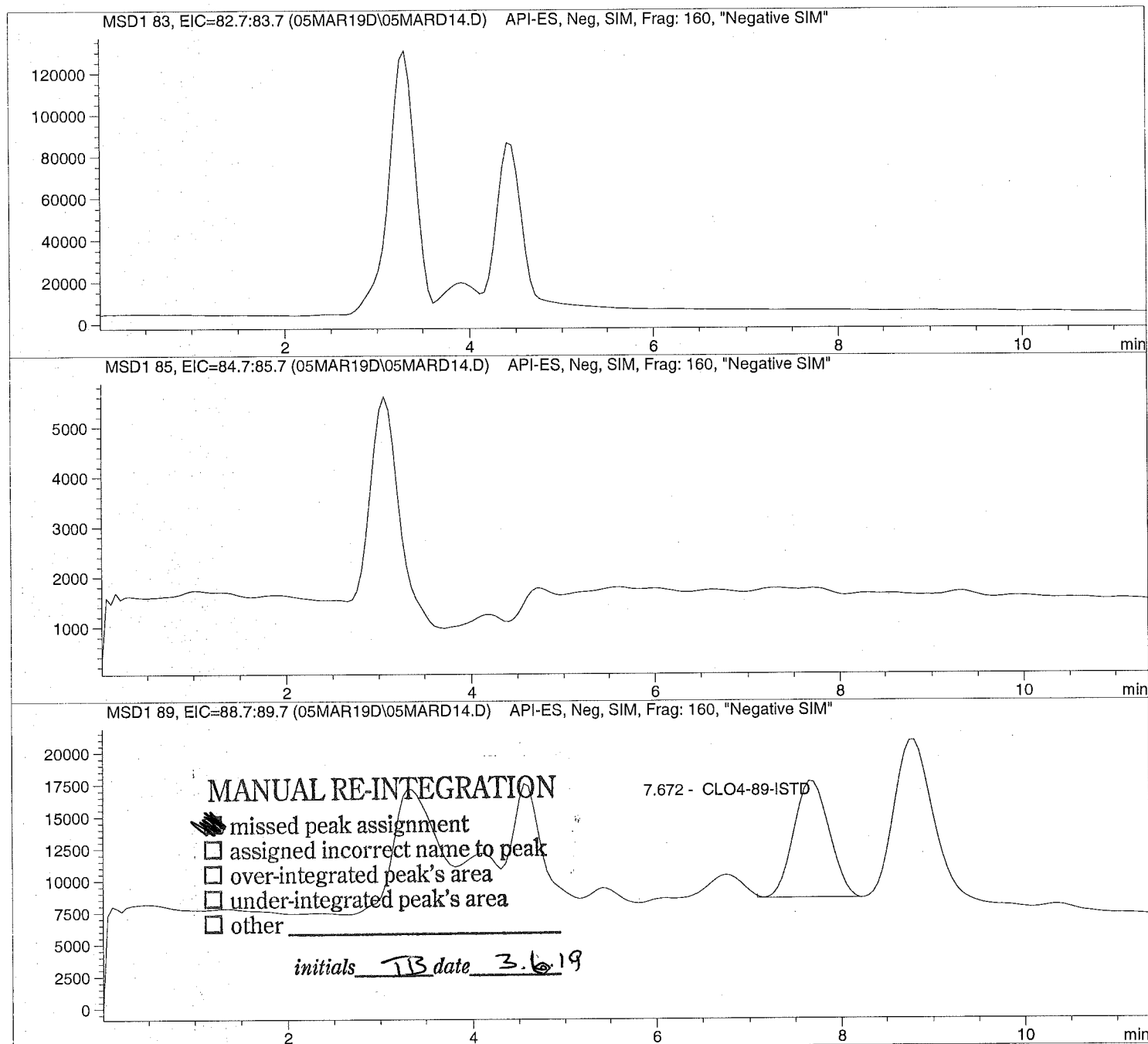
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D Sample Name: 1906112009

=====
 Injection Date: 3/05/2019 11:39:24 Seq Line: 14
 Sample Name: 1906112009 Location: Vial 84
 Acq Operator: TNB Inj. No.: 1
 Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
 Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
 Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D Sample Name: 1906112009

```

=====
Injection Date: 3/05/2019 11:39:24      Seq Line: 14
Sample Name: 1906112009                 Location: Vial 84
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

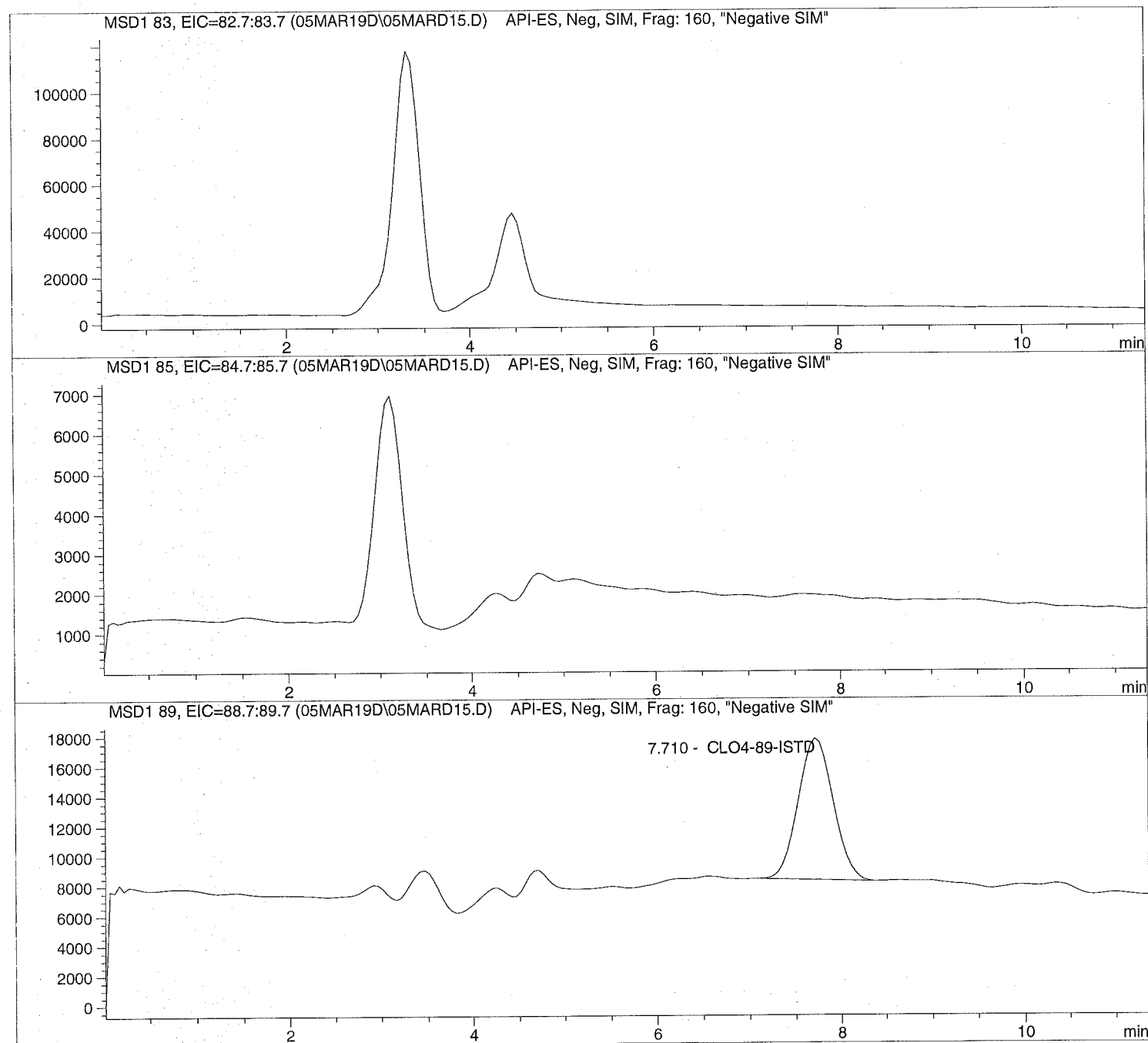
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD15.D Sample Name: 1906112010

```
=====
Injection Date: 3/05/2019 11:52:27      Seq Line: 15
Sample Name: 1906112010                 Location: Vial 85
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD15.D Sample Name: 1906112010

```

=====
Injection Date: 3/05/2019 11:52:27      Seq Line:          15
Sample Name:    1906112010              Location:          Vial 85
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.710	BBA	251865.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD16.D

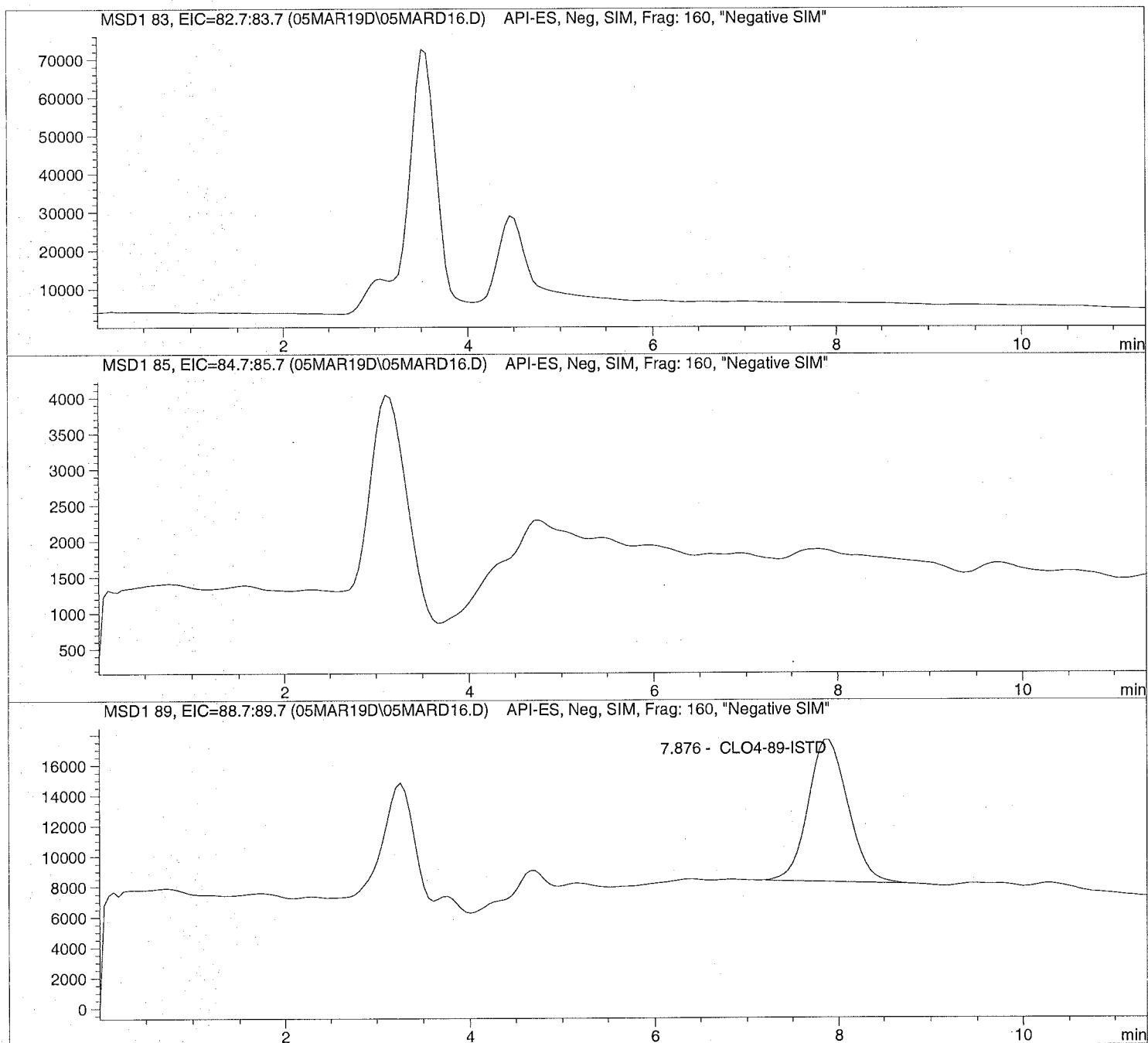
Sample Name: 1906112011

Injection Date: 3/05/2019 12:05:39
Sample Name: 1906112011
Acq Operator: TNB

Seq Line: 16
Location: Vial 86
Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD16.D Sample Name: 1906112011

```

=====
Injection Date: 3/05/2019 12:05:39      Seq Line:          16
Sample Name:    1906112011              Location:         Vial 86
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.876	BBA	280792.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

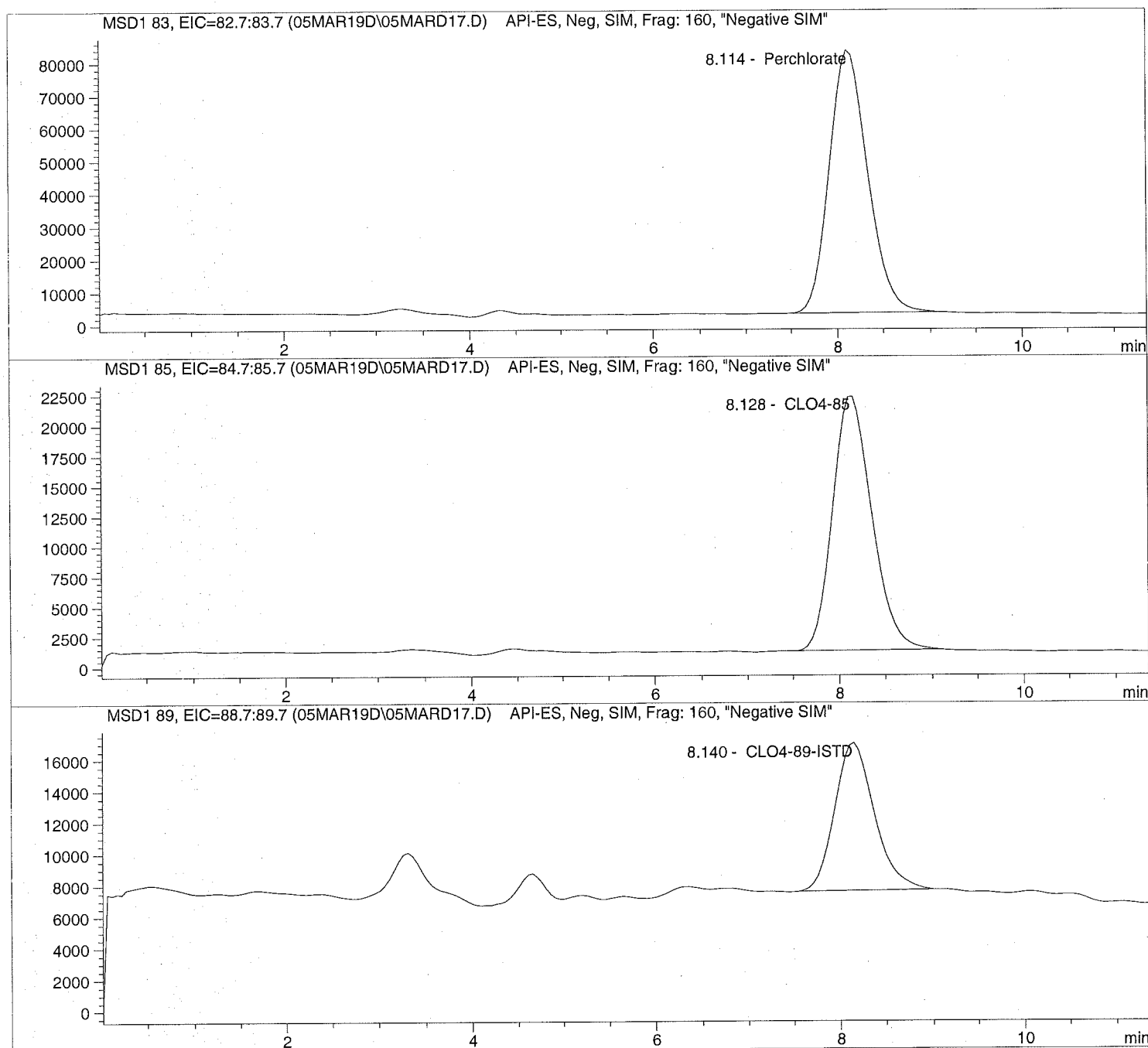
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD17.D Sample Name: 642101 CCV@25

```
=====
Injection Date: 3/05/2019 12:18:41      Seq Line:      17
Sample Name:    642101  CCV@25          Location:      Vial 71
Acq Operator:  TNB                    Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD17.D Sample Name: 642101 CCV@25

```

=====
Injection Date: 3/05/2019 12:18:41      Seq Line:          17
Sample Name:    642101  CCV@25          Location:          Vial 71
Acq Operator:   TNB                    Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	2336112.7	25.0199	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.128	PBA	628961.2	25.6341	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.140	PBA	283460.4	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD19.D

Sample Name: 1906332001

Injection Date: 3/05/2019 12:49:07

Seq Line: 19

Sample Name: 1906332001

Location: Vial 88

Acq Operator: TNB

Inj. No.: 1

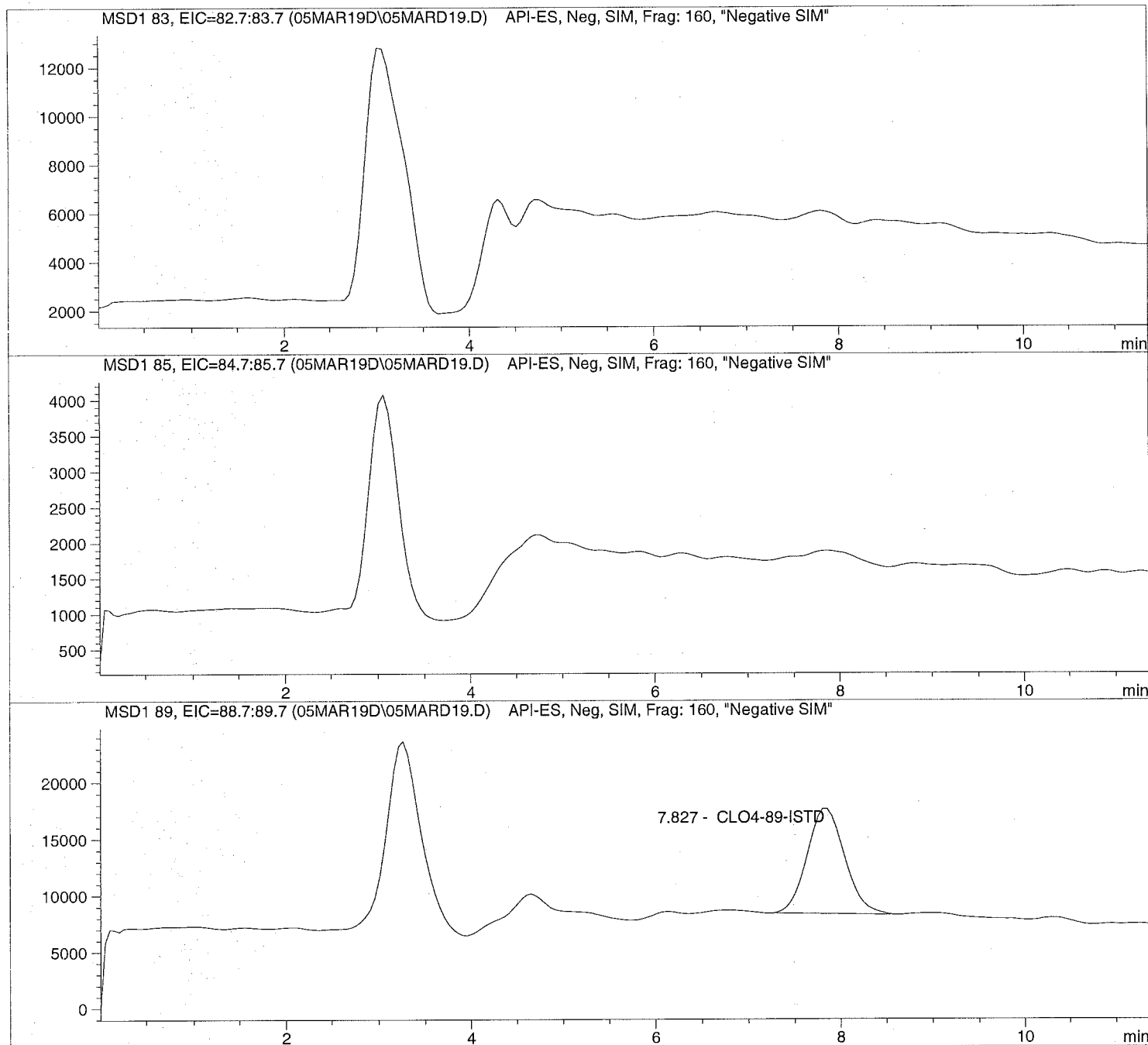
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M

Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD19.D Sample Name: 1906332001

```

=====
Injection Date: 3/05/2019 12:49:07      Seq Line: 19
Sample Name: 1906332001                  Location: Vial 88
Acq Operator: TNB                        Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.827	PBA	264674.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD20.D

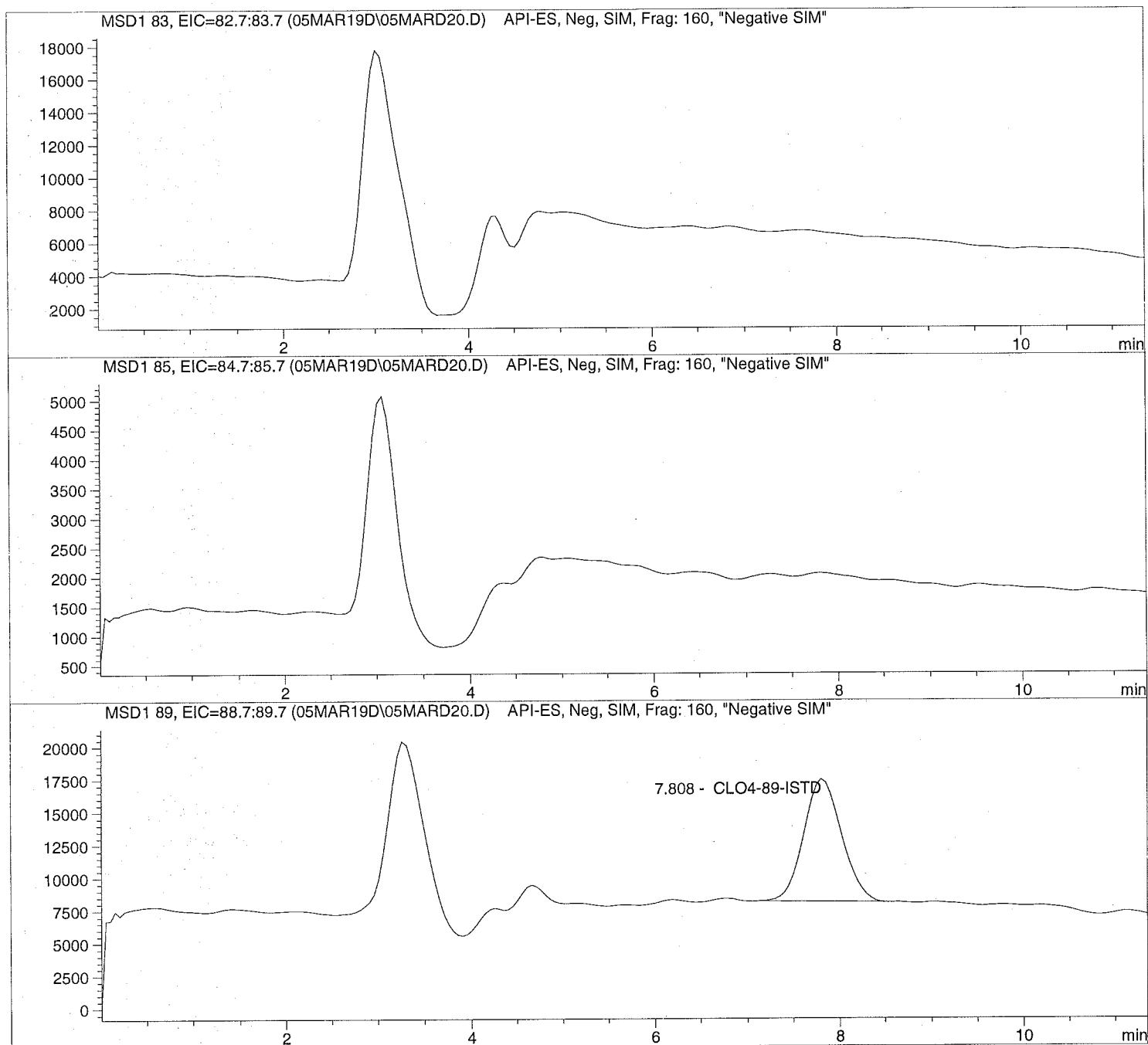
Sample Name: 1906334001

Injection Date: 3/05/2019 13:02:15
Sample Name: 1906334001
Acq Operator: TNB

Seq Line: 20
Location: Vial 89
Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD20.D Sample Name: 1906334001

```

=====
Injection Date: 3/05/2019 13:02:15      Seq Line:          20
Sample Name:    1906334001              Location:          Vial 89
Acq Operator:   TNB                      Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.808	PBA	265662.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

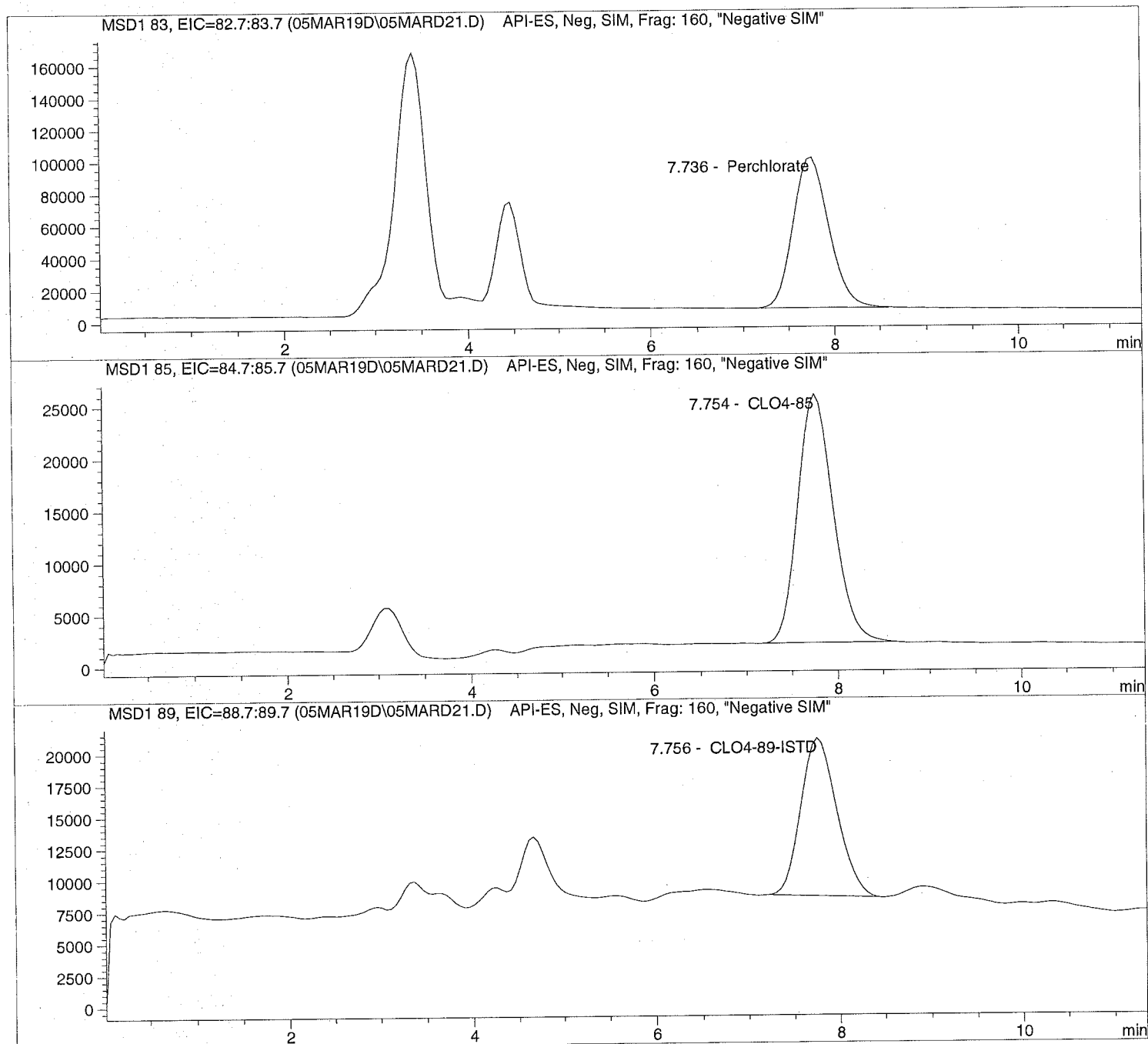
```


Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD21.D Sample Name: 1906112004 10X

```
=====
Injection Date: 3/05/2019 13:15:18      Seq Line:      21
Sample Name:    1906112004 10X          Location:      Vial 90
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:   20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD21.D Sample Name: 1906112004 10X

```

=====
Injection Date: 3/05/2019 13:15:18      Seq Line: 21
Sample Name: 1906112004 10X            Location: Vial 90
Acq Operator: TNB                      Inj. No.: 1
                                         Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 10.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.736	PBA	2531259.3	226.4815	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.754	PBA	646870.4	220.9191	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	PB	341607.9	50.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

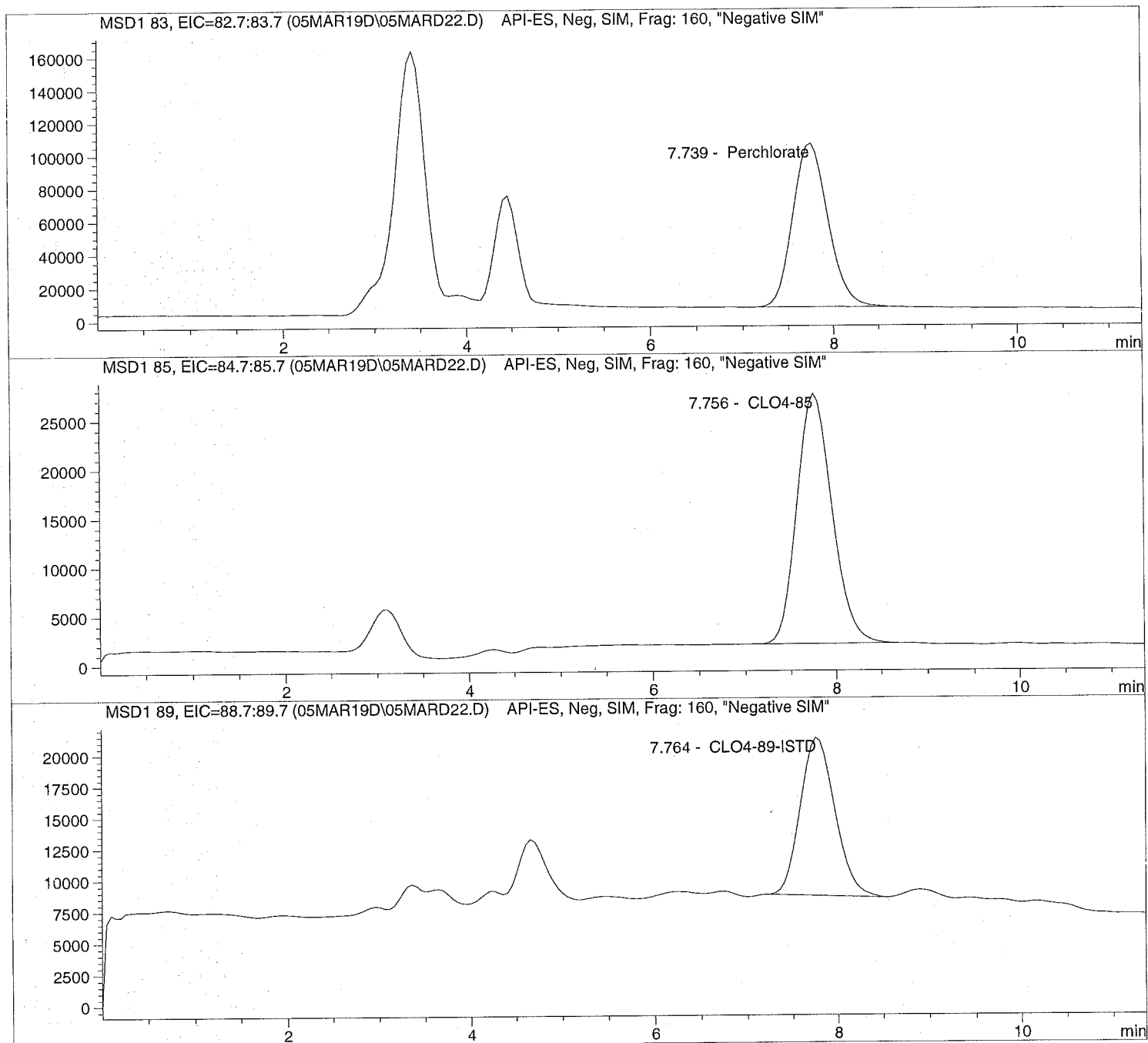
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD22.D Sample Name: 1906112005 10X

```
=====
Injection Date: 3/05/2019 13:28:32      Seq Line: 22
Sample Name:    1906112005 10X          Location:  Vial 91
Acq Operator:  TNB                      Inj. No.:  1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD22.D Sample Name: 1906112005 10X

```

=====
Injection Date: 3/05/2019 13:28:32      Seq Line:           22
Sample Name:    1906112005 10X          Location:           Vial 91
Acq Operator:   TNB                     Inj. No.:          1
                                           Inj. Vol.:         20 µl

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       10.000000
Sample Amount:  0.000

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.739	PBA	2682370.5	241.0371	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.756	BBA	695643.1	238.4467	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.764	PB	338724.1	50.0000	CLO4-89-ISTD

```

=====
*** End of Report ***

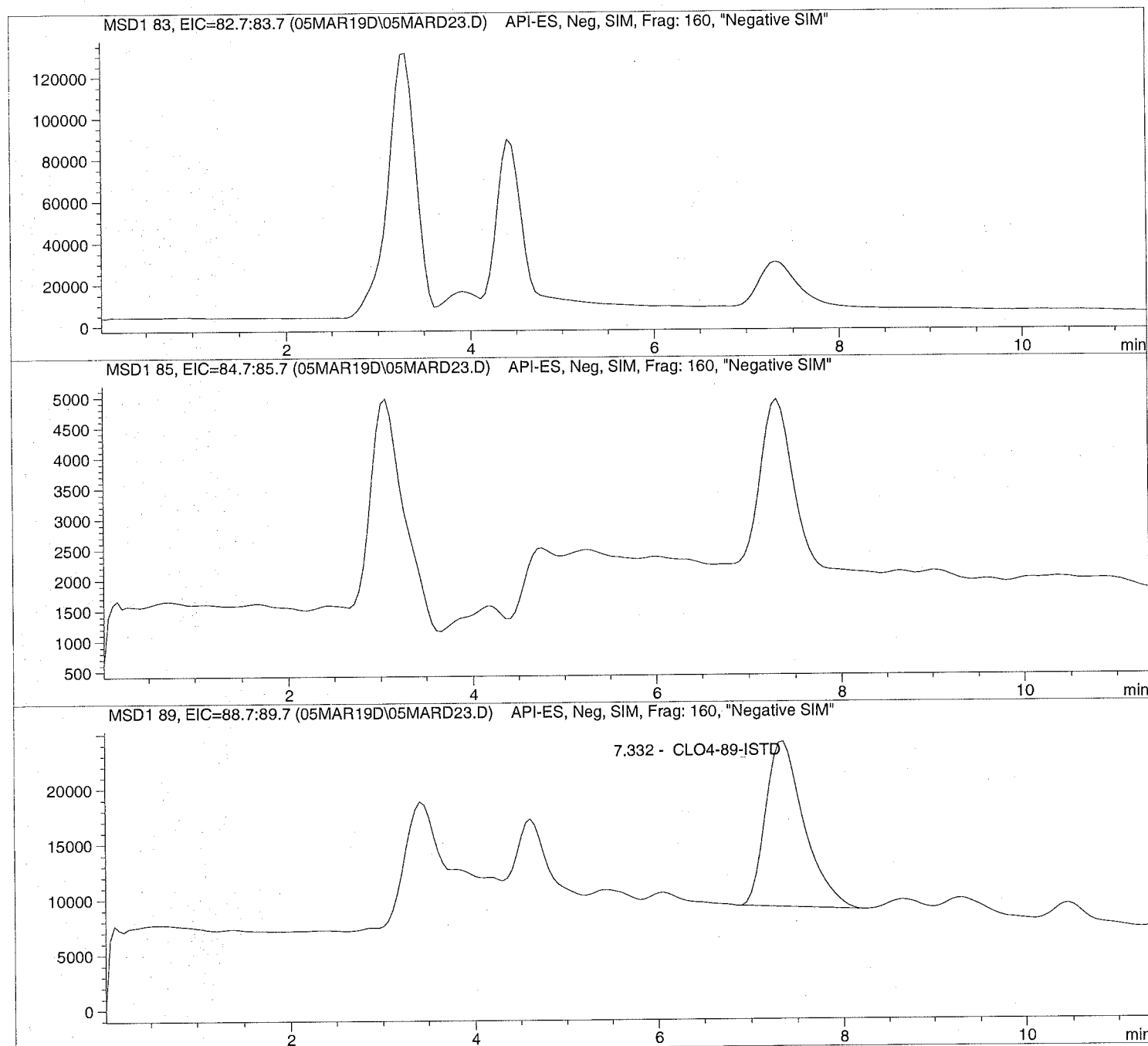
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD23.D Sample Name: 1906112007 RE

=====
Injection Date: 3/05/2019 13:41:34 Seq Line: 23
Sample Name: 1906112007 RE Location: Vial 82
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis
=====



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD23.D Sample Name: 1906112007 RE

```

=====
Injection Date: 3/05/2019 13:41:34      Seq Line:          23
Sample Name:    1906112007 RE           Location:          Vial 82
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.332	PB	431134.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

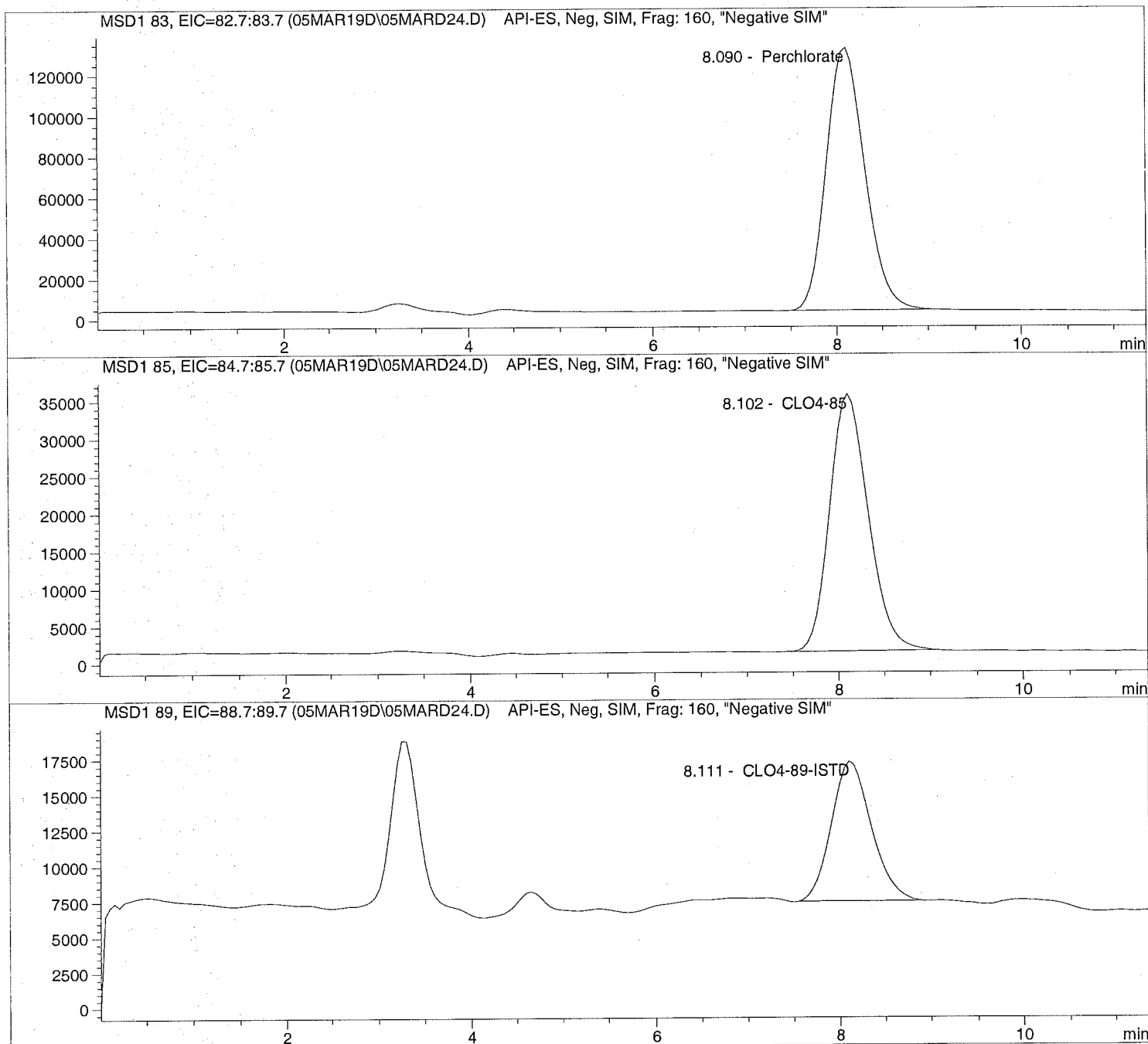
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD24.D Sample Name: 1906330001 100

```
=====
Injection Date: 3/05/2019 13:54:34      Seq Line:      24
Sample Name:    1906330001 100          Location:      Vial 92
Acq Operator:   TNB                    Inj. No.:     1
                                           Inj. Vol.:    20 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD24.D Sample Name: 1906330001 100

```

=====
Injection Date: 3/05/2019 13:54:34      Seq Line:          24
Sample Name:   1906330001 100          Location:         Vial 92
Acq Operator:  TNB                    Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      100.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.090	PBA	3773628.7	3708.6567	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.102	PBA	1010205.2	3771.6748	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.111	PBA	298984.6	500.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

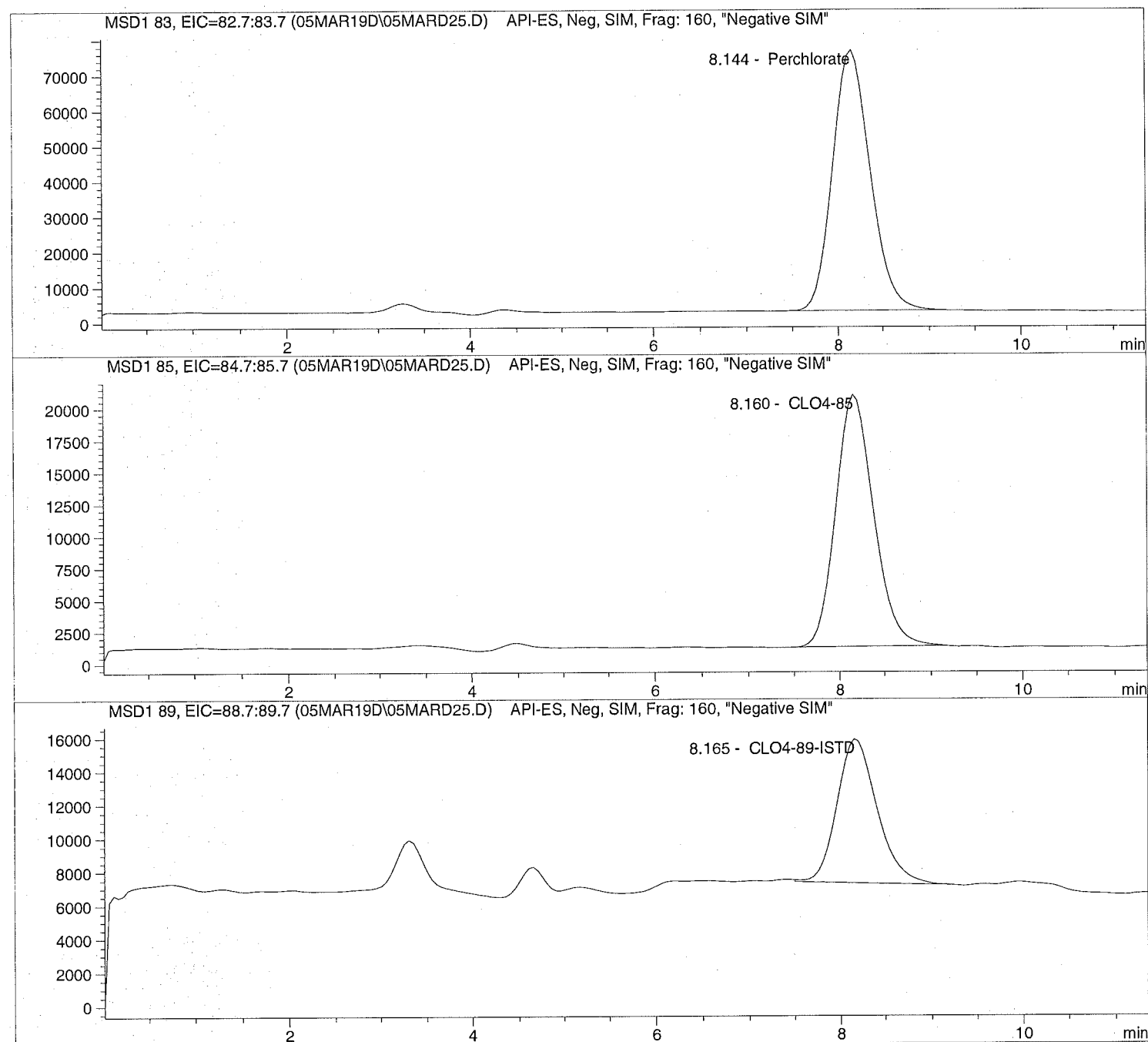
```


Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD25.D Sample Name: 642102 CCV@25

```
=====
Injection Date: 3/05/2019 14:07:36      Seq Line: 25
Sample Name:    642102  CCV@25          Location:  Vial 71
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method:  CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD25.D Sample Name: 642102 CCV@25

=====
Injection Date: 3/05/2019 14:07:36 Seq Line: 25
Sample Name: 642102 CCV@25 Location: Vial 71
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis

=====
Sample Information
=====

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 25.000

=====
LCMS Results
=====

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.144	PBA	2157871.5	24.4553	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.160	PBA	583711.5	25.1665	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.165	BBA	268305.4	5.0000	CLO4-89-ISTD

=====
*** End of Report ***



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

**Initial
Calibration**

Batch Review Method:

C:\HPCHEM\1\METHODS\CLO4-DP1.M

['#' ==> Run has not been reprocessed with Batch Review Method
 '*' ==> Run has been saved with batch file]

#*	Sample	Location	Inj	SampleType	Run	Perchlorate Area	Perchlorat RT	Perchlorate Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	8.94006e4	7.889	9.89924e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	1.97443e5	8.114	2.26028
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	4.79370e5	7.828	4.65688
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	9.30136e5	7.904	9.14998
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	2.81067e6	7.793	25.52636
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	5.66830e6	7.976	51.07439
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	8.69624e6	7.886	74.30603
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	1.01141e6	7.988	9.46019

#*	Sample	Location	Inj	SampleType	Run	CLO4-85 Area	CLO4-85 RT	CLO4-85 Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.26121e4	7.914	9.98836e-1
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	5.53134e4	8.127	2.11360
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	1.39247e5	7.842	4.91261
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	2.54396e5	7.923	9.39034
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	7.35969e5	7.811	25.48268
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	1.47152e6	7.993	50.35774
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	2.32809e6	7.900	74.72233
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	2.81230e5	8.007	9.87858

#*	Sample	Location	Inj	SampleType	Run	CLO4-89-ISTD Area	CLO4-89-IS RT	CLO4-89-ISTD Amount
#*	CLO4@ 1.0ug/L	Vial 73	1	Control	3	3.41443e5	7.900	5.00000
#*	CLO4@ 2.0ug/L	Vial 74	1	Control	4	2.99651e5	8.132	5.00000
#*	CLO4@ 5.0ug/L	Vial 75	1	Control	5	3.38646e5	7.853	5.00000
#*	CLO4@ 10.ug/L	Vial 76	1	Control	6	3.25154e5	7.925	5.00000
#*	CLO4@ 25.ug/L	Vial 77	1	Control	7	3.33799e5	7.819	5.00000
#*	CLO4@ 50.ug/L	Vial 78	1	Control	8	3.14712e5	7.999	5.00000
#*	CLO4@ 75.ug/L	Vial 79	1	Control	9	3.13909e5	7.908	5.00000
#*	ICAL Verf@10ug/L	Vial 80	1	Control	10	3.41503e5	8.005	5.00000

*** End of Report ***

Sequence Table:

Method and Injection Info Part:

Line	Location	SampleName	Method	Inj	SampleType	InjVolume	DataFile
====	=====	=====	=====	===	=====	=====	=====
1	Vial 71	CLO4@ .20ug/L	CLO4-AQN	1	Ctrl Samp		
2	Vial 72	CLO4@ 0.5ug/L	CLO4-AQN	1	Ctrl Samp		
3	Vial 73	CLO4@ 1.0ug/L	CLO4-AQN	1	Ctrl Samp		
4	Vial 74	CLO4@ 2.0ug/L	CLO4-AQN	1	Ctrl Samp		
5	Vial 75	CLO4@ 5.0ug/L	CLO4-AQN	1	Ctrl Samp		
6	Vial 76	CLO4@ 10.ug/L	CLO4-AQN	1	Ctrl Samp		
7	Vial 77	CLO4@ 25.ug/L	CLO4-AQN	1	Ctrl Samp		
8	Vial 78	CLO4@ 50.ug/L	CLO4-AQN	1	Ctrl Samp		
9	Vial 79	CLO4@ 75.ug/L	CLO4-AQN	1	Ctrl Samp		
10	Vial 80	ICAL Verf@10ug/L	CLO4-AQN	1	Ctrl Samp		

Data file: C:\HPCHEM\1\DATA\15FEB19T\15FEBI03.D

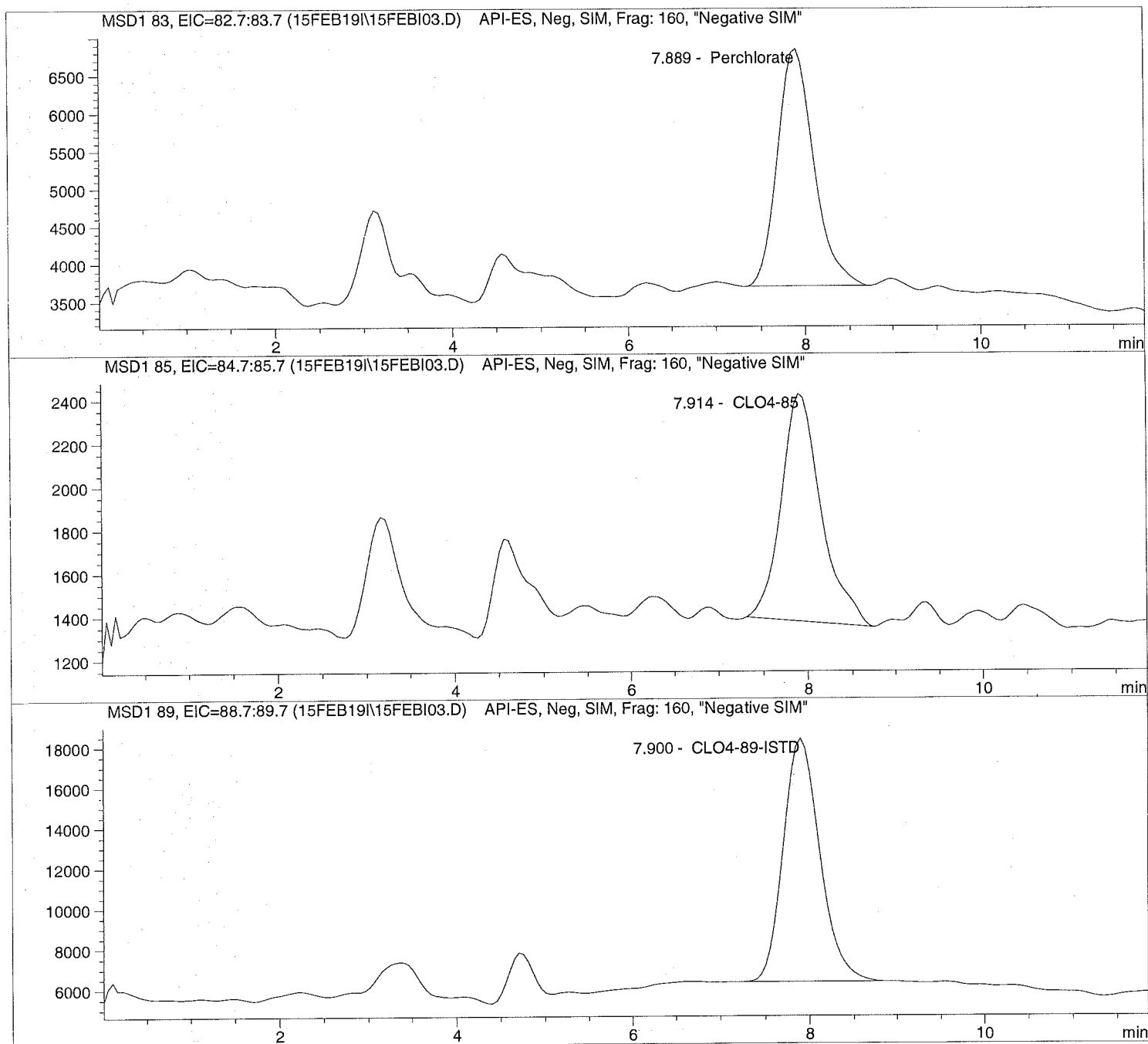
Sample Name: CLO4@ 1.0ug/L

Injection Date: 2/15/2019 09:51:42
Sample Name: CLO4@ 1.0ug/L
Acq Operator: TNB

Seq Line: 3
Location: Vial 73
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI03.D Sample Name: CLO4@ 1.0ug/L

```

=====
Injection Date: 2/15/2019 09:51:42      Seq Line: 3
Sample Name:    CLO4@ 1.0ug/L           Location:  Vial 73
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 1.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.889	PBA	89400.6	0.9899	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.914	BBA	32612.1	0.9988	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	BBA	341443.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D

Sample Name: CLO4@ 2.0ug/L

Injection Date: 2/15/2019 10:05:24

Seq Line: 4

Sample Name: CLO4@ 2.0ug/L

Location: Vial 74

Acq Operator: TNB

Inj. No.: 1

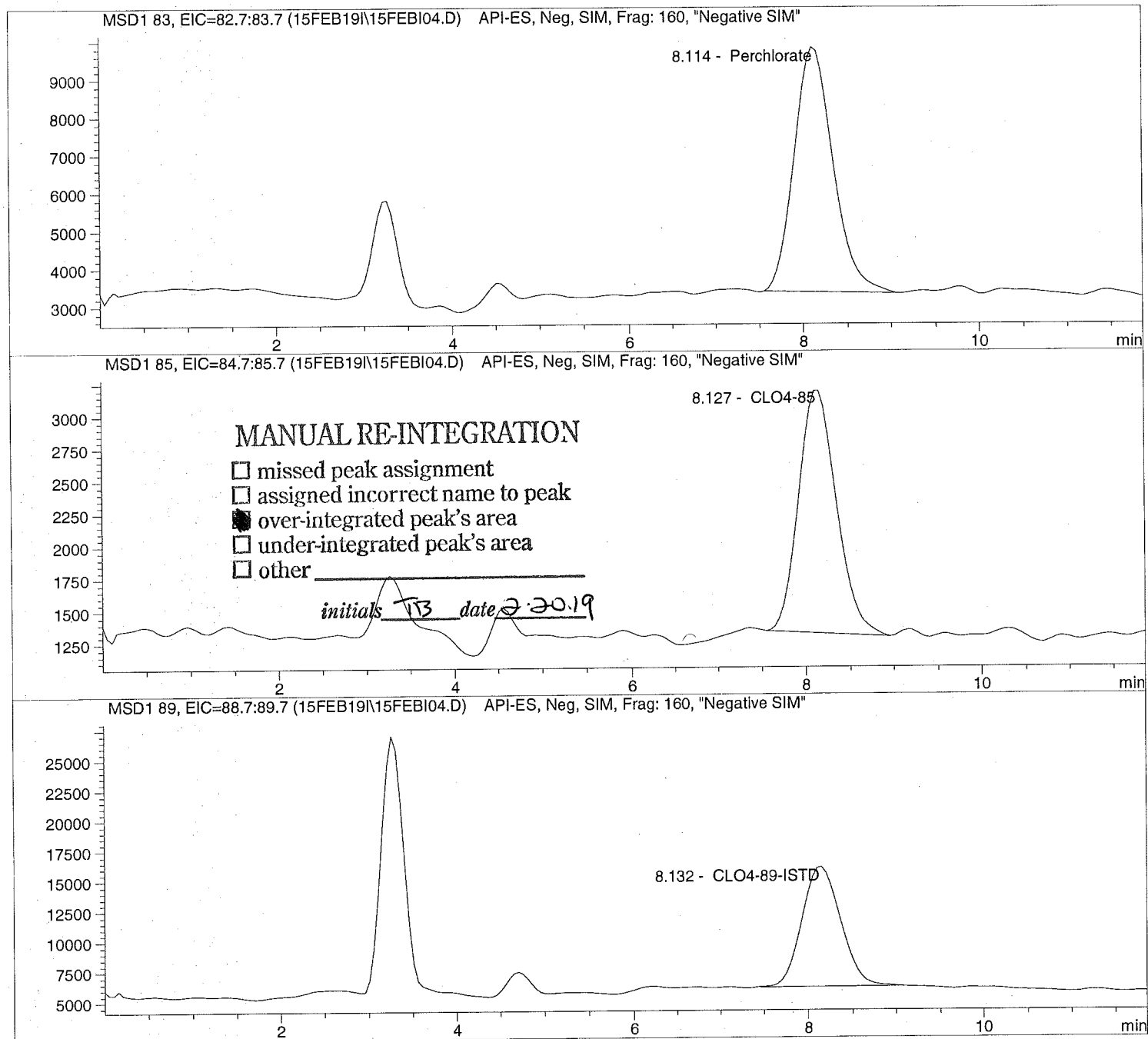
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M

Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 2/15/2019 10:05:24      Seq Line: 4
Sample Name:    CLO4@ 2.0ug/L           Location:  Vial 74
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 2.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	MM	55313.4	2.1136	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

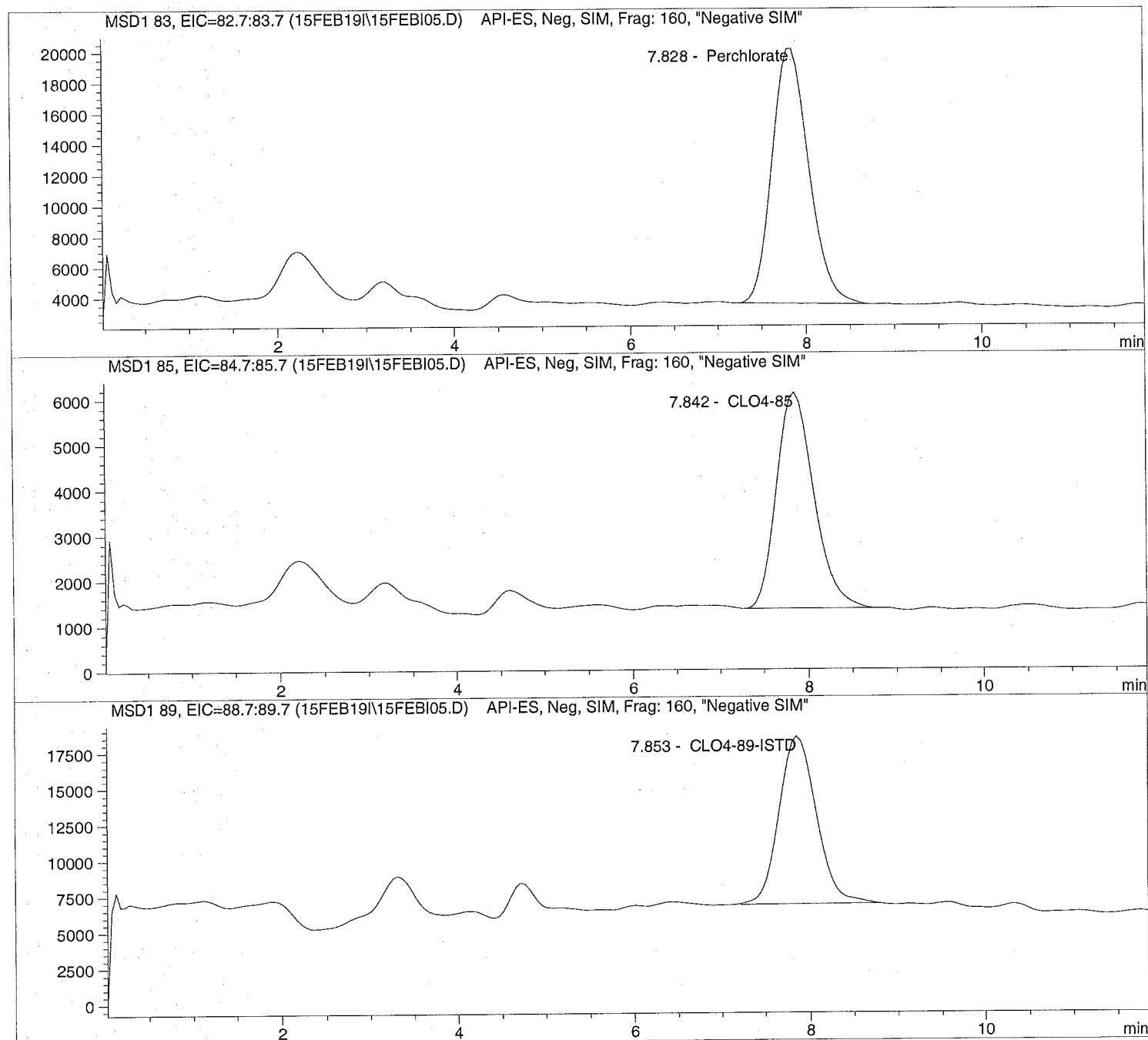
```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI05.D Sample Name: CLO4@ 5.0ug/L

```
=====
Injection Date: 2/15/2019 11:42:56      Seq Line:      5
Sample Name:    CLO4@ 5.0ug/L           Location:      Vial 75
Acq Operator:  TNB                      Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI05.D Sample Name: CLO4@ 5.0ug/L

```

=====
Injection Date: 2/15/2019 11:42:56      Seq Line: 5
Sample Name: CLO4@ 5.0ug/L              Location: Vial 75
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 5.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.828	PBA	479370.4	4.6569	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.842	PBA	139246.9	4.9126	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.853	PBA	338646.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

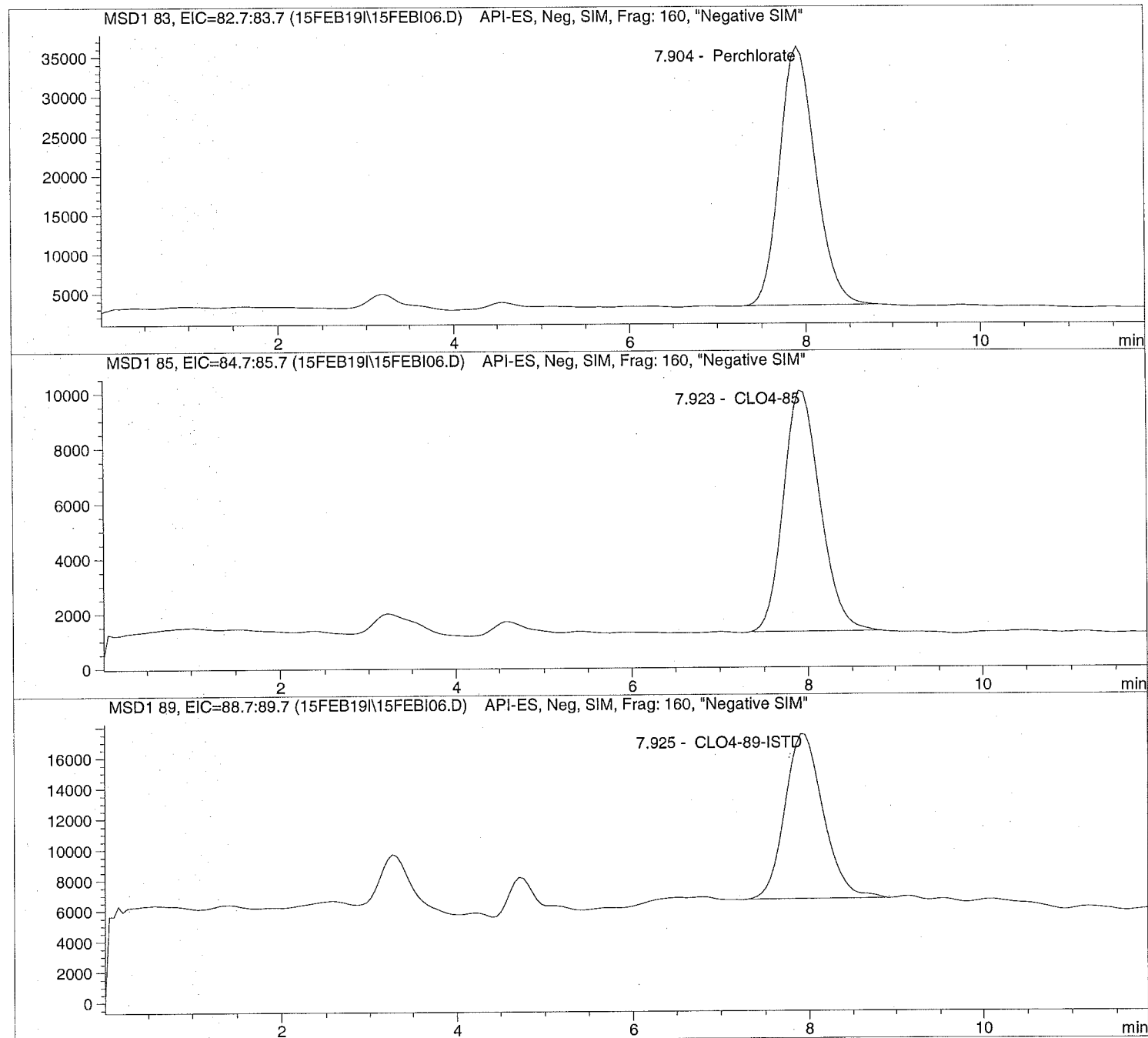
```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI06.D Sample Name: CLO4@ 10.ug/L

```
=====
Injection Date: 2/15/2019 11:56:38      Seq Line: 6
Sample Name:    CLO4@ 10.ug/L           Location:  Vial 76
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI06.D Sample Name: CLO4@ 10.ug/L

```
=====
Injection Date: 2/15/2019 11:56:38      Seq Line: 6
Sample Name: CLO4@ 10.ug/L              Location: Vial 76
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20
=====
```

Perchlorate analysis

Sample Information

```
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 10.000
=====
```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.904	PBA	930135.8	9.1500	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.923	BBA	254395.6	9.3903	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.925	PBA	325154.4	5.0000	CLO4-89-ISTD

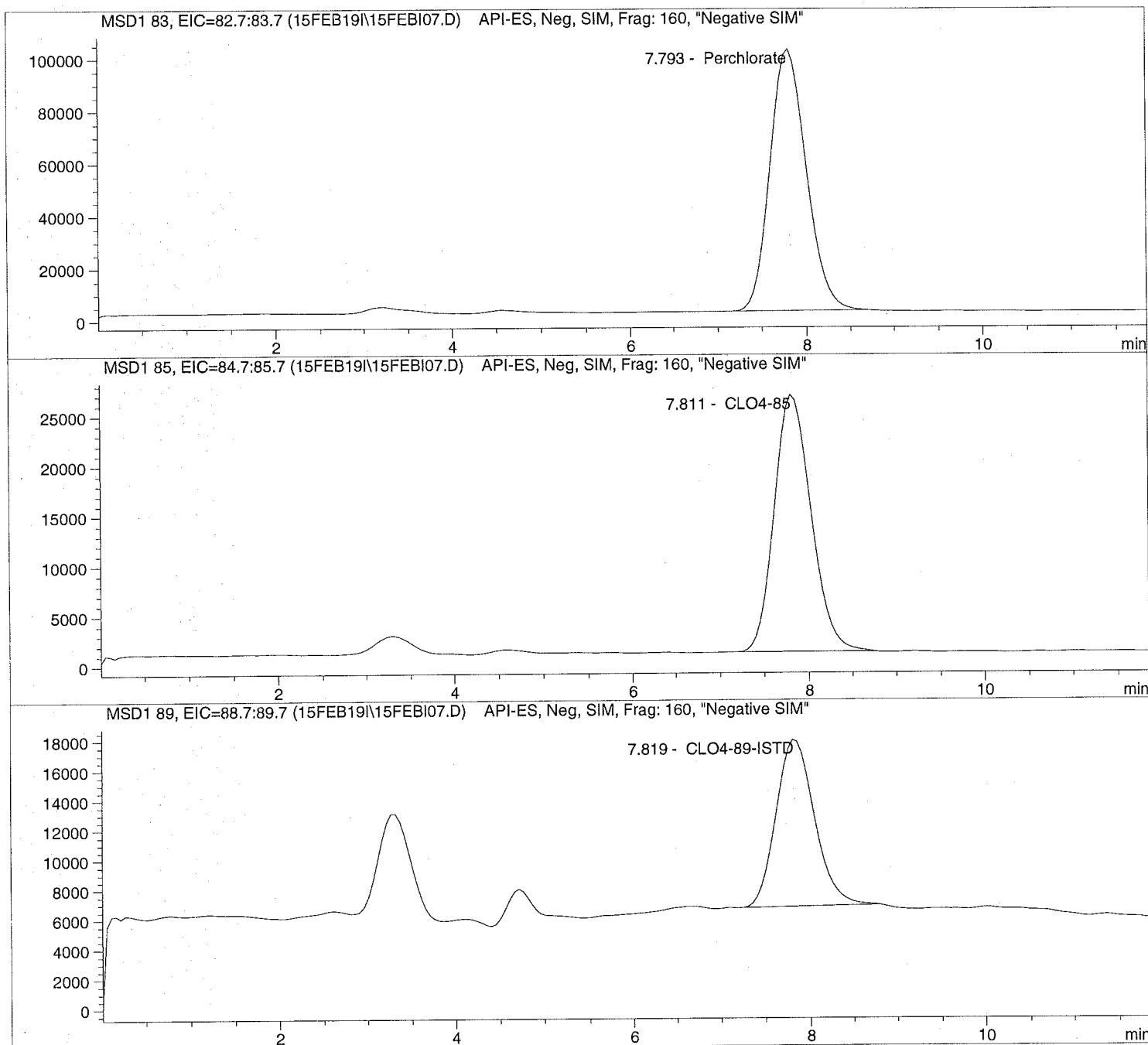
*** End of Report ***

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI07.D Sample Name: CLO4@ 25.ug/L

```
=====
Injection Date: 2/15/2019 12:10:22      Seq Line:          7
Sample Name:    CLO4@ 25.ug/L           Location:          Vial 77
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI07.D Sample Name: CLO4@ 25.ug/L

```

=====
Injection Date: 2/15/2019 12:10:22      Seq Line: 7
Sample Name:    CLO4@ 25.ug/L           Location:  Vial 77
Acq Operator:   TNB                     Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  25.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.793	PBA	2810669.2	25.5264	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.811	BBA	735968.9	25.4827	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.819	PBA	333799.0	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI08.D

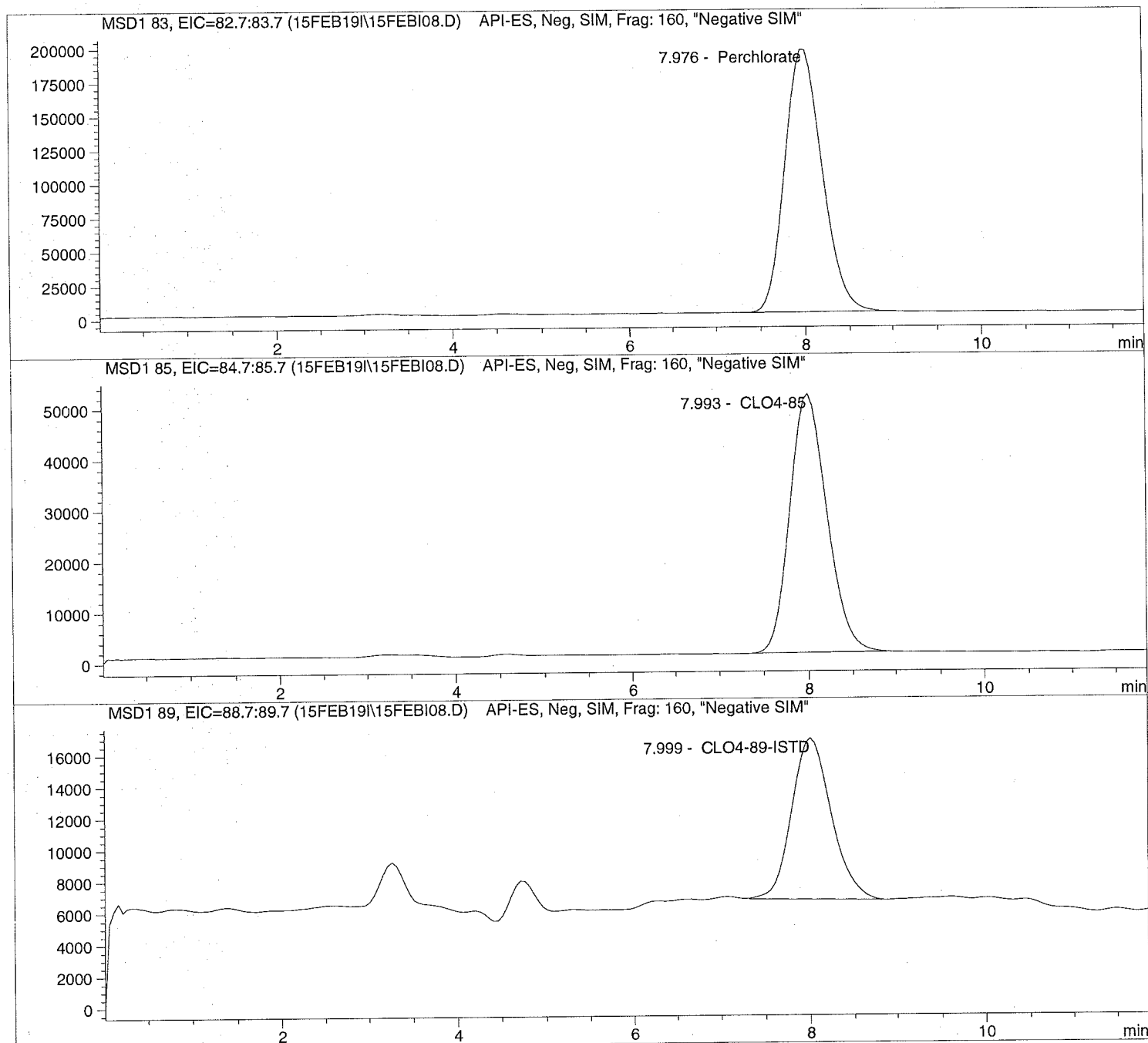
Sample Name: CLO4@ 50.ug/L

=====
Injection Date: 2/15/2019 12:24:06
Sample Name: CLO4@ 50.ug/L
Acq Operator: TNB

Seq Line: 8
Location: Vial 78
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI08.D Sample Name: CLO4@ 50.ug/L

```

=====
Injection Date: 2/15/2019 12:24:06      Seq Line: 8
Sample Name: CLO4@ 50.ug/L      Location: Vial 78
Acq Operator: TNB      Inj. No.: 1
                                 Inj. Vol.: 25 µl

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:09:20

```

Perchlorate analysis

Sample Information

```

=====
Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 50.000

```

LCMS Results

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.976	PBA	5668301.5	51.0744	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.993	PBA	1471522.9	50.3577	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.999	BBA	314711.8	5.0000	CLO4-89-ISTD

*** End of Report ***

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI09.D

Sample Name: CLO4@ 75.ug/L

Injection Date: 2/15/2019 12:37:48

Seq Line: 9

Sample Name: CLO4@ 75.ug/L

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

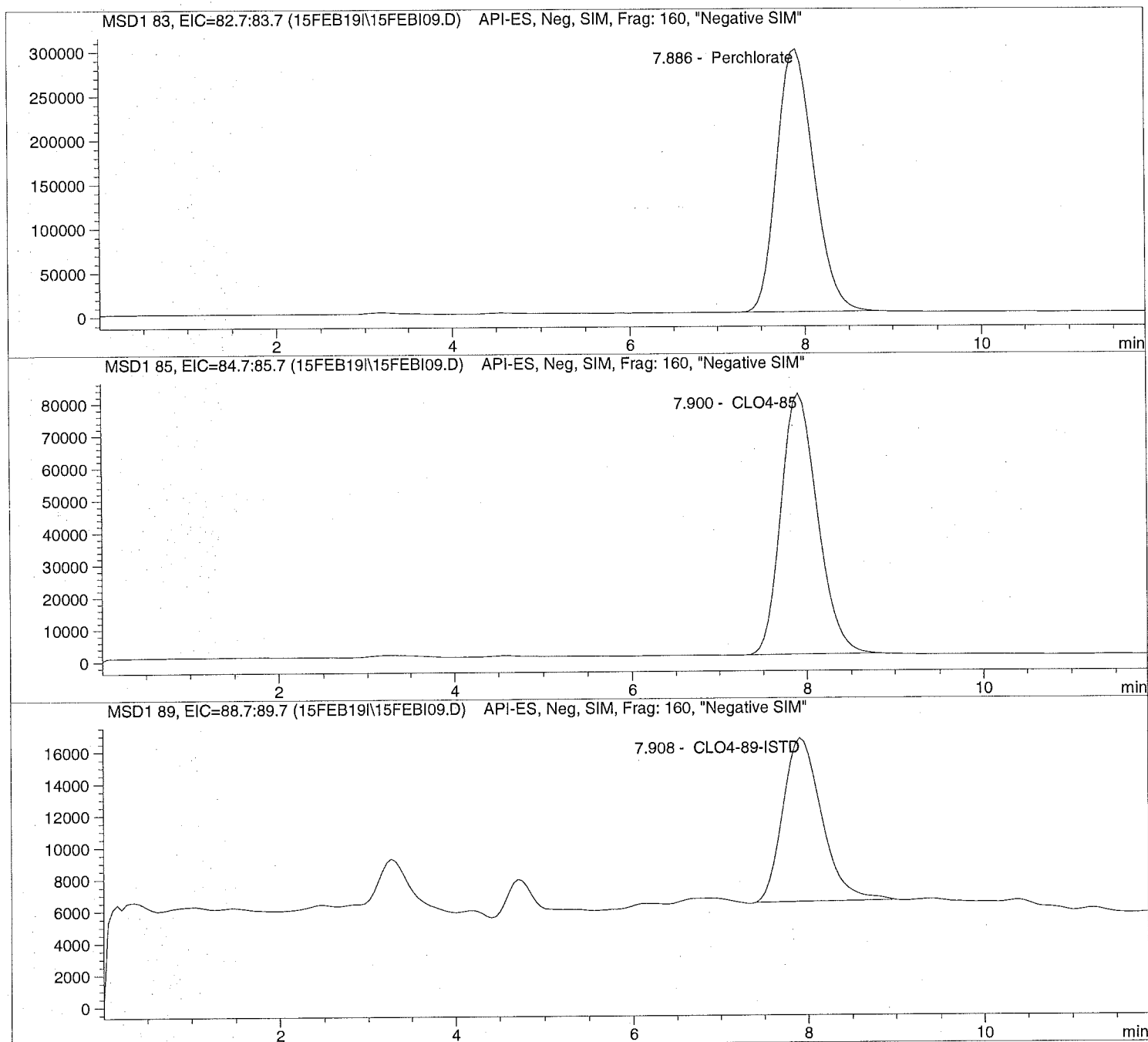
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M

Last Changed: 2/19/2019 09:09:20

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI09.D Sample Name: CLO4@ 75.ug/L

```

=====
Injection Date: 2/15/2019 12:37:48      Seq Line:          9
Sample Name:    CLO4@ 75.ug/L           Location:          Vial 79
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:       25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:     1.000000
Dilution:       1.000000
Sample Amount:  75.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.886	PBA	8696239.0	74.3060	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.900	PBA	2328089.5	74.7223	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.908	PBA	313908.9	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

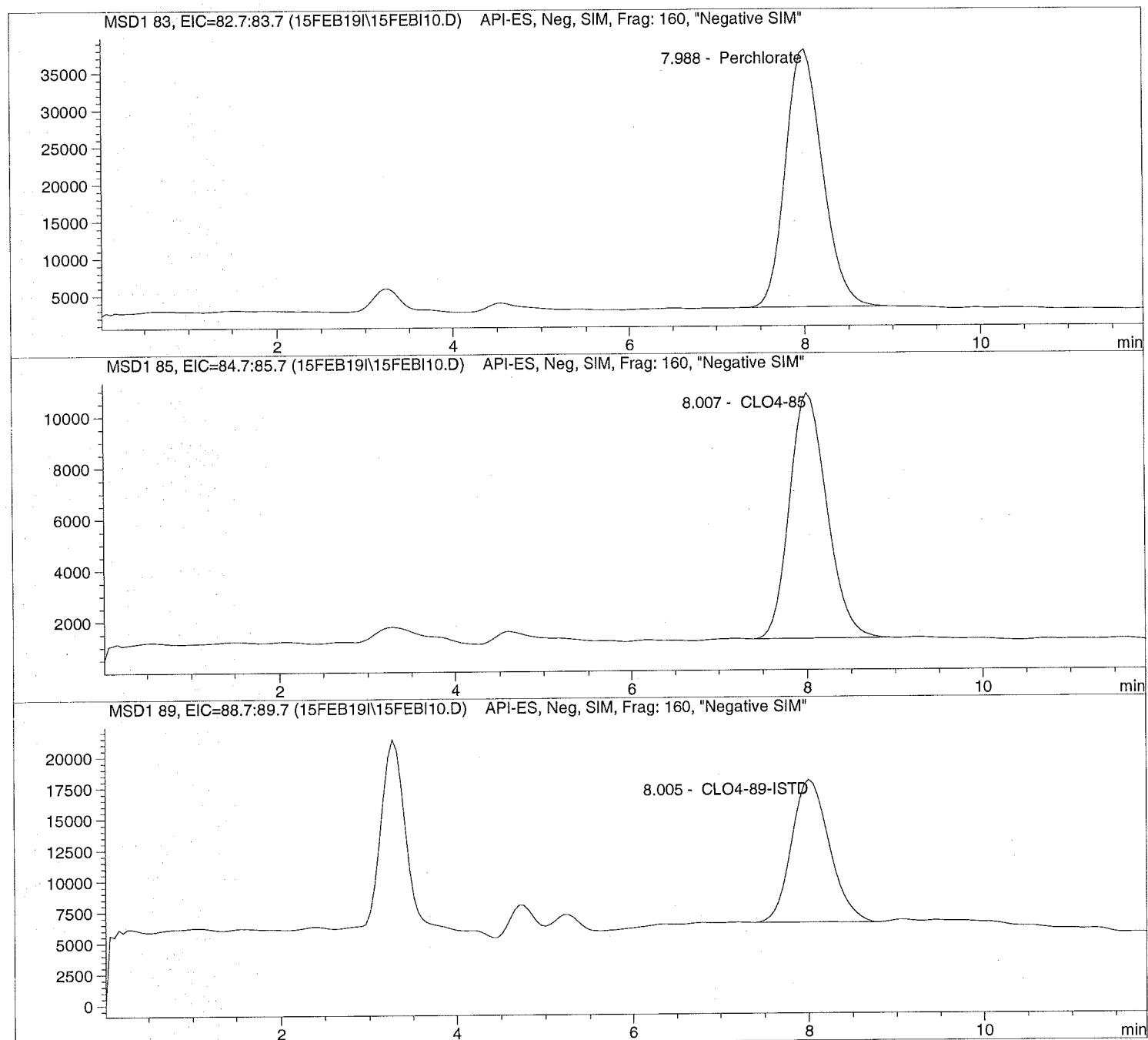
```

Data file: C:\HPCHEM\1\DATA\15FEB19\15FEB10.D Sample Name: ICAL Verf@10ug/L

```
=====
Injection Date: 2/15/2019 12:51:29      Seq Line:      10
Sample Name:    ICAL Verf@10ug/L        Location:      Vial 80
Acq Operator:   TNB                     Inj. No.:     1
                                           Inj. Vol.:    25 µl
=====
```

```
Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI10.D Sample Name: ICAL Verf@10ug/L

```

=====
Injection Date: 2/15/2019 12:51:29      Seq Line:          10
Sample Name:    ICAL Verf@10ug/L        Location:          Vial 80
Acq Operator:   TNB                     Inj. No.:         1
                                           Inj. Vol.:        25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:09:20
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 10.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.988	BBA	1011409.8	9.4602	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.007	BBA	281229.9	9.8786	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.005	BBA	341503.2	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```



ALS Laboratory Group
ANALYTICAL CHEMISTRY & TESTING SERVICES

Environmental Division

Raw Data

Unmodified

Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D

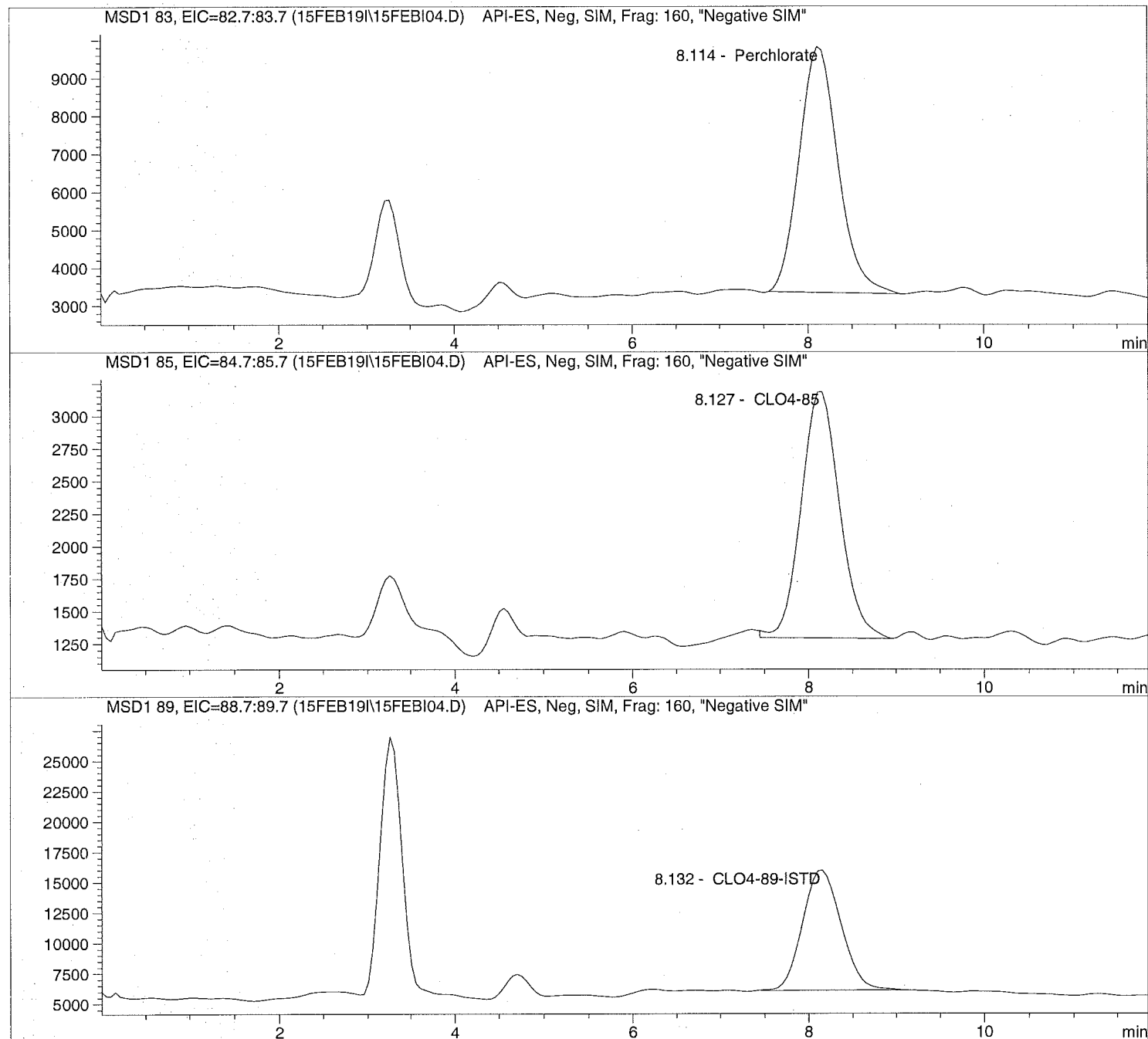
Sample Name: CLO4@ 2.0ug/L

Injection Date: 2/15/2019 10:05:24
Sample Name: CLO4@ 2.0ug/L
Acq Operator: TNB

Seq Line: 4
Location: Vial 74
Inj. No.: 1
Inj. Vol.: 25 µl

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 09:12:36

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\15FEB19I\15FEBI04.D Sample Name: CLO4@ 2.0ug/L

```

=====
Injection Date: 2/15/2019 10:05:24      Seq Line: 4
Sample Name:    CLO4@ 2.0ug/L           Location:  Vial 74
Acq Operator:  TNB                      Inj. No.: 1
                                           Inj. Vol.: 25 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 09:12:36
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 2.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.114	PBA	197442.9	2.2603	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.127	BBA	57206.1	2.1923	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.132	BBA	299650.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

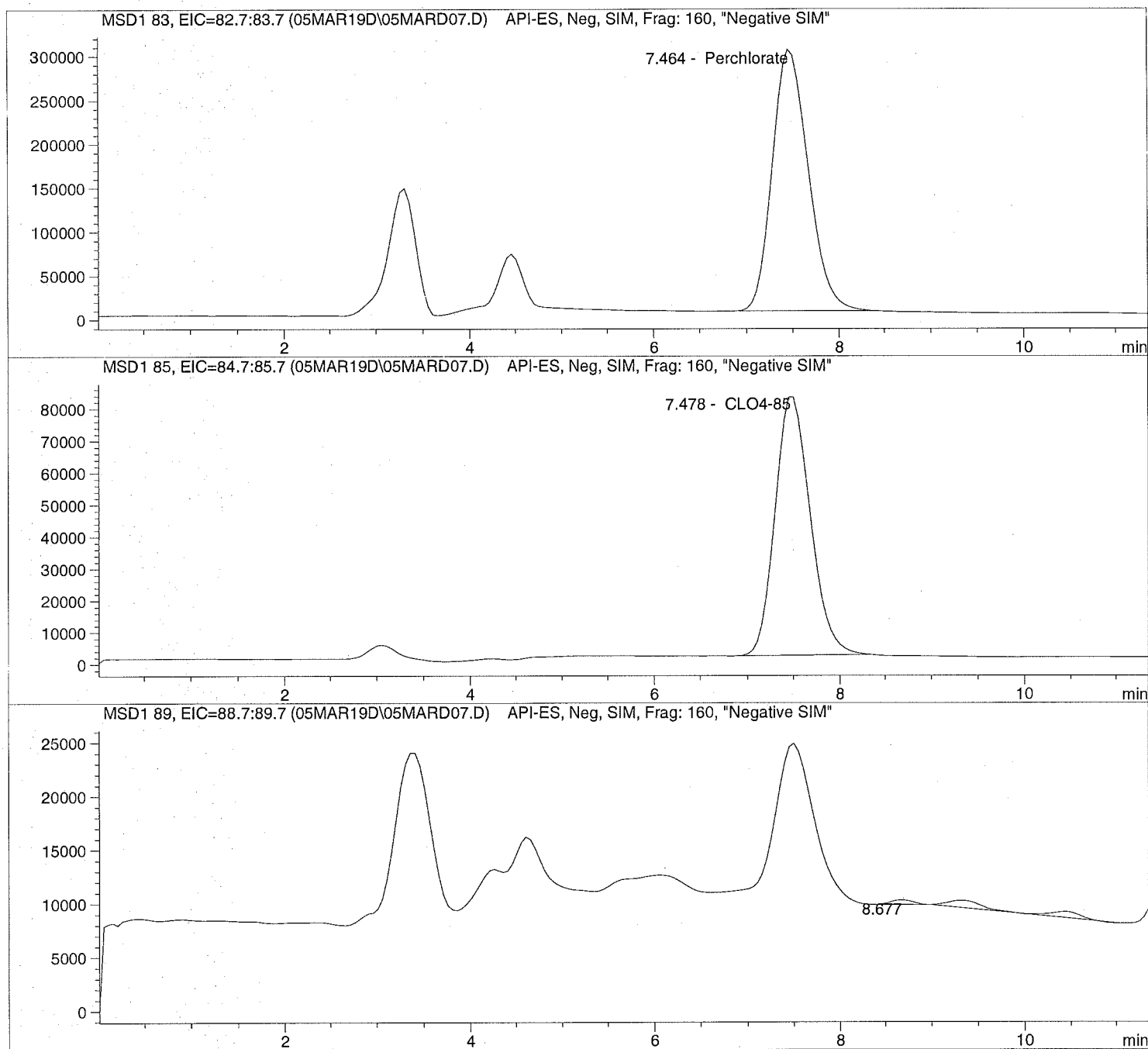
```


Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

```
=====
Injection Date: 3/05/2019 10:07:11      Seq Line: 7
Sample Name: 1906112002 MS              Location: Vial 77
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====
```

```
Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
```

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD07.D Sample Name: 1906112002 MS

```

=====
Injection Date: 3/05/2019 10:07:11      Seq Line:          7
Sample Name:   1906112002  MS           Location:         Vial 77
Acq Operator:  TNB                      Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.464	PBA	8026959.0	419.8794	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.478	PBA	2146365.0	402.6179	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.677	BB	7208.6	0.0000	
9.316	VBA	28561.1	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

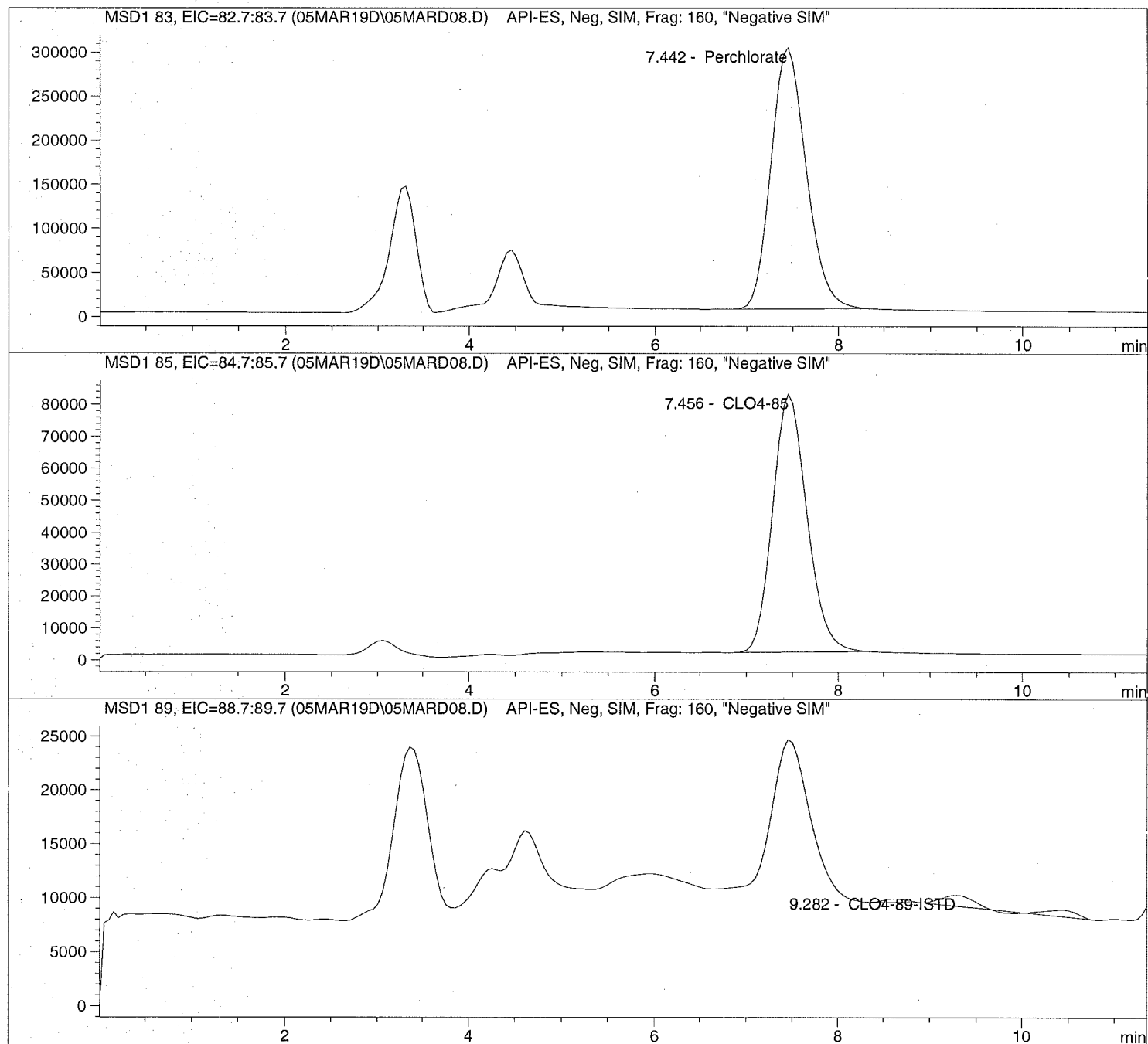
```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

Injection Date: 3/05/2019 10:20:17 Seq Line: 8
Sample Name: 1906112003 MSD Location: Vial 78
Acq Operator: TNB Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD08.D Sample Name: 1906112003 MSD

```

=====
Injection Date: 3/05/2019 10:20:17      Seq Line:      8
Sample Name:   1906112003  MSD          Location:     Vial 78
Acq Operator:  TNB                    Inj. No.:    1
                                           Inj. Vol.:   20 µl
=====

```

```

Acq. Method:   CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:  2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.442	PBA	7942422.5	359.0996	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.456	PBA	2109911.2	344.2233	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
8.621	VB	5769.9	0.0000	
9.282	VBA	35831.6	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D

Sample Name: 1906112004

Injection Date: 3/05/2019 10:33:21

Seq Line: 9

Sample Name: 1906112004

Location: Vial 79

Acq Operator: TNB

Inj. No.: 1

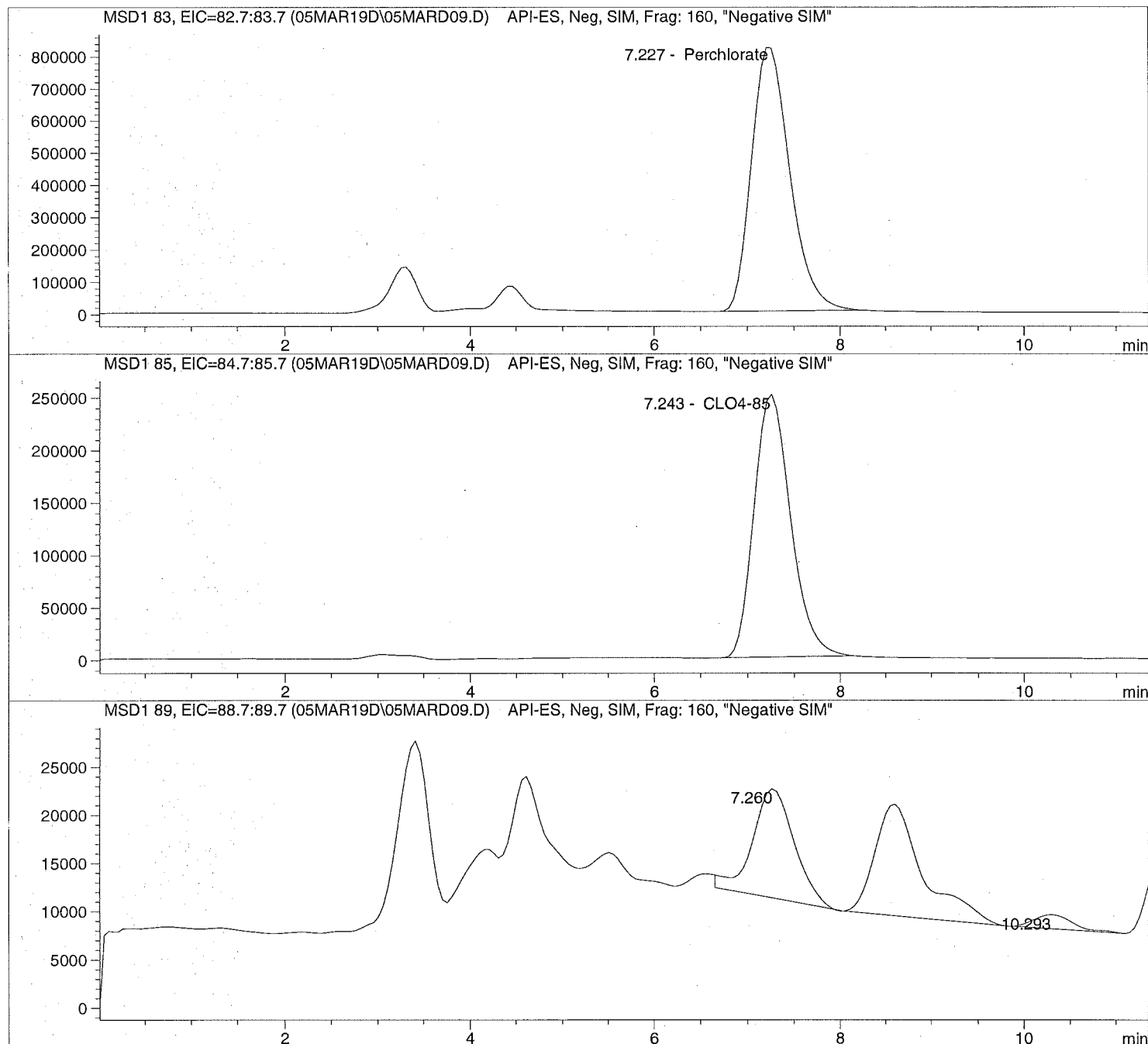
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M

Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD09.D Sample Name: 1906112004

```

=====
Injection Date: 3/05/2019 10:33:21      Seq Line:          9
Sample Name:    1906112004              Location:         Vial 79
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.227	PBA	23301694.0	131.1742	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.243	PBA	6796677.5	138.8050	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.260	BB	352149.7	0.0000	
8.589	VBA	421141.9	5.0000	CLO4-89-ISTD
10.293	BBA	41603.7	0.0000	

```

=====
*** End of Report ***
=====

```

Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D

Sample Name: 1906112005

Injection Date: 3/05/2019 10:46:26

Seq Line: 10

Sample Name: 1906112005

Location: Vial 80

Acq Operator: TNB

Inj. No.: 1

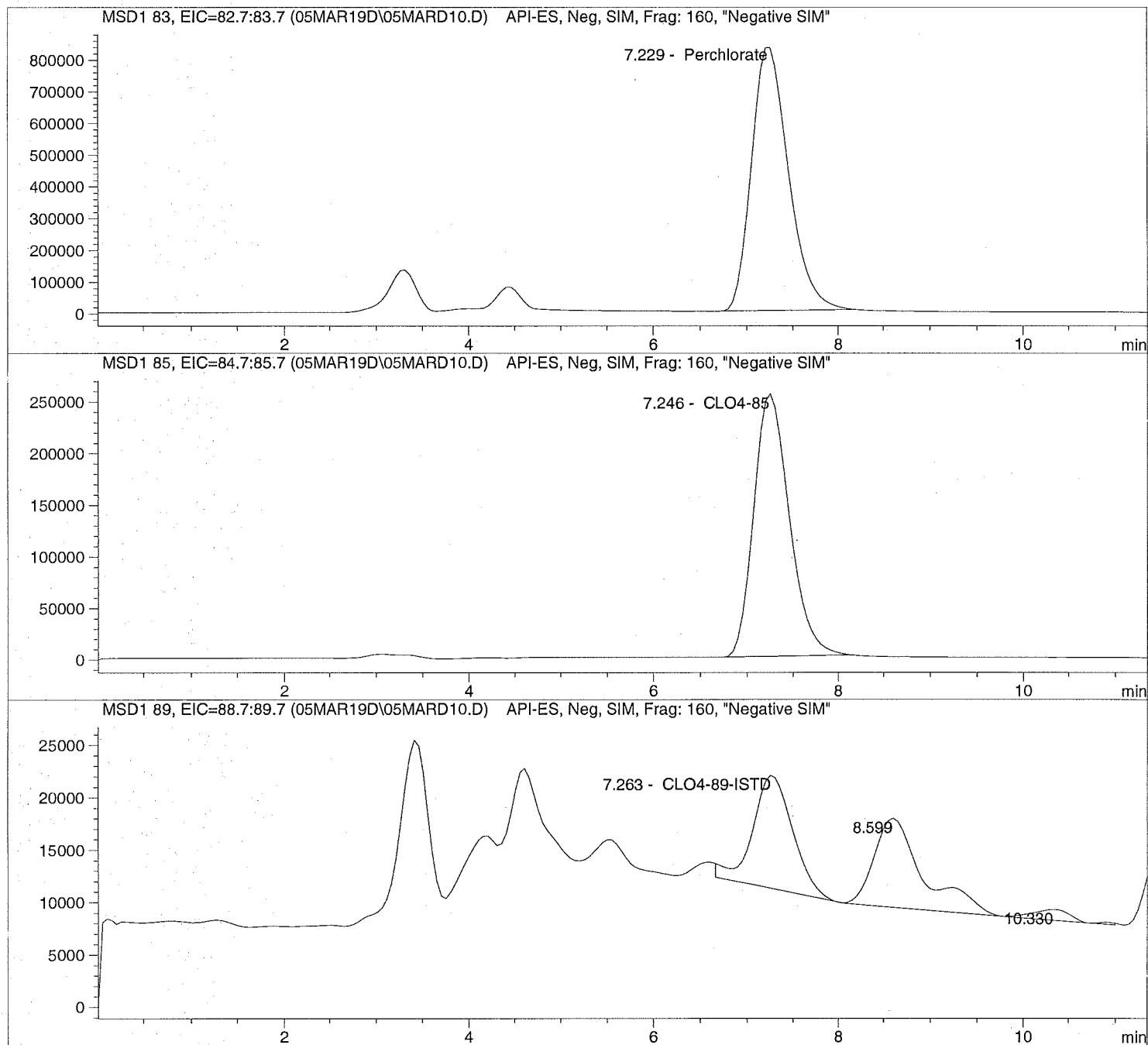
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M

Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M

Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD10.D Sample Name: 1906112005

```

=====
Injection Date: 3/05/2019 10:46:26      Seq Line:          10
Sample Name:    1906112005              Location:         Vial 80
Acq Operator:   TNB                     Inj. No.:        1
                                           Inj. Vol.:       20 µl
=====

```

```

Acq. Method:    CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed:   2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
                          Sample Information
=====

```

```

Sorted By:      Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier:    1.000000
Dilution:      1.000000
Sample Amount: 0.000
=====

```

```

=====
                          LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.229	PBA	23099082.0	159.0418	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.246	PBA	6828341.5	168.6882	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.263	BB	325841.0	5.0000	CLO4-89-ISTD
8.599	VB	308921.9	0.0000	
10.330	VBA	30210.3	0.0000	

```

=====
*** End of Report ***
=====

```


Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D

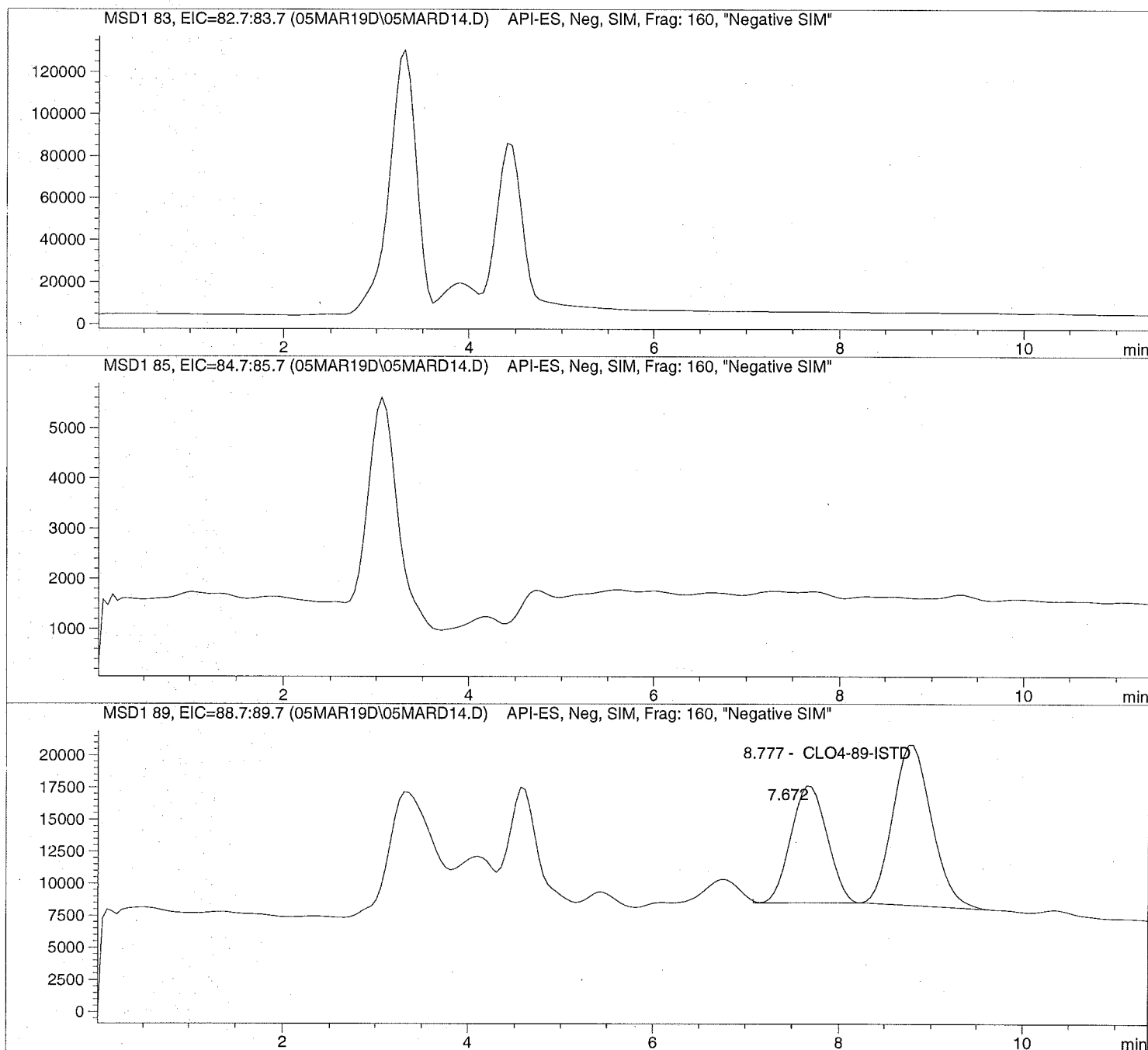
Sample Name: 1906112009

Injection Date: 3/05/2019 11:39:24
Sample Name: 1906112009
Acq Operator: TNB

Seq Line: 14
Location: Vial 84
Inj. No.: 1
Inj. Vol.: 20 μ l

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46

Perchlorate analysis



Data file: C:\HPCHEM\1\DATA\05MAR19D\05MARD14.D Sample Name: 1906112009

```

=====
Injection Date: 3/05/2019 11:39:24      Seq Line: 14
Sample Name: 1906112009                 Location: Vial 84
Acq Operator: TNB                       Inj. No.: 1
                                           Inj. Vol.: 20 µl
=====

```

```

Acq. Method: CLO4-AQN.M
Analysis Method: C:\HPCHEM\1\METHODS\CLO4-DP1.M
Last Changed: 2/19/2019 12:13:46
=====

```

Perchlorate analysis

```

=====
Sample Information
=====

```

```

Sorted By: Signal
Calib. Data Modified: Tue, 19. Feb. 2019,09:07:33 am
Multiplier: 1.000000
Dilution: 1.000000
Sample Amount: 0.000
=====

```

```

=====
LCMS Results
=====

```

Signal1: MSD1 83, EIC=82.7:83.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	Perchlorate

Signal2: MSD1 85, EIC=84.7:85.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
0.000		0.0	0.0000	CLO4-85

Signal3: MSD1 89, EIC=88.7:89.7

RT [min]	Type	Area	Amount [ug/sample]	Compound Name
7.672	BB	243675.3	0.0000	
8.777	VBA	362717.3	5.0000	CLO4-89-ISTD

```

=====
*** End of Report ***
=====

```

GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

APPENDIX E
QUALITY CONTROL SUMMARY REPORT

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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**QUARTERLY CONTROL SUMMARY REPORT
1st QUARTER (JANUARY - MARCH) 2019
GROUNDWATER TREATMENT PLANT
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS**

May 2019

Prepared For:



**Longhorn Army Ammunition Plant
Karnack, Texas**

Under Contract To:



**U.S. Army Corps of Engineers
Tulsa District
Tulsa, Oklahoma**

Contract Number: W9128F-13-D-0012

Task Order Number: W912BV17F0150

Prepared By:



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1 INTRODUCTION

Bhate reviewed 17 data packages from ALS Environmental, Houston, Texas. Groundwater samples were collected from February 21, 2019, through March 27, 2019, at the Groundwater Treatment Plant (GWTP) and Site LHAAP-16 at Longhorn Army Ammunition Plant (LHAAP), Karnack, Texas. No samples were collected during the month of January due to mechanical issues at the GWTP. Data were reviewed for conformance to the requirements of the following guidance documents: *USEPA Contract Laboratory Program [CLP] National Functional Guidelines for Superfund Organic Methods Data Review* (USEPA, January 2017); *USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review* (USEPA, January 2017); and the *Final Basewide Uniform Federal Policy [UFP] – Quality Assurance Project Plan [QAPP] Longhorn Army Ammunition Plant* which is in Appendix C of the *Final Installation-Wide Work Plan for Longhorn Army Ammunition Plant Karnack, Texas* (Bhate, May 2018).

1.1 Intended Use of Data

The objective of sampling at the GWTP is to monitor effluent streams to confirm compliance with discharge limits.

Analyses performed include:

- SW6850 – Perchlorate
- E350.3/SM4500 NH₃ – Nitrogen, Ammonia
- E365.3/SM4500 P – Orthophosphate
- E415.1/SM5310 – Total Organic Carbon
- SW8260C – Volatile Organic Compounds (VOCs)
- SW8270D Selected Ion Monitoring (SIM) - 1,4-Dioxane
- SW6020A/7470A – Metals
- SW9056A – Chloride (Cl) and Sulfate (SO₄)
- SW7196A – Hexavalent Chromium
- E410.4 – Chemical Oxygen Demand
- E1664A – Oil and Grease
- TO-15 – Volatiles in Air

Table 1 lists the sample identification numbers (IDs) and their associated laboratory package.

Table 2 lists qualified results with the qualification flag and reason code.

The following narrative is a brief synopsis of data that required qualification due to quality control discrepancies.

1.2 Preservation and Holding Times

Sample identification data were evaluated for agreement with the chain-of-custody (COC). All samples were received in appropriate containers, within the proper temperature range, in good condition, and within the required hold time.

1.3 Calibrations

All analytes reported must be present in the initial and continuing calibration. The calibrations must meet the acceptance criteria specified in Worksheet 24 (Analytical Instrument Calibration) of the QAPP. All results reported must be within the calibration range. Samples were diluted, if necessary, to bring analyte responses within the calibration range.

1.3.1 Continuing Calibration Verifications (CCV)

The calibrations must meet the following criteria otherwise the compound is qualified J or UJ: The continuing calibration verification (CCV) criteria are 20 percent difference (%D) for VOCs and Semi-Volatile Organic Compounds (SVOCs) and 50% for closing CCVs. Metals and general chemistry - 10%D; perchlorate - 15%D; and volatiles in air - 30%D.

The CCV for benzene and toluene was exceeded for the bi-weekly sample LH18/24-SP650_032119. The CCV for 1,2-dibromoethane, bromoform, and dibromochloromethane were also outside control limits for quarterly influent sample LH18/24-SP140_032719. These compounds were qualified as estimated non-detects and flagged "UJ".

1.4 Blanks

If the analyte result for an associated sample was less than 5X (10X for common laboratory contaminants) the analyte concentration in the blank, that result was qualified "UB" and considered an artifact of blank contamination. Where the sample result for the affected analyte was non-detect or greater than 5X the amount in the blank, no qualifier was applied.

1.4.1 SW6020A

The method blank for quarterly effluent and influent samples LH18/24-SP650_032719 and LH18/24-SP140_032719, respectively reported a low level detection of vanadium. Since the sample results were < 5X the blank result, they were considered artifacts of blank contamination and flagged "UB".

1.5 Surrogates

Surrogates were evaluated using limits defined by each method in the project-specific QAPP Worksheet 28.

1.5.1 SW8270D SIM

The monthly effluent sample LH18/24-SP650_031419 reported 2-fluorobiphenyl (193%) above QC limits. 1,4-Dioxane was considered as estimated and flagged "J".

1.6 Laboratory Control Sample (LCS)/Laboratory Control Sample Duplicate (LCSD)

LCS/LCSD recoveries were evaluated using limits defined in the project-specific QAPP Worksheet 15.

All sample recoveries were within control limits.

1.7 Matrix Spike (MS)/Matrix Spike Duplicate Sample (MSD)

MS/MSD recoveries were evaluated using limits defined in Worksheet 15 of the project-specific QAPP. When sample results were greater than 4X the spike amount, control limits were not applicable and require no qualification. Furthermore, if a MS/MSD analyses was performed on a batched (unrelated) sample, no qualification was made to the sample data.

1.7.1 SW6850

Site LHAAP-16 reported the MS and MSD recoveries of perchlorate below control limits. This compound was qualified as estimated, "J", in the sample used for the spike analysis (16EW01_022619).

1.7.2 SW8260C

Site LHAAP-16 reported the MSD recovery of cis-1,2-dichloroethene below control limits. This compound was qualified as estimated, "J", in the sample used for the spike analysis (16EW01_022619).

1.7.3 SW6850

The MS/MSD recoveries for GWTP sample LH18/24-SP650_032719_BIX were above control limits. The perchlorate result was considered estimated and flagged "J".

1.8 Internal Standards

If the percent recovery (%R) for an internal standard in a sample is not within the limit, the associated sample is qualified for those analytes associated with the internal standard(s) outside of the limit.

Internal standards were within acceptance criteria for the associated compounds.

1.9 Field Precision

Precision is the measure of variability of individual sample measurements. Evaluation of field duplicates for precision was done using the relative percent difference (RPD) calculation. The RPD is defined as the difference between two duplicate samples divided by the mean and expressed as a percent. Field duplicate RPD limits were set at <30% for groundwater and air matrices.

1.9.1 SW8260C

The LHAAP-16 RPD, between 16EW05_022619 and its duplicate, was exceeded for vinyl chloride. This compound was qualified as estimated, "J", in both sample and duplicate.

2 DATA USABILITY SUMMARY

The data are usable for the intended purposes of the project (see Table 3). The data quality objectives have been met for the project.

Table 1: Field Sample Identification and Laboratory Packages

Client Sample ID	Lab Package	SW6850	E350.3	E365.3	EA15.1	SM SW8270D	SW8260C	SW6020A	SW9056A	SW7196A	EA10.4	E1664A	TO-15
GWTP Samples													
LH18/24-SP650_022119/BIX	HS19021158	X	X	X	X								
LH18/24-SP650_022119	HS19021165						X		X				
LH18/24-SP650_022819/BIX	HS19030014	X	X	X	X								
LH18/24-SP650_022819/BIX	HS19030012	X				X	X	X		X			
LH18/24-SP140_022819	HS19030011	X						X		X			
LH18/24-SP650_030619/BIX	HS19030298	X	X	X	X								
LH18/24-SP650_030619	HS19030301						X		X				
LH18/24-SP650_031419/BIX	HS19030761	X				X	X	X		X			
LH18/24-SP650_031419/BIX	HS19030749	X	X	X	X								
LH18/24-SP140_031419	HS19030763	X						X		X			
LH18/24-SP650_032119/BIX	HS19031160	X	X	X	X								
LH18/24-SP650_032119	HS19031189						X		X				
LH18/24-SP650_032719/BIX	HS19031492	X	X	X	X								
LH18/24-SP140_032719	HS19031508	X				X	X	X	X		X	X	
LH18/24-SP650_032719/BIX	HS19031511	X				X	X	X	X		X	X	
Air Samples													
LH18/24-Air_022719_Stripper	P1901179												X
LH18/24-Air_022719_Stripper_a	P1901179												X
LH18/24-Air_022719_GWTP	P1901179												X
LH18/24-Air_022719_Downwind	P1901179												X
LHAAP-16													
16EW01_022619	HS19021428	X					X						
16EW02_022619	HS19021428	X					X						
16EW03_022619	HS19021428	X					X						
16EW04_022619	HS19021428	X					X						
16EW05_022619	HS19021428	X					X						
16EW05_022619_a	HS19021428	X					X						
16EW06_022619	HS19021428	X					X						
16EW07_022619	HS19021428	X					X						
16EW08_022619	HS19021428	X					X						
Notes: MW – Monitoring Well SM – Standard Method SW-846 - Test Methods for Evaluating Solid Waste, Physical/Chemical Methods. E – U.S. Environmental Protection Agency Method HS – Houston BIX – before Ion exchange (perchlorate analysis)													

Table 2: Qualified Analytical Data

Client Sample ID Laboratory	Laboratory Package	Analyte Name	Data Validation Qualifier	Reason for Qualification
GWTP				
LH18/24-SP650_031419	HS19030761	1,4-Dioxane	3.1 J	Surr >
LH18/24-SP650_032119	HS19031189	Benzene Toluene	0.50 UJ 0.50 UJ	CCV >
LH18/24-SP650_032719_BIX	HS19031492	Perchlorate	2.6 J	MS/MSD >
LH18/24-SP650_032719	HS19031511	Vanadium	0.00382 UB	MB
LH18/24-SP140_032719	HS19031508	Vanadium	0.00237 UB	MB
		1,2-Dibromoethane	5.0 UJ	CCV
		Bromoform	5.0 UJ	CCV
		Dibromochloromethane	5.0 UJ	CCV
LHAAP-16				
16EW01_022619	HS19021428	Perchlorate cis-1,2-Dichloroethene	51 J 3,700 J	MS/MSD < MSD <
16EW05_022619	HS19021428	Vinyl chloride	260 J	Dup RPD
16EW05_022619_a	HS19021428	Vinyl chloride	420 J	Dup RPD
Notes:				
ID – identification				
J – Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.				
UJ – The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.				
UB – considered an artifact of blank contamination				
MS/MSD </>– Matrix spike/duplicate recovery below/above control limits				
Dup RPD – the relative percent difference between sample and duplicate was outside control limits				
MB – Method blank contamination				
CCV – continuing calibration verification outside control limits				
HS - Houston				

Table 3: Completeness by Method

Method	No. of Rejected Results	% Completeness
SW6850	0	100
E350.3/SM4500 NH3	0	100
E365.3/SM4500 P	0	100
E415.1/SM5310	0	100
SW8270D	0	100
SW8260C	0	100
SW6020A	0	100
SW9056A	0	100
SW7196A	0	100
E410.4	0	100
E1664A	0	100
TO-15	0	100

GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

APPENDIX F
AIR MONITORING ANALYTICAL LABORATORY REPORTS
(PROVIDED ON CD ONLY)

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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LABORATORY REPORT

March 25, 2019

Marcia Olive
Bhate Environmental Associates, Inc.
445 Union Blvd Suite 129
Lakewood, CO 80228

RE: LHAAP GWTP / NW01312.0150

Dear Marcia:

Enclosed are the results of the samples submitted to our laboratory on March 5, 2019. For your reference, these analyses have been assigned our service request number P1901179.

All analyses were performed according to our laboratory's NELAP and DoD-ELAP-approved quality assurance program. The test results meet requirements of the current NELAP and DoD-ELAP standards, where applicable, and except as noted in the laboratory case narrative provided. For a specific list of NELAP and DoD-ELAP-accredited analytes, refer to the certifications section at www.alsglobal.com. Results are intended to be considered in their entirety and apply only to the samples analyzed and reported herein.

If you have any questions, please call me at (805) 526-7161.

Respectfully submitted,

ALS | Environmental

By Michael Conejo at 9:24 am, Mar 25, 2019

Michael Conejo
Project Manager



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www.alsglobal.com

Client: Bhate Environmental Associates, Inc.
 Project: LHAAP GWTP / NW01312.0150

Service Request No: P1901179

CASE NARRATIVE

The samples were received intact under chain of custody on March 5, 2019 and were stored in accordance with the analytical method requirements. Please refer to the sample acceptance check form for additional information. The results reported herein are applicable only to the condition of the samples at the time of sample receipt.

Volatile Organic Compound Analysis

The samples were analyzed for volatile organic compounds and in accordance with EPA Method TO-15 from the Compendium of Methods for the Determination of Toxic Organic Compounds in Ambient Air, Second Edition (EPA/625/R-96/010b), January, 1999. This procedure is described in laboratory SOP VOA-TO15. The analytical system was comprised of a gas chromatograph / mass spectrometer (GC/MS) interfaced to a whole-air preconcentrator. This method is included on the laboratory's NELAP and DoD-ELAP scope of accreditation. Any analytes flagged with an X are not included on the NELAP or DoD-ELAP accreditation.

The containers were cleaned, prior to sampling, down to the method reporting limit (MRL) reported for this project. For projects requiring DoD QSM 5.1 compliance canisters were cleaned to <1/2 the MRL. Please note, projects which require reporting below the MRL could have results between the MRL and method detection limit (MDL) that are biased high.

The results of analyses are given in the attached laboratory report. All results are intended to be considered in their entirety, and ALS Environmental (ALS) is not responsible for utilization of less than the complete report.

Use of ALS Environmental (ALS)'s Name. Client shall not use ALS's name or trademark in any marketing or reporting materials, press releases or in any other manner ("Materials") whatsoever and shall not attribute to ALS any test result, tolerance or specification derived from ALS's data ("Attribution") without ALS's prior written consent, which may be withheld by ALS for any reason in its sole discretion. To request ALS's consent, Client shall provide copies of the proposed Materials or Attribution and describe in writing Client's proposed use of such Materials or Attribution. If ALS has not provided written approval of the Materials or Attribution within ten (10) days of receipt from Client, Client's request to use ALS's name or trademark in any Materials or Attribution shall be deemed denied. ALS may, in its discretion, reasonably charge Client for its time in reviewing Materials or Attribution requests. Client acknowledges and agrees that the unauthorized use of ALS's name or trademark may cause ALS to incur irreparable harm for which the recovery of money damages will be inadequate. Accordingly, Client acknowledges and agrees that a violation shall justify preliminary injunctive relief. For questions contact the laboratory.



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ALS Environmental – Simi Valley

CERTIFICATIONS, ACCREDITATIONS, AND REGISTRATIONS

Agency	Web Site	Number
Alaska DEC	http://dec.alaska.gov/eh/lab.aspx	17-019
Arizona DHS	http://www.azdhs.gov/preparedness/state-laboratory/lab-licensure-certification/index.php#laboratory-licensure-home	AZ0694
Florida DOH (NELAP)	http://www.floridahealth.gov/licensing-and-regulation/environmental-laboratories/index.html	E871020
Louisiana DEQ (NELAP)	http://www.deq.louisiana.gov/page/la-lab-accreditation	05071
Maine DHHS	http://www.maine.gov/dhhs/mecdc/environmental-health/dwp/professionals/labCert.shtml	2018027
Minnesota DOH (NELAP)	http://www.health.state.mn.us/accreditation	1521096
New Jersey DEP (NELAP)	http://www.nj.gov/dep/enforcement/oqa.html	CA009
New York DOH (NELAP)	http://www.wadsworth.org/labcert/elap/elap.html	11221
Oregon PHD (NELAP)	http://www.oregon.gov/oha/ph/LaboratoryServices/EnvironmentalLaboratoryAccreditation/Pages/index.aspx	4068-006
Pennsylvania DEP	http://www.dep.pa.gov/Business/OtherPrograms/Labs/Pages/Laboratory-Accreditation-Program.aspx	68-03307 (Registration)
PJLA (DoD ELAP)	http://www.pjlabs.com/search-accredited-labs	65818 (Testing)
Texas CEQ (NELAP)	http://www.tceq.texas.gov/agency/qa/env_lab_accreditation.html	T104704413- 18-9
Utah DOH (NELAP)	http://health.utah.gov/lab/lab_cert_env	CA01627201 8-9
Washington DOE	http://www.ecy.wa.gov/programs/eap/labs/lab-accreditation.html	C946

Analyses were performed according to our laboratory's NELAP and DoD-ELAP approved quality assurance program. A complete listing of specific NELAP and DoD-ELAP certified analytes can be found in the certifications section at www.alsglobal.com, or at the accreditation body's website.

Each of the certifications listed above have an explicit Scope of Accreditation that applies to specific matrices/methods/analytes; therefore, please contact the laboratory for information corresponding to a particular certification.

ALS ENVIRONMENTAL

DETAIL SUMMARY REPORT

Client: Bhate Environmental Associates, Inc.
 Project ID: LHAAP GWTP / NW01312.0150

Service Request: P1901179

Date Received: 3/5/2019

Time Received: 09:20

TO-15 - VOC Cans

Client Sample ID	Lab Code	Matrix	Date Collected	Time Collected	Container ID	Pi1 (psig)	Pf1 (psig)	TO-15 - VOC Cans
LH18/24-Air_022719_Stripper	P1901179-001	Air	2/27/2019	14:00	AS00629	-0.03	3.53	X
LH18/24-Air_022719_Stripper_a	P1901179-002	Air	2/27/2019	14:00	AS00210	-0.15	3.90	X
LH18/24-Air_022719_GWTP	P1901179-003	Air	2/27/2019	14:15	AS01007	-0.47	3.56	X
LH18/24-Air_022719_Downwind	P1901179-004	Air	2/28/2019	06:30	AS01290	1.24	3.77	X

**ALS Environmental
Sample Acceptance Check Form**

Client: Bhate Environmental Associates, Inc. Work order: P1901179
 Project: LHAAP GWTP / NW01312.0150
 Sample(s) received on: 3/5/19 Date opened: 3/5/19 by: ADAVID

Note: This form is used for all samples received by ALS. The use of this form for custody seals is strictly meant to indicate presence/absence and not as an indication of compliance or nonconformity. Thermal preservation and pH will only be evaluated either at the request of the client and/or as required by the method/SOP.

- | | Yes | No | N/A |
|---|-------------------------------------|-------------------------------------|-------------------------------------|
| 1 Were sample containers properly marked with client sample ID? | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 2 Did sample containers arrive in good condition? | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 3 Were chain-of-custody papers used and filled out? | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 4 Did sample container labels and/or tags agree with custody papers? | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 5 Was sample volume received adequate for analysis? | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 6 Are samples within specified holding times? | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> |
| 7 Was proper temperature (thermal preservation) of cooler at receipt adhered to? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 8 Were custody seals on outside of cooler/Box/Container? | <input type="checkbox"/> | <input checked="" type="checkbox"/> | <input type="checkbox"/> |
| Location of seal(s)? _____ Sealing Lid? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Were signature and date included? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Were seals intact? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 9 Do containers have appropriate preservation , according to method/SOP or Client specified information? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Is there a client indication that the submitted samples are pH preserved? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Were VOA vials checked for presence/absence of air bubbles? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Does the client/method/SOP require that the analyst check the sample pH and <u>if necessary</u> alter it? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 10 Tubes: Are the tubes capped and intact? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| 11 Badges: Are the badges properly capped and intact? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |
| Are dual bed badges separated and individually capped and intact? | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> |

Lab Sample ID	Container Description	Required pH *	Received pH	Adjusted pH	VOA Headspace (Presence/Absence)	Receipt / Preservation Comments
P1901179-001.01	6.0 L Silonite Can					
P1901179-002.01	6.0 L Silonite Can					
P1901179-003.01	6.0 L Silonite Can					
P1901179-004.01	6.0 L Silonite Can					

Explain any discrepancies: (include lab sample ID numbers): _____

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RESULTS OF ANALYSIS

Page 1 of 3

Client: **Bhate Environmental Associates, Inc.**Client Sample ID: **LH18/24-Air_022719_Stripper**Client Project ID: **LHAAP GWTP / NW01312.0150**

ALS Project ID: P1901179

ALS Sample ID: P1901179-001

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00629

Initial Pressure (psig): -0.03 Final Pressure (psig): 3.53

Container Dilution Factor: 1.24

CAS #	Compound	Result	MRL	Result	MRL	Data Qualifier
		µg/m ³	µg/m ³	ppbV	ppbV	
115-07-1	Propene	ND	64	ND	37	
75-71-8	Dichlorodifluoromethane (CFC 12)	ND	64	ND	13	
74-87-3	Chloromethane	ND	62	ND	30	
76-14-2	1,2-Dichloro-1,1,2,2-tetrafluoroethane (CFC 114)	ND	63	ND	9.1	
75-01-4	Vinyl Chloride	ND	66	ND	26	
106-99-0	1,3-Butadiene	ND	64	ND	29	
74-83-9	Bromomethane	ND	62	ND	16	
75-00-3	Chloroethane	ND	63	ND	24	
64-17-5	Ethanol	ND	630	ND	340	
75-05-8	Acetonitrile	ND	64	ND	38	
107-02-8	Acrolein	ND	120	ND	54	
67-64-1	Acetone	ND	670	ND	280	
75-69-4	Trichlorofluoromethane (CFC 11)	ND	66	ND	12	
67-63-0	2-Propanol (Isopropyl Alcohol)	ND	260	ND	110	
107-13-1	Acrylonitrile	ND	64	ND	30	
75-35-4	1,1-Dichloroethene	ND	67	ND	17	
75-09-2	Methylene Chloride	580	67	170	19	
107-05-1	3-Chloro-1-propene (Allyl Chloride)	ND	66	ND	21	
76-13-1	Trichlorotrifluoroethane (CFC 113)	1,800	66	240	8.6	
75-15-0	Carbon Disulfide	ND	140	ND	44	
156-60-5	trans-1,2-Dichloroethene	ND	66	ND	17	
75-34-3	1,1-Dichloroethane	ND	64	ND	16	
1634-04-4	Methyl tert-Butyl Ether	ND	67	ND	19	
108-05-4	Vinyl Acetate	ND	660	ND	190	
78-93-3	2-Butanone (MEK)	ND	120	ND	42	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

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Client: **Bhate Environmental Associates, Inc.**Client Sample ID: **LH18/24-Air_022719_Stripper**Client Project ID: **LHAAP GWTP / NW01312.0150**

ALS Project ID: P1901179

ALS Sample ID: P1901179-001

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00629

Initial Pressure (psig): -0.03 Final Pressure (psig): 3.53

Container Dilution Factor: 1.24

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
156-59-2	cis-1,2-Dichloroethene	2,700	66	690	17	
141-78-6	Ethyl Acetate	ND	140	ND	38	
110-54-3	n-Hexane	ND	67	ND	19	
67-66-3	Chloroform	ND	67	ND	14	
109-99-9	Tetrahydrofuran (THF)	ND	66	ND	22	
107-06-2	1,2-Dichloroethane	140	66	34	16	
71-55-6	1,1,1-Trichloroethane	ND	67	ND	12	
71-43-2	Benzene	ND	64	ND	20	
56-23-5	Carbon Tetrachloride	ND	64	ND	10	
110-82-7	Cyclohexane	ND	120	ND	36	
78-87-5	1,2-Dichloropropane	ND	67	ND	14	
75-27-4	Bromodichloromethane	ND	66	ND	9.8	
79-01-6	Trichloroethene	11,000	66	2,100	12	
123-91-1	1,4-Dioxane	ND	66	ND	18	
80-62-6	Methyl Methacrylate	ND	140	ND	33	
142-82-5	n-Heptane	ND	67	ND	16	
10061-01-5	cis-1,3-Dichloropropene	ND	69	ND	15	
108-10-1	4-Methyl-2-pentanone	ND	66	ND	16	
10061-02-6	trans-1,3-Dichloropropene	ND	66	ND	14	
79-00-5	1,1,2-Trichloroethane	ND	67	ND	12	
108-88-3	Toluene	ND	66	ND	17	
591-78-6	2-Hexanone	ND	67	ND	16	
124-48-1	Dibromochloromethane	ND	67	ND	7.9	
106-93-4	1,2-Dibromoethane	ND	67	ND	8.7	
123-86-4	n-Butyl Acetate	ND	67	ND	14	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

Page 3 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_Stripper**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-001

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00629

Initial Pressure (psig): -0.03 Final Pressure (psig): 3.53

Container Dilution Factor: 1.24

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
111-65-9	n-Octane	ND	67	ND	14	
127-18-4	Tetrachloroethene	ND	66	ND	9.7	
108-90-7	Chlorobenzene	ND	66	ND	14	
100-41-4	Ethylbenzene	ND	64	ND	15	
179601-23-1	m,p-Xylenes	ND	140	ND	31	
75-25-2	Bromoform	ND	66	ND	6.4	
100-42-5	Styrene	ND	66	ND	15	
95-47-6	o-Xylene	ND	66	ND	15	
111-84-2	n-Nonane	ND	67	ND	13	
79-34-5	1,1,2,2-Tetrachloroethane	ND	66	ND	9.6	
98-82-8	Cumene	ND	66	ND	13	
80-56-8	alpha-Pinene	ND	64	ND	12	
103-65-1	n-Propylbenzene	ND	67	ND	14	
622-96-8	4-Ethyltoluene	ND	66	ND	13	
108-67-8	1,3,5-Trimethylbenzene	ND	66	ND	13	
95-63-6	1,2,4-Trimethylbenzene	ND	66	ND	13	
100-44-7	Benzyl Chloride	ND	140	ND	26	
541-73-1	1,3-Dichlorobenzene	ND	67	ND	11	
106-46-7	1,4-Dichlorobenzene	ND	67	ND	11	
95-50-1	1,2-Dichlorobenzene	ND	67	ND	11	
5989-27-5	d-Limonene	ND	63	ND	11	
96-12-8	1,2-Dibromo-3-chloropropane	ND	64	ND	6.7	
120-82-1	1,2,4-Trichlorobenzene	ND	66	ND	8.9	
91-20-3	Naphthalene	ND	63	ND	12	
87-68-3	Hexachlorobutadiene	ND	66	ND	6.2	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

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Client: **Bhate Environmental Associates, Inc.**Client Sample ID: **LH18/24-Air_022719_Stripper_a**Client Project ID: **LHAAP GWTP / NW01312.0150**

ALS Project ID: P1901179

ALS Sample ID: P1901179-002

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00210

Initial Pressure (psig): -0.15 Final Pressure (psig): 3.90

Container Dilution Factor: 1.28

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
115-07-1	Propene	ND	67	ND	39	
75-71-8	Dichlorodifluoromethane (CFC 12)	ND	67	ND	13	
74-87-3	Chloromethane	ND	64	ND	31	
76-14-2	1,2-Dichloro-1,1,2,2-tetrafluoroethane (CFC 114)	ND	65	ND	9.3	
75-01-4	Vinyl Chloride	ND	68	ND	27	
106-99-0	1,3-Butadiene	ND	67	ND	30	
74-83-9	Bromomethane	ND	64	ND	16	
75-00-3	Chloroethane	ND	65	ND	25	
64-17-5	Ethanol	ND	650	ND	350	
75-05-8	Acetonitrile	ND	67	ND	40	
107-02-8	Acrolein	ND	130	ND	56	
67-64-1	Acetone	ND	690	ND	290	
75-69-4	Trichlorofluoromethane (CFC 11)	ND	68	ND	12	
67-63-0	2-Propanol (Isopropyl Alcohol)	ND	270	ND	110	
107-13-1	Acrylonitrile	ND	67	ND	31	
75-35-4	1,1-Dichloroethene	ND	69	ND	17	
75-09-2	Methylene Chloride	630	69	180	20	
107-05-1	3-Chloro-1-propene (Allyl Chloride)	ND	68	ND	22	
76-13-1	Trichlorotrifluoroethane (CFC 113)	1,900	68	240	8.9	
75-15-0	Carbon Disulfide	ND	140	ND	45	
156-60-5	trans-1,2-Dichloroethene	ND	68	ND	17	
75-34-3	1,1-Dichloroethane	ND	67	ND	16	
1634-04-4	Methyl tert-Butyl Ether	ND	69	ND	19	
108-05-4	Vinyl Acetate	ND	680	ND	190	
78-93-3	2-Butanone (MEK)	ND	130	ND	43	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

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Client: **Bhate Environmental Associates, Inc.**Client Sample ID: **LH18/24-Air_022719_Stripper_a**Client Project ID: **LHAAP GWTP / NW01312.0150**

ALS Project ID: P1901179

ALS Sample ID: P1901179-002

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00210

Initial Pressure (psig): -0.15 Final Pressure (psig): 3.90

Container Dilution Factor: 1.28

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
156-59-2	cis-1,2-Dichloroethene	2,800	68	710	17	
141-78-6	Ethyl Acetate	ND	140	ND	39	
110-54-3	n-Hexane	ND	69	ND	20	
67-66-3	Chloroform	ND	69	ND	14	
109-99-9	Tetrahydrofuran (THF)	ND	68	ND	23	
107-06-2	1,2-Dichloroethane	140	68	35	17	
71-55-6	1,1,1-Trichloroethane	ND	69	ND	13	
71-43-2	Benzene	ND	67	ND	21	
56-23-5	Carbon Tetrachloride	ND	67	ND	11	
110-82-7	Cyclohexane	ND	130	ND	37	
78-87-5	1,2-Dichloropropane	ND	69	ND	15	
75-27-4	Bromodichloromethane	ND	68	ND	10	
79-01-6	Trichloroethene	12,000	68	2,200	13	
123-91-1	1,4-Dioxane	ND	68	ND	19	
80-62-6	Methyl Methacrylate	ND	140	ND	34	
142-82-5	n-Heptane	ND	69	ND	17	
10061-01-5	cis-1,3-Dichloropropene	ND	72	ND	16	
108-10-1	4-Methyl-2-pentanone	ND	68	ND	17	
10061-02-6	trans-1,3-Dichloropropene	ND	68	ND	15	
79-00-5	1,1,2-Trichloroethane	ND	69	ND	13	
108-88-3	Toluene	ND	68	ND	18	
591-78-6	2-Hexanone	ND	69	ND	17	
124-48-1	Dibromochloromethane	ND	69	ND	8.1	
106-93-4	1,2-Dibromoethane	ND	69	ND	9.0	
123-86-4	n-Butyl Acetate	ND	69	ND	15	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

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RESULTS OF ANALYSIS

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Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_Stripper_a**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-002

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00210

Initial Pressure (psig): -0.15 Final Pressure (psig): 3.90

Container Dilution Factor: 1.28

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
111-65-9	n-Octane	ND	69	ND	15	
127-18-4	Tetrachloroethene	ND	68	ND	10	
108-90-7	Chlorobenzene	ND	68	ND	15	
100-41-4	Ethylbenzene	ND	67	ND	15	
179601-23-1	m,p-Xylenes	ND	140	ND	32	
75-25-2	Bromoform	ND	68	ND	6.6	
100-42-5	Styrene	ND	68	ND	16	
95-47-6	o-Xylene	ND	68	ND	16	
111-84-2	n-Nonane	ND	69	ND	13	
79-34-5	1,1,2,2-Tetrachloroethane	ND	68	ND	9.9	
98-82-8	Cumene	ND	68	ND	14	
80-56-8	alpha-Pinene	ND	67	ND	12	
103-65-1	n-Propylbenzene	ND	69	ND	14	
622-96-8	4-Ethyltoluene	ND	68	ND	14	
108-67-8	1,3,5-Trimethylbenzene	ND	68	ND	14	
95-63-6	1,2,4-Trimethylbenzene	ND	68	ND	14	
100-44-7	Benzyl Chloride	ND	140	ND	27	
541-73-1	1,3-Dichlorobenzene	ND	69	ND	12	
106-46-7	1,4-Dichlorobenzene	ND	69	ND	12	
95-50-1	1,2-Dichlorobenzene	ND	69	ND	12	
5989-27-5	d-Limonene	ND	65	ND	12	
96-12-8	1,2-Dibromo-3-chloropropane	ND	67	ND	6.9	
120-82-1	1,2,4-Trichlorobenzene	ND	68	ND	9.1	
91-20-3	Naphthalene	ND	65	ND	12	
87-68-3	Hexachlorobutadiene	ND	68	ND	6.4	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

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Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_GWTP**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-003

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 1.00 Liter(s)

Test Notes:

Container ID: AS01007

Initial Pressure (psig): -0.47 Final Pressure (psig): 3.56

Container Dilution Factor: 1.28

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
115-07-1	Propene	ND	0.67	ND	0.39	
75-71-8	Dichlorodifluoromethane (CFC 12)	2.5	0.67	0.52	0.13	
74-87-3	Chloromethane	ND	0.64	ND	0.31	
76-14-2	1,2-Dichloro-1,1,2,2-tetrafluoroethane (CFC 114)	ND	0.65	ND	0.093	
75-01-4	Vinyl Chloride	ND	0.68	ND	0.27	
106-99-0	1,3-Butadiene	ND	0.67	ND	0.30	
74-83-9	Bromomethane	ND	0.64	ND	0.16	
75-00-3	Chloroethane	ND	0.65	ND	0.25	
64-17-5	Ethanol	ND	6.5	ND	3.5	
75-05-8	Acetonitrile	ND	0.67	ND	0.40	
107-02-8	Acrolein	ND	1.3	ND	0.56	
67-64-1	Acetone	9.0	6.9	3.8	2.9	
75-69-4	Trichlorofluoromethane (CFC 11)	1.4	0.68	0.24	0.12	
67-63-0	2-Propanol (Isopropyl Alcohol)	ND	2.7	ND	1.1	
107-13-1	Acrylonitrile	ND	0.67	ND	0.31	
75-35-4	1,1-Dichloroethene	ND	0.69	ND	0.17	
75-09-2	Methylene Chloride	2.8	0.69	0.82	0.20	
107-05-1	3-Chloro-1-propene (Allyl Chloride)	ND	0.68	ND	0.22	
76-13-1	Trichlorotrifluoroethane (CFC 113)	19	0.68	2.5	0.089	
75-15-0	Carbon Disulfide	ND	1.4	ND	0.45	
156-60-5	trans-1,2-Dichloroethene	ND	0.68	ND	0.17	
75-34-3	1,1-Dichloroethane	ND	0.67	ND	0.16	
1634-04-4	Methyl tert-Butyl Ether	ND	0.69	ND	0.19	
108-05-4	Vinyl Acetate	ND	6.8	ND	1.9	
78-93-3	2-Butanone (MEK)	1.5	1.3	0.52	0.43	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

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Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_GWTP**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-003

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 1.00 Liter(s)

Test Notes:

Container ID: AS01007

Initial Pressure (psig): -0.47 Final Pressure (psig): 3.56

Container Dilution Factor: 1.28

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
156-59-2	cis-1,2-Dichloroethene	12	0.68	3.1	0.17	
141-78-6	Ethyl Acetate	5.7	1.4	1.6	0.39	
110-54-3	n-Hexane	1.0	0.69	0.29	0.20	
67-66-3	Chloroform	ND	0.69	ND	0.14	
109-99-9	Tetrahydrofuran (THF)	ND	0.68	ND	0.23	
107-06-2	1,2-Dichloroethane	ND	0.68	ND	0.17	
71-55-6	1,1,1-Trichloroethane	ND	0.69	ND	0.13	
71-43-2	Benzene	0.80	0.67	0.25	0.21	
56-23-5	Carbon Tetrachloride	ND	0.67	ND	0.11	
110-82-7	Cyclohexane	ND	1.3	ND	0.37	
78-87-5	1,2-Dichloropropane	ND	0.69	ND	0.15	
75-27-4	Bromodichloromethane	ND	0.68	ND	0.10	
79-01-6	Trichloroethene	55	0.68	10	0.13	
123-91-1	1,4-Dioxane	ND	0.68	ND	0.19	
80-62-6	Methyl Methacrylate	ND	1.4	ND	0.34	
142-82-5	n-Heptane	ND	0.69	ND	0.17	
10061-01-5	cis-1,3-Dichloropropene	ND	0.72	ND	0.16	
108-10-1	4-Methyl-2-pentanone	ND	0.68	ND	0.17	
10061-02-6	trans-1,3-Dichloropropene	ND	0.68	ND	0.15	
79-00-5	1,1,2-Trichloroethane	ND	0.69	ND	0.13	
108-88-3	Toluene	1.2	0.68	0.31	0.18	
591-78-6	2-Hexanone	ND	0.69	ND	0.17	
124-48-1	Dibromochloromethane	ND	0.69	ND	0.081	
106-93-4	1,2-Dibromoethane	ND	0.69	ND	0.090	
123-86-4	n-Butyl Acetate	ND	0.69	ND	0.15	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

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Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_GWTP**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-003

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 1.00 Liter(s)

Test Notes:

Container ID: AS01007

Initial Pressure (psig): -0.47 Final Pressure (psig): 3.56

Container Dilution Factor: 1.28

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
111-65-9	n-Octane	ND	0.69	ND	0.15	
127-18-4	Tetrachloroethene	ND	0.68	ND	0.10	
108-90-7	Chlorobenzene	ND	0.68	ND	0.15	
100-41-4	Ethylbenzene	ND	0.67	ND	0.15	
179601-23-1	m,p-Xylenes	ND	1.4	ND	0.32	
75-25-2	Bromoform	ND	0.68	ND	0.066	
100-42-5	Styrene	ND	0.68	ND	0.16	
95-47-6	o-Xylene	ND	0.68	ND	0.16	
111-84-2	n-Nonane	ND	0.69	ND	0.13	
79-34-5	1,1,2,2-Tetrachloroethane	ND	0.68	ND	0.099	
98-82-8	Cumene	ND	0.68	ND	0.14	
80-56-8	alpha-Pinene	ND	0.67	ND	0.12	
103-65-1	n-Propylbenzene	ND	0.69	ND	0.14	
622-96-8	4-Ethyltoluene	ND	0.68	ND	0.14	
108-67-8	1,3,5-Trimethylbenzene	ND	0.68	ND	0.14	
95-63-6	1,2,4-Trimethylbenzene	ND	0.68	ND	0.14	
100-44-7	Benzyl Chloride	ND	1.4	ND	0.27	
541-73-1	1,3-Dichlorobenzene	ND	0.69	ND	0.12	
106-46-7	1,4-Dichlorobenzene	ND	0.69	ND	0.12	
95-50-1	1,2-Dichlorobenzene	ND	0.69	ND	0.12	
5989-27-5	d-Limonene	ND	0.65	ND	0.12	
96-12-8	1,2-Dibromo-3-chloropropane	ND	0.67	ND	0.069	
120-82-1	1,2,4-Trichlorobenzene	ND	0.68	ND	0.091	
91-20-3	Naphthalene	ND	0.65	ND	0.12	
87-68-3	Hexachlorobutadiene	ND	0.68	ND	0.064	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

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Client: **Bhate Environmental Associates, Inc.**Client Sample ID: **LH18/24-Air_022719_Downwind**Client Project ID: **LHAAP GWTP / NW01312.0150**

ALS Project ID: P1901179

ALS Sample ID: P1901179-004

Test Code: EPA TO-15

Date Collected: 2/28/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 1.00 Liter(s)

Test Notes:

Container ID: AS01290

Initial Pressure (psig): 1.24 Final Pressure (psig): 3.77

Container Dilution Factor: 1.16

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
115-07-1	Propene	ND	0.60	ND	0.35	
75-71-8	Dichlorodifluoromethane (CFC 12)	2.6	0.60	0.52	0.12	
74-87-3	Chloromethane	ND	0.58	ND	0.28	
76-14-2	1,2-Dichloro-1,1,2,2-tetrafluoroethane (CFC 114)	ND	0.59	ND	0.085	
75-01-4	Vinyl Chloride	ND	0.61	ND	0.24	
106-99-0	1,3-Butadiene	ND	0.60	ND	0.27	
74-83-9	Bromomethane	ND	0.58	ND	0.15	
75-00-3	Chloroethane	ND	0.59	ND	0.22	
64-17-5	Ethanol	ND	5.9	ND	3.1	
75-05-8	Acetonitrile	ND	0.60	ND	0.36	
107-02-8	Acrolein	ND	1.2	ND	0.51	
67-64-1	Acetone	ND	6.3	ND	2.6	
75-69-4	Trichlorofluoromethane (CFC 11)	1.3	0.61	0.24	0.11	
67-63-0	2-Propanol (Isopropyl Alcohol)	ND	2.4	ND	0.99	
107-13-1	Acrylonitrile	ND	0.60	ND	0.28	
75-35-4	1,1-Dichloroethene	ND	0.63	ND	0.16	
75-09-2	Methylene Chloride	1.4	0.63	0.40	0.18	
107-05-1	3-Chloro-1-propene (Allyl Chloride)	ND	0.61	ND	0.20	
76-13-1	Trichlorotrifluoroethane (CFC 113)	7.0	0.61	0.91	0.080	
75-15-0	Carbon Disulfide	ND	1.3	ND	0.41	
156-60-5	trans-1,2-Dichloroethene	ND	0.61	ND	0.16	
75-34-3	1,1-Dichloroethane	ND	0.60	ND	0.15	
1634-04-4	Methyl tert-Butyl Ether	ND	0.63	ND	0.17	
108-05-4	Vinyl Acetate	ND	6.1	ND	1.7	
78-93-3	2-Butanone (MEK)	ND	1.2	ND	0.39	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

Page 2 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_Downwind**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-004

Test Code: EPA TO-15

Date Collected: 2/28/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 1.00 Liter(s)

Test Notes:

Container ID: AS01290

Initial Pressure (psig): 1.24 Final Pressure (psig): 3.77

Container Dilution Factor: 1.16

CAS #	Compound	Result	MRL	Result	MRL	Data Qualifier
		$\mu\text{g}/\text{m}^3$	$\mu\text{g}/\text{m}^3$	ppbV	ppbV	
156-59-2	cis-1,2-Dichloroethene	0.64	0.61	0.16	0.16	
141-78-6	Ethyl Acetate	2.7	1.3	0.76	0.35	
110-54-3	n-Hexane	0.83	0.63	0.23	0.18	
67-66-3	Chloroform	ND	0.63	ND	0.13	
109-99-9	Tetrahydrofuran (THF)	ND	0.61	ND	0.21	
107-06-2	1,2-Dichloroethane	ND	0.61	ND	0.15	
71-55-6	1,1,1-Trichloroethane	ND	0.63	ND	0.11	
71-43-2	Benzene	0.69	0.60	0.22	0.19	
56-23-5	Carbon Tetrachloride	ND	0.60	ND	0.096	
110-82-7	Cyclohexane	ND	1.2	ND	0.34	
78-87-5	1,2-Dichloropropane	ND	0.63	ND	0.14	
75-27-4	Bromodichloromethane	ND	0.61	ND	0.092	
79-01-6	Trichloroethene	2.1	0.61	0.39	0.11	
123-91-1	1,4-Dioxane	ND	0.61	ND	0.17	
80-62-6	Methyl Methacrylate	ND	1.3	ND	0.31	
142-82-5	n-Heptane	ND	0.63	ND	0.15	
10061-01-5	cis-1,3-Dichloropropene	ND	0.65	ND	0.14	
108-10-1	4-Methyl-2-pentanone	ND	0.61	ND	0.15	
10061-02-6	trans-1,3-Dichloropropene	ND	0.61	ND	0.14	
79-00-5	1,1,2-Trichloroethane	ND	0.63	ND	0.11	
108-88-3	Toluene	0.76	0.61	0.20	0.16	
591-78-6	2-Hexanone	ND	0.63	ND	0.15	
124-48-1	Dibromochloromethane	ND	0.63	ND	0.074	
106-93-4	1,2-Dibromoethane	ND	0.63	ND	0.082	
123-86-4	n-Butyl Acetate	ND	0.63	ND	0.13	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

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Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_Downwind**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-004

Test Code: EPA TO-15

Date Collected: 2/28/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 1.00 Liter(s)

Test Notes:

Container ID: AS01290

Initial Pressure (psig): 1.24 Final Pressure (psig): 3.77

Container Dilution Factor: 1.16

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
111-65-9	n-Octane	ND	0.63	ND	0.13	
127-18-4	Tetrachloroethene	ND	0.61	ND	0.091	
108-90-7	Chlorobenzene	ND	0.61	ND	0.13	
100-41-4	Ethylbenzene	ND	0.60	ND	0.14	
179601-23-1	m,p-Xylenes	ND	1.3	ND	0.29	
75-25-2	Bromoform	ND	0.61	ND	0.059	
100-42-5	Styrene	ND	0.61	ND	0.14	
95-47-6	o-Xylene	ND	0.61	ND	0.14	
111-84-2	n-Nonane	ND	0.63	ND	0.12	
79-34-5	1,1,2,2-Tetrachloroethane	ND	0.61	ND	0.090	
98-82-8	Cumene	ND	0.61	ND	0.13	
80-56-8	alpha-Pinene	ND	0.60	ND	0.11	
103-65-1	n-Propylbenzene	ND	0.63	ND	0.13	
622-96-8	4-Ethyltoluene	ND	0.61	ND	0.13	
108-67-8	1,3,5-Trimethylbenzene	ND	0.61	ND	0.13	
95-63-6	1,2,4-Trimethylbenzene	ND	0.61	ND	0.13	
100-44-7	Benzyl Chloride	ND	1.3	ND	0.25	
541-73-1	1,3-Dichlorobenzene	ND	0.63	ND	0.10	
106-46-7	1,4-Dichlorobenzene	ND	0.63	ND	0.10	
95-50-1	1,2-Dichlorobenzene	ND	0.63	ND	0.10	
5989-27-5	d-Limonene	ND	0.59	ND	0.11	
96-12-8	1,2-Dibromo-3-chloropropane	ND	0.60	ND	0.062	
120-82-1	1,2,4-Trichlorobenzene	ND	0.61	ND	0.083	
91-20-3	Naphthalene	ND	0.59	ND	0.11	
87-68-3	Hexachlorobutadiene	ND	0.61	ND	0.058	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

Page 1 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** Method Blank**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P190311-MB

Test Code: EPA TO-15

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Analyst: Wida Ang

Sample Type: 6.0 L Silonite Canister

Test Notes:

Date Collected: NA

Date Received: NA

Date Analyzed: 3/11/19

Volume(s) Analyzed: 1.00 Liter(s)

Container Dilution Factor: 1.00

CAS #	Compound	Result	MRL	Result	MRL	Data Qualifier
		µg/m ³	µg/m ³	ppbV	ppbV	
115-07-1	Propene	ND	0.52	ND	0.30	
75-71-8	Dichlorodifluoromethane (CFC 12)	ND	0.52	ND	0.11	
74-87-3	Chloromethane	ND	0.50	ND	0.24	
76-14-2	1,2-Dichloro-1,1,2,2-tetrafluoroethane (CFC 114)	ND	0.51	ND	0.073	
75-01-4	Vinyl Chloride	ND	0.53	ND	0.21	
106-99-0	1,3-Butadiene	ND	0.52	ND	0.24	
74-83-9	Bromomethane	ND	0.50	ND	0.13	
75-00-3	Chloroethane	ND	0.51	ND	0.19	
64-17-5	Ethanol	ND	5.1	ND	2.7	
75-05-8	Acetonitrile	ND	0.52	ND	0.31	
107-02-8	Acrolein	ND	1.0	ND	0.44	
67-64-1	Acetone	ND	5.4	ND	2.3	
75-69-4	Trichlorofluoromethane (CFC 11)	ND	0.53	ND	0.094	
67-63-0	2-Propanol (Isopropyl Alcohol)	ND	2.1	ND	0.85	
107-13-1	Acrylonitrile	ND	0.52	ND	0.24	
75-35-4	1,1-Dichloroethene	ND	0.54	ND	0.14	
75-09-2	Methylene Chloride	ND	0.54	ND	0.16	
107-05-1	3-Chloro-1-propene (Allyl Chloride)	ND	0.53	ND	0.17	
76-13-1	Trichlorotrifluoroethane (CFC 113)	ND	0.53	ND	0.069	
75-15-0	Carbon Disulfide	ND	1.1	ND	0.35	
156-60-5	trans-1,2-Dichloroethene	ND	0.53	ND	0.13	
75-34-3	1,1-Dichloroethane	ND	0.52	ND	0.13	
1634-04-4	Methyl tert-Butyl Ether	ND	0.54	ND	0.15	
108-05-4	Vinyl Acetate	ND	5.3	ND	1.5	
78-93-3	2-Butanone (MEK)	ND	1.0	ND	0.34	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

Page 2 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** Method Blank**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P190311-MB

Test Code: EPA TO-15
 Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13
 Analyst: Wida Ang
 Sample Type: 6.0 L Silonite Canister
 Test Notes:

Date Collected: NA
 Date Received: NA
 Date Analyzed: 3/11/19
 Volume(s) Analyzed: 1.00 Liter(s)

Container Dilution Factor: 1.00

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
156-59-2	cis-1,2-Dichloroethene	ND	0.53	ND	0.13	
141-78-6	Ethyl Acetate	ND	1.1	ND	0.31	
110-54-3	n-Hexane	ND	0.54	ND	0.15	
67-66-3	Chloroform	ND	0.54	ND	0.11	
109-99-9	Tetrahydrofuran (THF)	ND	0.53	ND	0.18	
107-06-2	1,2-Dichloroethane	ND	0.53	ND	0.13	
71-55-6	1,1,1-Trichloroethane	ND	0.54	ND	0.099	
71-43-2	Benzene	ND	0.52	ND	0.16	
56-23-5	Carbon Tetrachloride	ND	0.52	ND	0.083	
110-82-7	Cyclohexane	ND	1.0	ND	0.29	
78-87-5	1,2-Dichloropropane	ND	0.54	ND	0.12	
75-27-4	Bromodichloromethane	ND	0.53	ND	0.079	
79-01-6	Trichloroethene	ND	0.53	ND	0.099	
123-91-1	1,4-Dioxane	ND	0.53	ND	0.15	
80-62-6	Methyl Methacrylate	ND	1.1	ND	0.27	
142-82-5	n-Heptane	ND	0.54	ND	0.13	
10061-01-5	cis-1,3-Dichloropropene	ND	0.56	ND	0.12	
108-10-1	4-Methyl-2-pentanone	ND	0.53	ND	0.13	
10061-02-6	trans-1,3-Dichloropropene	ND	0.53	ND	0.12	
79-00-5	1,1,2-Trichloroethane	ND	0.54	ND	0.099	
108-88-3	Toluene	ND	0.53	ND	0.14	
591-78-6	2-Hexanone	ND	0.54	ND	0.13	
124-48-1	Dibromochloromethane	ND	0.54	ND	0.063	
106-93-4	1,2-Dibromoethane	ND	0.54	ND	0.070	
123-86-4	n-Butyl Acetate	ND	0.54	ND	0.11	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

Page 3 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** Method Blank**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P190311-MB

Test Code: EPA TO-15

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Analyst: Wida Ang

Sample Type: 6.0 L Silonite Canister

Test Notes:

Date Collected: NA

Date Received: NA

Date Analyzed: 3/11/19

Volume(s) Analyzed: 1.00 Liter(s)

Container Dilution Factor: 1.00

CAS #	Compound	Result µg/m ³	MRL µg/m ³	Result ppbV	MRL ppbV	Data Qualifier
111-65-9	n-Octane	ND	0.54	ND	0.12	
127-18-4	Tetrachloroethene	ND	0.53	ND	0.078	
108-90-7	Chlorobenzene	ND	0.53	ND	0.12	
100-41-4	Ethylbenzene	ND	0.52	ND	0.12	
179601-23-1	m,p-Xylenes	ND	1.1	ND	0.25	
75-25-2	Bromoform	ND	0.53	ND	0.051	
100-42-5	Styrene	ND	0.53	ND	0.12	
95-47-6	o-Xylene	ND	0.53	ND	0.12	
111-84-2	n-Nonane	ND	0.54	ND	0.10	
79-34-5	1,1,2,2-Tetrachloroethane	ND	0.53	ND	0.077	
98-82-8	Cumene	ND	0.53	ND	0.11	
80-56-8	alpha-Pinene	ND	0.52	ND	0.093	
103-65-1	n-Propylbenzene	ND	0.54	ND	0.11	
622-96-8	4-Ethyltoluene	ND	0.53	ND	0.11	
108-67-8	1,3,5-Trimethylbenzene	ND	0.53	ND	0.11	
95-63-6	1,2,4-Trimethylbenzene	ND	0.53	ND	0.11	
100-44-7	Benzyl Chloride	ND	1.1	ND	0.21	
541-73-1	1,3-Dichlorobenzene	ND	0.54	ND	0.090	
106-46-7	1,4-Dichlorobenzene	ND	0.54	ND	0.090	
95-50-1	1,2-Dichlorobenzene	ND	0.54	ND	0.090	
5989-27-5	d-Limonene	ND	0.51	ND	0.092	
96-12-8	1,2-Dibromo-3-chloropropane	ND	0.52	ND	0.054	
120-82-1	1,2,4-Trichlorobenzene	ND	0.53	ND	0.071	
91-20-3	Naphthalene	ND	0.51	ND	0.097	
87-68-3	Hexachlorobutadiene	ND	0.53	ND	0.050	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

MRL = Method Reporting Limit - The minimum quantity of a target analyte that can be confidently determined by the referenced method.

ALS ENVIRONMENTAL

SURROGATE SPIKE RECOVERY RESULTS

Page 1 of 1

Client: Bhate Environmental Associates, Inc.
Client Project ID: LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

Test Code: EPA TO-15
Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13
Analyst: Wida Ang
Sample Type: 6.0 L Silonite Canister(s)
Test Notes:

Date(s) Collected: 2/27 - 2/28/19

Date(s) Received: 3/5/19

Date(s) Analyzed: 3/11/19

Client Sample ID	ALS Sample ID	1,2-Dichloroethane-d4	Toluene-d8	Bromofluorobenzene	Acceptance Limits	Data Qualifier
		Percent Recovered	Percent Recovered	Percent Recovered		
Method Blank	P190311-MB	112	96	99	70-130	
Lab Control Sample	P190311-LCS	108	94	99	70-130	
LH18/24-Air_022719_Stripper	P1901179-001	112	95	98	70-130	
LH18/24-Air_022719_Stripper_a	P1901179-002	113	94	98	70-130	
LH18/24-Air_022719_Stripper_a	P1901179-002DUP	114	94	98	70-130	
LH18/24-Air_022719_GWTP	P1901179-003	112	94	98	70-130	
LH18/24-Air_022719_Downwind	P1901179-004	112	94	98	70-130	

Surrogate percent recovery is verified and accepted based on the on-column result.

Reported results are shown in concentration units and as a result of the calculation, may vary slightly from the on-column percent recovery.

ALS ENVIRONMENTAL

LABORATORY CONTROL SAMPLE SUMMARY

Page 1 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** Lab Control Sample**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P190311-LCS

Test Code: EPA TO-15

Date Collected: NA

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: NA

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.125 Liter(s)

Test Notes:

CAS #	Compound	Spike Amount µg/m ³	Result µg/m ³	% Recovery	ALS	Data Qualifier
					Acceptance Limits	
115-07-1	Propene	211	215	102	53-112	
75-71-8	Dichlorodifluoromethane (CFC 12)	210	210	100	62-103	
74-87-3	Chloromethane	211	234	111	51-121	
76-14-2	1,2-Dichloro-1,1,2,2-tetrafluoroethane (CFC 114)	211	206	98	56-111	
75-01-4	Vinyl Chloride	214	236	110	57-117	
106-99-0	1,3-Butadiene	210	243	116	53-134	
74-83-9	Bromomethane	212	217	102	65-110	
75-00-3	Chloroethane	214	218	102	64-111	
64-17-5	Ethanol	1,020	1080	106	57-124	
75-05-8	Acetonitrile	206	227	110	57-126	
107-02-8	Acrolein	205	234	114	62-121	
67-64-1	Acetone	1,060	1030	97	60-113	
75-69-4	Trichlorofluoromethane (CFC 11)	211	211	100	63-104	
67-63-0	2-Propanol (Isopropyl Alcohol)	413	449	109	60-124	
107-13-1	Acrylonitrile	207	259	125	66-125	
75-35-4	1,1-Dichloroethene	218	218	100	68-107	
75-09-2	Methylene Chloride	217	228	105	66-105	
107-05-1	3-Chloro-1-propene (Allyl Chloride)	216	245	113	63-127	
76-13-1	Trichlorotrifluoroethane (CFC 113)	216	204	94	59-109	
75-15-0	Carbon Disulfide	218	203	93	67-109	
156-60-5	trans-1,2-Dichloroethene	214	242	113	70-115	
75-34-3	1,1-Dichloroethane	216	218	101	66-106	
1634-04-4	Methyl tert-Butyl Ether	214	217	101	67-109	
108-05-4	Vinyl Acetate	1,060	1140	108	68-136	
78-93-3	2-Butanone (MEK)	208	218	105	71-116	

Laboratory Control Sample percent recovery is verified and accepted based on the on-column result. Reported results are shown in concentration units and as a result of the calculation, may vary slightly.

ALS ENVIRONMENTAL

LABORATORY CONTROL SAMPLE SUMMARY

Page 2 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** Lab Control Sample**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P190311-LCS

Test Code: EPA TO-15

Date Collected: NA

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: NA

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.125 Liter(s)

Test Notes:

CAS #	Compound	Spike Amount µg/m ³	Result µg/m ³	% Recovery	ALS	Data Qualifier
					Acceptance Limits	
156-59-2	cis-1,2-Dichloroethene	211	225	107	67-110	
141-78-6	Ethyl Acetate	436	463	106	64-127	
110-54-3	n-Hexane	216	209	97	60-115	
67-66-3	Chloroform	217	218	100	66-105	
109-99-9	Tetrahydrofuran (THF)	216	210	97	65-110	
107-06-2	1,2-Dichloroethane	215	231	107	60-110	
71-55-6	1,1,1-Trichloroethane	215	215	100	64-108	
71-43-2	Benzene	211	202	96	67-106	
56-23-5	Carbon Tetrachloride	212	218	103	64-112	
110-82-7	Cyclohexane	416	405	97	67-110	
78-87-5	1,2-Dichloropropane	216	218	101	66-112	
75-27-4	Bromodichloromethane	215	229	107	67-113	
79-01-6	Trichloroethene	213	208	98	66-108	
123-91-1	1,4-Dioxane	214	216	101	70-116	
80-62-6	Methyl Methacrylate	431	463	107	73-118	
142-82-5	n-Heptane	215	213	99	66-110	
10061-01-5	cis-1,3-Dichloropropene	214	225	105	75-120	
108-10-1	4-Methyl-2-pentanone	209	229	110	65-124	
10061-02-6	trans-1,3-Dichloropropene	213	233	109	77-123	
79-00-5	1,1,2-Trichloroethane	215	218	101	68-112	
108-88-3	Toluene	212	174	82	62-111	
591-78-6	2-Hexanone	214	225	105	59-128	
124-48-1	Dibromochloromethane	213	214	100	67-123	
106-93-4	1,2-Dibromoethane	216	214	99	66-122	
123-86-4	n-Butyl Acetate	219	218	100	64-128	

Laboratory Control Sample percent recovery is verified and accepted based on the on-column result.

Reported results are shown in concentration units and as a result of the calculation, may vary slightly.

ALS ENVIRONMENTAL

LABORATORY CONTROL SAMPLE SUMMARY

Page 3 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** Lab Control Sample**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P190311-LCS

Test Code: EPA TO-15

Date Collected: NA

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: NA

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.125 Liter(s)

Test Notes:

CAS #	Compound	Spike Amount µg/m ³	Result µg/m ³	% Recovery	ALS	Data Qualifier
					Acceptance Limits	
111-65-9	n-Octane	217	194	89	65-114	
127-18-4	Tetrachloroethene	213	182	85	55-120	
108-90-7	Chlorobenzene	215	184	86	61-114	
100-41-4	Ethylbenzene	212	182	86	64-113	
179601-23-1	m,p-Xylenes	426	368	86	64-114	
75-25-2	Bromoform	213	216	101	65-132	
100-42-5	Styrene	212	199	94	67-124	
95-47-6	o-Xylene	214	184	86	65-114	
111-84-2	n-Nonane	215	198	92	64-117	
79-34-5	1,1,2,2-Tetrachloroethane	214	194	91	66-119	
98-82-8	Cumene	214	180	84	61-116	
80-56-8	alpha-Pinene	211	195	92	65-120	
103-65-1	n-Propylbenzene	218	188	86	63-117	
622-96-8	4-Ethyltoluene	214	189	88	63-124	
108-67-8	1,3,5-Trimethylbenzene	214	175	82	60-117	
95-63-6	1,2,4-Trimethylbenzene	215	185	86	61-122	
100-44-7	Benzyl Chloride	217	219	101	77-142	
541-73-1	1,3-Dichlorobenzene	216	189	88	61-125	
106-46-7	1,4-Dichlorobenzene	216	183	85	59-123	
95-50-1	1,2-Dichlorobenzene	216	189	88	61-126	
5989-27-5	d-Limonene	211	206	98	66-124	
96-12-8	1,2-Dibromo-3-chloropropane	209	195	93	67-138	
120-82-1	1,2,4-Trichlorobenzene	214	195	91	62-141	
91-20-3	Naphthalene	203	208	102	62-145	
87-68-3	Hexachlorobutadiene	209	162	78	49-131	

Laboratory Control Sample percent recovery is verified and accepted based on the on-column result.
Reported results are shown in concentration units and as a result of the calculation, may vary slightly.

ALS ENVIRONMENTAL

LABORATORY DUPLICATE SUMMARY RESULTS

Page 1 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_Stripper_a**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-002DUP

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00210

Initial Pressure (psig): -0.15

Final Pressure (psig): 3.90

Container Dilution Factor: 1.28

Compound	Sample Result		Duplicate Sample Result		Average µg/m ³	% RPD	RPD Limit	Data Qualifier
	µg/m ³	ppbV	µg/m ³	ppbV				
Propene	ND	ND	ND	ND	-	-	25	
Dichlorodifluoromethane (CFC 12)	ND	ND	ND	ND	-	-	25	
Chloromethane	ND	ND	ND	ND	-	-	25	
1,2-Dichloro-1,1,2,2-tetrafluoroethane (CFC 114)	ND	ND	ND	ND	-	-	25	
Vinyl Chloride	ND	ND	ND	ND	-	-	25	
1,3-Butadiene	ND	ND	ND	ND	-	-	25	
Bromomethane	ND	ND	ND	ND	-	-	25	
Chloroethane	ND	ND	ND	ND	-	-	25	
Ethanol	ND	ND	ND	ND	-	-	25	
Acetonitrile	ND	ND	ND	ND	-	-	25	
Acrolein	ND	ND	ND	ND	-	-	25	
Acetone	ND	ND	ND	ND	-	-	25	
Trichlorofluoromethane	ND	ND	ND	ND	-	-	25	
2-Propanol (Isopropyl Alcohol)	ND	ND	ND	ND	-	-	25	
Acrylonitrile	ND	ND	ND	ND	-	-	25	
1,1-Dichloroethene	ND	ND	ND	ND	-	-	25	
Methylene Chloride	632	182	606	174	619	4	25	
3-Chloro-1-propene (Allyl Chloride)	ND	ND	ND	ND	-	-	25	
Trichlorotrifluoroethane	1,870	244	1,900	248	1885	2	25	
Carbon Disulfide	ND	ND	ND	ND	-	-	25	
trans-1,2-Dichloroethene	ND	ND	ND	ND	-	-	25	
1,1-Dichloroethane	ND	ND	ND	ND	-	-	25	
Methyl tert-Butyl Ether	ND	ND	ND	ND	-	-	25	
Vinyl Acetate	ND	ND	ND	ND	-	-	25	
2-Butanone (MEK)	ND	ND	ND	ND	-	-	25	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

ALS ENVIRONMENTAL

LABORATORY DUPLICATE SUMMARY RESULTS

Page 2 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_Stripper_a**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-002DUP

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00210

Initial Pressure (psig): -0.15

Final Pressure (psig): 3.90

Container Dilution Factor: 1.28

Compound	Sample Result		Duplicate Sample Result		Average µg/m ³	% RPD	RPD Limit	Data Qualifier
	µg/m ³	ppbV	µg/m ³	ppbV				
cis-1,2-Dichloroethene	2,830	714	2,900	731	2865	2	25	
Ethyl Acetate	ND	ND	ND	ND	-	-	25	
n-Hexane	ND	ND	ND	ND	-	-	25	
Chloroform	ND	ND	ND	ND	-	-	25	
Tetrahydrofuran (THF)	ND	ND	ND	ND	-	-	25	
1,2-Dichloroethane	142	35.1	143	35.2	142.5	0.7	25	
1,1,1-Trichloroethane	ND	ND	ND	ND	-	-	25	
Benzene	ND	ND	ND	ND	-	-	25	
Carbon Tetrachloride	ND	ND	ND	ND	-	-	25	
Cyclohexane	ND	ND	ND	ND	-	-	25	
1,2-Dichloropropane	ND	ND	ND	ND	-	-	25	
Bromodichloromethane	ND	ND	ND	ND	-	-	25	
Trichloroethene	11,700	2,170	11,900	2,220	11800	2	25	
1,4-Dioxane	ND	ND	ND	ND	-	-	25	
Methyl Methacrylate	ND	ND	ND	ND	-	-	25	
n-Heptane	ND	ND	ND	ND	-	-	25	
cis-1,3-Dichloropropene	ND	ND	ND	ND	-	-	25	
4-Methyl-2-pentanone	ND	ND	ND	ND	-	-	25	
trans-1,3-Dichloropropene	ND	ND	ND	ND	-	-	25	
1,1,2-Trichloroethane	ND	ND	ND	ND	-	-	25	
Toluene	ND	ND	ND	ND	-	-	25	
2-Hexanone	ND	ND	ND	ND	-	-	25	
Dibromochloromethane	ND	ND	ND	ND	-	-	25	
1,2-Dibromoethane	ND	ND	ND	ND	-	-	25	
n-Butyl Acetate	ND	ND	ND	ND	-	-	25	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

ALS ENVIRONMENTAL

LABORATORY DUPLICATE SUMMARY RESULTS

Page 3 of 3

Client: Bhate Environmental Associates, Inc.**Client Sample ID:** LH18/24-Air_022719_Stripper_a**Client Project ID:** LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

ALS Sample ID: P1901179-002DUP

Test Code: EPA TO-15

Date Collected: 2/27/19

Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13

Date Received: 3/5/19

Analyst: Wida Ang

Date Analyzed: 3/11/19

Sample Type: 6.0 L Silonite Canister

Volume(s) Analyzed: 0.010 Liter(s)

Test Notes:

Container ID: AS00210

Initial Pressure (psig): -0.15

Final Pressure (psig): 3.90

Container Dilution Factor: 1.28

Compound	Sample Result		Duplicate Sample Result		Average µg/m ³	% RPD	RPD Limit	Data Qualifier
	µg/m ³	ppbV	µg/m ³	ppbV				
n-Octane	ND	ND	ND	ND	-	-	25	
Tetrachloroethene	ND	ND	ND	ND	-	-	25	
Chlorobenzene	ND	ND	ND	ND	-	-	25	
Ethylbenzene	ND	ND	ND	ND	-	-	25	
m,p-Xylenes	ND	ND	ND	ND	-	-	25	
Bromoform	ND	ND	ND	ND	-	-	25	
Styrene	ND	ND	ND	ND	-	-	25	
o-Xylene	ND	ND	ND	ND	-	-	25	
n-Nonane	ND	ND	ND	ND	-	-	25	
1,1,2,2-Tetrachloroethane	ND	ND	ND	ND	-	-	25	
Cumene	ND	ND	ND	ND	-	-	25	
alpha-Pinene	ND	ND	ND	ND	-	-	25	
n-Propylbenzene	ND	ND	ND	ND	-	-	25	
4-Ethyltoluene	ND	ND	ND	ND	-	-	25	
1,3,5-Trimethylbenzene	ND	ND	ND	ND	-	-	25	
1,2,4-Trimethylbenzene	ND	ND	ND	ND	-	-	25	
Benzyl Chloride	ND	ND	ND	ND	-	-	25	
1,3-Dichlorobenzene	ND	ND	ND	ND	-	-	25	
1,4-Dichlorobenzene	ND	ND	ND	ND	-	-	25	
1,2-Dichlorobenzene	ND	ND	ND	ND	-	-	25	
d-Limonene	ND	ND	ND	ND	-	-	25	
1,2-Dibromo-3-chloropropane	ND	ND	ND	ND	-	-	25	
1,2,4-Trichlorobenzene	ND	ND	ND	ND	-	-	25	
Naphthalene	ND	ND	ND	ND	-	-	25	
Hexachlorobutadiene	ND	ND	ND	ND	-	-	25	

ND = Compound was analyzed for, but not detected above the laboratory reporting limit.

ALS ENVIRONMENTAL

RESULTS OF ANALYSIS

Page 1 of 1

Client: Bhate Environmental Associates, Inc.
Client Project ID: LHAAP GWTP / NW01312.0150

ALS Project ID: P1901179

Internal Standard Area and RT Summary

Test Code: EPA TO-15
Instrument ID: Tekmar AUTOCAN/Agilent 5975Binert/6890N/MS13
Analyst: Wida Ang
Sample Type: 6.0 L Silonite Canister(s)
Test Notes:

Lab File ID: 03111901.D
Date Analyzed: 3/11/19
Time Analyzed: 06:51

	IS1 (BCM)		IS2 (DFB)		IS3 (CBZ)	
	AREA #	RT #	AREA #	RT #	AREA #	RT #
24 Hour Standard	106667	10.98	447071	13.10	192180	17.42
Upper Limit	149334	11.31	625899	13.43	269052	17.75
Lower Limit	64000	10.65	268243	12.77	115308	17.09

Client Sample ID		IS1 (BCM)		IS2 (DFB)		IS3 (CBZ)	
		AREA #	RT #	AREA #	RT #	AREA #	RT #
01	Method Blank	90918	10.97	407856	13.10	175696	17.42
02	Lab Control Sample	100369	10.98	428145	13.11	184579	17.43
03	LH18/24-Air_022719_Stripper	95258	10.96	409116	13.10	177684	17.42
04	LH18/24-Air_022719_Stripper_a	93387	10.96	405288	13.10	175195	17.42
05	LH18/24-Air_022719_Stripper_a (Lab Dup)	88966	10.97	384241	13.10	167241	17.42
06	LH18/24-Air_022719_GWTP	93047	10.97	409377	13.10	180111	17.42
07	LH18/24-Air_022719_Downwind	93346	10.96	407063	13.10	179457	17.42
08							
09							
10							
11							
12							
13							
14							
15							
16							
17							
18							
19							
20							

IS1 (BCM) = Bromochloromethane

IS2 (DFB) = 1,4-Difluorobenzene

IS3 (CBZ) = Chlorobenzene-d5

AREA UPPER LIMIT = 140% of internal standard area

AREA LOWER LIMIT = 60% of internal standard area

RT UPPER LIMIT = 0.33 minutes of internal standard RT

RT LOWER LIMIT = 0.33 minutes of internal standard RT

Column used to flag values outside QC limits with an I.

I = Internal standard not within the specified limits.

Data File : I:\MS13\DATA\2019_03\11\03111914.D
 Acq On : 11 Mar 2019 14:37
 Sample : P1901179-001 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:05:30 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

DA 3/14/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	95258	12.500	ng	-0.03
37) 1,4-Difluorobenzene (IS2)	13.10	114	409116	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	177684	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	113684	13.997	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	112.00%	
57) Toluene-d8 (SS2)	15.54	98	450131	11.830	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	94.64%	
73) Bromofluorobenzene (SS3)	18.83	174	168136	12.273	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	98.16%	

Target Compounds

						Qvalue
2) Propene	0.00	42	0	N.D.		
3) Dichlorodifluoromethan...	4.19	85	117	N.D.		
4) Chloromethane	0.00	50	0	N.D.		
5) 1,2-Dichloro-1,1,2,2-t...	0.00	135	0	N.D.		
6) Vinyl Chloride	4.88	62	4957	0.470	ng	84
7) 1,3-Butadiene	0.00	54	0	N.D.		
8) Bromomethane	0.00	94	0	N.D.		
9) Chloroethane	0.00	64	0	N.D.		
10) Ethanol	0.00	45	0	N.D.		
11) Acetonitrile	0.00	41	0	N.D.		
12) Acrolein	0.00	56	0	N.D.		
13) Acetone	6.94	58	1870	N.D.		
14) Trichlorofluoromethane	0.00	101	0	N.D.		
15) 2-Propanol (Isopropanol)	0.00	45	0	N.D.		
16) Acrylonitrile	0.00	53	0	N.D.		
17) 1,1-Dichloroethene	8.03	96	4083	0.516	ng	96
18) 2-Methyl-2-Propanol (t...	0.00	59	0	N.D.		
19) Methylene Chloride	8.24	84	35520	4.667	ng	100
20) 3-Chloro-1-propene (Al...	0.00	41	0	N.D.	d	
21) Trichlorotrifluoroethane	8.66	151	122516	14.632	ng	98
22) Carbon Disulfide	8.55	76	1242	N.D.		
23) trans-1,2-Dichloroethene	0.00	61	0	N.D.		
24) 1,1-Dichloroethane	9.77	63	1589	N.D.		
25) Methyl tert-Butyl Ether	0.00	73	0	N.D.		
26) Vinyl Acetate	0.00	86	0	N.D.		
27) 2-Butanone (MEK)	0.00	72	0	N.D.		
28) cis-1,2-Dichloroethene	10.79	61	211966	22.114	ng	96
29) Diisopropyl Ether	0.00	87	0	N.D.		
30) Ethyl Acetate	11.13	61	236	N.D.		
31) n-Hexane	0.00	57	0	N.D.		
32) Chloroform	11.14	83	2156	N.D.		
34) Tetrahydrofuran (THF)	0.00	72	0	N.D.		
35) Ethyl tert-Butyl Ether	0.00	87	0	N.D.		
36) 1,2-Dichloroethane	11.95	62	8843	1.124	ng	94
38) 1,1,1-Trichloroethane	0.00	97	0	N.D.		
39) Isopropyl Acetate	0.00	61	0	N.D.		
40) 1-Butanol	0.00	56	0	N.D.		
41) Benzene	12.71	78	1076	N.D.		
42) Carbon Tetrachloride	12.86	117	632	N.D.		
43) Cyclohexane	0.00	84	0	N.D.		
44) tert-Amyl Methyl Ether	0.00	73	0	N.D.		
45) 1,2-Dichloropropane	0.00	63	0	N.D.		
46) Bromodichloromethane	0.00	83	0	N.D.	d	
47) Trichloroethene	13.80	130	904294	92.605	ng	99
48) 1,4-Dioxane	0.00	88	0	N.D.		
49) 2,2,4-Trimethylpentane...	0.00	57	0	N.D.		
50) Methyl Methacrylate	0.00	100	0	N.D.	d	

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Data File : I:\MS13\DATA\2019_03\11\03111914.D
 Acq On : 11 Mar 2019 14:37
 Sample : P1901179-001 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:05:30 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

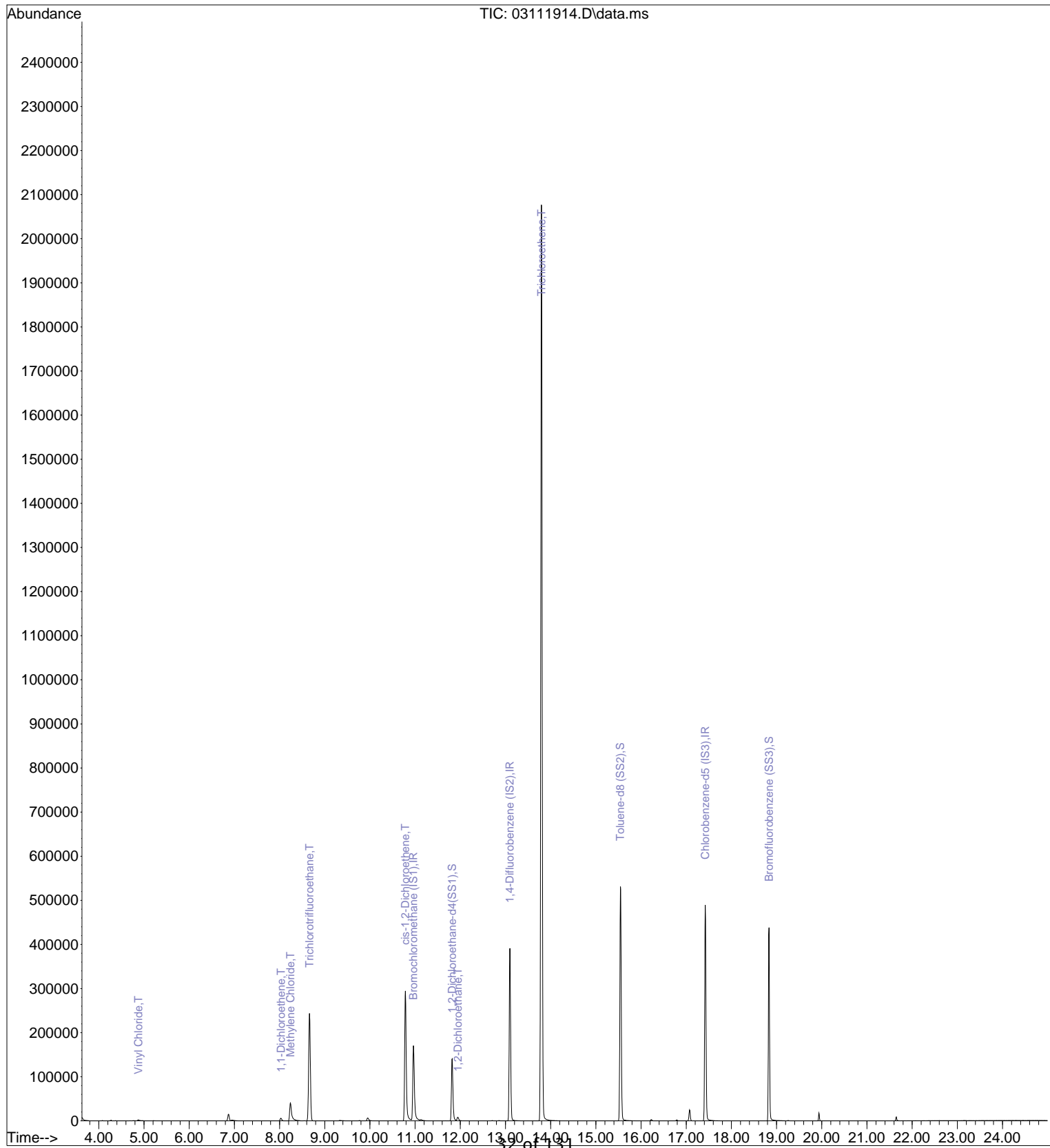
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	0.00	71	0	N.D.		
52) cis-1,3-Dichloropropene	0.00	75	0	N.D.		
53) 4-Methyl-2-pentanone	0.00	58	0	N.D.		
54) trans-1,3-Dichloropropene	0.00	75	0	N.D.		
55) 1,1,2-Trichloroethane	0.00	97	0	N.D.		
58) Toluene	15.66	91	537	N.D.		
59) 2-Hexanone	0.00	43	0	N.D.		
60) Dibromochloromethane	0.00	129	0	N.D.		
61) 1,2-Dibromoethane	0.00	107	0	N.D.		
62) n-Butyl Acetate	0.00	43	0	N.D.		
63) n-Octane	0.00	57	0	N.D.		
64) Tetrachloroethene	16.80	166	1091	N.D.		
65) Chlorobenzene	0.00	112	0	N.D.		
66) Ethylbenzene	0.00	91	0	N.D.		
67) m- & p-Xylenes	0.00	91	0	N.D.		
68) Bromoform	0.00	173	0	N.D.		
69) Styrene	0.00	104	0	N.D.		
70) o-Xylene	0.00	91	0	N.D.		
71) n-Nonane	0.00	43	0	N.D.		
72) 1,1,2,2-Tetrachloroethane	0.00	83	0	N.D.		
74) Cumene	0.00	105	0	N.D.		
75) alpha-Pinene	0.00	93	0	N.D.		
76) n-Propylbenzene	0.00	91	0	N.D.		
77) 3-Ethyltoluene	0.00	105	0	N.D.		
78) 4-Ethyltoluene	0.00	105	0	N.D.		
79) 1,3,5-Trimethylbenzene	0.00	105	0	N.D.		
80) alpha-Methylstyrene	0.00	118	0	N.D.		
81) 2-Ethyltoluene	0.00	105	0	N.D.		
82) 1,2,4-Trimethylbenzene	0.00	105	0	N.D.		
83) n-Decane	0.00	57	0	N.D.		
84) Benzyl Chloride	0.00	91	0	N.D.		
85) 1,3-Dichlorobenzene	0.00	146	0	N.D.		
86) 1,4-Dichlorobenzene	0.00	146	0	N.D.		
87) sec-Butylbenzene	0.00	105	0	N.D.		
88) 4-Isopropyltoluene (p-...	0.00	119	0	N.D.		
89) 1,2,3-Trimethylbenzene	0.00	105	0	N.D.		
90) 1,2-Dichlorobenzene	0.00	146	0	N.D.		
91) d-Limonene	0.00	68	0	N.D.		
92) 1,2-Dibromo-3-Chloropr...	0.00	157	0	N.D.		
93) n-Undecane	0.00	57	0	N.D.		
94) 1,2,4-Trichlorobenzene	0.00	180	0	N.D.		
95) Naphthalene	0.00	128	0	N.D.		
96) n-Dodecane	0.00	57	0	N.D.		
97) Hexachlorobutadiene	0.00	225	0	N.D.		
98) Cyclohexanone	0.00	55	0	N.D.		
99) tert-Butylbenzene	19.94	119	312	N.D.		
100) n-Butylbenzene	0.00	91	0	N.D.		

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111914.D
 Acq On : 11 Mar 2019 14:37
 Sample : P1901179-001 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:05:30 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M



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Data File : I:\MS13\DATA\2019_03\11\03111914.D
 Acq On : 11 Mar 2019 14:37
 Sample : P1901179-001 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:05:30 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

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Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	95258	12.500	ng	-0.03
37) 1,4-Difluorobenzene (IS2)	13.10	114	409116	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	177684	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	113684	13.997	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	112.00%	
57) Toluene-d8 (SS2)	15.54	98	450131	11.830	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	94.64%	
73) Bromofluorobenzene (SS3)	18.83	174	168136	12.273	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	98.16%	

Target Compounds

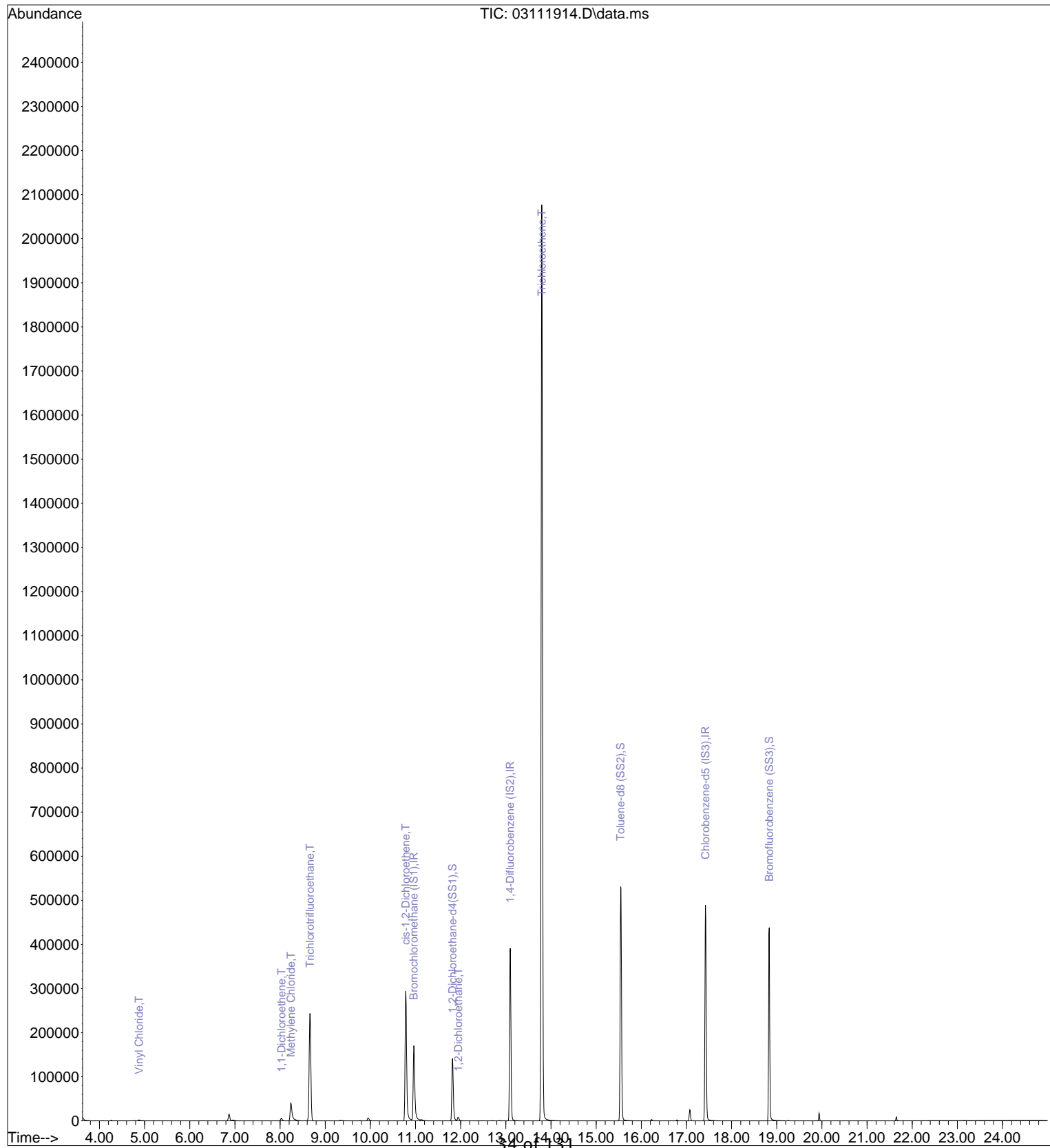
	R.T.	QIon	Response	Conc	Units	Qvalue
6) Vinyl Chloride	4.88	62	4957	0.470	ng	84
17) 1,1-Dichloroethene	8.03	96	4083	0.516	ng	96
19) Methylene Chloride	8.24	84	35520	4.667	ng	100
21) Trichlorotrifluoroethane	8.66	151	122516	14.632	ng	98
28) cis-1,2-Dichloroethene	10.79	61	211966	22.114	ng	96
36) 1,2-Dichloroethane	11.95	62	8843	1.124	ng	94
47) Trichloroethene	13.80	130	904294	92.605	ng	99

(#) = qualifier out of range (m) = manual integration (+) = signals summed

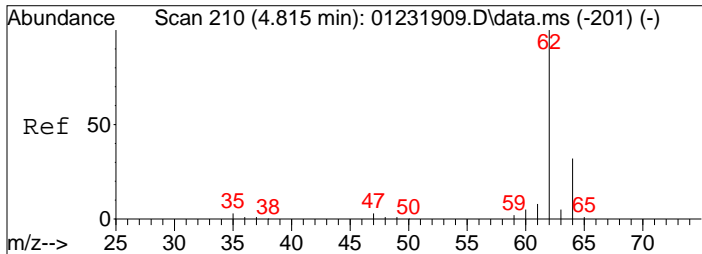
Data File : I:\MS13\DATA\2019_03\11\03111914.D
 Acq On : 11 Mar 2019 14:37
 Sample : P1901179-001 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:05:30 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

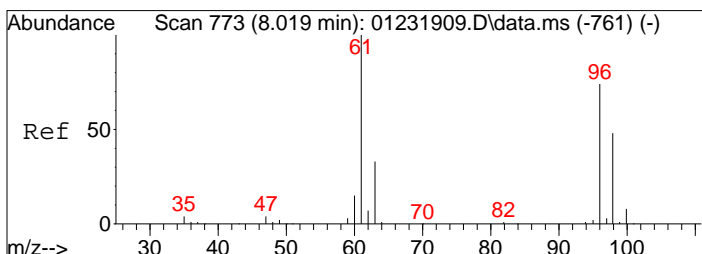
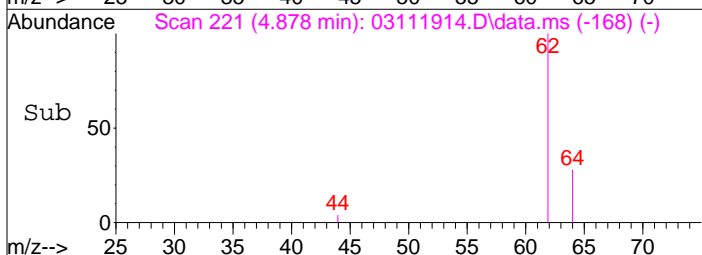
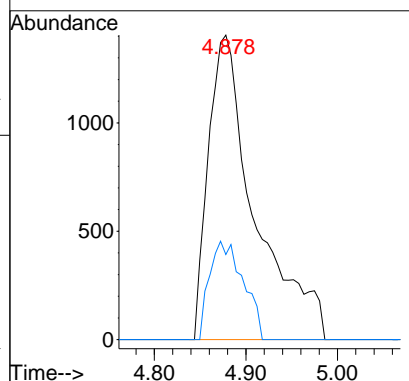
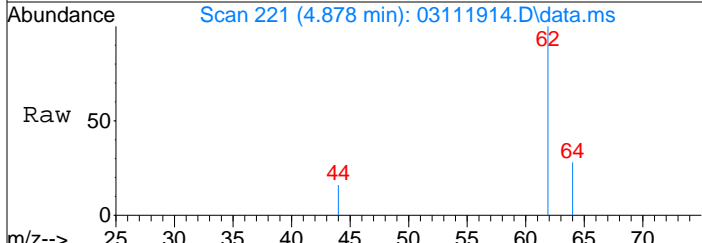


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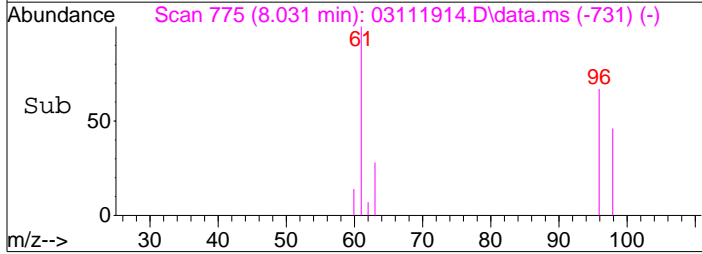
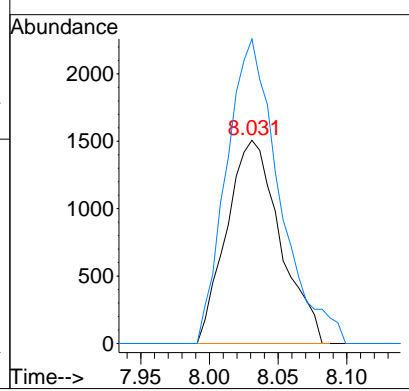
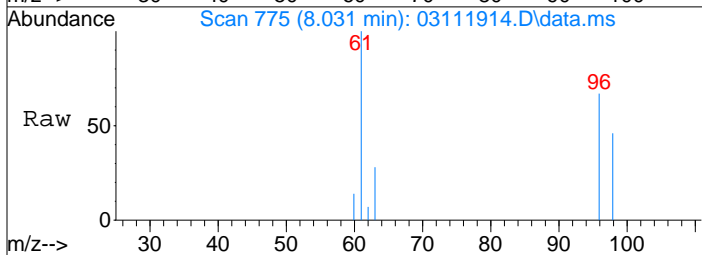
#6
 Vinyl Chloride
 Concen: 0.47 ng
 RT: 4.88 min Scan# 221
 Delta R.T. 0.051 min
 Lab File: 03111914.D
 Acq: 11 Mar 2019 14:37

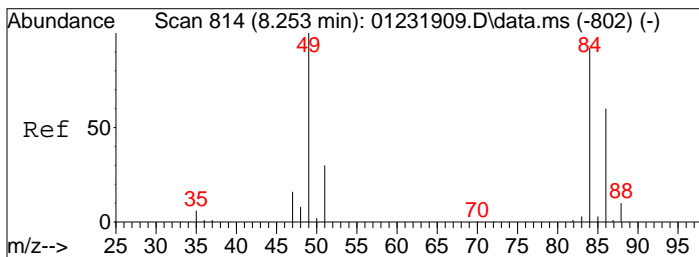
Tgt Ion	Resp	Lower	Upper
62	4957		
64	100	23.5	52.4



#17
 1,1-Dichloroethene
 Concen: 0.52 ng
 RT: 8.03 min Scan# 775
 Delta R.T. 0.000 min
 Lab File: 03111914.D
 Acq: 11 Mar 2019 14:37

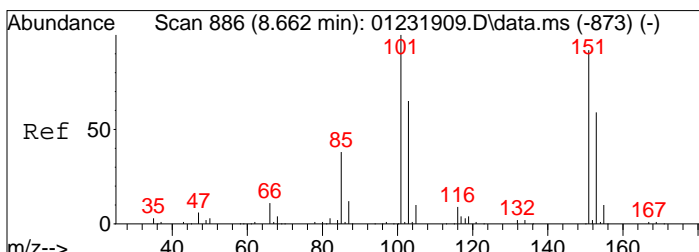
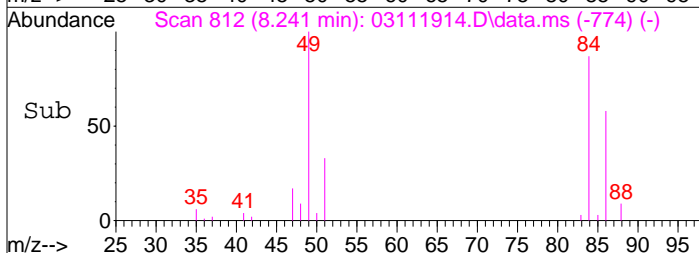
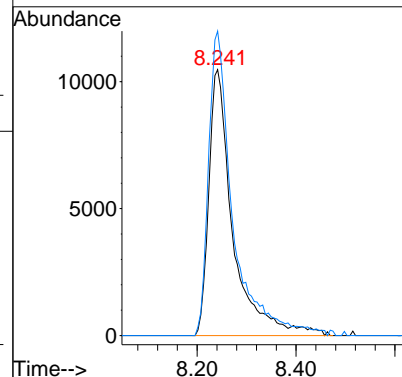
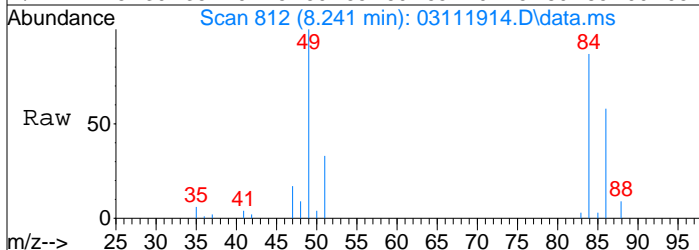
Tgt Ion	Resp	Lower	Upper
96	4083		
61	148.3	132.9	172.9





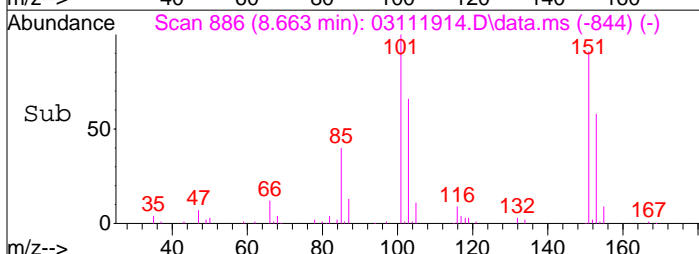
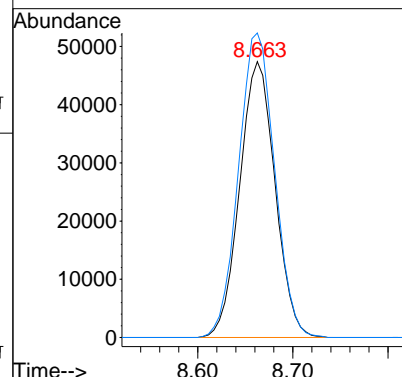
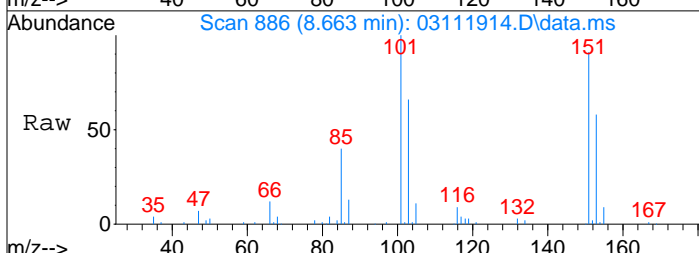
#19
 Methylene Chloride
 Concen: 4.67 ng
 RT: 8.24 min Scan# 812
 Delta R.T. -0.034 min
 Lab File: 03111914.D
 Acq: 11 Mar 2019 14:37

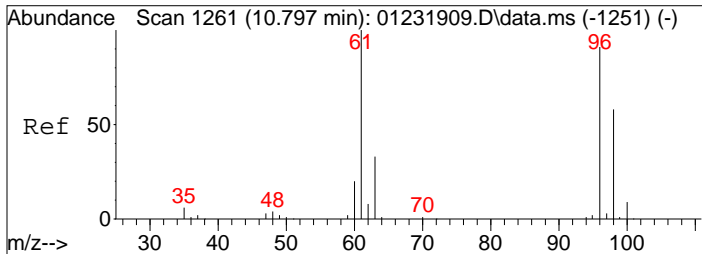
Tgt Ion: 84 Resp: 35520
 Ion Ratio Lower Upper
 84 100
 49 115.6 90.3 140.3



#21
 Trichlorotrifluoroethane
 Concen: 14.63 ng
 RT: 8.66 min Scan# 886
 Delta R.T. -0.011 min
 Lab File: 03111914.D
 Acq: 11 Mar 2019 14:37

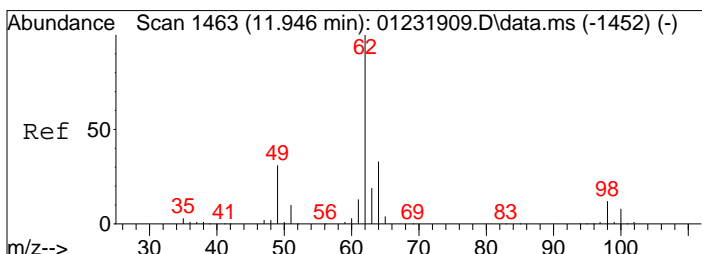
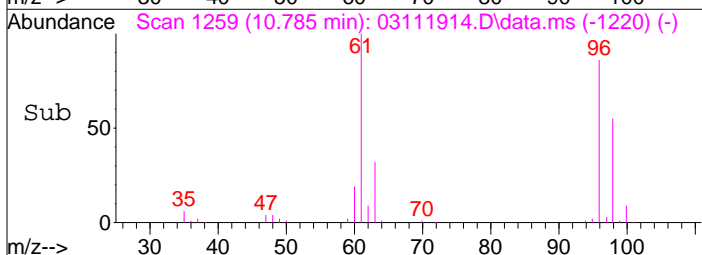
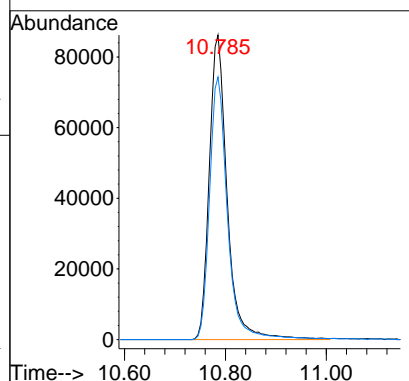
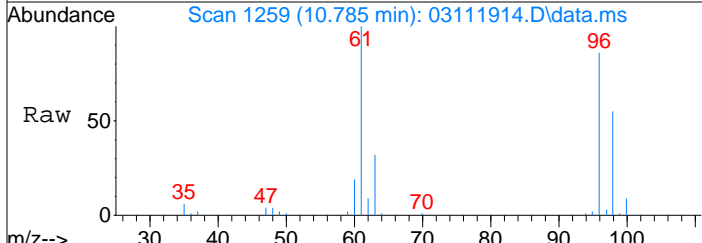
Tgt Ion: 151 Resp: 122516
 Ion Ratio Lower Upper
 151 100
 101 112.6 95.1 135.1





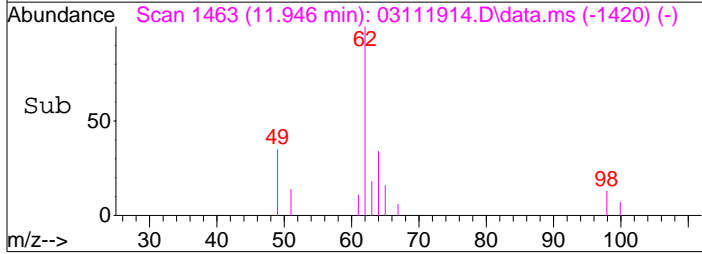
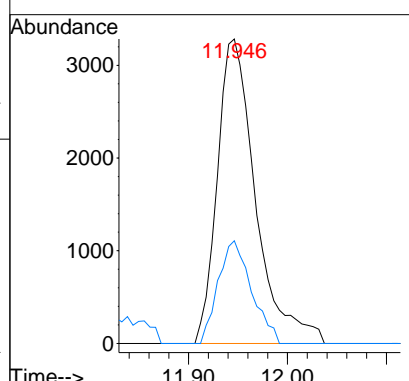
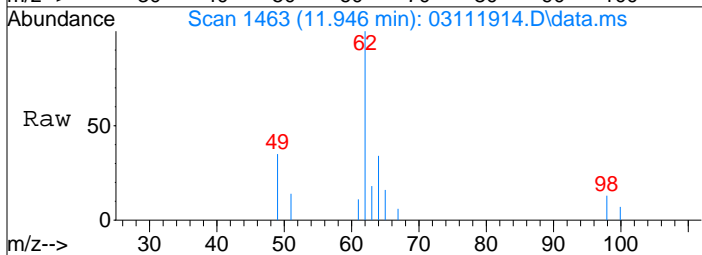
#28
 cis-1,2-Dichloroethene
 Concen: 22.11 ng
 RT: 10.79 min Scan# 1259
 Delta R.T. -0.028 min
 Lab File: 03111914.D
 Acq: 11 Mar 2019 14:37

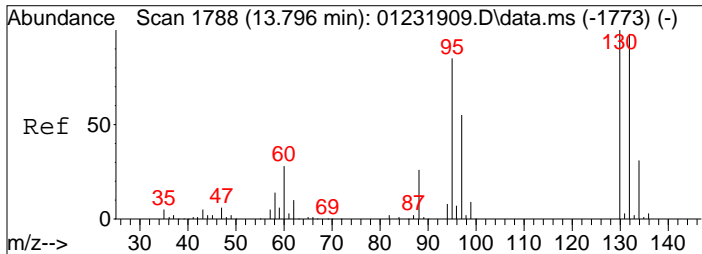
Tgt Ion	Resp	Lower	Upper
61	100		
96	86.6	63.3	103.3



#36
 1,2-Dichloroethane
 Concen: 1.12 ng
 RT: 11.95 min Scan# 1463
 Delta R.T. -0.006 min
 Lab File: 03111914.D
 Acq: 11 Mar 2019 14:37

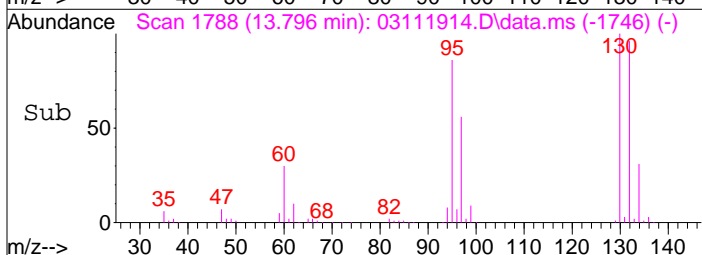
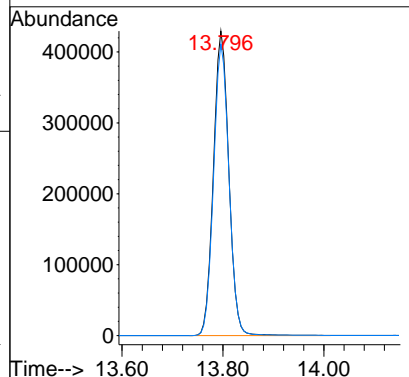
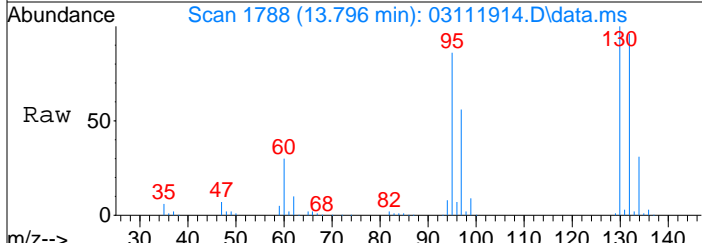
Tgt Ion	Resp	Lower	Upper
62	100		
64	29.4	12.6	52.6





#47
Trichloroethene
Concen: 92.60 ng
RT: 13.80 min Scan# 1788
Delta R.T. -0.011 min
Lab File: 03111914.D
Acq: 11 Mar 2019 14:37

Tgt Ion:130 Resp: 904294
Ion Ratio Lower Upper
130 100
132 96.2 76.7 116.7



Data File : I:\MS13\DATA\2019_03\11\03111915.D
 Acq On : 11 Mar 2019 15:20
 Sample : P1901179-002 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:53 2019

Quant Method : I:\MS13\METHODS\R13012319.M

Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)

QLast Update : Wed Jan 23 11:21:29 2019

Response via : Initial Calibration

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DataAcq Meth:TO15.M

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	93387	12.500	ng	-0.03
37) 1,4-Difluorobenzene (IS2)	13.10	114	405288	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	175195	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	112078	14.076	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	112.64%	
57) Toluene-d8 (SS2)	15.54	98	442470	11.794	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	94.32%	
73) Bromofluorobenzene (SS3)	18.83	174	165126	12.224	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	97.76%	

Target Compounds

						Qvalue
2) Propene	4.02	42	3124	N.D.		
3) Dichlorodifluoromethan...	4.16	85	699	N.D.		
4) Chloromethane	0.00	50	0	N.D.		
5) 1,2-Dichloro-1,1,2,2-t...	0.00	135	0	N.D.		
6) Vinyl Chloride	4.87	62	4432	N.D.		
7) 1,3-Butadiene	5.02	54	801	N.D.		
8) Bromomethane	0.00	94	0	N.D.		
9) Chloroethane	0.00	64	0	N.D.		
10) Ethanol	0.00	45	0	N.D.	d	
11) Acetonitrile	0.00	41	0	N.D.		
12) Acrolein	0.00	56	0	N.D.		
13) Acetone	6.92	58	3637	0.618	ng	93
14) Trichlorofluoromethane	0.00	101	0	N.D.		
15) 2-Propanol (Isopropanol)	7.50	45	62	N.D.		
16) Acrylonitrile	0.00	53	0	N.D.		
17) 1,1-Dichloroethene	8.03	96	4181	0.538	ng	90
18) 2-Methyl-2-Propanol (t...	8.39	59	53	N.D.		
19) Methylene Chloride	8.24	84	36847	4.938	ng	100
20) 3-Chloro-1-propene (Al...	8.24	41	1131	N.D.		
21) Trichlorotrifluoroethane	8.66	151	119725	14.586	ng	98
22) Carbon Disulfide	8.54	76	2387	N.D.		
23) trans-1,2-Dichloroethene	0.00	61	0	N.D.		
24) 1,1-Dichloroethane	9.77	63	1561	N.D.		
25) Methyl tert-Butyl Ether	0.00	73	0	N.D.		
26) Vinyl Acetate	0.00	86	0	N.D.		
27) 2-Butanone (MEK)	0.00	72	0	N.D.		
28) cis-1,2-Dichloroethene	10.78	61	207802	22.114	ng	98
29) Diisopropyl Ether	0.00	87	0	N.D.		
30) Ethyl Acetate	0.00	61	0	N.D.		
31) n-Hexane	11.09	57	430	N.D.		
32) Chloroform	11.13	83	2122	N.D.		
34) Tetrahydrofuran (THF)	11.64	72	193	N.D.		
35) Ethyl tert-Butyl Ether	0.00	87	0	N.D.		
36) 1,2-Dichloroethane	11.94	62	8571	1.111	ng	95
38) 1,1,1-Trichloroethane	0.00	97	0	N.D.		
39) Isopropyl Acetate	0.00	61	0	N.D.		
40) 1-Butanol	0.00	56	0	N.D.		
41) Benzene	12.71	78	2576	N.D.		
42) Carbon Tetrachloride	12.85	117	593	N.D.		
43) Cyclohexane	12.99	84	185	N.D.		
44) tert-Amyl Methyl Ether	0.00	73	0	N.D.		
45) 1,2-Dichloropropane	0.00	63	0	N.D.		
46) Bromodichloromethane	0.00	83	0	N.D.	d	
47) Trichloroethene	13.80	130	880723	91.043	ng	100
48) 1,4-Dioxane	0.00	88	0	N.D.		
49) 2,2,4-Trimethylpentane...	13.86	57	945	N.D.		
50) Methyl Methacrylate	0.00	100	0	N.D.	d	

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Data File : I:\MS13\DATA\2019_03\11\03111915.D
 Acq On : 11 Mar 2019 15:20
 Sample : P1901179-002 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:53 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

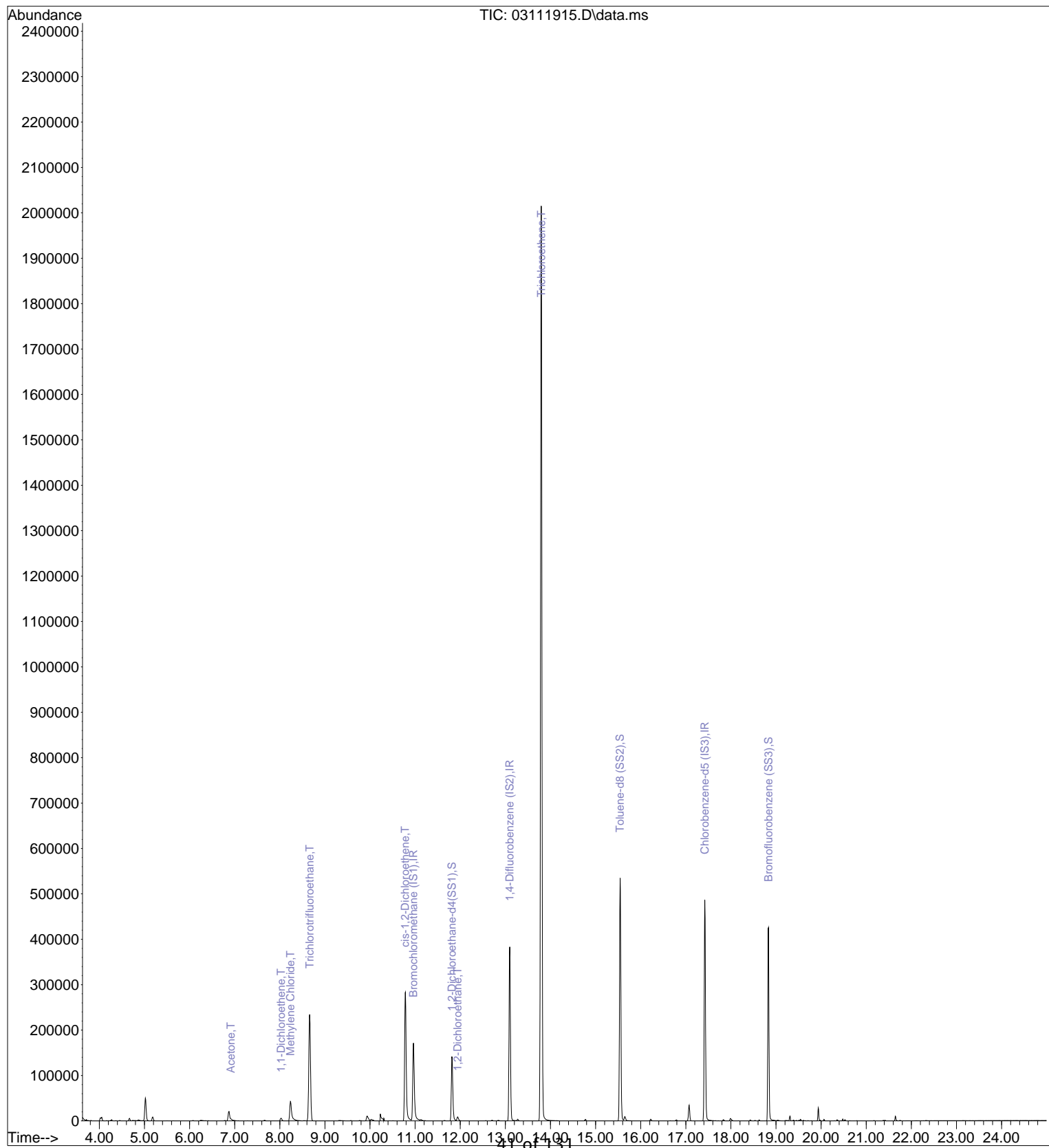
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	0.00	71	0	N.D.		
52) cis-1,3-Dichloropropene	0.00	75	0	N.D.		
53) 4-Methyl-2-pentanone	0.00	58	0	N.D.		
54) trans-1,3-Dichloropropene	0.00	75	0	N.D.		
55) 1,1,2-Trichloroethane	0.00	97	0	N.D.		
58) Toluene	15.65	91	9407	N.D.		
59) 2-Hexanone	16.05	43	164	N.D.		
60) Dibromochloromethane	0.00	129	0	N.D.		
61) 1,2-Dibromoethane	0.00	107	0	N.D.		
62) n-Butyl Acetate	16.65	43	316	N.D.		
63) n-Octane	0.00	57	0	N.D.		
64) Tetrachloroethene	16.79	166	1180	N.D.		
65) Chlorobenzene	17.47	112	205	N.D.		
66) Ethylbenzene	17.84	91	3597	N.D.		
67) m- & p-Xylenes	17.99	91	6088	N.D.		
68) Bromoform	0.00	173	0	N.D.		
69) Styrene	0.00	104	0	N.D.		
70) o-Xylene	18.42	91	2201	N.D.		
71) n-Nonane	18.63	43	1097	N.D.		
72) 1,1,2,2-Tetrachloroethane	0.00	83	0	N.D.		
74) Cumene	18.96	105	499	N.D.		
75) alpha-Pinene	19.31	93	4899	N.D.		
76) n-Propylbenzene	19.38	91	2876	N.D.		
77) 3-Ethyltoluene	19.50	105	1336	N.D.		
78) 4-Ethyltoluene	19.54	105	631	N.D.		
79) 1,3,5-Trimethylbenzene	19.61	105	735	N.D.		
80) alpha-Methylstyrene	19.94	118	281	N.D.		
81) 2-Ethyltoluene	19.78	105	516	N.D.		
82) 1,2,4-Trimethylbenzene	19.98	105	1383	N.D.		
83) n-Decane	20.06	57	1241	N.D.		
84) Benzyl Chloride	20.29	91	113	N.D.		
85) 1,3-Dichlorobenzene	0.00	146	0	N.D.		
86) 1,4-Dichlorobenzene	0.00	146	0	N.D.		
87) sec-Butylbenzene	20.35	105	332	N.D.		
88) 4-Isopropyltoluene (p-...	20.35	119	1688	N.D.		
89) 1,2,3-Trimethylbenzene	20.35	105	332	N.D.		
90) 1,2-Dichlorobenzene	0.00	146	0	N.D.		
91) d-Limonene	20.48	68	1222	N.D.		
92) 1,2-Dibromo-3-Chloropr...	0.00	157	0	N.D.		
93) n-Undecane	21.17	57	465	N.D.		
94) 1,2,4-Trichlorobenzene	0.00	180	0	N.D.		
95) Naphthalene	0.00	128	0	N.D.		
96) n-Dodecane	0.00	57	0	N.D.		
97) Hexachlorobutadiene	0.00	225	0	N.D.		
98) Cyclohexanone	0.00	55	0	N.D.		
99) tert-Butylbenzene	19.94	119	514	N.D.		
100) n-Butylbenzene	20.53	91	177	N.D.		

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111915.D
 Acq On : 11 Mar 2019 15:20
 Sample : P1901179-002 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:53 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_03\11\03111915.D
 Acq On : 11 Mar 2019 15:20
 Sample : P1901179-002 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:53 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

WA 3/14/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	93387	12.500	ng	-0.03
37) 1,4-Difluorobenzene (IS2)	13.10	114	405288	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	175195	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	112078	14.076	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	112.64%	
57) Toluene-d8 (SS2)	15.54	98	442470	11.794	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	94.32%	
73) Bromofluorobenzene (SS3)	18.83	174	165126	12.224	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	97.76%	

Target Compounds

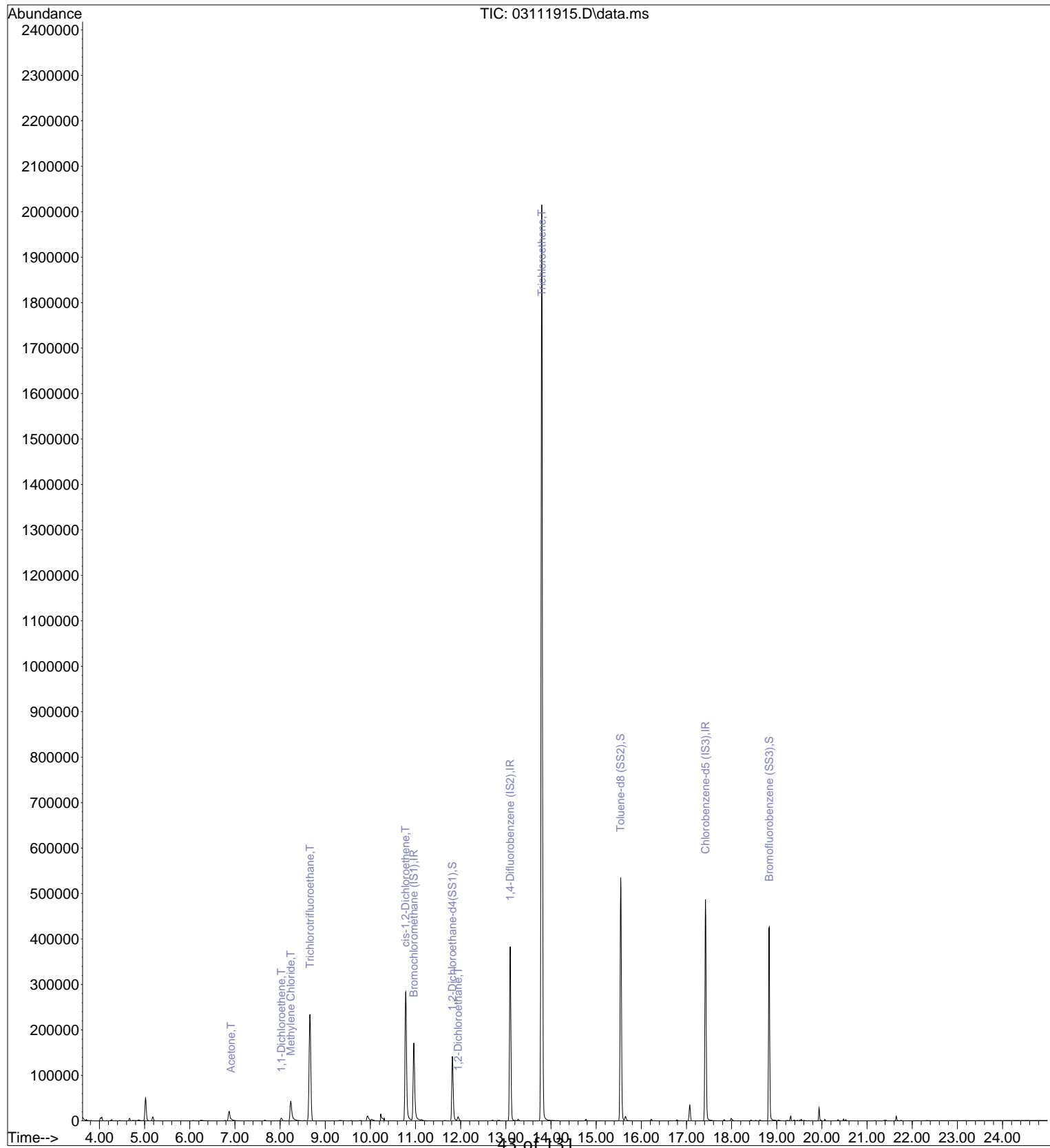
	R.T.	QIon	Response	Conc	Units	Qvalue
13) Acetone	6.92	58	3637	0.618	ng	93
17) 1,1-Dichloroethene	8.03	96	4181	0.538	ng	90
19) Methylene Chloride	8.24	84	36847	4.938	ng	100
21) Trichlorotrifluoroethane	8.66	151	119725	14.586	ng	98
28) cis-1,2-Dichloroethene	10.78	61	207802	22.114	ng	98
36) 1,2-Dichloroethane	11.94	62	8571	1.111	ng	95
47) Trichloroethene	13.80	130	880723	91.043	ng	100

(#) = qualifier out of range (m) = manual integration (+) = signals summed

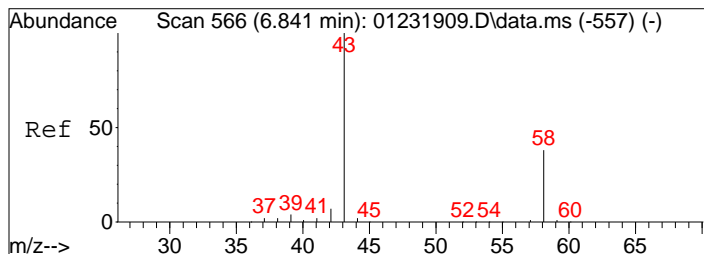
Data File : I:\MS13\DATA\2019_03\11\03111915.D
 Acq On : 11 Mar 2019 15:20
 Sample : P1901179-002 (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:53 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

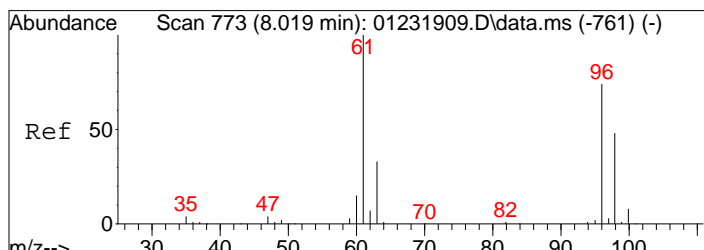
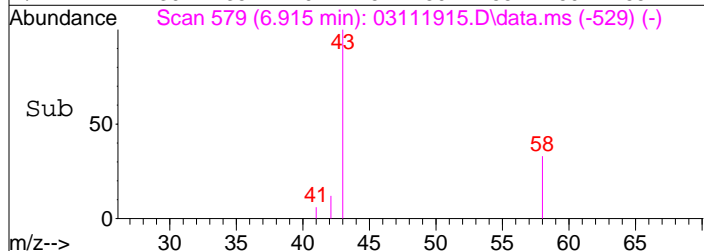
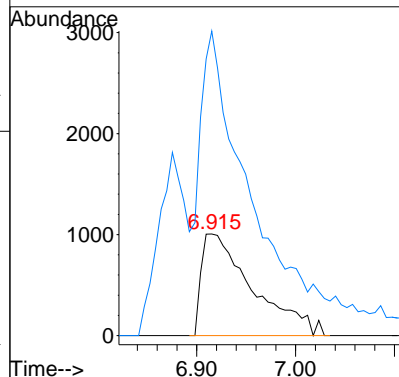
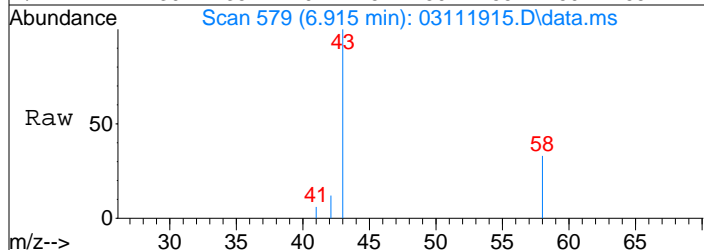


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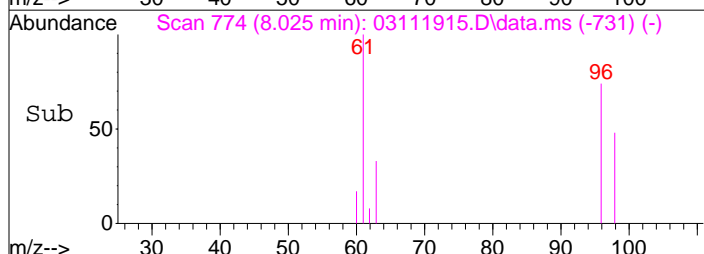
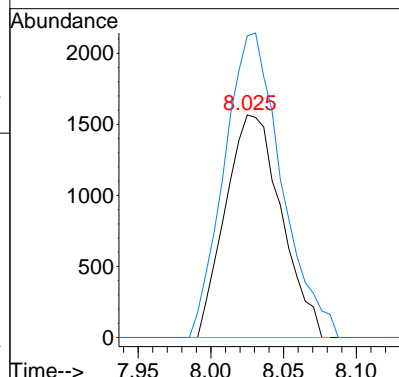
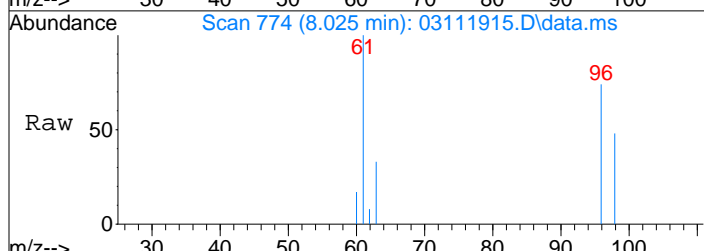
#13
 Acetone
 Concen: 0.62 ng
 RT: 6.92 min Scan# 579
 Delta R.T. 0.034 min
 Lab File: 03111915.D
 Acq: 11 Mar 2019 15:20

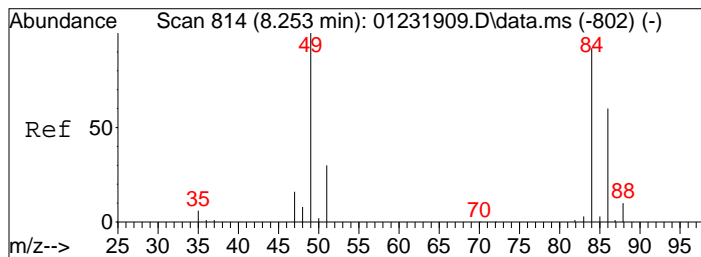
Tgt Ion	Resp	Lower	Upper
58	100		
43	287.3	271.8	331.8



#17
 1,1-Dichloroethene
 Concen: 0.54 ng
 RT: 8.03 min Scan# 774
 Delta R.T. -0.006 min
 Lab File: 03111915.D
 Acq: 11 Mar 2019 15:20

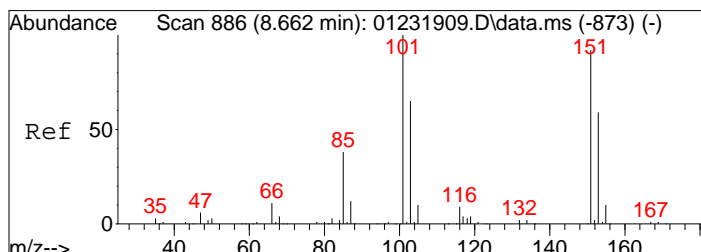
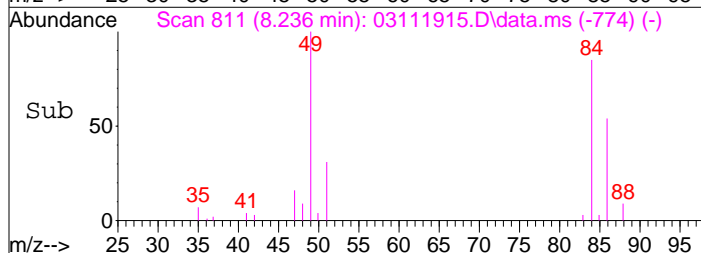
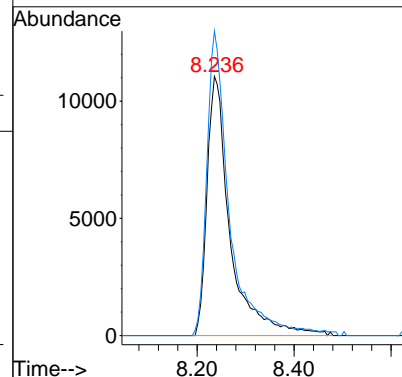
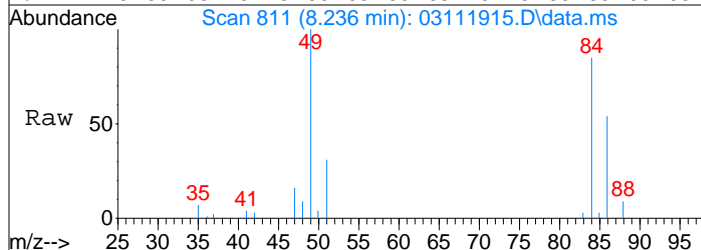
Tgt Ion	Resp	Lower	Upper
96	100		
61	140.4	132.9	172.9





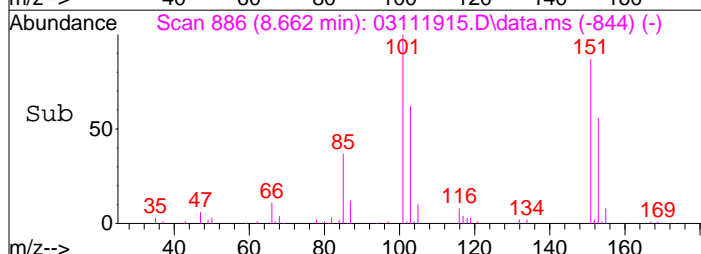
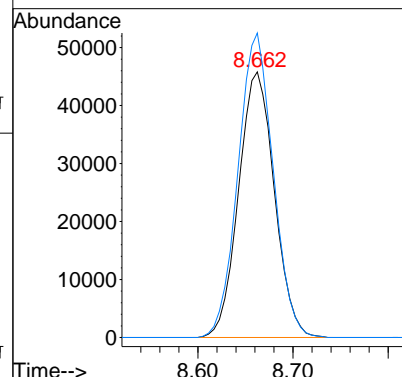
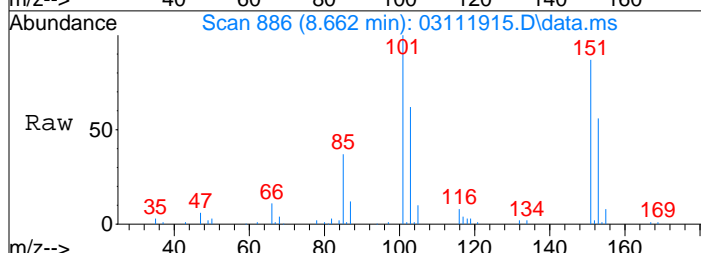
#19
 Methylene Chloride
 Concen: 4.94 ng
 RT: 8.24 min Scan# 811
 Delta R.T. -0.040 min
 Lab File: 03111915.D
 Acq: 11 Mar 2019 15:20

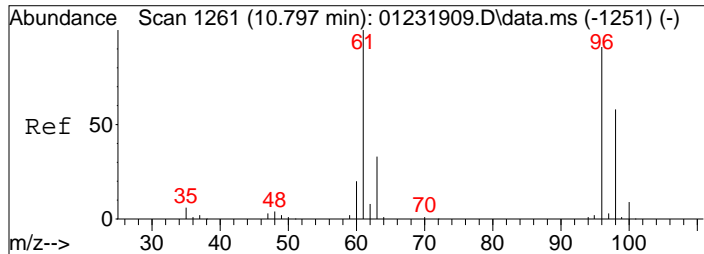
Tgt Ion: 84 Resp: 36847
 Ion Ratio Lower Upper
 84 100
 49 115.7 90.3 140.3



#21
 Trichlorotrifluoroethane
 Concen: 14.59 ng
 RT: 8.66 min Scan# 886
 Delta R.T. -0.012 min
 Lab File: 03111915.D
 Acq: 11 Mar 2019 15:20

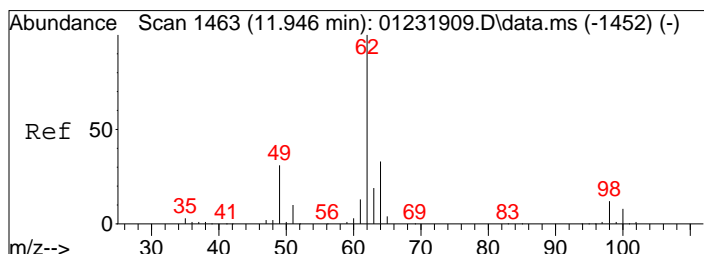
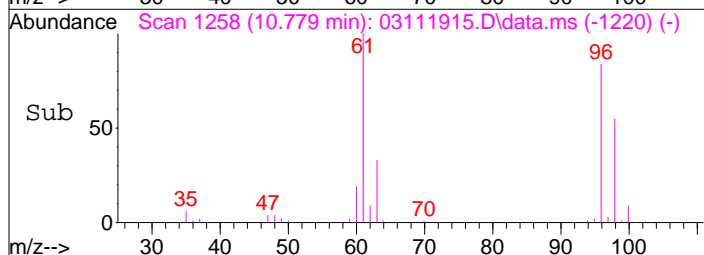
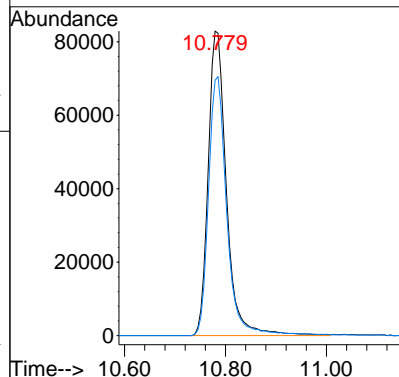
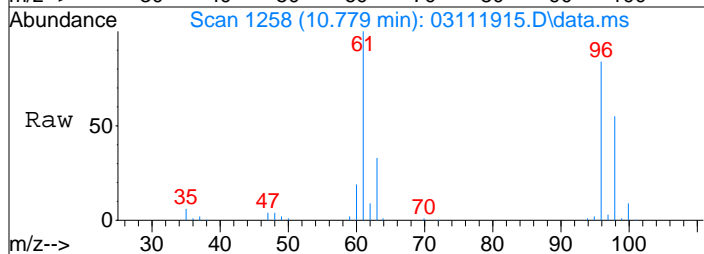
Tgt Ion: 151 Resp: 119725
 Ion Ratio Lower Upper
 151 100
 101 112.6 95.1 135.1





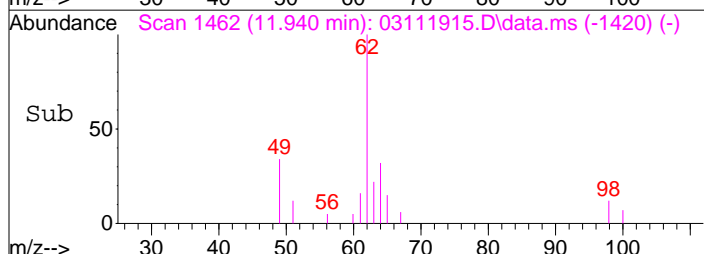
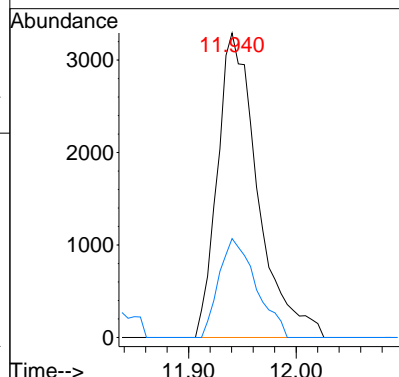
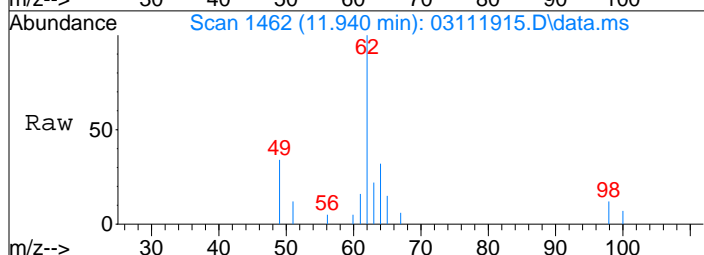
#28
 cis-1,2-Dichloroethene
 Concen: 22.11 ng
 RT: 10.78 min Scan# 1258
 Delta R.T. -0.034 min
 Lab File: 03111915.D
 Acq: 11 Mar 2019 15:20

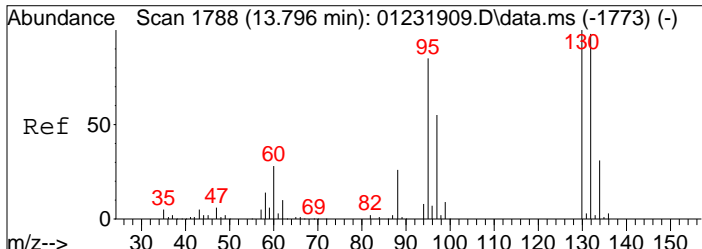
Tgt Ion: 61 Resp: 207802
 Ion Ratio Lower Upper
 61 100
 96 85.3 63.3 103.3



#36
 1,2-Dichloroethane
 Concen: 1.11 ng
 RT: 11.94 min Scan# 1462
 Delta R.T. -0.012 min
 Lab File: 03111915.D
 Acq: 11 Mar 2019 15:20

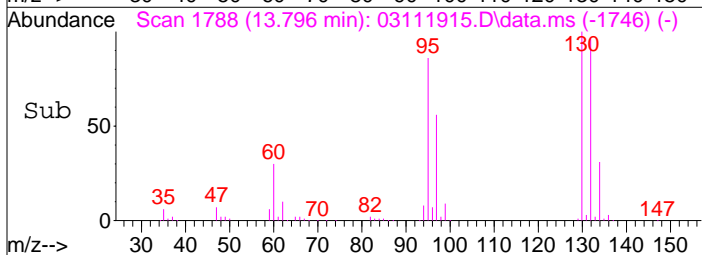
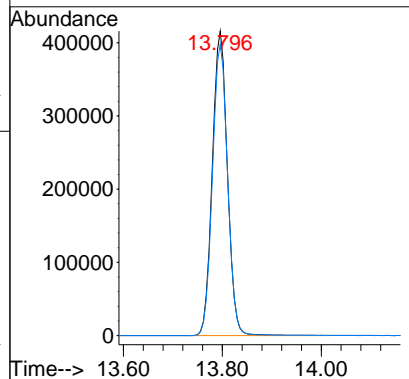
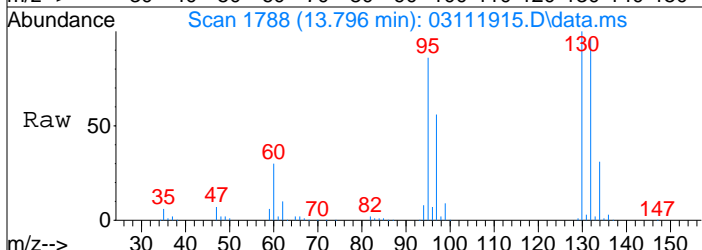
Tgt Ion: 62 Resp: 8571
 Ion Ratio Lower Upper
 62 100
 64 30.0 12.6 52.6





#47
Trichloroethene
Concen: 91.04 ng
RT: 13.80 min Scan# 1788
Delta R.T. -0.011 min
Lab File: 03111915.D
Acq: 11 Mar 2019 15:20

Tgt Ion	Resp	Lower	Upper
130	100		
132	96.3	76.7	116.7



Data File : I:\MS13\DATA\2019_03\11\03111921.D
 Acq On : 11 Mar 2019 18:42
 Sample : P1901179-003 (1000mL)
 Misc : S31-02211904

Vial: 1
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:09:24 2019

Quant Method : I:\MS13\METHODS\R13012319.M

Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)

QLast Update : Wed Jan 23 11:21:29 2019

Response via : Initial Calibration

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DataAcq Meth:TO15.M

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.97	130	93047	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.10	114	409377	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	180111	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	110943	13.984	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	111.84%	
57) Toluene-d8 (SS2)	15.54	98	450833	11.689	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	93.52%	
73) Bromofluorobenzene (SS3)	18.83	174	169718	12.221	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	97.76%	

Target Compounds

						Qvalue
2) Propene	0.00	42	0	N.D.	d	
3) Dichlorodifluoromethan...	4.15	85	27542	1.991	ng	99
4) Chloromethane	4.45	50	3587	N.D.		
5) 1,2-Dichloro-1,1,2,2-t...	4.68	135	701	N.D.		
6) Vinyl Chloride	4.85	62	3197	N.D.		
7) 1,3-Butadiene	0.00	54	0	N.D.		
8) Bromomethane	0.00	94	0	N.D.		
9) Chloroethane	0.00	64	0	N.D.		
10) Ethanol	6.21	45	15665	2.838	ng	98
11) Acetonitrile	6.54	41	1992	N.D.		
12) Acrolein	6.69	56	774	N.D.		
13) Acetone	6.86	58	41157	7.018	ng	86
14) Trichlorofluoromethane	7.07	101	12455	1.075	ng	100
15) 2-Propanol (Isopropanol)	7.42	45	4246	N.D.		
16) Acrylonitrile	7.66	53	225	N.D.		
17) 1,1-Dichloroethene	8.03	96	2510	N.D.		
18) 2-Methyl-2-Propanol (t...	8.33	59	647	N.D.		
19) Methylene Chloride	8.24	84	16541	2.225	ng	97
20) 3-Chloro-1-propene (Al...	8.24	41	523	N.D.		
21) Trichlorotrifluoroethane	8.66	151	122773	15.012	ng	98
22) Carbon Disulfide	8.53	76	18566	0.611	ng	91
23) trans-1,2-Dichloroethene	0.00	61	0	N.D.		
24) 1,1-Dichloroethane	9.77	63	627	N.D.		
25) Methyl tert-Butyl Ether	9.93	73	356	N.D.		
26) Vinyl Acetate	0.00	86	0	N.D.	d	
27) 2-Butanone (MEK)	10.32	72	6464	1.202	ng	# 69
28) cis-1,2-Dichloroethene	10.79	61	88520	9.455	ng	96
29) Diisopropyl Ether	11.13	87	109	N.D.		
30) Ethyl Acetate	11.12	61	11883	4.464	ng	86
31) n-Hexane	11.08	57	10028	0.812	ng	# 100
32) Chloroform	11.13	83	1697	N.D.		
34) Tetrahydrofuran (THF)	11.62	72	1862	N.D.		
35) Ethyl tert-Butyl Ether	0.00	87	0	N.D.		
36) 1,2-Dichloroethane	11.95	62	3715	0.483	ng	97
38) 1,1,1-Trichloroethane	0.00	97	0	N.D.		
39) Isopropyl Acetate	0.00	61	0	N.D.		
40) 1-Butanol	0.00	56	0	N.D.	d	
41) Benzene	12.70	78	21228	0.628	ng	98
42) Carbon Tetrachloride	12.86	117	4001	N.D.		
43) Cyclohexane	12.99	84	3145	N.D.		
44) tert-Amyl Methyl Ether	13.46	73	183	N.D.		
45) 1,2-Dichloropropane	0.00	63	0	N.D.		
46) Bromodichloromethane	13.80	83	4188	N.D.		
47) Trichloroethene	13.80	130	420086	42.992	ng	100
48) 1,4-Dioxane	13.84	88	456	N.D.		
49) 2,2,4-Trimethylpentane...	13.86	57	4992	N.D.		
50) Methyl Methacrylate	14.13	100	848	N.D.		

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Data File : I:\MS13\DATA\2019_03\11\03111921.D
 Acq On : 11 Mar 2019 18:42
 Sample : P1901179-003 (1000mL)
 Misc : S31-02211904

Vial: 1
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:09:24 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

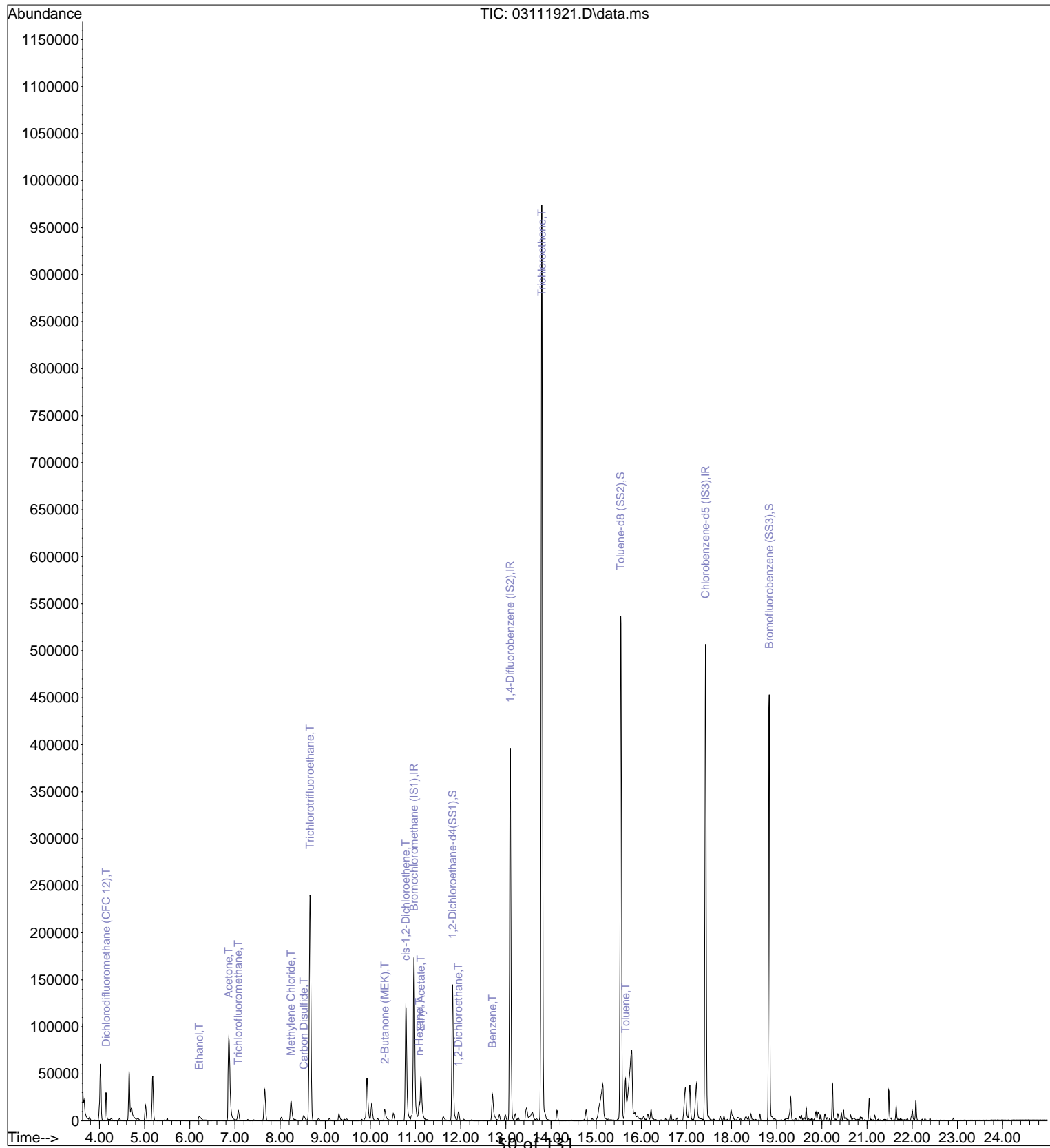
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	3089	N.D.		
52) cis-1,3-Dichloropropene	0.00	75	0	N.D.		
53) 4-Methyl-2-pentanone	14.73	58	395	N.D.		
54) trans-1,3-Dichloropropene	0.00	75	0	N.D.		
55) 1,1,2-Trichloroethane	0.00	97	0	N.D.		
58) Toluene	15.65	91	37664	0.919	ng	100
59) 2-Hexanone	15.92	43	1969	N.D.		
60) Dibromochloromethane	0.00	129	0	N.D.		
61) 1,2-Dibromoethane	0.00	107	0	N.D.		
62) n-Butyl Acetate	16.54	43	2949	N.D.		
63) n-Octane	16.65	57	1378	N.D.		
64) Tetrachloroethene	16.79	166	1285	N.D.		
65) Chlorobenzene	17.47	112	435	N.D.		
66) Ethylbenzene	17.83	91	4884	N.D.		
67) m- & p-Xylenes	17.99	91	10875	N.D.		
68) Bromoform	0.00	173	0	N.D.		
69) Styrene	18.33	104	2348	N.D.		
70) o-Xylene	18.43	91	4136	N.D.		
71) n-Nonane	18.63	43	3326	N.D.		
72) 1,1,2,2-Tetrachloroethane	0.00	83	0	N.D.		
74) Cumene	18.96	105	931	N.D.		
75) alpha-Pinene	19.31	93	9761	N.D.		
76) n-Propylbenzene	19.41	91	2393	N.D.		
77) 3-Ethyltoluene	19.50	105	3024	N.D.		
78) 4-Ethyltoluene	19.54	105	1887	N.D.		
79) 1,3,5-Trimethylbenzene	19.61	105	1372	N.D.		
80) alpha-Methylstyrene	19.75	118	606	N.D.		
81) 2-Ethyltoluene	19.78	105	1417	N.D.		
82) 1,2,4-Trimethylbenzene	19.97	105	3248	N.D.		
83) n-Decane	20.06	57	3329	N.D.		
84) Benzyl Chloride	20.10	91	865	N.D.		
85) 1,3-Dichlorobenzene	20.11	146	682	N.D.		
86) 1,4-Dichlorobenzene	20.17	146	1208	N.D.		
87) sec-Butylbenzene	20.22	105	604	N.D.		
88) 4-Isopropyltoluene (p-...	20.35	119	3260	N.D.		
89) 1,2,3-Trimethylbenzene	20.35	105	1323	N.D.		
90) 1,2-Dichlorobenzene	20.47	146	692	N.D.		
91) d-Limonene	20.48	68	2429	N.D.		
92) 1,2-Dibromo-3-Chloropr...	20.86	157	457	N.D.		
93) n-Undecane	21.17	57	2469	N.D.		
94) 1,2,4-Trichlorobenzene	21.98	180	1896	N.D.		
95) Naphthalene	22.08	128	11945	N.D.		
96) n-Dodecane	22.08	57	3644	N.D.		
97) Hexachlorobutadiene	22.39	225	861	N.D.		
98) Cyclohexanone	18.14	55	1964	N.D.		
99) tert-Butylbenzene	19.97	119	662	N.D.		
100) n-Butylbenzene	20.72	91	1555	N.D.		

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111921.D
 Acq On : 11 Mar 2019 18:42
 Sample : P1901179-003 (1000mL)
 Misc : S31-02211904

Vial: 1
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:09:24 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M



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Data File : I:\MS13\DATA\2019_03\11\03111921.D
 Acq On : 11 Mar 2019 18:42
 Sample : P1901179-003 (1000mL)
 Misc : S31-02211904

Vial: 1
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Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.97	130	93047	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.10	114	409377	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	180111	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	110943	13.984	ng	-0.02
Spiked Amount	12.500	Range	70 - 130	Recovery	=	111.84%
57) Toluene-d8 (SS2)	15.54	98	450833	11.689	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	93.52%
73) Bromofluorobenzene (SS3)	18.83	174	169718	12.221	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	97.76%

Target Compounds

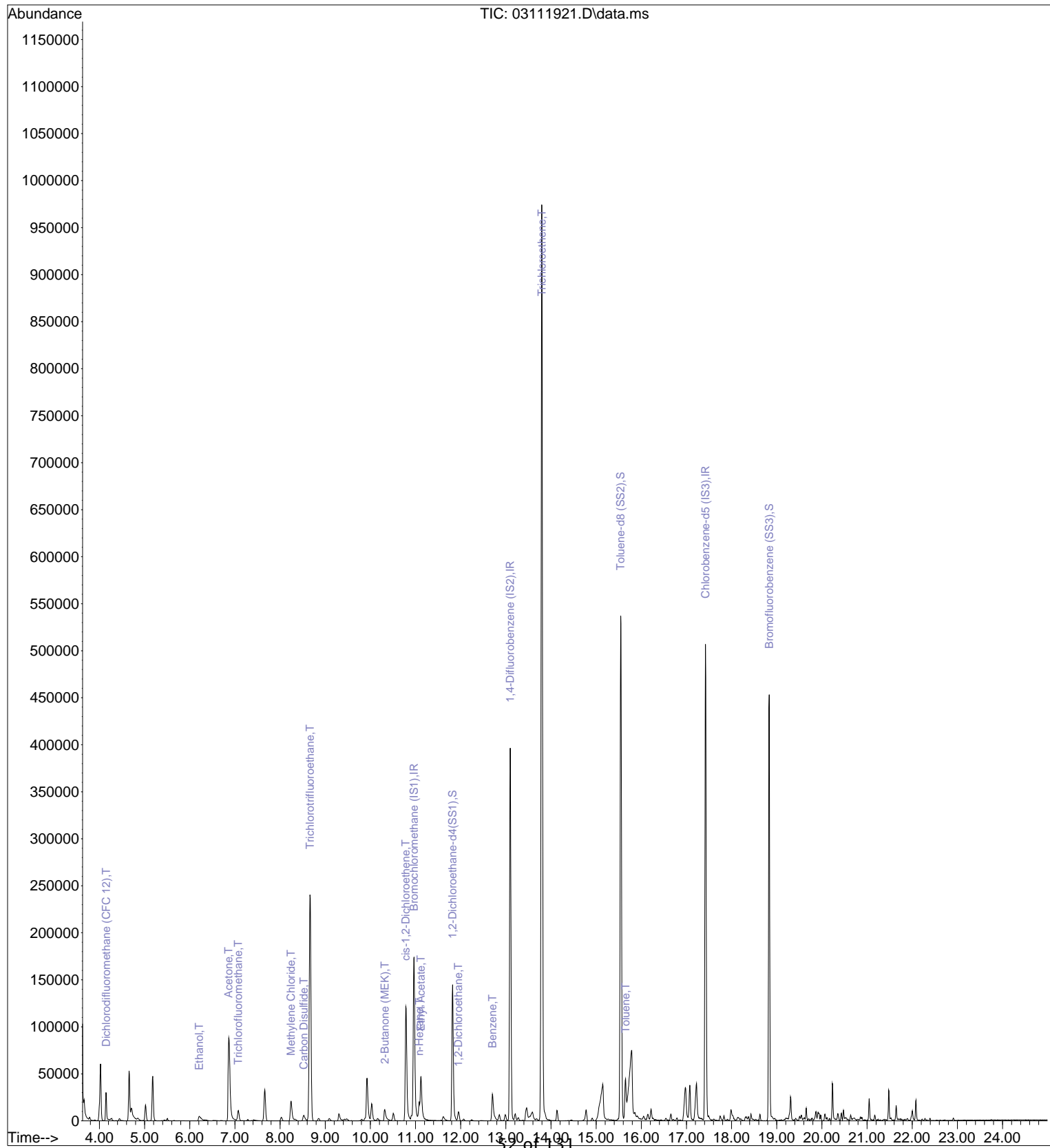
						Qvalue
3) Dichlorodifluoromethan...	4.15	85	27542	1.991	ng	99
10) Ethanol	6.21	45	15665	2.838	ng	98
13) Acetone	6.86	58	41157	7.018	ng	86
14) Trichlorofluoromethane	7.07	101	12455	1.075	ng	100
19) Methylene Chloride	8.24	84	16541	2.225	ng	97
21) Trichlorotrifluoroethane	8.66	151	122773	15.012	ng	98
22) Carbon Disulfide	8.53	76	18566	0.611	ng	91
27) 2-Butanone (MEK)	10.32	72	6464	1.202	ng	# 69
28) cis-1,2-Dichloroethene	10.79	61	88520	9.455	ng	96
30) Ethyl Acetate	11.12	61	11883	4.464	ng	86
31) n-Hexane	11.08	57	10028	0.812	ng	# 100
36) 1,2-Dichloroethane	11.95	62	3715	0.483	ng	97
41) Benzene	12.70	78	21228	0.628	ng	98
47) Trichloroethene	13.80	130	420086	42.992	ng	100
58) Toluene	15.65	91	37664	0.919	ng	100

(#) = qualifier out of range (m) = manual integration (+) = signals summed

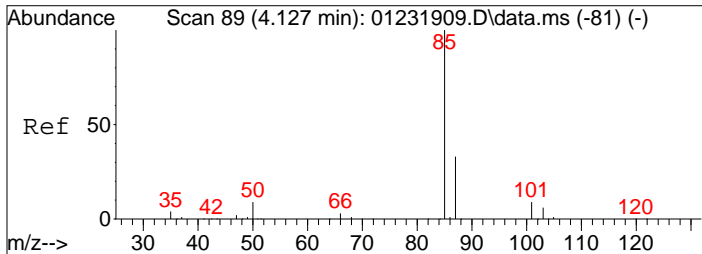
Data File : I:\MS13\DATA\2019_03\11\03111921.D
 Acq On : 11 Mar 2019 18:42
 Sample : P1901179-003 (1000mL)
 Misc : S31-02211904

Vial: 1
 Operator: WA
 Inst : MS13

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 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

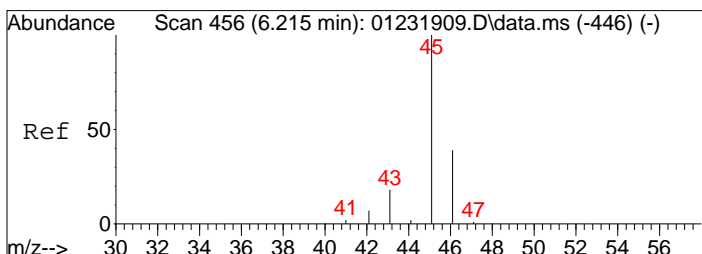
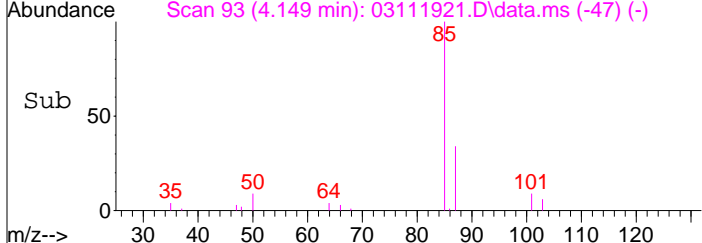
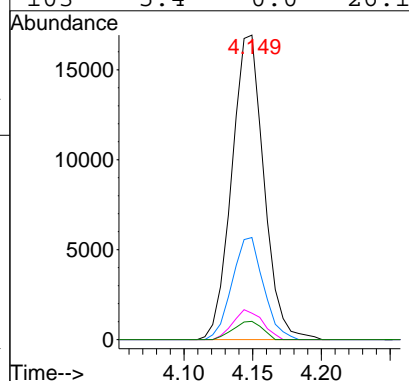
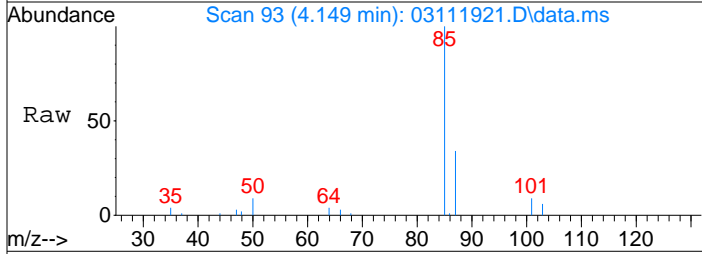


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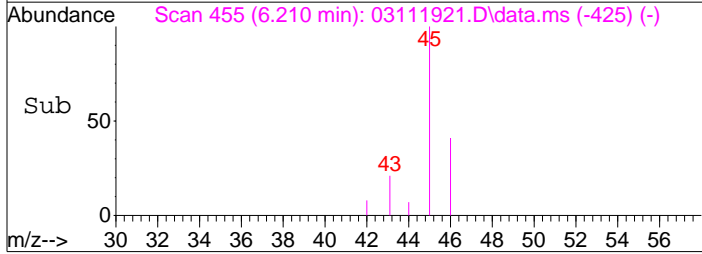
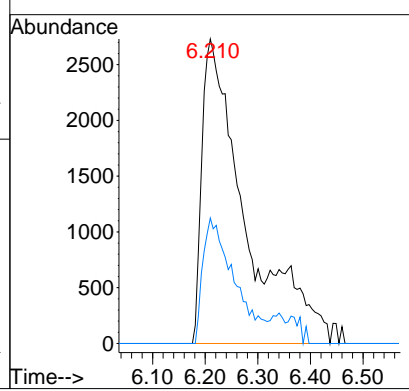
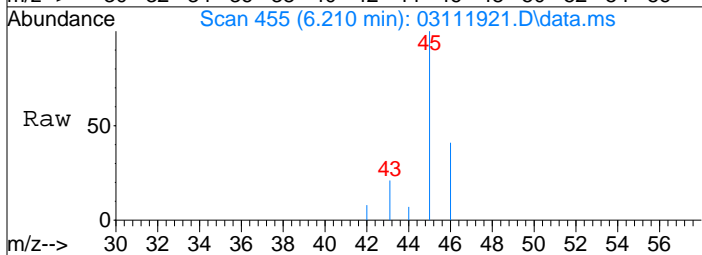
#3
 Dichlorodifluoromethane (CFC 12)
 Concen: 1.99 ng
 RT: 4.15 min Scan# 93
 Delta R.T. 0.011 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

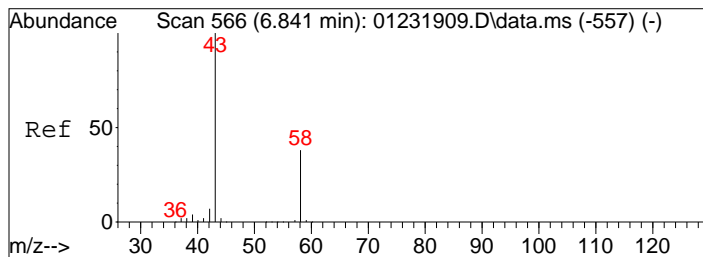
Tgt Ion	Resp	Lower	Upper
85	100		
87	32.7	12.6	52.6
101	9.0	0.0	29.5
103	5.4	0.0	26.1



#10
 Ethanol
 Concen: 2.84 ng
 RT: 6.21 min Scan# 455
 Delta R.T. -0.080 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

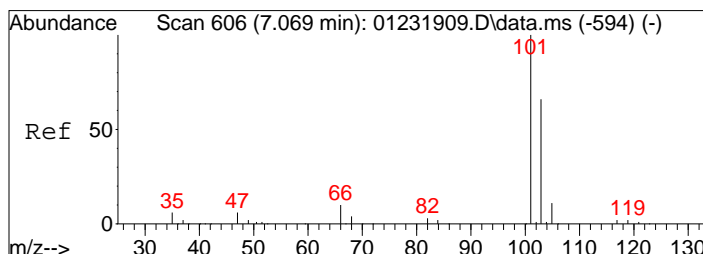
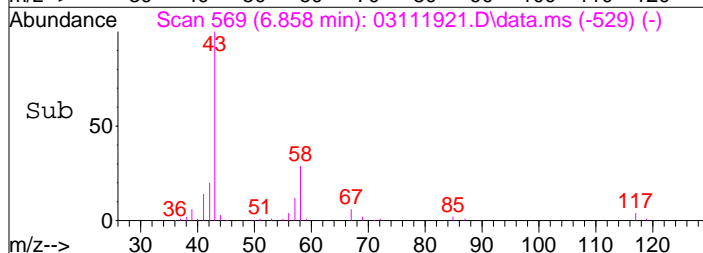
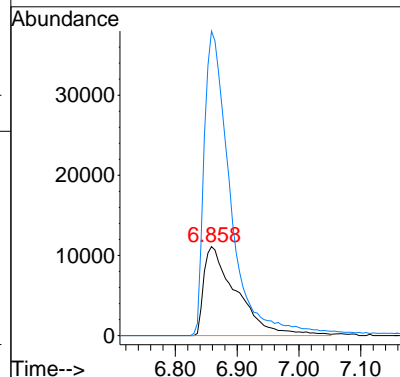
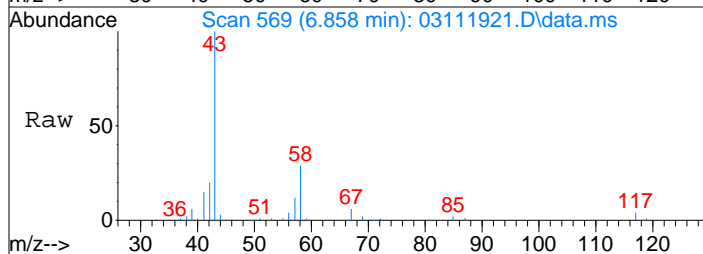
Tgt Ion	Resp	Lower	Upper
45	100		
46	35.7	16.8	56.8





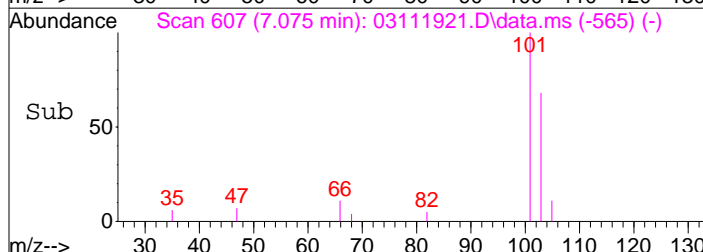
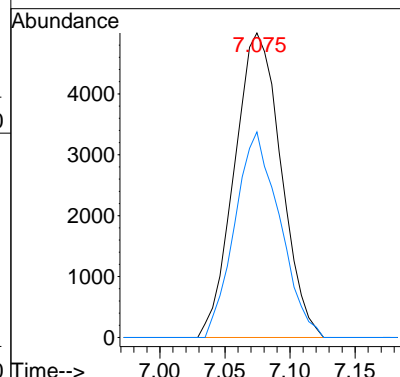
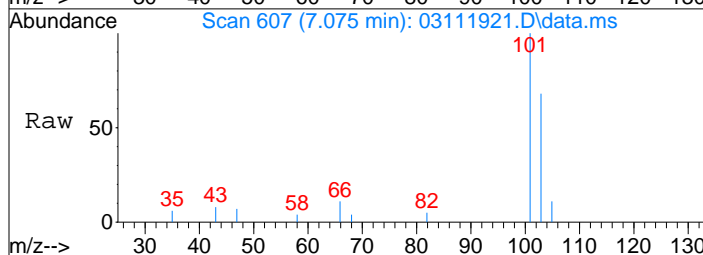
#13
 Acetone
 Concen: 7.02 ng
 RT: 6.86 min Scan# 569
 Delta R.T. -0.023 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

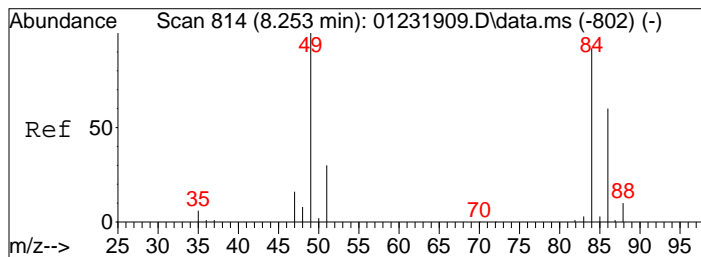
Tgt Ion: 58 Resp: 41157
 Ion Ratio Lower Upper
 58 100
 43 275.0 271.8 331.8



#14
 Trichlorofluoromethane
 Concen: 1.07 ng
 RT: 7.07 min Scan# 607
 Delta R.T. -0.012 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

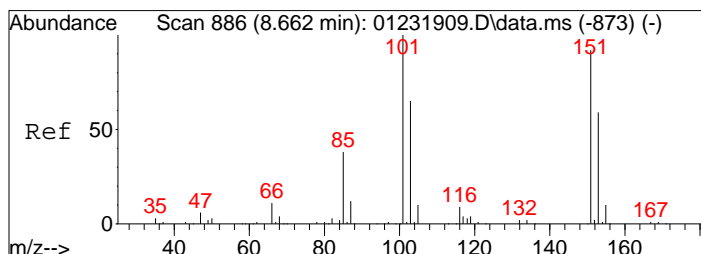
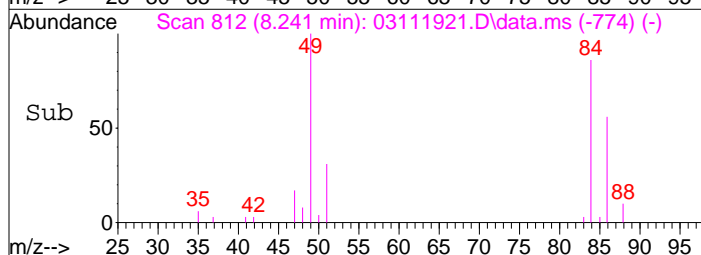
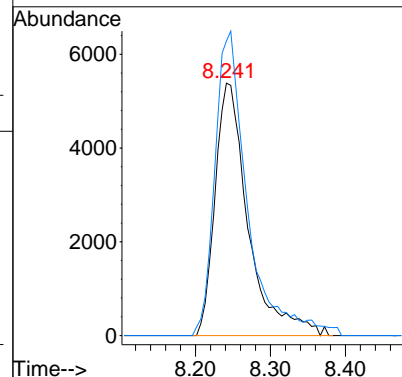
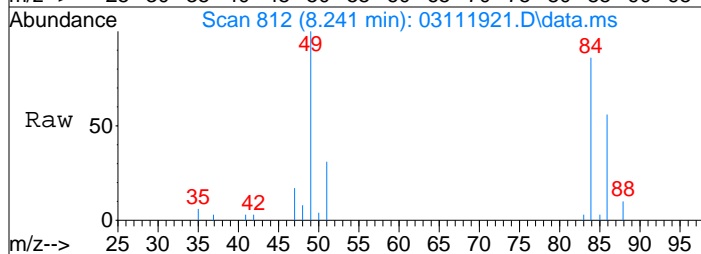
Tgt Ion: 101 Resp: 12455
 Ion Ratio Lower Upper
 101 100
 103 64.9 44.6 84.6





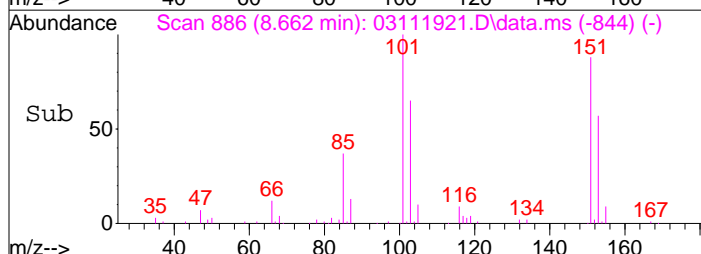
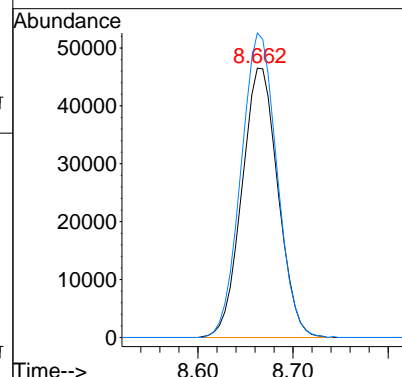
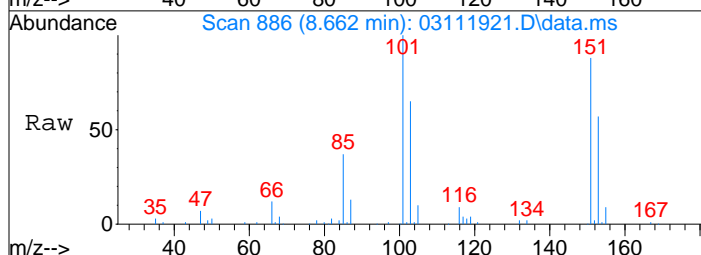
#19
 Methylene Chloride
 Concen: 2.22 ng
 RT: 8.24 min Scan# 812
 Delta R.T. -0.034 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

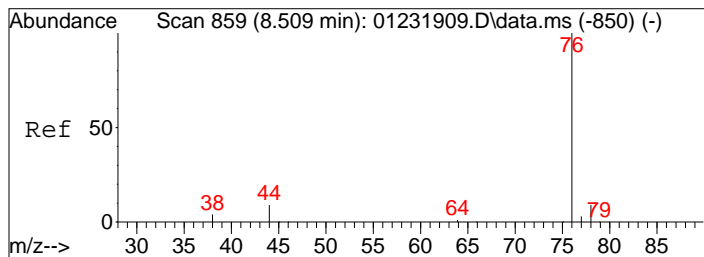
Tgt Ion: 84 Resp: 16541
 Ion Ratio Lower Upper
 84 100
 49 119.0 90.3 140.3



#21
 Trichlorotrifluoroethane
 Concen: 15.01 ng
 RT: 8.66 min Scan# 886
 Delta R.T. -0.012 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

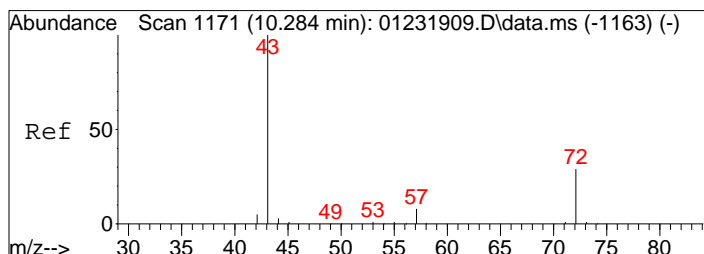
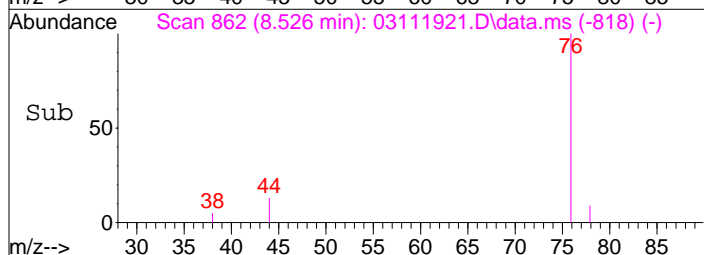
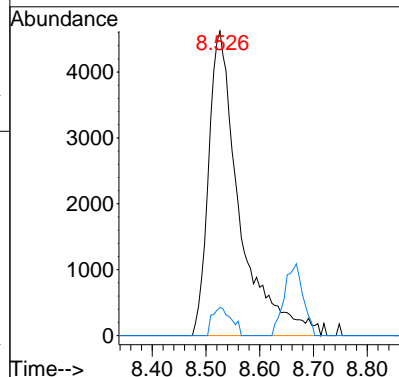
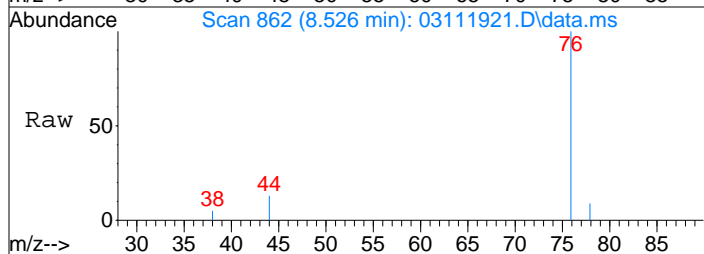
Tgt Ion: 151 Resp: 122773
 Ion Ratio Lower Upper
 151 100
 101 112.4 95.1 135.1





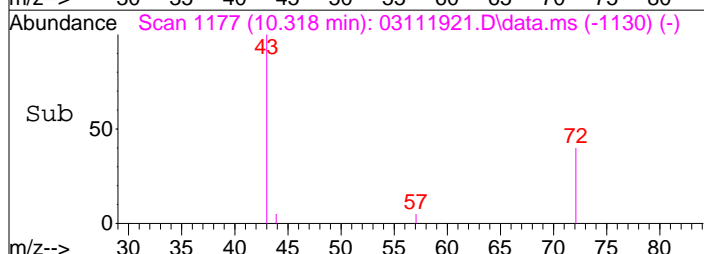
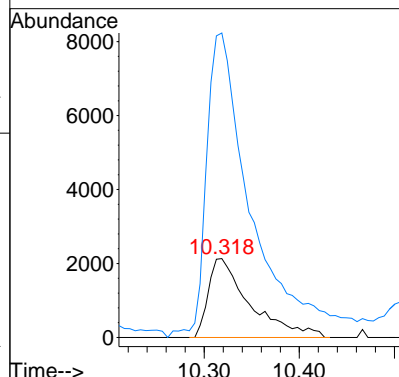
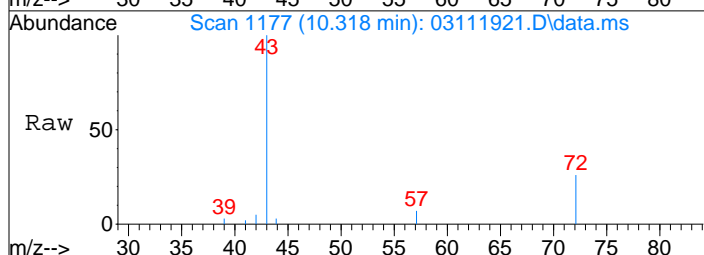
#22
 Carbon Disulfide
 Concen: 0.61 ng
 RT: 8.53 min Scan# 862
 Delta R.T. -0.000 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

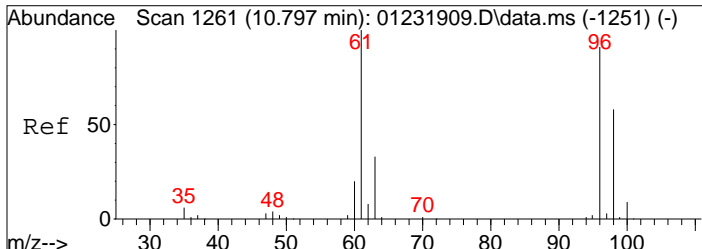
Tgt Ion: 76 Resp: 18566
 Ion Ratio Lower Upper
 76 100
 78 5.6 0.0 29.0



#27
 2-Butanone (MEK)
 Concen: 1.20 ng
 RT: 10.32 min Scan# 1177
 Delta R.T. 0.016 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

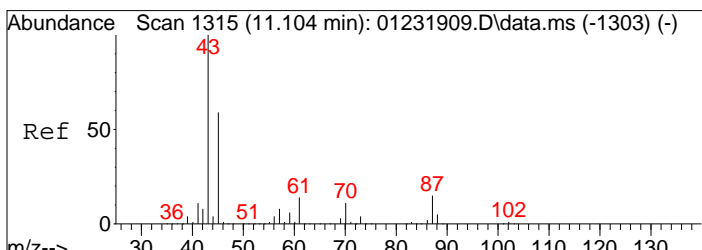
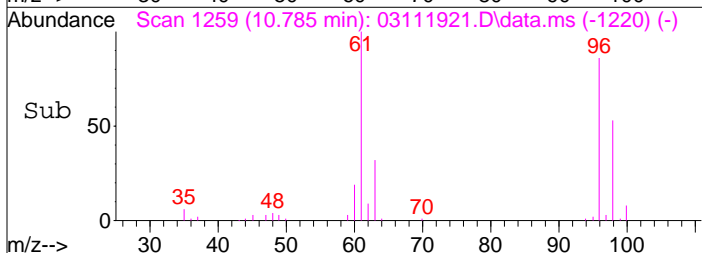
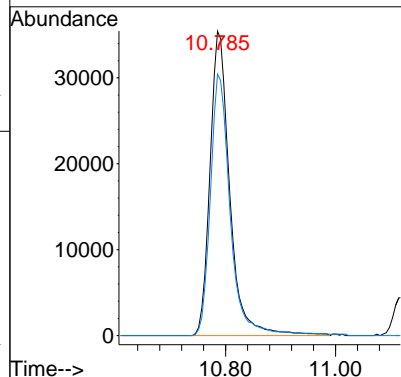
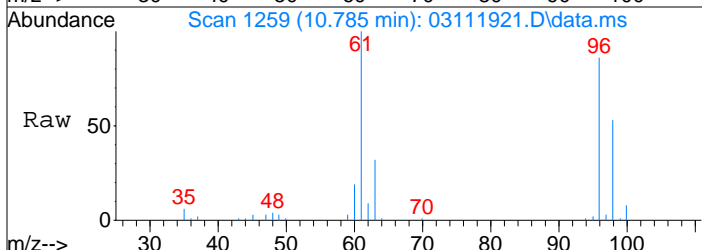
Tgt Ion: 72 Resp: 6464
 Ion Ratio Lower Upper
 72 100
 43 431.1 343.2 383.2#





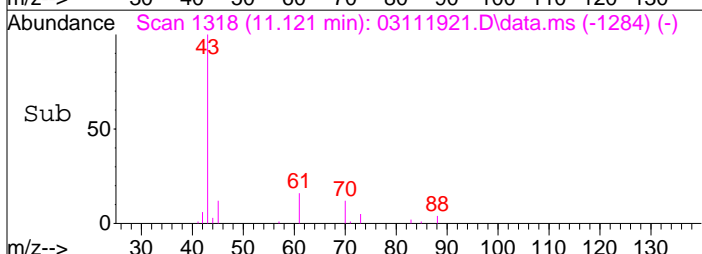
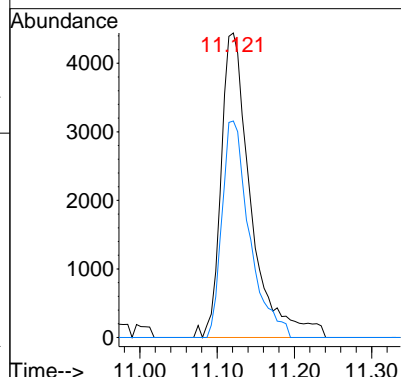
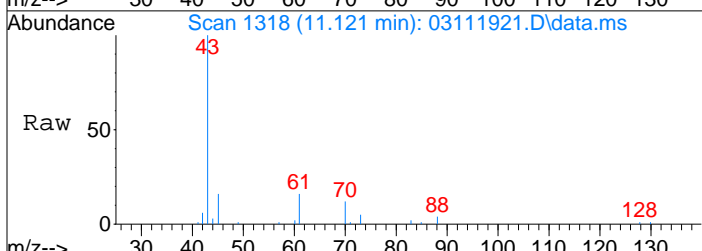
#28
 cis-1,2-Dichloroethene
 Concen: 9.45 ng
 RT: 10.79 min Scan# 1259
 Delta R.T. -0.029 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

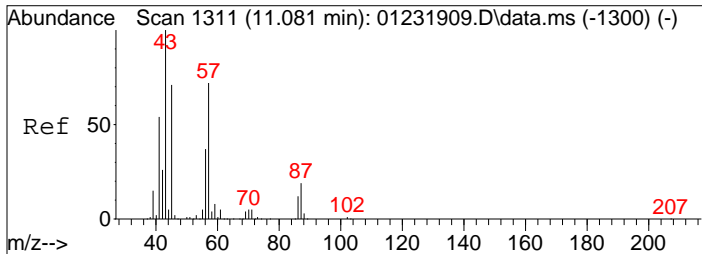
Tgt Ion	Resp	Lower	Upper
61	100		
96	86.6	63.3	103.3



#30
 Ethyl Acetate
 Concen: 4.46 ng
 RT: 11.12 min Scan# 1318
 Delta R.T. -0.006 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

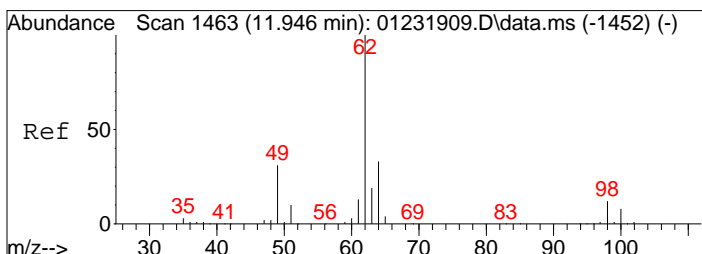
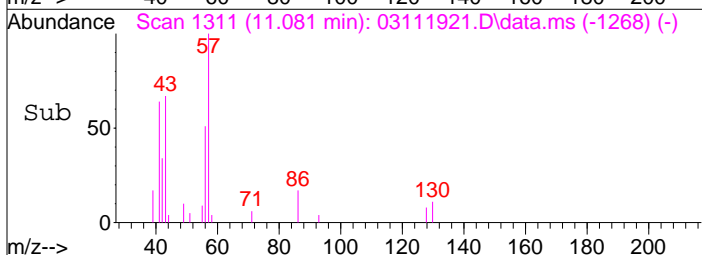
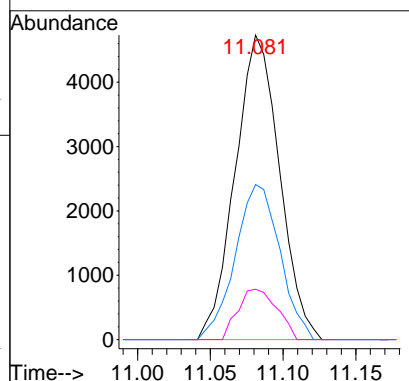
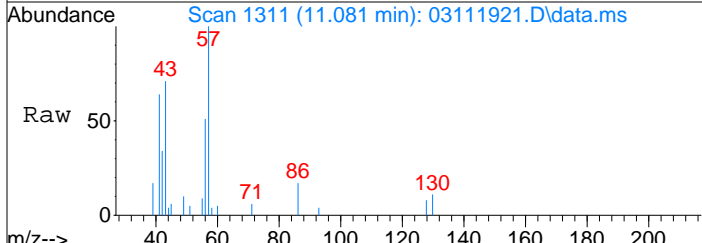
Tgt Ion	Resp	Lower	Upper
61	100		
70	65.7	57.7	97.7





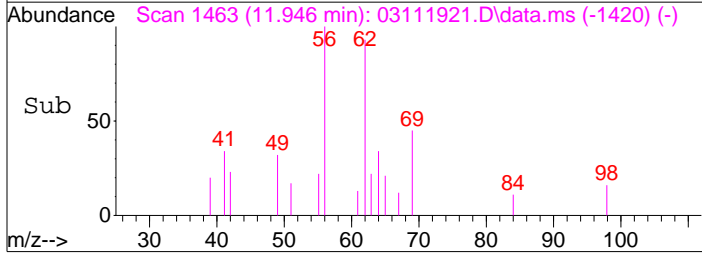
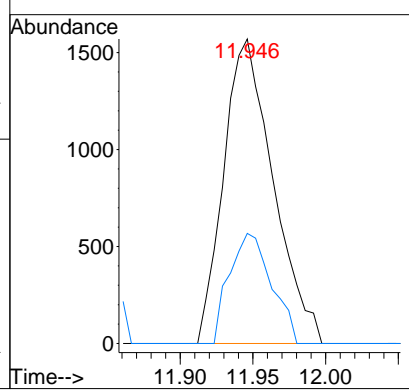
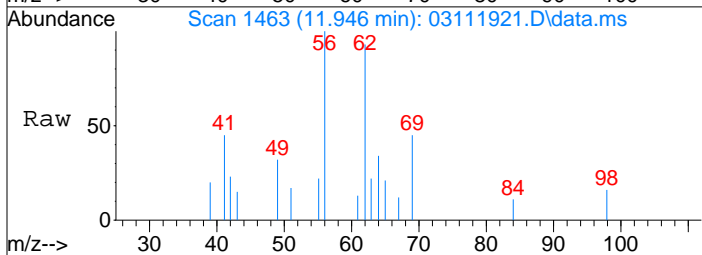
#31
 n-Hexane
 Concen: 0.81 ng
 RT: 11.08 min Scan# 1311
 Delta R.T. -0.006 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

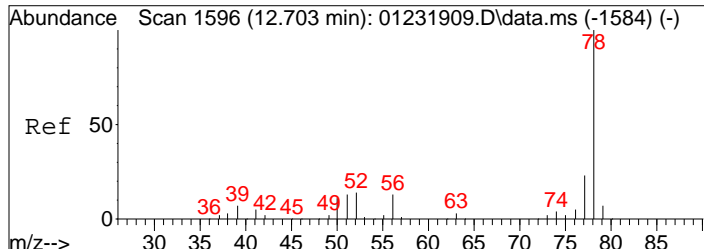
Tgt Ion	Resp	Lower	Upper
57	10028		
56	51.2	0.0	0.0#
86	14.6	0.0	0.0#



#36
 1,2-Dichloroethane
 Concen: 0.48 ng
 RT: 11.95 min Scan# 1463
 Delta R.T. -0.006 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

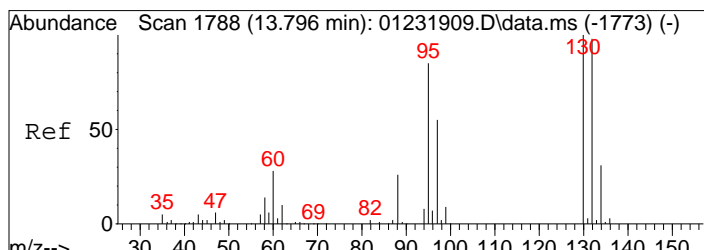
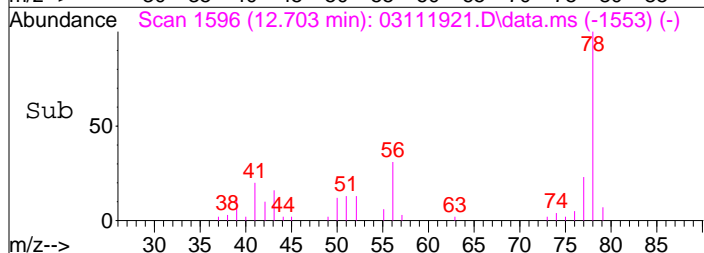
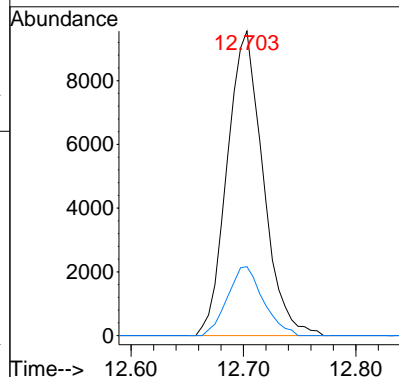
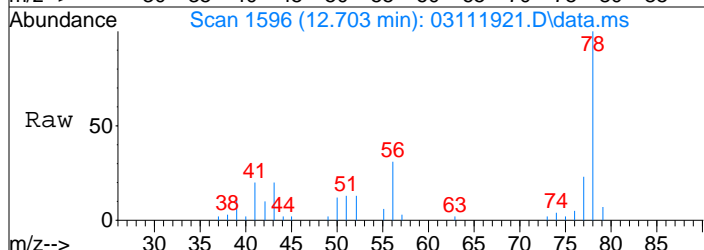
Tgt Ion	Resp	Lower	Upper
62	3715		
64	30.7	12.6	52.6





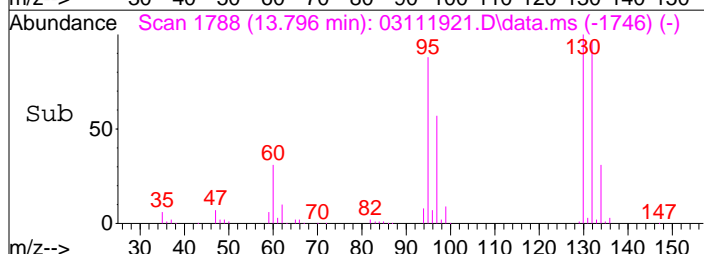
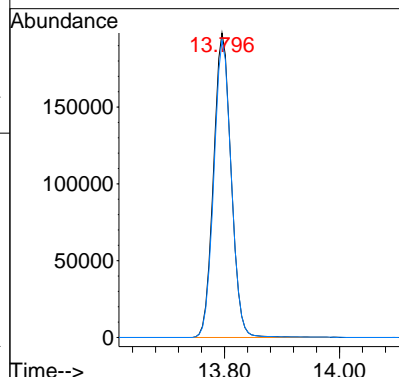
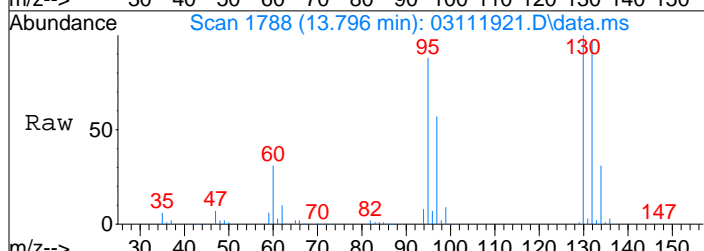
#41
Benzene
Concen: 0.63 ng
RT: 12.70 min Scan# 1596
Delta R.T. -0.006 min
Lab File: 03111921.D
Acq: 11 Mar 2019 18:42

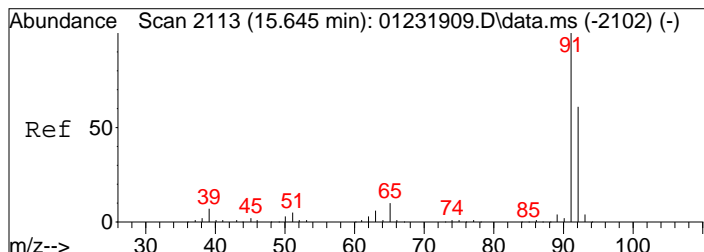
Tgt Ion	Resp	Lower	Upper
78	21228	100	
77	22.5	3.5	43.5



#47
Trichloroethene
Concen: 42.99 ng
RT: 13.80 min Scan# 1788
Delta R.T. -0.012 min
Lab File: 03111921.D
Acq: 11 Mar 2019 18:42

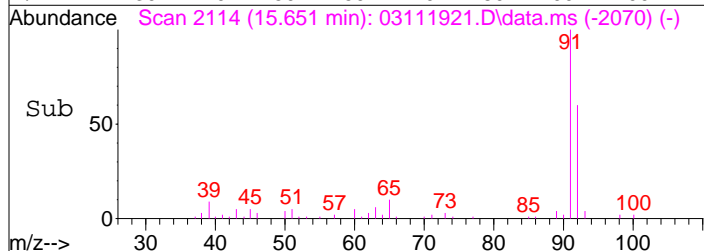
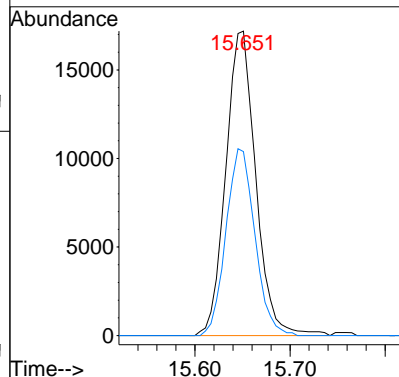
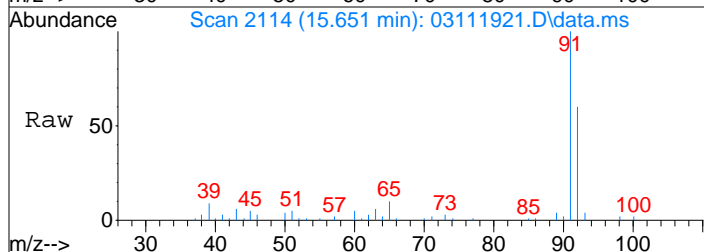
Tgt Ion	Resp	Lower	Upper
130	420086	100	
132	96.9	76.7	116.7





#58
 Toluene
 Concen: 0.92 ng
 RT: 15.65 min Scan# 2114
 Delta R.T. -0.000 min
 Lab File: 03111921.D
 Acq: 11 Mar 2019 18:42

Tgt Ion	Resp	Lower	Upper
91	100		
92	59.2	39.5	79.5



Data File : I:\MS13\DATA\2019_03\11\03111922.D
 Acq On : 11 Mar 2019 19:15
 Sample : P1901179-004 (1000mL)
 Misc : S31-02211904

Vial: 5
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:10:04 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

DA 3/14/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	93346	12.500	ng	-0.03
37) 1,4-Difluorobenzene (IS2)	13.10	114	407063	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	179457	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	111465	14.005	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	112.08%	
57) Toluene-d8 (SS2)	15.54	98	449465	11.696	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	93.60%	
73) Bromofluorobenzene (SS3)	18.83	174	169599	12.257	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	98.08%	

Target Compounds

						Qvalue
2) Propene	0.00	42	0	N.D.	d	
3) Dichlorodifluoromethan...	4.14	85	31011	2.235	ng	98
4) Chloromethane	4.43	50	3311	N.D.		
5) 1,2-Dichloro-1,1,2,2-t...	4.68	135	801	N.D.		
6) Vinyl Chloride	0.00	62	0	N.D.		
7) 1,3-Butadiene	0.00	54	0	N.D.		
8) Bromomethane	0.00	94	0	N.D.		
9) Chloroethane	0.00	64	0	N.D.		
10) Ethanol	6.23	45	6211	1.122	ng	79
11) Acetonitrile	6.51	41	65	N.D.		
12) Acrolein	6.69	56	60	N.D.		
13) Acetone	6.86	58	23740	4.035	ng	# 74
14) Trichlorofluoromethane	7.07	101	13262	1.141	ng	99
15) 2-Propanol (Isopropanol)	7.42	45	2710	N.D.		
16) Acrylonitrile	7.66	53	61	N.D.		
17) 1,1-Dichloroethene	0.00	96	0	N.D.		
18) 2-Methyl-2-Propanol (t...	8.34	59	675	N.D.		
19) Methylene Chloride	8.25	84	9009	1.208	ng	95
20) 3-Chloro-1-propene (Al...	0.00	41	0	N.D.		
21) Trichlorotrifluoroethane	8.66	151	49381	6.019	ng	97
22) Carbon Disulfide	8.53	76	2046	N.D.		
23) trans-1,2-Dichloroethene	0.00	61	0	N.D.		
24) 1,1-Dichloroethane	0.00	63	0	N.D.		
25) Methyl tert-Butyl Ether	0.00	73	0	N.D.		
26) Vinyl Acetate	10.03	86	768	N.D.		
27) 2-Butanone (MEK)	10.34	72	1507	N.D.		
28) cis-1,2-Dichloroethene	10.80	61	5209	0.555	ng	97
29) Diisopropyl Ether	0.00	87	0	N.D.		
30) Ethyl Acetate	11.13	61	6320	2.366	ng	88
31) n-Hexane	11.08	57	8837	0.713	ng	# 100
32) Chloroform	11.13	83	853	N.D.		
34) Tetrahydrofuran (THF)	11.64	72	478	N.D.		
35) Ethyl tert-Butyl Ether	0.00	87	0	N.D.		
36) 1,2-Dichloroethane	11.94	62	598	N.D.		
38) 1,1,1-Trichloroethane	0.00	97	0	N.D.		
39) Isopropyl Acetate	0.00	61	0	N.D.		
40) 1-Butanol	12.74	56	1513	N.D.		
41) Benzene	12.70	78	19912	0.592	ng	98
42) Carbon Tetrachloride	12.86	117	3770	N.D.		
43) Cyclohexane	12.99	84	2850	N.D.		
44) tert-Amyl Methyl Ether	0.00	73	0	N.D.		
45) 1,2-Dichloropropane	0.00	63	0	N.D.		
46) Bromodichloromethane	0.00	83	0	N.D.		
47) Trichloroethene	13.80	130	17610	1.812	ng	100
48) 1,4-Dioxane	0.00	88	0	N.D.		
49) 2,2,4-Trimethylpentane...	13.86	57	2408	N.D.		
50) Methyl Methacrylate	14.14	100	644	N.D.		

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Data File : I:\MS13\DATA\2019_03\11\03111922.D
 Acq On : 11 Mar 2019 19:15
 Sample : P1901179-004 (1000mL)
 Misc : S31-02211904

Vial: 5
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:10:04 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

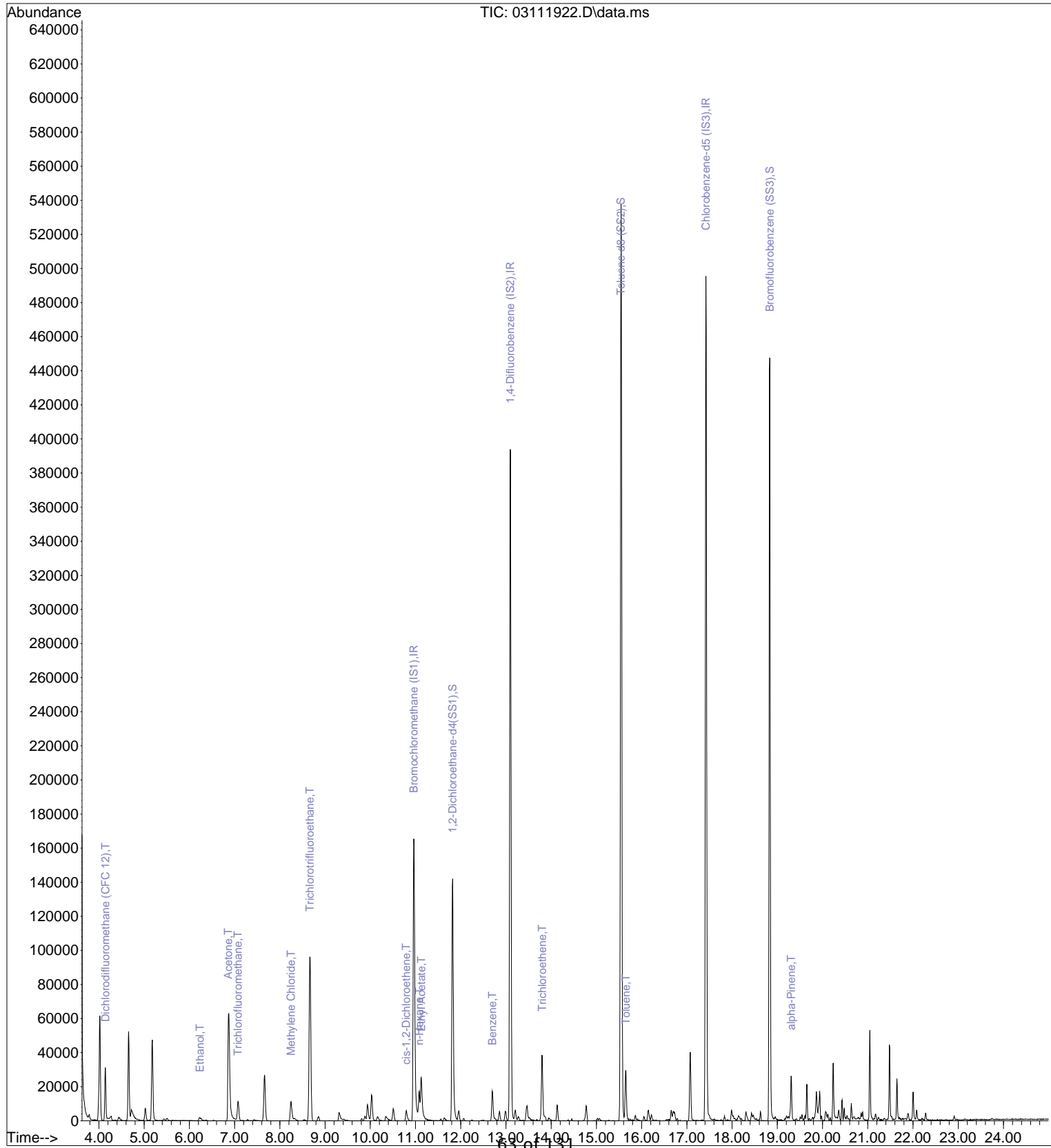
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.14	71	2616	N.D.		
52) cis-1,3-Dichloropropene	0.00	75	0	N.D.		
53) 4-Methyl-2-pentanone	0.00	58	0	N.D.		
54) trans-1,3-Dichloropropene	0.00	75	0	N.D.		
55) 1,1,2-Trichloroethane	0.00	97	0	N.D.		
58) Toluene	15.65	91	26877	0.658	ng	99
59) 2-Hexanone	15.90	43	1527	N.D.		
60) Dibromochloromethane	0.00	129	0	N.D.		
61) 1,2-Dibromoethane	0.00	107	0	N.D.		
62) n-Butyl Acetate	16.54	43	1758	N.D.		
63) n-Octane	16.66	57	1309	N.D.		
64) Tetrachloroethene	16.79	166	557	N.D.		
65) Chlorobenzene	0.00	112	0	N.D.		
66) Ethylbenzene	17.83	91	2885	N.D.		
67) m- & p-Xylenes	17.99	91	6109	N.D.		
68) Bromoform	0.00	173	0	N.D.		
69) Styrene	18.34	104	1242	N.D.		
70) o-Xylene	18.43	91	2615	N.D.		
71) n-Nonane	18.63	43	2470	N.D.		
72) 1,1,2,2-Tetrachloroethane	0.00	83	0	N.D.		
74) Cumene	18.96	105	405	N.D.		
75) alpha-Pinene	19.30	93	10401	0.471	ng	# 55
76) n-Propylbenzene	19.41	91	863	N.D.		
77) 3-Ethyltoluene	19.50	105	1577	N.D.		
78) 4-Ethyltoluene	19.54	105	637	N.D.		
79) 1,3,5-Trimethylbenzene	19.61	105	562	N.D.		
80) alpha-Methylstyrene	19.75	118	306	N.D.		
81) 2-Ethyltoluene	19.78	105	698	N.D.		
82) 1,2,4-Trimethylbenzene	19.97	105	1535	N.D.		
83) n-Decane	20.07	57	3245	N.D.		
84) Benzyl Chloride	19.91	91	658	N.D.		
85) 1,3-Dichlorobenzene	20.18	146	319	N.D.		
86) 1,4-Dichlorobenzene	20.18	146	319	N.D.		
87) sec-Butylbenzene	20.21	105	113	N.D.		
88) 4-Isopropyltoluene (p-...	20.35	119	1972	N.D.		
89) 1,2,3-Trimethylbenzene	20.35	105	528	N.D.		
90) 1,2-Dichlorobenzene	0.00	146	0	N.D.		
91) d-Limonene	20.48	68	1748	N.D.		
92) 1,2-Dibromo-3-Chloropr...	0.00	157	0	N.D.		
93) n-Undecane	21.17	57	1342	N.D.		
94) 1,2,4-Trichlorobenzene	21.99	180	384	N.D.		
95) Naphthalene	22.09	128	3238	N.D.		
96) n-Dodecane	22.08	57	1726	N.D.		
97) Hexachlorobutadiene	0.00	225	0	N.D.		
98) Cyclohexanone	18.14	55	1522	N.D.		
99) tert-Butylbenzene	19.97	119	193	N.D.		
100) n-Butylbenzene	20.70	91	687	N.D.		

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111922.D
Acq On : 11 Mar 2019 19:15
Sample : P1901179-004 (1000mL)
Misc : S31-02211904

Vial: 5
Operator: WA
Inst : MS13

Quant Time: Mar 14 11:10:04 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 11:21:29 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



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Data File : I:\MS13\DATA\2019_03\11\03111922.D
 Acq On : 11 Mar 2019 19:15
 Sample : P1901179-004 (1000mL)
 Misc : S31-02211904

Vial: 5
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:10:04 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

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Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	93346	12.500	ng	-0.03
37) 1,4-Difluorobenzene (IS2)	13.10	114	407063	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	179457	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	111465	14.005	ng	-0.02
Spiked Amount	12.500	Range	70 - 130	Recovery	=	112.08%
57) Toluene-d8 (SS2)	15.54	98	449465	11.696	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	93.60%
73) Bromofluorobenzene (SS3)	18.83	174	169599	12.257	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	98.08%

Target Compounds

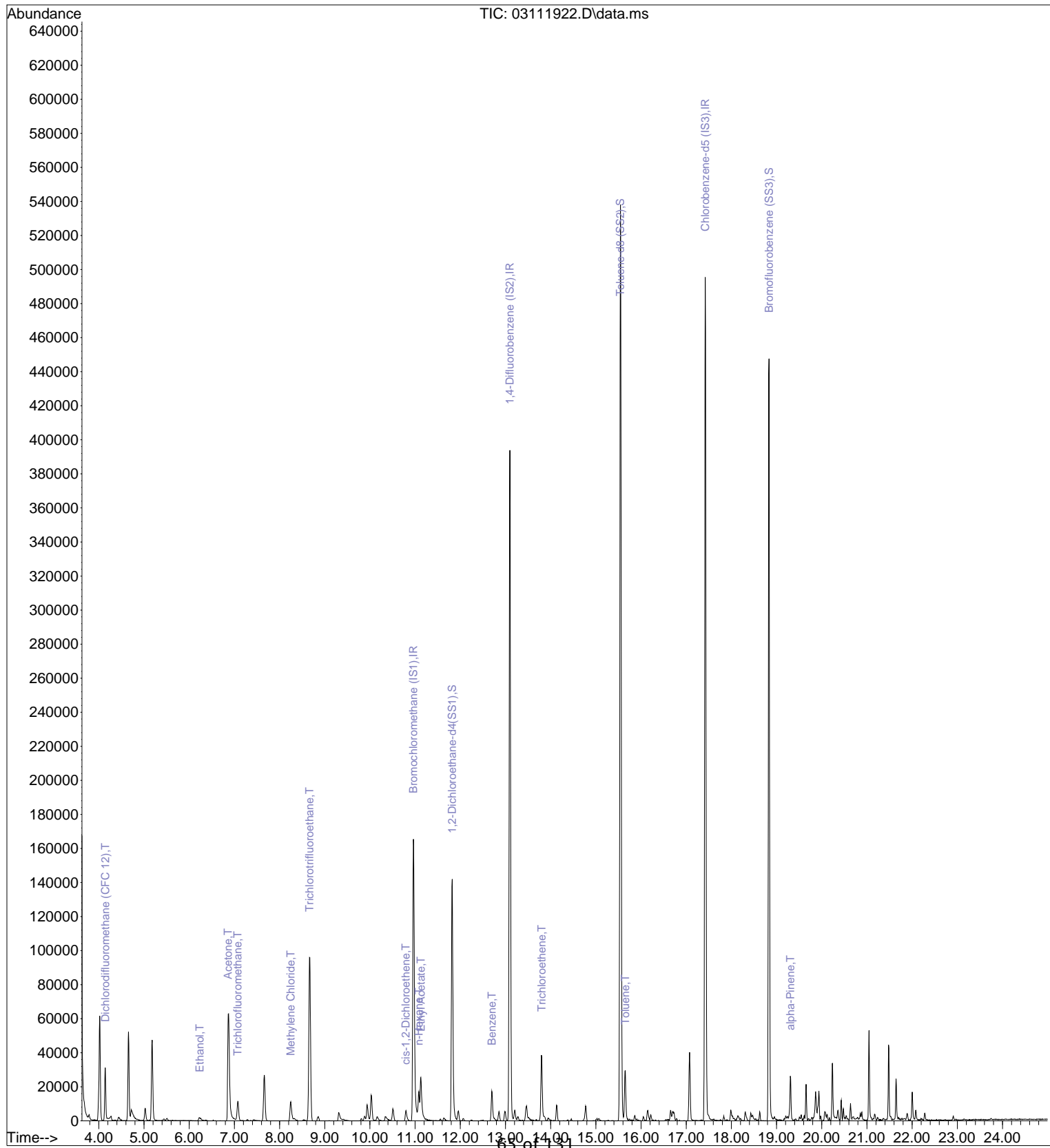
	R.T.	QIon	Response	Conc	Units	Qvalue
3) Dichlorodifluoromethan...	4.14	85	31011	2.235	ng	98
10) Ethanol	6.23	45	6211	1.122	ng	79
13) Acetone	6.86	58	23740	4.035	ng	# 74
14) Trichlorofluoromethane	7.07	101	13262	1.141	ng	99
19) Methylene Chloride	8.25	84	9009	1.208	ng	95
21) Trichlorotrifluoroethane	8.66	151	49381	6.019	ng	97
28) cis-1,2-Dichloroethene	10.80	61	5209	0.555	ng	97
30) Ethyl Acetate	11.13	61	6320	2.366	ng	88
31) n-Hexane	11.08	57	8837	0.713	ng	# 100
41) Benzene	12.70	78	19912	0.592	ng	98
47) Trichloroethene	13.80	130	17610	1.812	ng	100
58) Toluene	15.65	91	26877	0.658	ng	99
75) alpha-Pinene	19.30	93	10401	0.471	ng	# 55

(#) = qualifier out of range (m) = manual integration (+) = signals summed

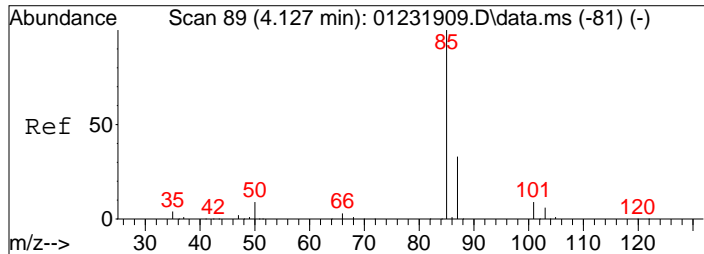
Data File : I:\MS13\DATA\2019_03\11\03111922.D
 Acq On : 11 Mar 2019 19:15
 Sample : P1901179-004 (1000mL)
 Misc : S31-02211904

Vial: 5
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:10:04 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

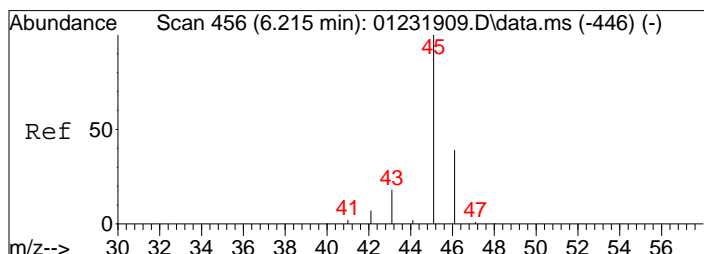
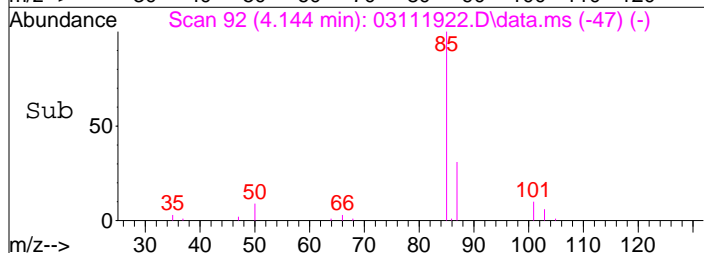
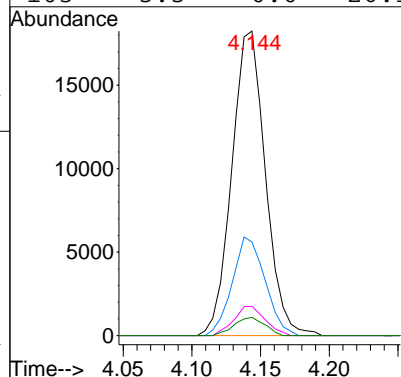
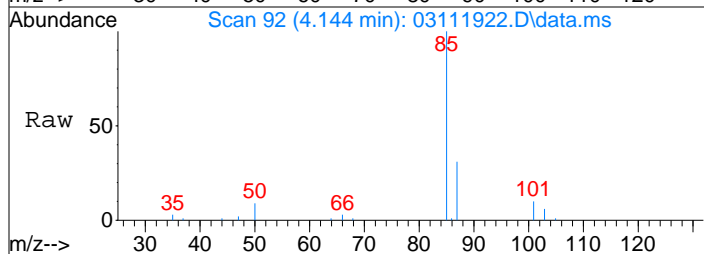


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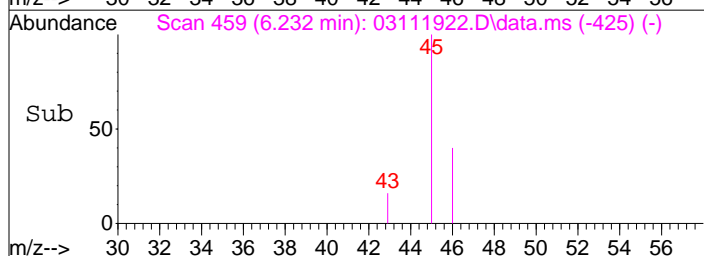
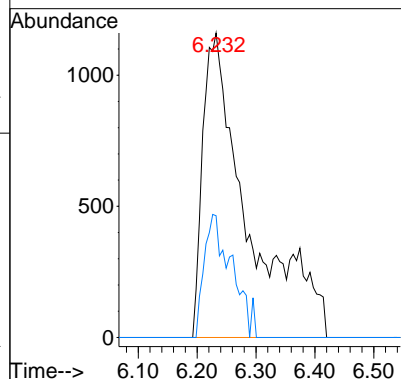
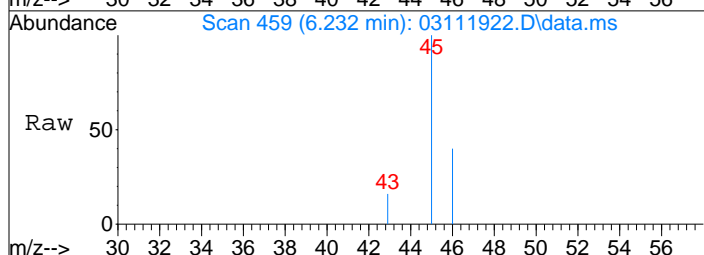
#3
 Dichlorodifluoromethane (CFC 12)
 Concen: 2.23 ng
 RT: 4.14 min Scan# 92
 Delta R.T. 0.005 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

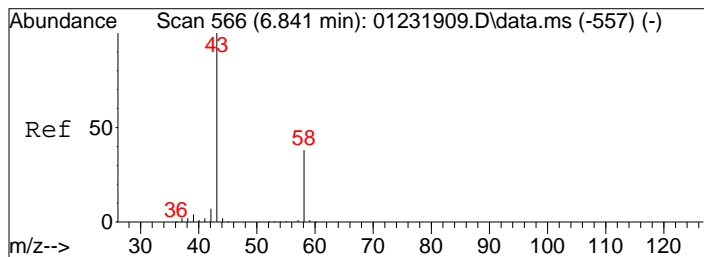
Tgt Ion	Resp	Lower	Upper
85	100		
87	31.6	12.6	52.6
101	9.0	0.0	29.5
103	5.5	0.0	26.1



#10
 Ethanol
 Concen: 1.12 ng
 RT: 6.23 min Scan# 459
 Delta R.T. -0.057 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

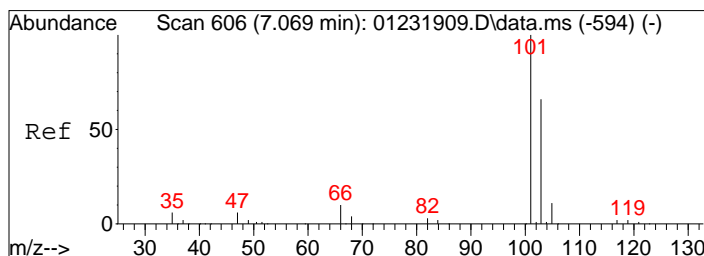
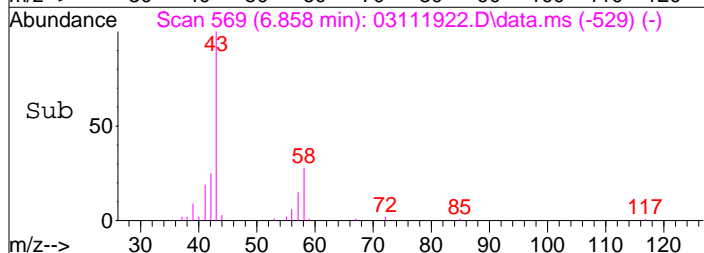
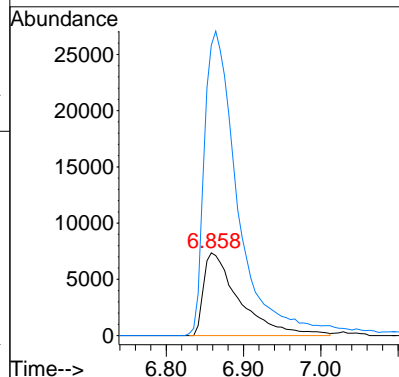
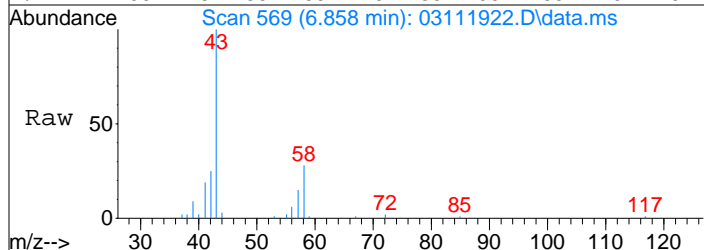
Tgt Ion	Resp	Lower	Upper
45	100		
46	24.6	16.8	56.8





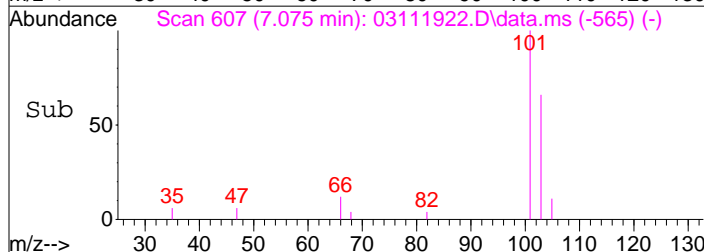
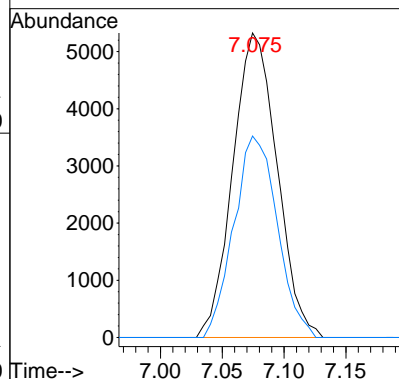
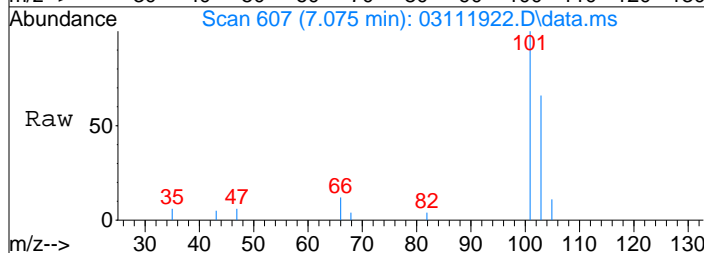
#13
 Acetone
 Concen: 4.04 ng
 RT: 6.86 min Scan# 569
 Delta R.T. -0.023 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

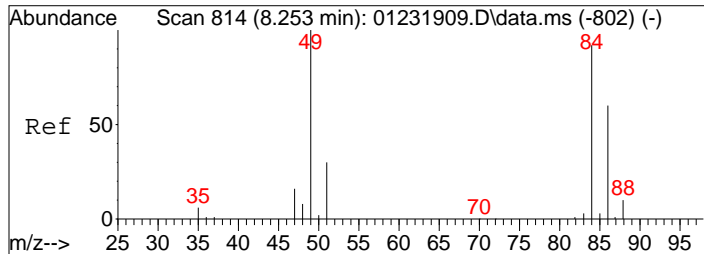
Tgt Ion	Resp	Lower	Upper
58	100		
43	352.5	271.8	331.8#



#14
 Trichlorofluoromethane
 Concen: 1.14 ng
 RT: 7.07 min Scan# 607
 Delta R.T. -0.012 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

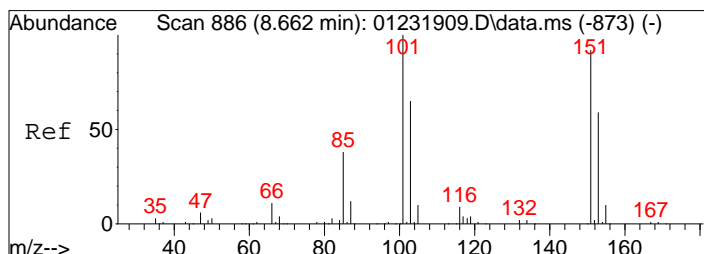
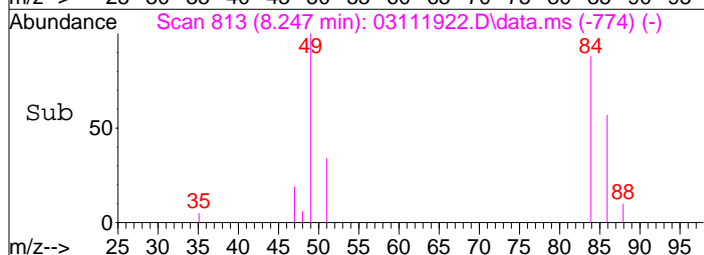
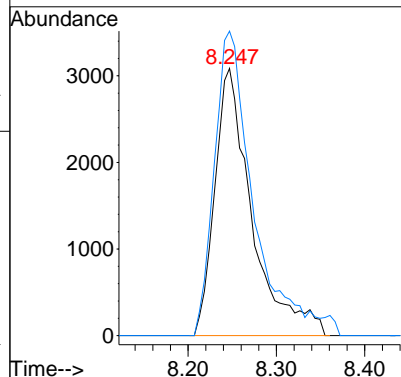
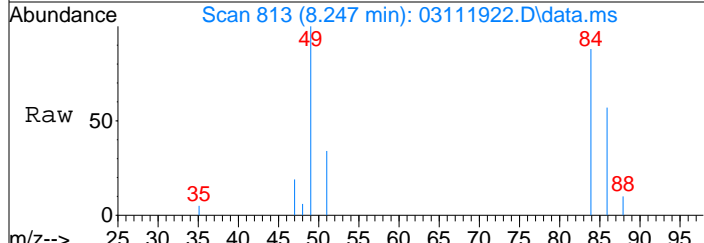
Tgt Ion	Resp	Lower	Upper
101	100		
103	65.0	44.6	84.6





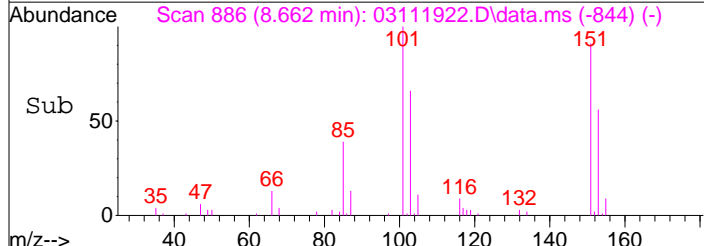
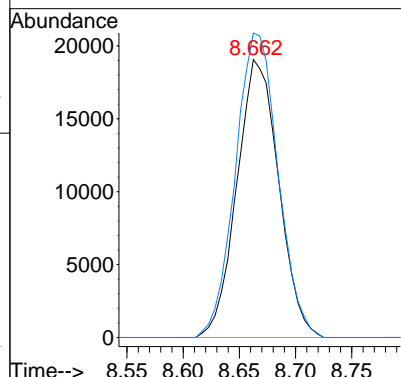
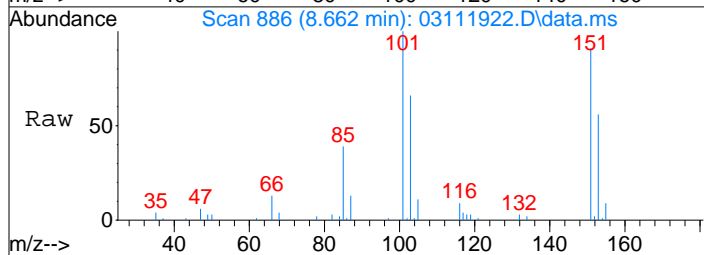
#19
 Methylene Chloride
 Concen: 1.21 ng
 RT: 8.25 min Scan# 813
 Delta R.T. -0.029 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

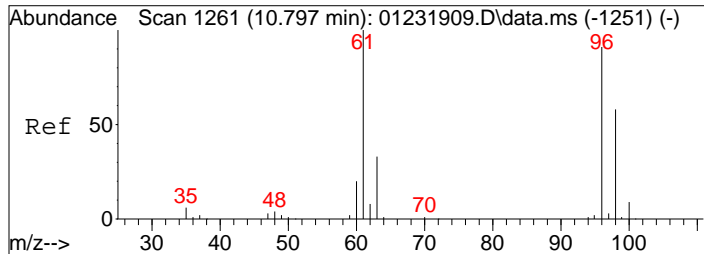
Tgt Ion: 84 Resp: 9009
 Ion Ratio Lower Upper
 84 100
 49 120.7 90.3 140.3



#21
 Trichlorotrifluoroethane
 Concen: 6.02 ng
 RT: 8.66 min Scan# 886
 Delta R.T. -0.012 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

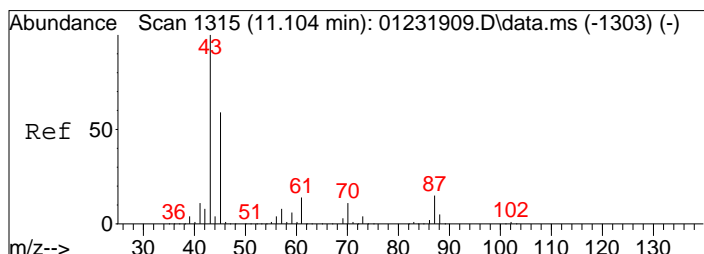
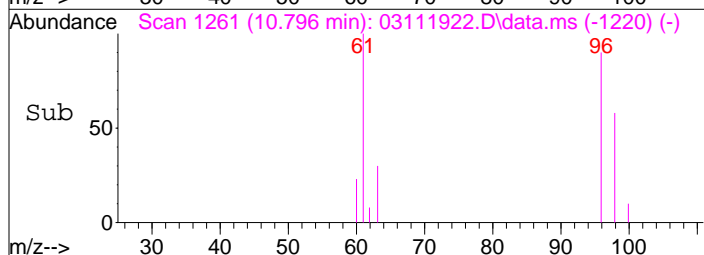
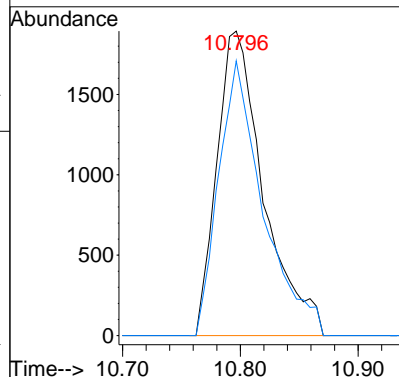
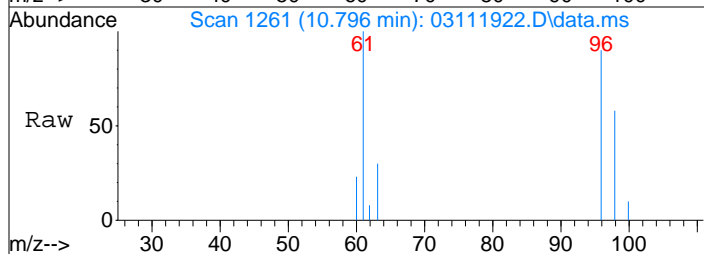
Tgt Ion: 151 Resp: 49381
 Ion Ratio Lower Upper
 151 100
 101 111.8 95.1 135.1





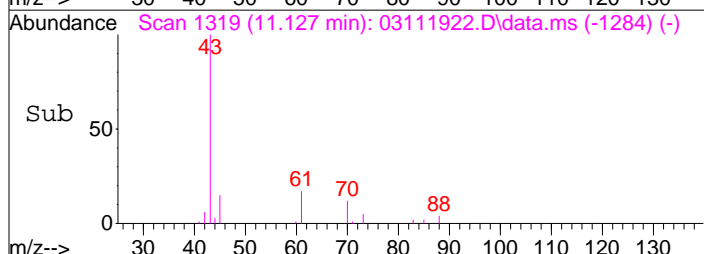
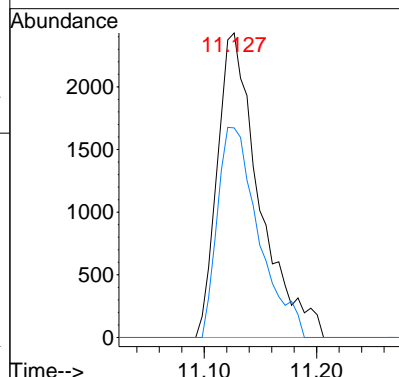
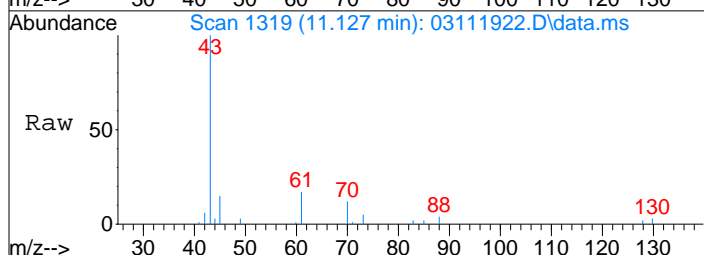
#28
 cis-1,2-Dichloroethene
 Concen: 0.55 ng
 RT: 10.80 min Scan# 1261
 Delta R.T. -0.017 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

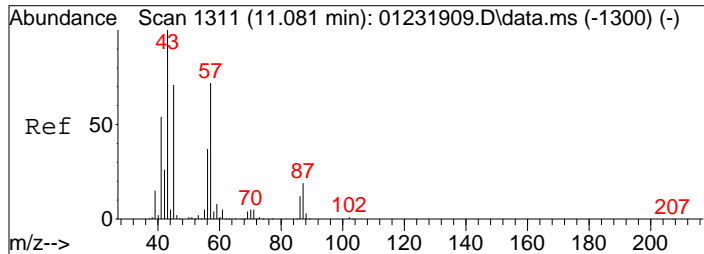
Tgt Ion	Resp	Lower	Upper
61	100		
96	85.7	63.3	103.3



#30
 Ethyl Acetate
 Concen: 2.37 ng
 RT: 11.13 min Scan# 1319
 Delta R.T. -0.000 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

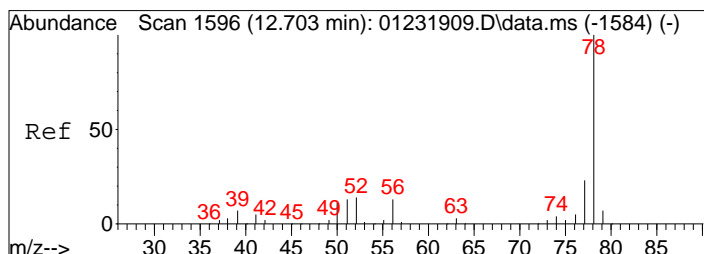
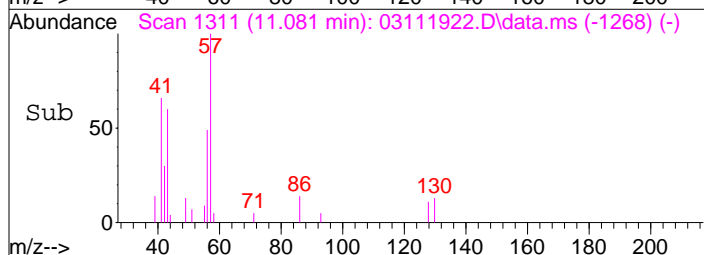
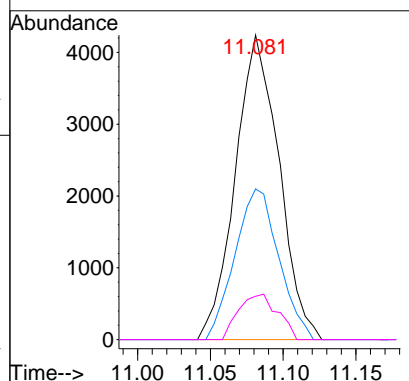
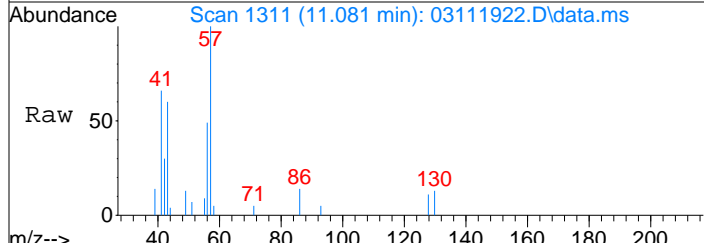
Tgt Ion	Resp	Lower	Upper
61	100		
70	67.6	57.7	97.7





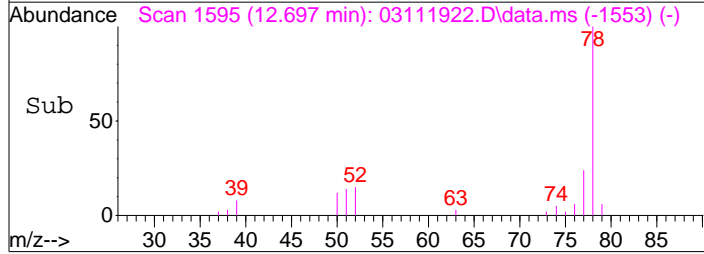
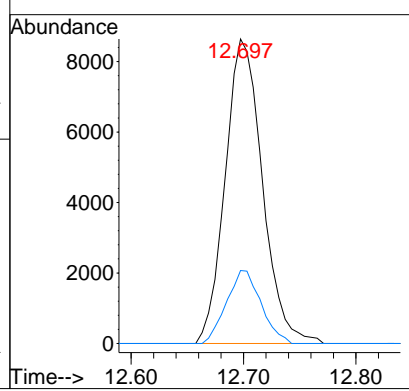
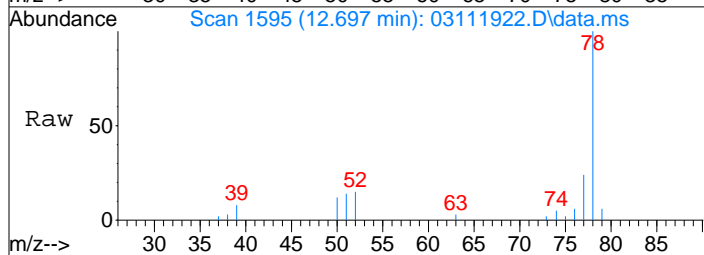
#31
 n-Hexane
 Concen: 0.71 ng
 RT: 11.08 min Scan# 1311
 Delta R.T. -0.006 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

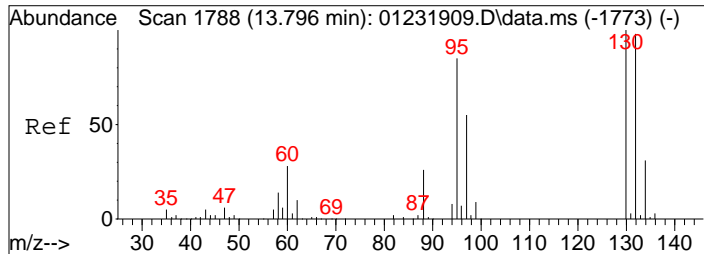
Tgt Ion	Resp	Lower	Upper
57	100		
56	49.7	0.0	0.0#
86	13.4	0.0	0.0#



#41
 Benzene
 Concen: 0.59 ng
 RT: 12.70 min Scan# 1595
 Delta R.T. -0.012 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

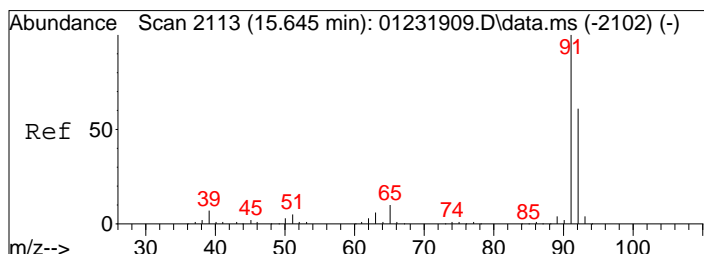
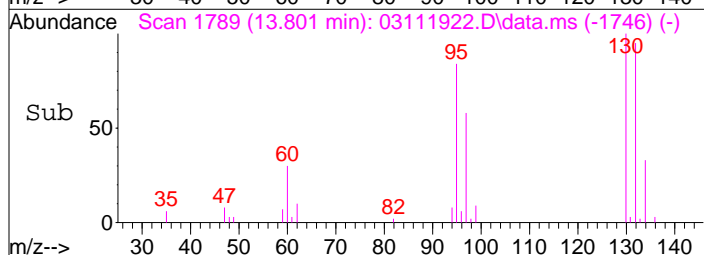
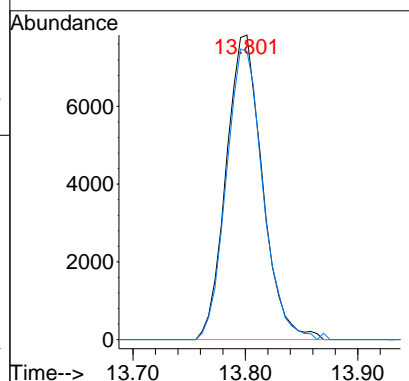
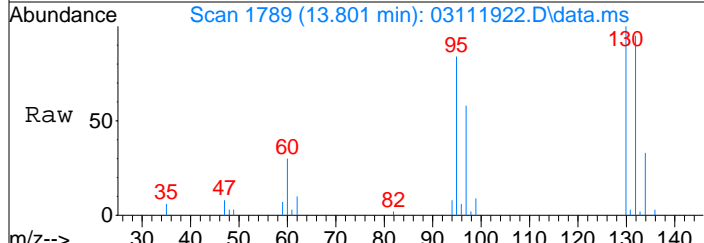
Tgt Ion	Resp	Lower	Upper
78	100		
77	22.3	3.5	43.5





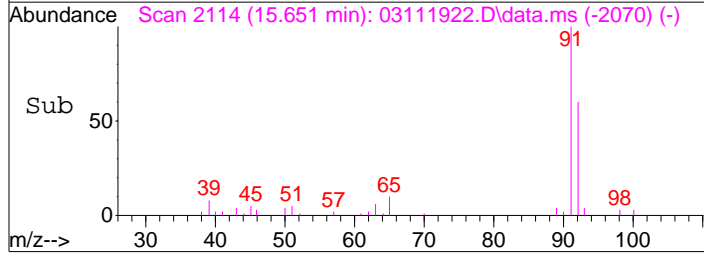
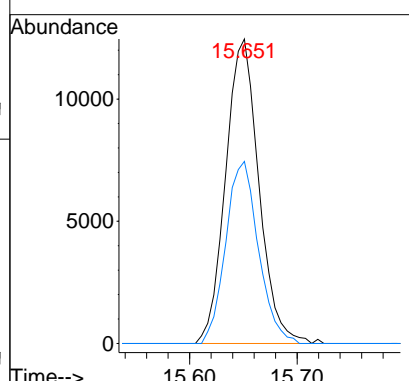
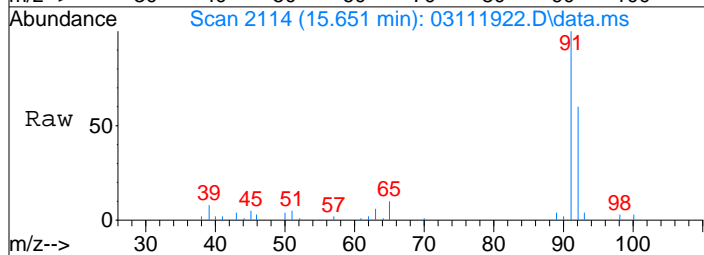
#47
 Trichloroethene
 Concen: 1.81 ng
 RT: 13.80 min Scan# 1789
 Delta R.T. -0.006 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

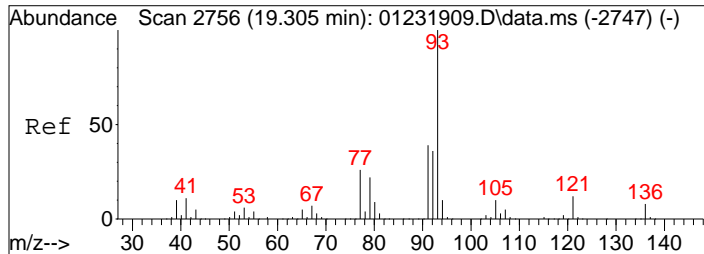
Tgt Ion: 130 Resp: 17610
 Ion Ratio Lower Upper
 130 100
 132 96.3 76.7 116.7



#58
 Toluene
 Concen: 0.66 ng
 RT: 15.65 min Scan# 2114
 Delta R.T. -0.000 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

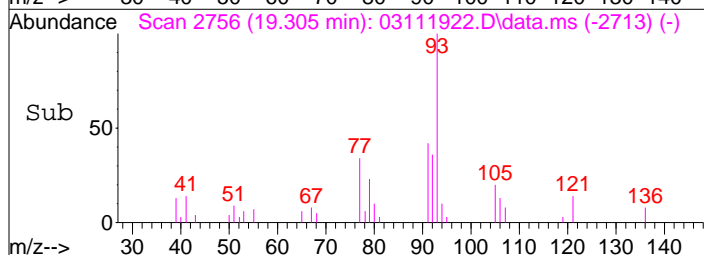
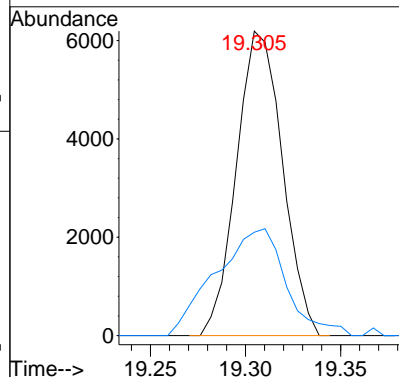
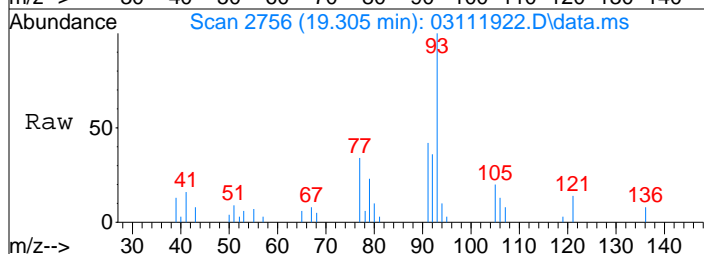
Tgt Ion: 91 Resp: 26877
 Ion Ratio Lower Upper
 91 100
 92 58.6 39.5 79.5





#75
 alpha-Pinene
 Concen: 0.47 ng
 RT: 19.30 min Scan# 2756
 Delta R.T. -0.006 min
 Lab File: 03111922.D
 Acq: 11 Mar 2019 19:15

Tgt Ion	Resp	Lower	Upper
93	10401		
77	53.8	9.6	49.6#



Data File : I:\MS13\DATA\2019_03\11\03111903.D
 Acq On : 11 Mar 2019 7:57
 Sample : MB R13031119_1000mL
 Misc : S31-02211904_AC00880

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 08:20:41 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

 3/11/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.97	130	90918	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.10	114	407856	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	175696	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	108894	14.047	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	112.40%	
57) Toluene-d8 (SS2)	15.54	98	452700	12.032	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	96.24%	
73) Bromofluorobenzene (SS3)	18.83	174	166974	12.326	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	98.64%	

Target Compounds

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Propene	0.00	42	0	N.D.		
3) Dichlorodifluoromethan...	0.00	85	0	N.D.		
4) Chloromethane	0.00	50	0	N.D.		
5) 1,2-Dichloro-1,1,2,2-t...	0.00	135	0	N.D.		
6) Vinyl Chloride	0.00	62	0	N.D.		
7) 1,3-Butadiene	0.00	54	0	N.D.		
8) Bromomethane	0.00	94	0	N.D.		
9) Chloroethane	0.00	64	0	N.D.		
10) Ethanol	0.00	45	0	N.D.		
11) Acetonitrile	0.00	41	0	N.D.		
12) Acrolein	0.00	56	0	N.D.		
13) Acetone	6.93	58	4163	0.727	ng	# 58
14) Trichlorofluoromethane	0.00	101	0	N.D.		
15) 2-Propanol (Isopropanol)	0.00	45	0	N.D.		
16) Acrylonitrile	0.00	53	0	N.D.		
17) 1,1-Dichloroethene	0.00	96	0	N.D.		
18) 2-Methyl-2-Propanol (t...	0.00	59	0	N.D.		
19) Methylene Chloride	0.00	84	0	N.D.		
20) 3-Chloro-1-propene (Al...	0.00	41	0	N.D.		
21) Trichlorotrifluoroethane	0.00	151	0	N.D.		
22) Carbon Disulfide	0.00	76	0	N.D.	d	
23) trans-1,2-Dichloroethene	0.00	61	0	N.D.		
24) 1,1-Dichloroethane	0.00	63	0	N.D.		
25) Methyl tert-Butyl Ether	0.00	73	0	N.D.		
26) Vinyl Acetate	0.00	86	0	N.D.		
27) 2-Butanone (MEK)	10.18	72	367	0.070	ng	# 87
28) cis-1,2-Dichloroethene	0.00	61	0	N.D.		
29) Diisopropyl Ether	0.00	87	0	N.D.		
30) Ethyl Acetate	0.00	61	0	N.D.		
31) n-Hexane	0.00	57	0	N.D.		
32) Chloroform	0.00	83	0	N.D.		
34) Tetrahydrofuran (THF)	0.00	72	0	N.D.		
35) Ethyl tert-Butyl Ether	0.00	87	0	N.D.		
36) 1,2-Dichloroethane	0.00	62	0	N.D.		
38) 1,1,1-Trichloroethane	0.00	97	0	N.D.		
39) Isopropyl Acetate	0.00	61	0	N.D.		
40) 1-Butanol	0.00	56	0	N.D.		
41) Benzene	0.00	78	0	N.D.	d	
42) Carbon Tetrachloride	0.00	117	0	N.D.		
43) Cyclohexane	0.00	84	0	N.D.		
44) tert-Amyl Methyl Ether	0.00	73	0	N.D.		
45) 1,2-Dichloropropane	0.00	63	0	N.D.		
46) Bromodichloromethane	0.00	83	0	N.D.		
47) Trichloroethene	0.00	130	0	N.D.		
48) 1,4-Dioxane	0.00	88	0	N.D.		
49) 2,2,4-Trimethylpentane...	0.00	57	0	N.D.		
50) Methyl Methacrylate	0.00	100	0	N.D.		

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Data File : I:\MS13\DATA\2019_03\11\03111903.D
 Acq On : 11 Mar 2019 7:57
 Sample : MB R13031119_1000mL
 Misc : S31-02211904_AC00880

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 08:20:41 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

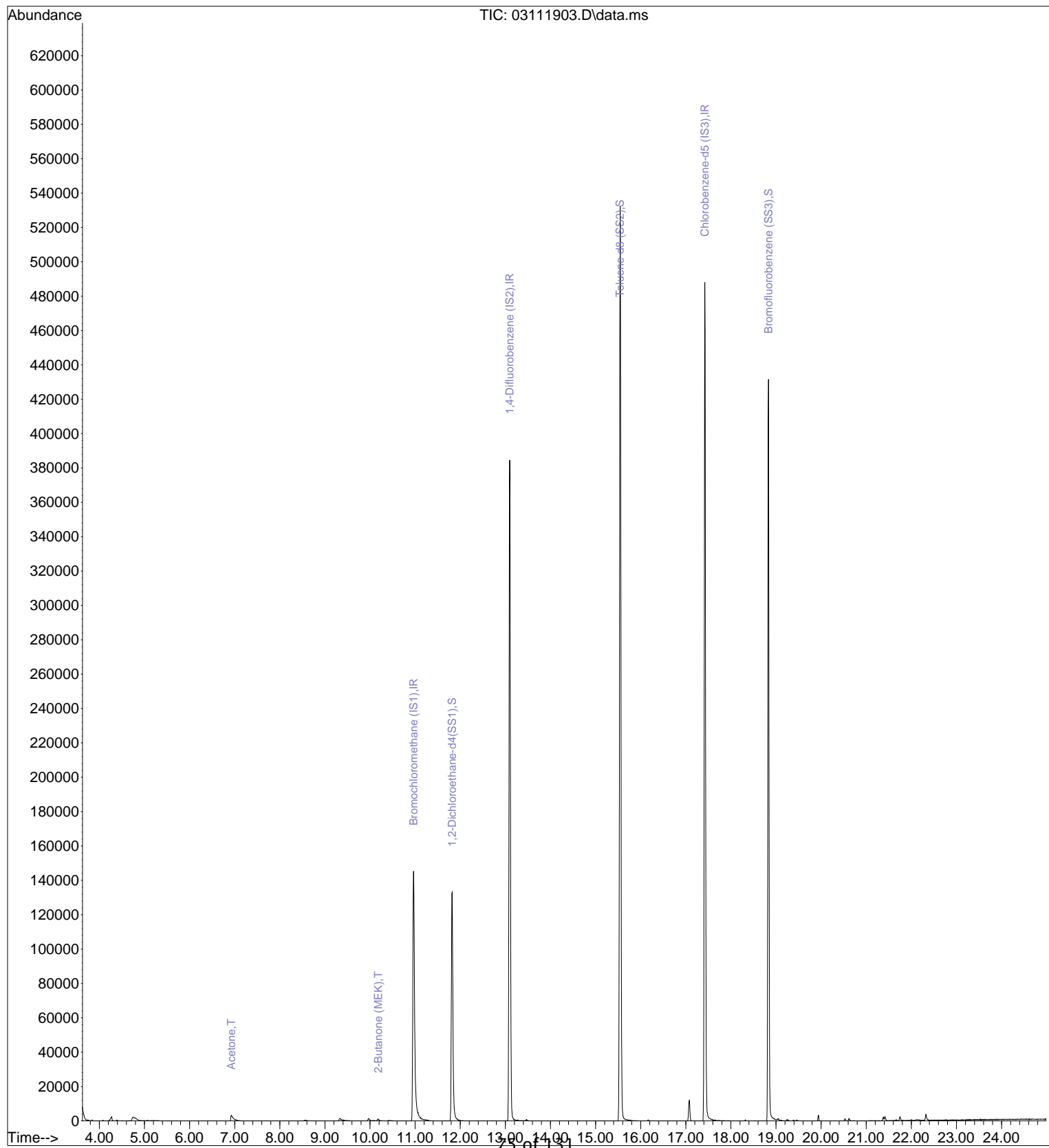
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	0.00	71	0	N.D.		
52) cis-1,3-Dichloropropene	0.00	75	0	N.D.		
53) 4-Methyl-2-pentanone	0.00	58	0	N.D.		
54) trans-1,3-Dichloropropene	0.00	75	0	N.D.		
55) 1,1,2-Trichloroethane	0.00	97	0	N.D.		
58) Toluene	0.00	91	0	N.D.		
59) 2-Hexanone	0.00	43	0	N.D.		
60) Dibromochloromethane	0.00	129	0	N.D.		
61) 1,2-Dibromoethane	0.00	107	0	N.D.		
62) n-Butyl Acetate	0.00	43	0	N.D.		
63) n-Octane	0.00	57	0	N.D.		
64) Tetrachloroethene	0.00	166	0	N.D.		
65) Chlorobenzene	0.00	112	0	N.D.		
66) Ethylbenzene	0.00	91	0	N.D.		
67) m- & p-Xylenes	0.00	91	0	N.D.		
68) Bromoform	0.00	173	0	N.D.		
69) Styrene	0.00	104	0	N.D.		
70) o-Xylene	0.00	91	0	N.D.		
71) n-Nonane	0.00	43	0	N.D.		
72) 1,1,2,2-Tetrachloroethane	0.00	83	0	N.D.		
74) Cumene	0.00	105	0	N.D.	d	
75) alpha-Pinene	0.00	93	0	N.D.		
76) n-Propylbenzene	0.00	91	0	N.D.	d	
77) 3-Ethyltoluene	0.00	105	0	N.D.		
78) 4-Ethyltoluene	0.00	105	0	N.D.		
79) 1,3,5-Trimethylbenzene	0.00	105	0	N.D.		
80) alpha-Methylstyrene	0.00	118	0	N.D.		
81) 2-Ethyltoluene	0.00	105	0	N.D.		
82) 1,2,4-Trimethylbenzene	0.00	105	0	N.D.		
83) n-Decane	0.00	57	0	N.D.		
84) Benzyl Chloride	0.00	91	0	N.D.		
85) 1,3-Dichlorobenzene	0.00	146	0	N.D.		
86) 1,4-Dichlorobenzene	0.00	146	0	N.D.		
87) sec-Butylbenzene	0.00	105	0	N.D.		
88) 4-Isopropyltoluene (p-...	0.00	119	0	N.D.		
89) 1,2,3-Trimethylbenzene	0.00	105	0	N.D.		
90) 1,2-Dichlorobenzene	0.00	146	0	N.D.		
91) d-Limonene	0.00	68	0	N.D.		
92) 1,2-Dibromo-3-Chloropr...	0.00	157	0	N.D.		
93) n-Undecane	0.00	57	0	N.D.		
94) 1,2,4-Trichlorobenzene	0.00	180	0	N.D.	d	
95) Naphthalene	0.00	128	0	N.D.	d	
96) n-Dodecane	0.00	57	0	N.D.		
97) Hexachlorobutadiene	0.00	225	0	N.D.		
98) Cyclohexanone	0.00	55	0	N.D.		
99) tert-Butylbenzene	0.00	119	0	N.D.		
100) n-Butylbenzene	0.00	91	0	N.D.		

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111903.D
 Acq On : 11 Mar 2019 7:57
 Sample : MB R13031119_1000mL
 Misc : S31-02211904_AC00880

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 08:20:41 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M



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Data File : I:\MS13\DATA\2019_03\11\03111904.D
 Acq On : 11 Mar 2019 8:31
 Sample : LCS R13031119_25ng
 Misc : S31-02211904/S31-03051901

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 10:35:42 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

WA 3/11/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.98	130	100369	12.500	ng	0.00
37) 1,4-Difluorobenzene (IS2)	13.11	114	428145	12.500	ng	0.00
56) Chlorobenzene-d5 (IS3)	17.43	82	184579	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.83	65	115998	13.555	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	108.40%
57) Toluene-d8 (SS2)	15.55	98	466873	11.812	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	94.48%
73) Bromofluorobenzene (SS3)	18.83	174	176829	12.425	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	99.44%

Target Compounds

						Qvalue
2) Propene	3.98	42	234695	26.874	ng	95
3) Dichlorodifluoromethan...	4.13	85	392578	26.312	ng	100
4) Chloromethane	4.41	50	339019	29.268	ng	100
5) 1,2-Dichloro-1,1,2,2-t...	4.67	135	247818	25.772	ng	100
6) Vinyl Chloride	4.82	62	327551	29.496	ng	100
7) 1,3-Butadiene	5.08	54	253702	30.370	ng	95
8) Bromomethane	5.51	94	216300	27.146	ng	99
9) Chloroethane	5.83	64	166872	27.303	ng	100
10) Ethanol	6.23	45	802859	134.853	ng	98
11) Acetonitrile	6.47	41	408383	28.348	ng	97
12) Acrolein	6.64	56	143986	29.261	ng	97
13) Acetone	6.85	58	817566	129.244	ng	86
14) Trichlorofluoromethane	7.07	101	329048	26.318	ng	99
15) 2-Propanol (Isopropanol)	7.35	45	1080771	56.103	ng	94
16) Acrylonitrile	7.60	53	298473	32.363	ng	98
17) 1,1-Dichloroethene	8.03	96	226949	27.196	ng	91
18) 2-Methyl-2-Propanol (t...	8.21	59	1068317	56.418	ng	96
19) Methylene Chloride	8.26	84	229024	28.559	ng	99
20) 3-Chloro-1-propene (Al...	8.41	41	329515	30.661	ng	95
21) Trichlorotrifluoroethane	8.67	151	224430	25.439	ng	97
22) Carbon Disulfide	8.51	76	831106	25.372	ng	99
23) trans-1,2-Dichloroethene	9.53	61	295699	30.281	ng	97
24) 1,1-Dichloroethane	9.78	63	377827	27.198	ng	100
25) Methyl tert-Butyl Ether	9.88	73	643597	27.122	ng	97
26) Vinyl Acetate	10.05	86	290762	142.887	ng	# 91
27) 2-Butanone (MEK)	10.29	72	158062	27.258	ng	100
28) cis-1,2-Dichloroethene	10.80	61	284545	28.175	ng	97
29) Diisopropyl Ether	11.10	87	223687	27.480	ng	# 69
30) Ethyl Acetate	11.11	61	166282	57.903	ng	100
31) n-Hexane	11.09	57	348093	26.115	ng	# 100
32) Chloroform	11.15	83	355762	27.244	ng	99
34) Tetrahydrofuran (THF)	11.55	72	154139	26.192	ng	94
35) Ethyl tert-Butyl Ether	11.70	87	266924	26.407	ng	98
36) 1,2-Dichloroethane	11.95	62	238941	28.825	ng	100
38) 1,1,1-Trichloroethane	12.23	97	303572	26.857	ng	98
39) Isopropyl Acetate	12.66	61	285422	52.564	ng	# 87
40) 1-Butanol	12.69	56	438762	59.338	ng	# 70
41) Benzene	12.70	78	891348	25.211	ng	99
42) Carbon Tetrachloride	12.86	117	283947	27.300	ng	99
43) Cyclohexane	12.99	84	703825	50.663	ng	96
44) tert-Amyl Methyl Ether	13.35	73	623967	27.040	ng	97
45) 1,2-Dichloropropane	13.56	63	219857	27.189	ng	100
46) Bromodichloromethane	13.75	83	280525	28.661	ng	99
47) Trichloroethene	13.80	130	265311	25.962	ng	99
48) 1,4-Dioxane	13.78	88	192767	27.061	ng	97
49) 2,2,4-Trimethylpentane...	13.87	57	898702	25.842	ng	91
50) Methyl Methacrylate	14.01	100	197910	57.896	ng	96

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Data File : I:\MS13\DATA\2019_03\11\03111904.D
 Acq On : 11 Mar 2019 8:31
 Sample : LCS R13031119_25ng
 Misc : S31-02211904/S31-03051901

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 10:35:42 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

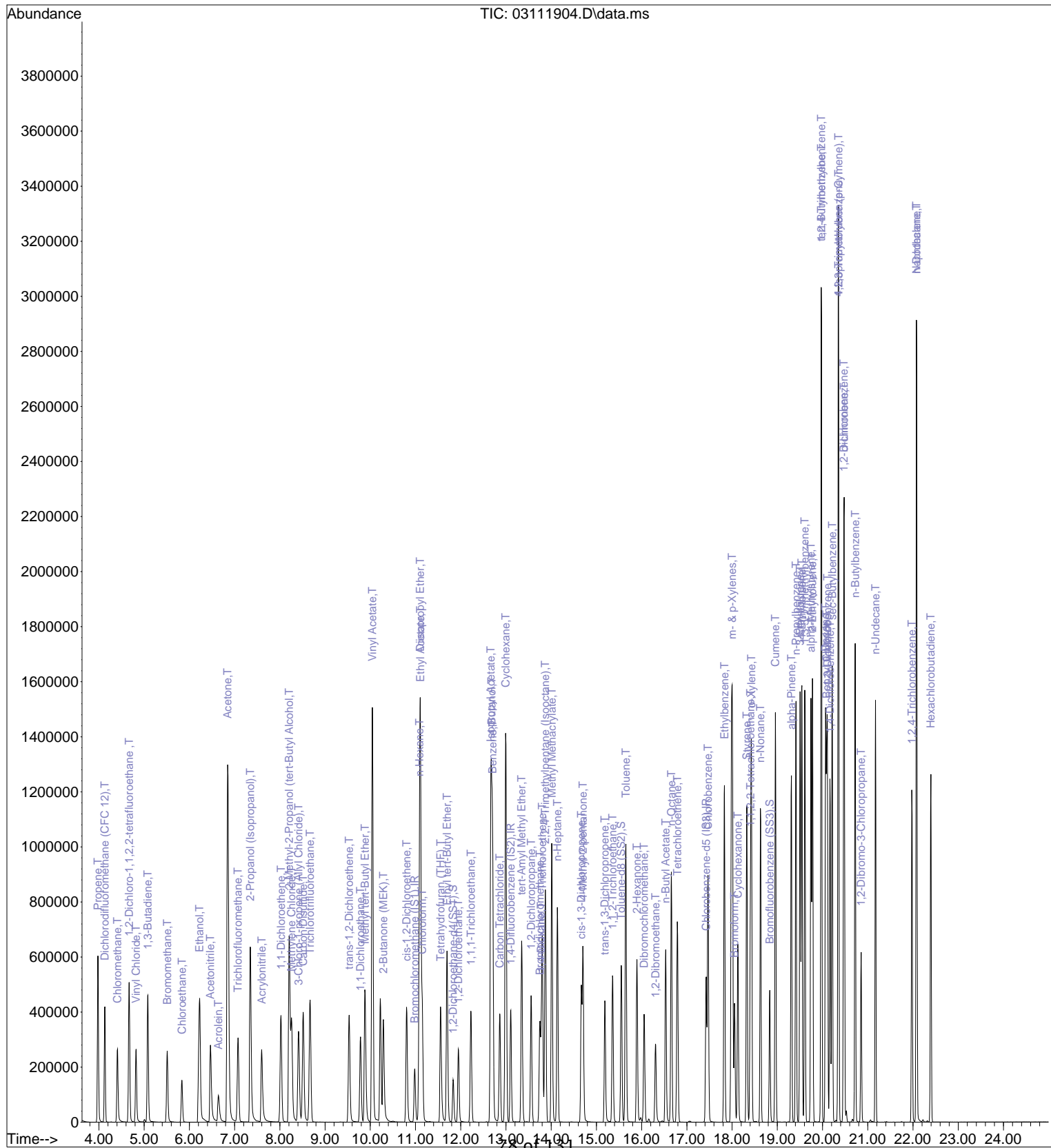
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.14	71	226505	26.580	ng	98
52) cis-1,3-Dichloropropene	14.67	75	359699	28.064	ng	100
53) 4-Methyl-2-pentanone	14.70	58	208089	28.563	ng	96
54) trans-1,3-Dichloropropene	15.18	75	314165	29.176	ng	100
55) 1,1,2-Trichloroethane	15.36	97	223513	27.203	ng	100
58) Toluene	15.65	91	915747	21.796	ng	99
59) 2-Hexanone	15.90	43	464734	28.149	ng	97
60) Dibromochloromethane	16.06	129	273371	26.707	ng	99
61) 1,2-Dibromoethane	16.31	107	254598	26.766	ng	100
62) n-Butyl Acetate	16.53	43	522898	27.292	ng	96
63) n-Octane	16.66	57	187215	24.250	ng	94
64) Tetrachloroethene	16.79	166	292558	22.752	ng	99
65) Chlorobenzene	17.47	112	640926	22.999	ng	100
66) Ethylbenzene	17.83	91	1025978	22.688	ng	97
67) m- & p-Xylenes	18.00	91	1607738	46.056	ng	100
68) Bromoform	18.05	173	252507	27.005	ng	99
69) Styrene	18.33	104	679796	24.912	ng	97
70) o-Xylene	18.43	91	800404	23.006	ng	98
71) n-Nonane	18.63	43	419987	24.697	ng	97
72) 1,1,2,2-Tetrachloroethane	18.41	83	395928	24.287	ng	100
74) Cumene	18.96	105	1067103	22.478	ng	100
75) alpha-Pinene	19.31	93	553597	24.360	ng	95
76) n-Propylbenzene	19.41	91	1252898	23.530	ng	99
77) 3-Ethyltoluene	19.50	105	1038302	23.011	ng	100
78) 4-Ethyltoluene	19.54	105	1063624	23.590	ng	100
79) 1,3,5-Trimethylbenzene	19.61	105	866239	21.889	ng	97
80) alpha-Methylstyrene	19.74	118	511886	26.204	ng	91
81) 2-Ethyltoluene	19.78	105	1054886	22.465	ng	99
82) 1,2,4-Trimethylbenzene	19.97	105	888629	23.147	ng	97
83) n-Decane	20.07	57	464807	24.880	ng	99
84) Benzyl Chloride	20.09	91	769286	27.400	ng	99
85) 1,3-Dichlorobenzene	20.11	146	569627	23.645	ng	100
86) 1,4-Dichlorobenzene	20.16	146	577843	22.849	ng	100
87) sec-Butylbenzene	20.22	105	1201166	22.538	ng	99
88) 4-Isopropyltoluene (p-...	20.35	119	1152722	23.172	ng	98
89) 1,2,3-Trimethylbenzene	20.35	105	915013	24.093	ng	97
90) 1,2-Dichlorobenzene	20.47	146	554361	23.576	ng	100
91) d-Limonene	20.48	68	345822	25.808	ng	94
92) 1,2-Dibromo-3-Chloropr...	20.85	157	207368	24.394	ng	90
93) n-Undecane	21.17	57	482071	27.983	ng	97
94) 1,2,4-Trichlorobenzene	21.97	180	436960	24.409	ng	100
95) Naphthalene	22.08	128	1268288	25.971	ng	100
96) n-Dodecane	22.08	57	461035	28.615	ng	98
97) Hexachlorobutadiene	22.39	225	269812	20.273	ng	100
98) Cyclohexanone	18.13	55	303795	23.876	ng	96
99) tert-Butylbenzene	19.97	119	895814	22.888	ng	100
100) n-Butylbenzene	20.72	91	932245	23.873	ng	99

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111904.D
Acq On : 11 Mar 2019 8:31
Sample : LCS R13031119_25ng
Misc : S31-02211904/S31-03051901

Vial: 2
Operator: WA
Inst : MS13

Quant Time: Mar 11 10:35:42 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 11:21:29 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_03\11\03111916.D
 Acq On : 11 Mar 2019 15:53
 Sample : P1901179-002dup (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:10 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

DA 3/14/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.97	130	88966	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.10	114	384241	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	167241	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	107717	14.201	ng	-0.02
Spiked Amount	12.500	Range	70 - 130	Recovery	=	113.60%
57) Toluene-d8 (SS2)	15.55	98	421843	11.779	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	94.24%
73) Bromofluorobenzene (SS3)	18.83	174	157571	12.220	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	97.76%

Target Compounds

						Qvalue
2) Propene	0.00	42	0	N.D.		
3) Dichlorodifluoromethan...	0.00	85	0	N.D.		
4) Chloromethane	0.00	50	0	N.D.		
5) 1,2-Dichloro-1,1,2,2-t...	0.00	135	0	N.D.		
6) Vinyl Chloride	4.87	62	4636	0.471	ng	85
7) 1,3-Butadiene	0.00	54	0	N.D.		
8) Bromomethane	0.00	94	0	N.D.		
9) Chloroethane	0.00	64	0	N.D.		
10) Ethanol	0.00	45	0	N.D.		
11) Acetonitrile	0.00	41	0	N.D.		
12) Acrolein	0.00	56	0	N.D.		
13) Acetone	6.95	58	2085	N.D.		
14) Trichlorofluoromethane	0.00	101	0	N.D.		
15) 2-Propanol (Isopropanol)	0.00	45	0	N.D.		
16) Acrylonitrile	0.00	53	0	N.D.		
17) 1,1-Dichloroethene	8.03	96	3747	0.507	ng	94
18) 2-Methyl-2-Propanol (t...	0.00	59	0	N.D.		
19) Methylene Chloride	8.24	84	33648	4.734	ng	98
20) 3-Chloro-1-propene (Al...	8.24	41	973	N.D.		
21) Trichlorotrifluoroethane	8.66	151	116096	14.846	ng	98
22) Carbon Disulfide	8.55	76	1652	N.D.		
23) trans-1,2-Dichloroethene	0.00	61	0	N.D.		
24) 1,1-Dichloroethane	9.77	63	1414	N.D.		
25) Methyl tert-Butyl Ether	0.00	73	0	N.D.		
26) Vinyl Acetate	0.00	86	0	N.D.		
27) 2-Butanone (MEK)	0.00	72	0	N.D.		
28) cis-1,2-Dichloroethene	10.79	61	202748	22.649	ng	98
29) Diisopropyl Ether	0.00	87	0	N.D.		
30) Ethyl Acetate	11.09	61	285	N.D.		
31) n-Hexane	0.00	57	0	N.D.		
32) Chloroform	11.13	83	2083	N.D.		
34) Tetrahydrofuran (THF)	0.00	72	0	N.D.		
35) Ethyl tert-Butyl Ether	0.00	87	0	N.D.		
36) 1,2-Dichloroethane	11.95	62	8186	1.114	ng	97
38) 1,1,1-Trichloroethane	0.00	97	0	N.D.		
39) Isopropyl Acetate	0.00	61	0	N.D.		
40) 1-Butanol	0.00	56	0	N.D.		
41) Benzene	12.71	78	958	N.D.		
42) Carbon Tetrachloride	12.85	117	588	N.D.		
43) Cyclohexane	0.00	84	0	N.D.		
44) tert-Amyl Methyl Ether	0.00	73	0	N.D.		
45) 1,2-Dichloropropane	0.00	63	0	N.D.		
46) Bromodichloromethane	0.00	83	0	N.D.	d	
47) Trichloroethene	13.80	130	854020	93.118	ng	100
48) 1,4-Dioxane	0.00	88	0	N.D.		
49) 2,2,4-Trimethylpentane...	0.00	57	0	N.D.		
50) Methyl Methacrylate	0.00	100	0	N.D.	d	

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Data File : I:\MS13\DATA\2019_03\11\03111916.D
 Acq On : 11 Mar 2019 15:53
 Sample : P1901179-002dup (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:10 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

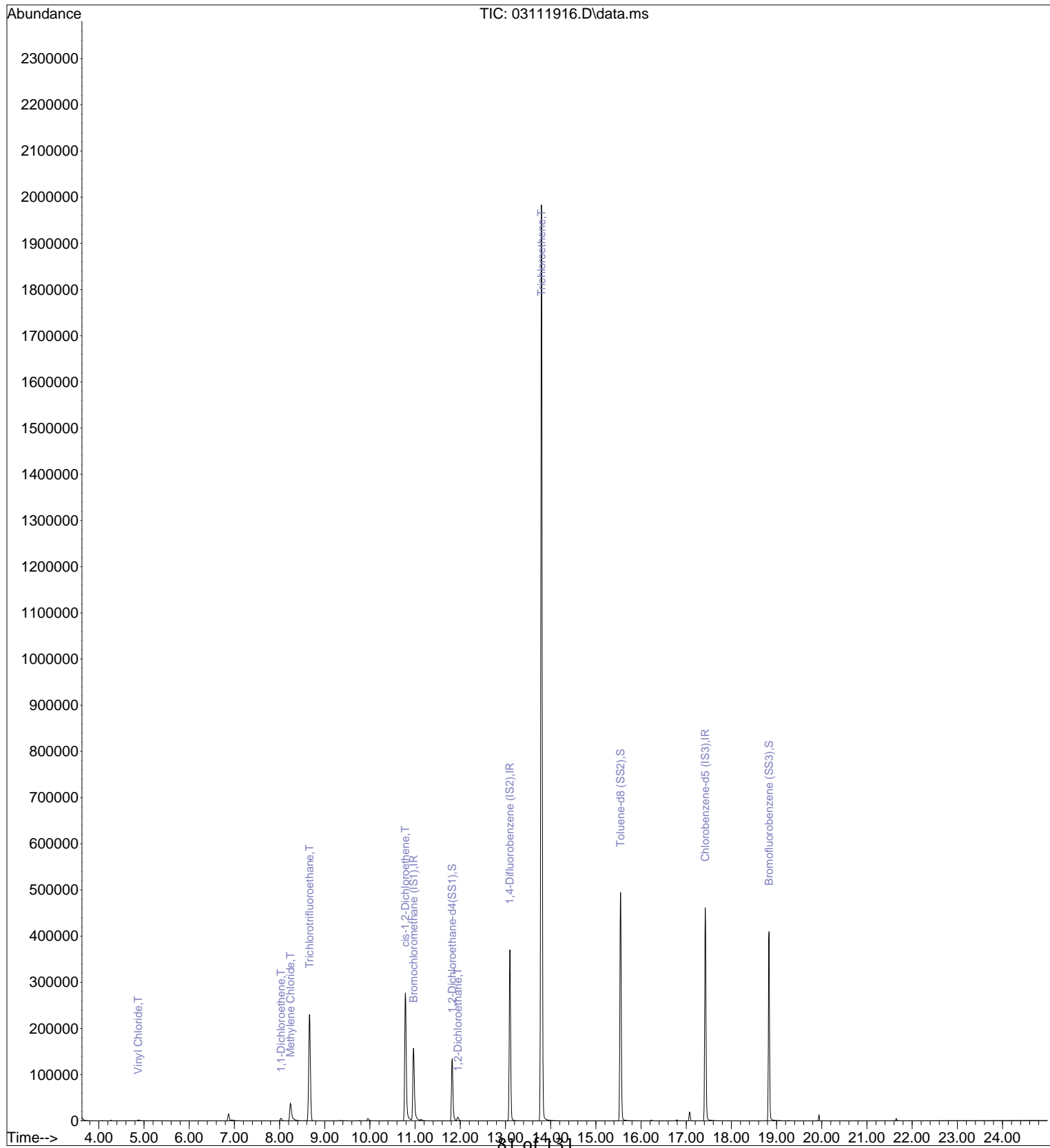
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	0.00	71	0	N.D.		
52) cis-1,3-Dichloropropene	0.00	75	0	N.D.		
53) 4-Methyl-2-pentanone	0.00	58	0	N.D.		
54) trans-1,3-Dichloropropene	0.00	75	0	N.D.		
55) 1,1,2-Trichloroethane	0.00	97	0	N.D.		
58) Toluene	15.66	91	518	N.D.		
59) 2-Hexanone	0.00	43	0	N.D.		
60) Dibromochloromethane	0.00	129	0	N.D.		
61) 1,2-Dibromoethane	0.00	107	0	N.D.		
62) n-Butyl Acetate	0.00	43	0	N.D.		
63) n-Octane	0.00	57	0	N.D.		
64) Tetrachloroethene	16.79	166	1013	N.D.		
65) Chlorobenzene	0.00	112	0	N.D.		
66) Ethylbenzene	0.00	91	0	N.D.		
67) m- & p-Xylenes	0.00	91	0	N.D.		
68) Bromoform	0.00	173	0	N.D.		
69) Styrene	0.00	104	0	N.D.		
70) o-Xylene	0.00	91	0	N.D.		
71) n-Nonane	0.00	43	0	N.D.		
72) 1,1,2,2-Tetrachloroethane	0.00	83	0	N.D.		
74) Cumene	0.00	105	0	N.D.		
75) alpha-Pinene	0.00	93	0	N.D.		
76) n-Propylbenzene	0.00	91	0	N.D.		
77) 3-Ethyltoluene	0.00	105	0	N.D.		
78) 4-Ethyltoluene	0.00	105	0	N.D.		
79) 1,3,5-Trimethylbenzene	0.00	105	0	N.D.		
80) alpha-Methylstyrene	0.00	118	0	N.D.		
81) 2-Ethyltoluene	0.00	105	0	N.D.		
82) 1,2,4-Trimethylbenzene	0.00	105	0	N.D.		
83) n-Decane	0.00	57	0	N.D.		
84) Benzyl Chloride	0.00	91	0	N.D.		
85) 1,3-Dichlorobenzene	0.00	146	0	N.D.		
86) 1,4-Dichlorobenzene	0.00	146	0	N.D.		
87) sec-Butylbenzene	0.00	105	0	N.D.		
88) 4-Isopropyltoluene (p-...	0.00	119	0	N.D.		
89) 1,2,3-Trimethylbenzene	0.00	105	0	N.D.		
90) 1,2-Dichlorobenzene	0.00	146	0	N.D.		
91) d-Limonene	0.00	68	0	N.D.		
92) 1,2-Dibromo-3-Chloropr...	0.00	157	0	N.D.		
93) n-Undecane	0.00	57	0	N.D.		
94) 1,2,4-Trichlorobenzene	0.00	180	0	N.D.		
95) Naphthalene	0.00	128	0	N.D.		
96) n-Dodecane	0.00	57	0	N.D.		
97) Hexachlorobutadiene	0.00	225	0	N.D.		
98) Cyclohexanone	0.00	55	0	N.D.		
99) tert-Butylbenzene	19.94	119	201	N.D.		
100) n-Butylbenzene	0.00	91	0	N.D.		

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111916.D
 Acq On : 11 Mar 2019 15:53
 Sample : P1901179-002dup (10mL)
 Misc : S31-02211904

Vial: 3
 Operator: WA
 Inst : MS13

Quant Time: Mar 14 11:07:10 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M



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Data File : I:\MS13\DATA\2019_03\11\03111916.D
 Acq On : 11 Mar 2019 15:53
 Sample : P1901179-002dup (10mL)
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 Response via : Initial Calibration
 DataAcq Meth:TO15.M

WA 3/14/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.97	130	88966	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.10	114	384241	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	167241	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	107717	14.201	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery	=	113.60%	
57) Toluene-d8 (SS2)	15.55	98	421843	11.779	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	94.24%	
73) Bromofluorobenzene (SS3)	18.83	174	157571	12.220	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	97.76%	

Target Compounds

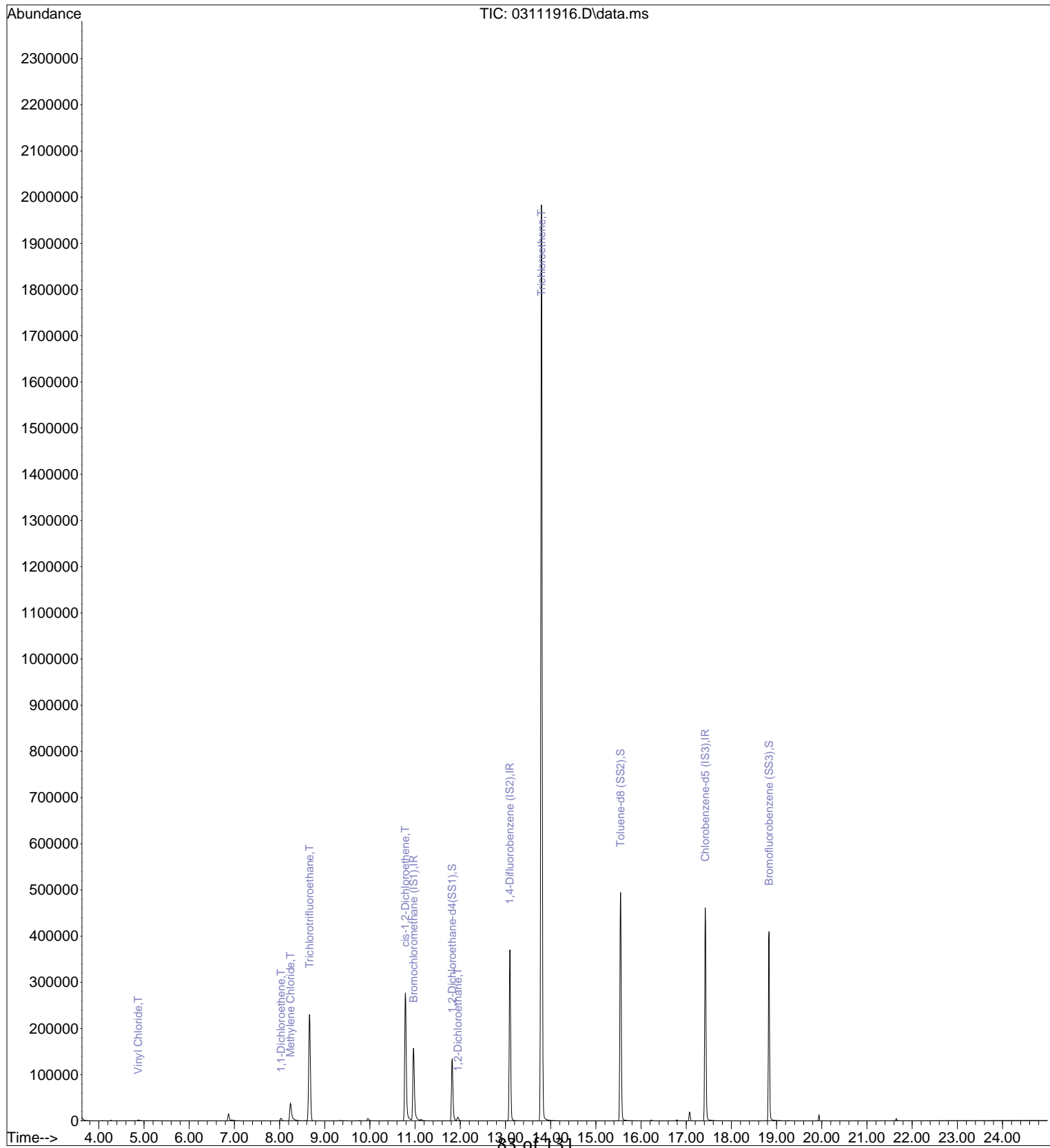
	R.T.	QIon	Response	Conc	Units	Qvalue
6) Vinyl Chloride	4.87	62	4636	0.471	ng	85
17) 1,1-Dichloroethene	8.03	96	3747	0.507	ng	94
19) Methylene Chloride	8.24	84	33648	4.734	ng	98
21) Trichlorotrifluoroethane	8.66	151	116096	14.846	ng	98
28) cis-1,2-Dichloroethene	10.79	61	202748	22.649	ng	98
36) 1,2-Dichloroethane	11.95	62	8186	1.114	ng	97
47) Trichloroethene	13.80	130	854020	93.118	ng	100

(#) = qualifier out of range (m) = manual integration (+) = signals summed

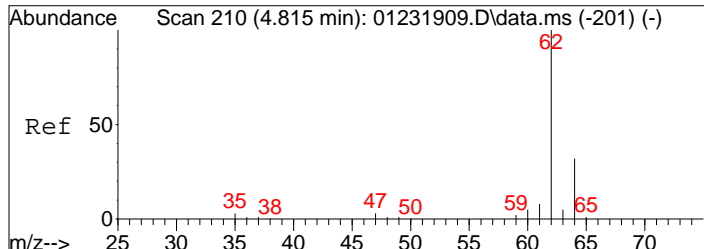
Data File : I:\MS13\DATA\2019_03\11\03111916.D
Acq On : 11 Mar 2019 15:53
Sample : P1901179-002dup (10mL)
Misc : S31-02211904

Vial: 3
Operator: WA
Inst : MS13

Quant Time: Mar 14 11:07:10 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 11:21:29 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M

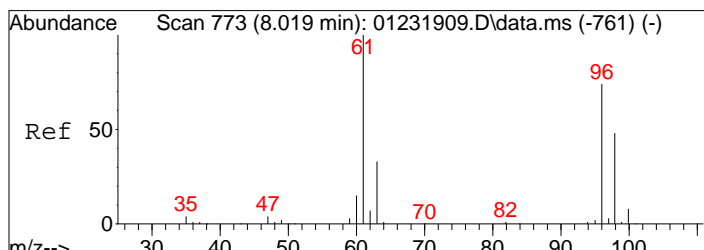
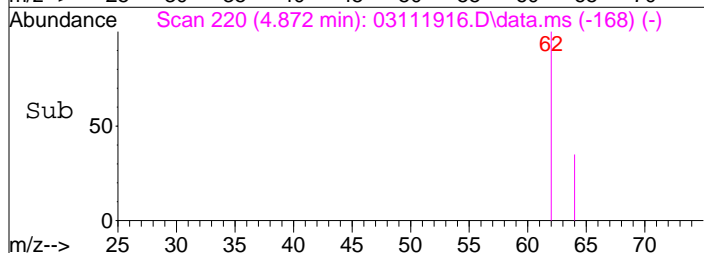
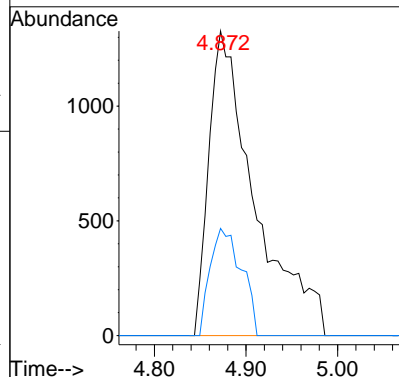
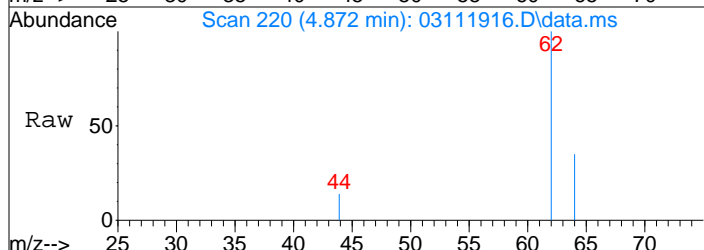


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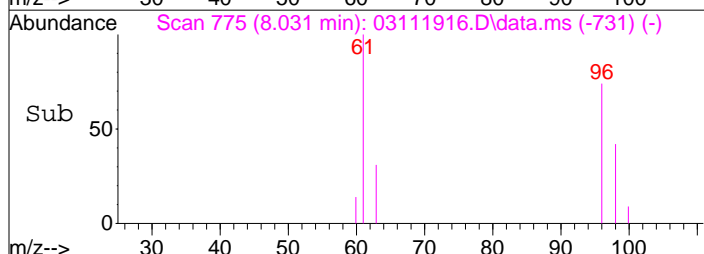
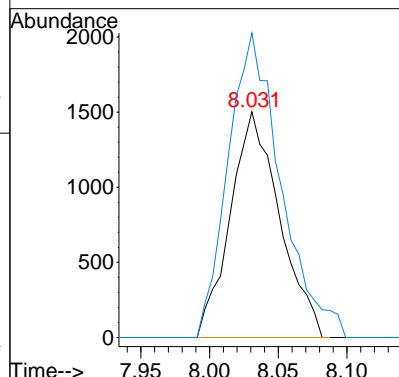
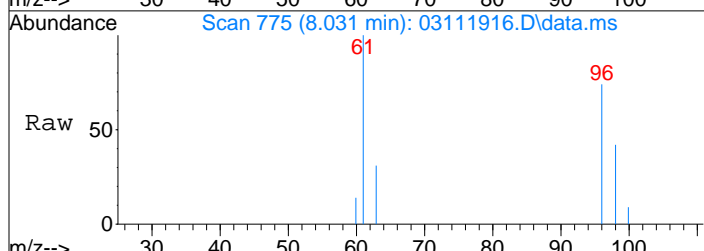
#6
 Vinyl Chloride
 Concen: 0.47 ng
 RT: 4.87 min Scan# 220
 Delta R.T. 0.045 min
 Lab File: 03111916.D
 Acq: 11 Mar 2019 15:53

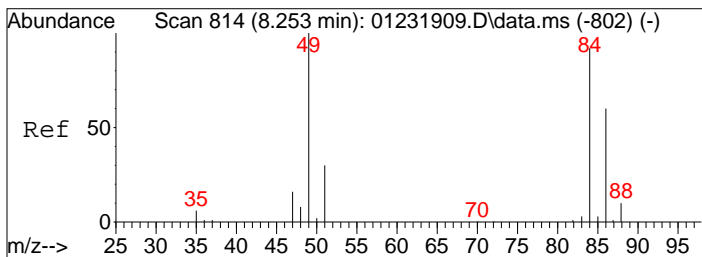
Tgt Ion	Resp	Lower	Upper
62	100		
64	24.0	12.4	52.4



#17
 1,1-Dichloroethene
 Concen: 0.51 ng
 RT: 8.03 min Scan# 775
 Delta R.T. -0.000 min
 Lab File: 03111916.D
 Acq: 11 Mar 2019 15:53

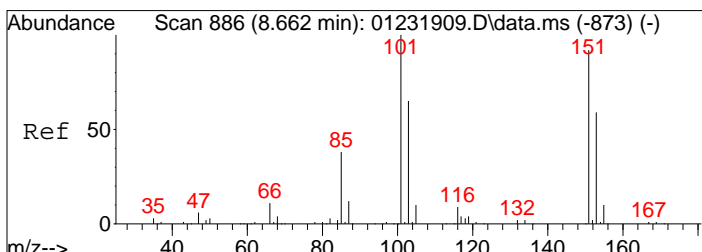
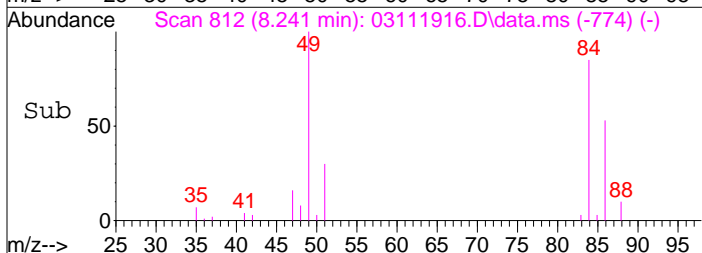
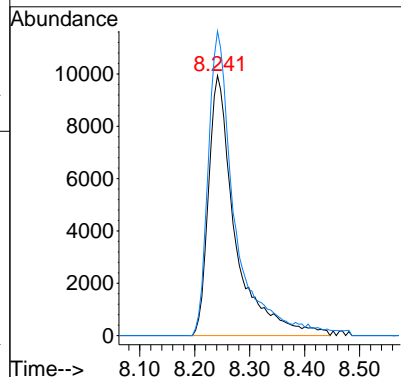
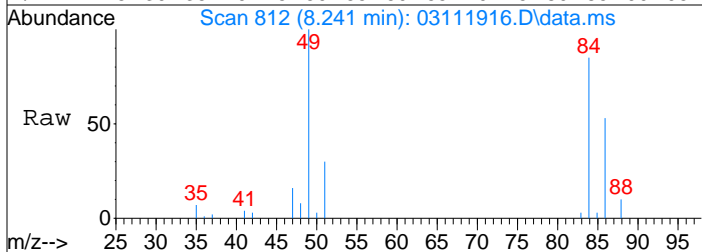
Tgt Ion	Resp	Lower	Upper
96	100		
61	144.7	132.9	172.9





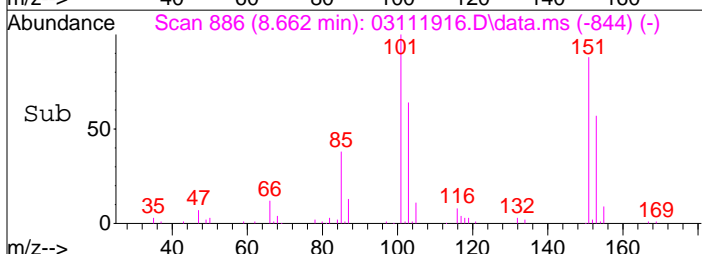
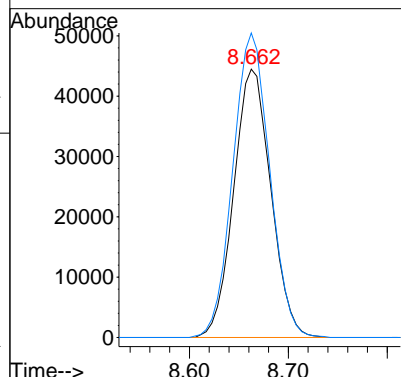
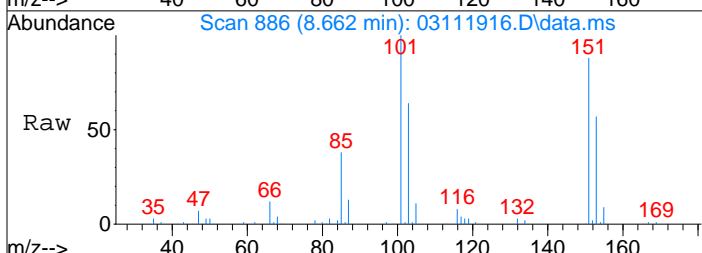
#19
 Methylene Chloride
 Concen: 4.73 ng
 RT: 8.24 min Scan# 812
 Delta R.T. -0.034 min
 Lab File: 03111916.D
 Acq: 11 Mar 2019 15:53

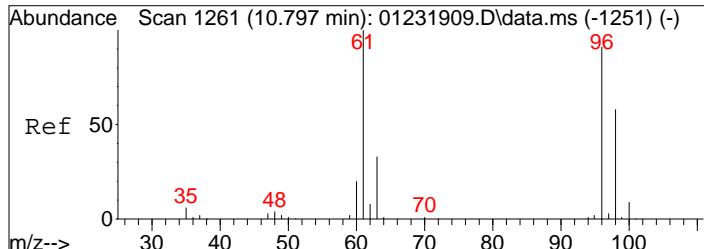
Tgt Ion: 84 Resp: 33648
 Ion Ratio Lower Upper
 84 100
 49 117.8 90.3 140.3



#21
 Trichlorotrifluoroethane
 Concen: 14.85 ng
 RT: 8.66 min Scan# 886
 Delta R.T. -0.011 min
 Lab File: 03111916.D
 Acq: 11 Mar 2019 15:53

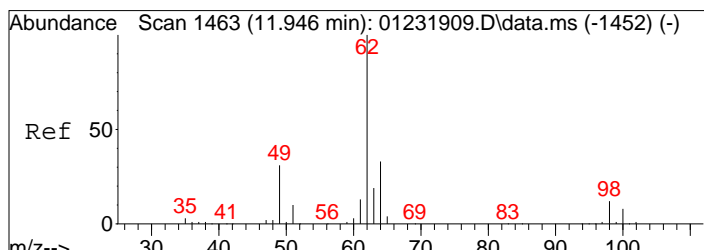
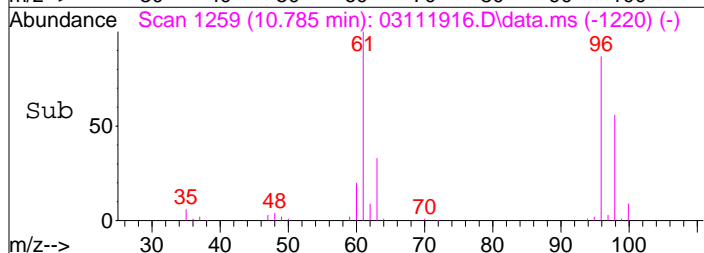
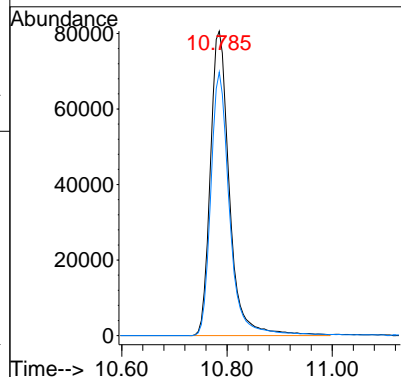
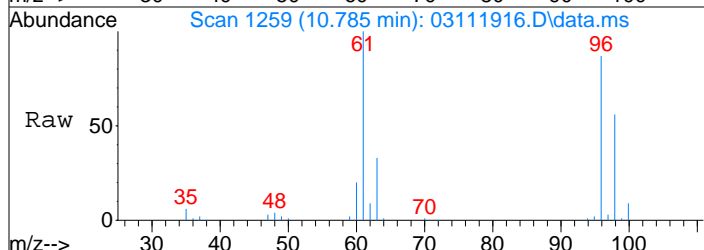
Tgt Ion: 151 Resp: 116096
 Ion Ratio Lower Upper
 151 100
 101 112.6 95.1 135.1





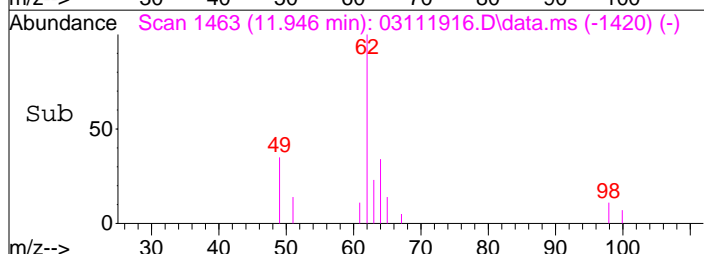
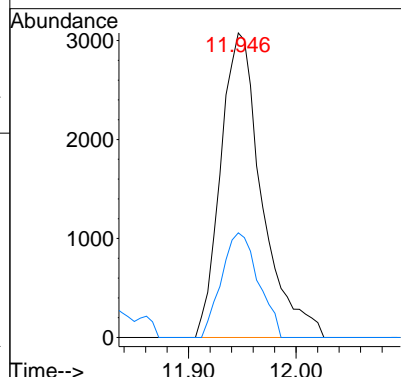
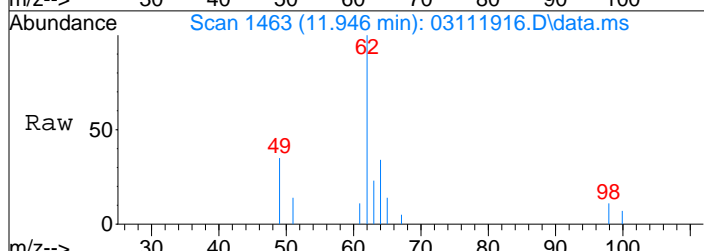
#28
 cis-1,2-Dichloroethene
 Concen: 22.65 ng
 RT: 10.79 min Scan# 1259
 Delta R.T. -0.029 min
 Lab File: 03111916.D
 Acq: 11 Mar 2019 15:53

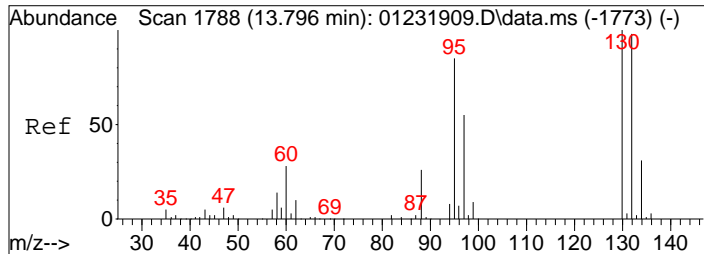
Tgt Ion	Resp	Lower	Upper
61	100		
96	85.3	63.3	103.3



#36
 1,2-Dichloroethane
 Concen: 1.11 ng
 RT: 11.95 min Scan# 1463
 Delta R.T. -0.006 min
 Lab File: 03111916.D
 Acq: 11 Mar 2019 15:53

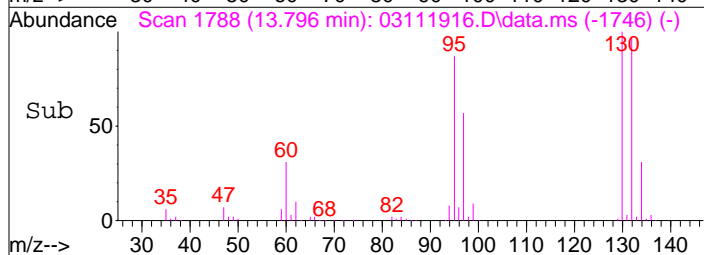
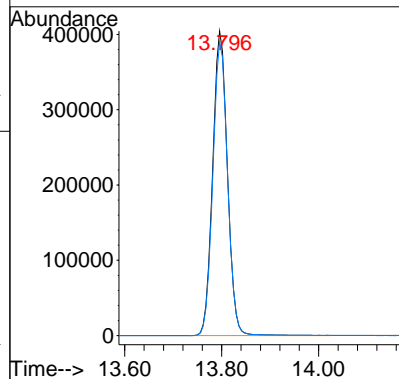
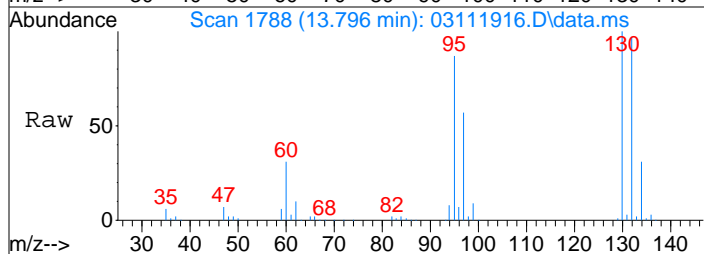
Tgt Ion	Resp	Lower	Upper
62	100		
64	30.8	12.6	52.6





#47
 Trichloroethene
 Concen: 93.12 ng
 RT: 13.80 min Scan# 1788
 Delta R.T. -0.011 min
 Lab File: 03111916.D
 Acq: 11 Mar 2019 15:53

Tgt Ion	Resp	Lower	Upper
130	100		
132	96.6	76.7	116.7



Method Path : I:\MS13\METHODS\
Method File : R13012319.M
Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
Last Update : Wed Jan 23 11:21:29 2019
Response Via : Initial Calibration

Calibration Files

0.1 =01231906.D 0.2 =01231907.D 0.5 =01231904.D 1.0 =01231905.D 5.0 =01231908.D 25 =01231909.D 50 =01231910.D
100 =01231911.D

Compound	0.1	0.2	0.5	1.0	5.0	25	50	100	AVG	%RSD
-----ISTD-----										
1) IR Bromochloromethane...	1.086	1.026	1.144	1.076	1.116	1.075	1.030	1.147	1.088	4.25
2) T Propene	1.809	1.904	1.983	1.928	1.929	1.844	1.767	1.701	1.858	5.11
3) T Dichlorodifluo...	1.367	1.297	1.705	1.481	1.555	1.501	1.434	1.201	1.443	10.85
4) T Chloromethane	1.133	1.207	1.294	1.236	1.216	1.182	1.162	1.150	1.198	4.37
5) T 1,2-Dichloro-1...	1.155	1.181	1.541	1.425	1.493	1.456	1.420	1.394	1.383	10.16
6) T Vinyl Chloride	0.839	0.766	1.099	1.063	1.193	1.139	1.130	1.094	1.040	14.71
7) T 1,3-Butadiene	0.974	0.896	1.022	1.000	1.047	1.027	0.990	0.981	0.992	4.64
8) T Bromomethane	0.661	0.806	0.783	0.799	0.786	0.755	0.739	0.761	0.761	6.58
9) T Chloroethane	0.911	0.684	0.807	0.723	0.751	0.725	0.683	0.649	0.741	11.27
10) T Ethanol	1.720	1.655	1.785	1.785	1.902	1.857	1.846	1.794	1.794	5.19
11) T Acetonitrile	0.478	0.578	0.638	0.677	0.745	0.656	0.654	0.613	0.613	12.08
12) T Acrolein	0.982	0.849	0.832	0.785	0.777	0.745	0.696	0.636	0.788	13.31
13) T Acetone	1.578	1.534	1.687	1.609	1.592	1.510	1.479	1.468	1.557	4.73
14) T Trichlorofluor...	2.391	2.376	2.589	2.458	2.558	2.520	2.371	1.931	2.399	8.63
15) T 2-Propanol (Is...	0.607	0.939	1.196	1.196	1.397	1.378	1.374	1.149	1.149	27.66
16) T Acrylonitrile	0.916	0.965	1.102	1.059	1.091	1.081	1.057	1.042	1.039	6.27
17) T 1,1-Dichloroet...	2.156	2.284	2.524	2.464	2.470	2.451	2.336	2.181	2.358	5.95
18) T 2-Methyl-2-Pro...	0.813	0.834	1.045	1.041	1.089	1.092	1.059	1.016	0.999	11.11
19) T Methylene Chlo...	1.150	1.232	1.363	1.307	1.435	1.433	1.410	1.377	1.338	7.65
20) T 3-Chloro-1-pro...	1.080	1.074	1.186	1.111	1.119	1.090	1.067	1.062	1.099	3.70
21) T Trichlorotrifl...	4.632	4.512	4.217	3.870	3.906	3.766	3.653	4.079	4.079	9.31
22) T Carbon Disulfide	0.797	0.909	1.278	1.306	1.364	1.394	1.357	1.323	1.216	18.81
23) T trans-1,2-Dich...	1.655	1.724	1.814	1.782	1.815	1.738	1.683	1.631	1.730	4.07
24) T 1,1-Dichloroet...	2.784	2.946	3.175	3.042	3.063	2.981	2.866	2.785	2.955	4.70
25) T Methyl tert-Bu...	0.227	0.242	0.266	0.266	0.276	0.263	0.247	0.253	0.253	7.11
26) T Vinyl Acetate	0.655	0.676	0.731	0.772	0.772	0.757	0.743	0.722	0.722	6.47
27) T 2-Butanone (MEK)	0.980	1.145	1.394	1.307	1.349	1.337	1.291	1.259	1.258	10.69
28) T cis-1,2-Dichlo...	0.828	1.007	1.104	1.080	1.089	1.048	1.001	0.953	1.014	8.96
29) T Diisopropyl Ether	0.354	0.361	0.386	0.376	0.376	0.352	0.317	0.358	0.358	6.66
30) T Ethyl Acetate	1.711	1.805	1.894	1.762	1.738	1.540	1.477	1.354	1.660	11.07
31) T n-Hexane	1.574	1.527	1.735	1.661	1.672	1.656	1.611	1.575	1.626	4.12
32) T Chloroform	1.071	1.072	1.080	1.077	1.066	1.048	1.051	1.061	1.066	1.09
33) S 1,2-Dichloroet...	0.744	0.728	0.765	0.749	0.745	0.731	0.707	0.695	0.733	3.15
34) T Tetrahydrofura...	1.153	1.191	1.333	1.335	1.301	1.285	1.248	1.225	1.259	5.27
35) T Ethyl tert-But...	0.877	0.970	1.089	1.066	1.085	1.092	1.051	1.028	1.032	7.24
36) T 1,2-Dichloroet...	-----ISTD-----									
37) IR 1,4-Difluorobenzen...	0.312	0.333	0.341	0.331	0.341	0.340	0.328	0.315	0.330	3.42
38) T 1,1,1-Trichlor...	0.156	0.166	0.173	0.164	0.164	0.160	0.149	0.135	0.159	7.45
39) T Isopropyl Acetate	0.226	0.166	0.225	0.207	0.226	0.243	0.227	0.207	0.216	10.78
40) T 1-Butanol	1.172	1.072	1.088	1.054	1.046	1.017	0.954	0.856	1.032	9.16
41) T Benzene	0.273	0.287	0.314	0.310	0.310	0.321	0.311	0.302	0.304	5.24
42) T Carbon Tetrach...	-----ISTD-----									

ADA 1/23/19

Method Path : I:\MS13\METHODS\
Method File : R13012319.M

Title	EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)										ISTD
43) T Cyclohexane	0.426	0.420	0.439	0.416	0.413	0.403	0.380	0.348	0.406	7.13	
44) T tert-Amyl Meth...	0.678	0.651	0.721	0.679	0.685	0.690	0.660	0.625	0.674	4.29	
45) T 1,2-Dichloropr...	0.231	0.230	0.253	0.241	0.245	0.241	0.230	0.218	0.236	4.61	
46) T Bromodichlorom...	0.259	0.263	0.291	0.282	0.299	0.307	0.298	0.287	0.286	6.04	
47) T Trichloroethene	0.276	0.295	0.318	0.297	0.306	0.308	0.299	0.288	0.298	4.31	
48) T 1,4-Dioxane	0.174	0.209	0.215	0.213	0.219	0.219	0.212	0.203	0.208	7.11	
49) T 2,2,4-Trimethy...	1.112	1.075	1.125	1.030	1.029	0.979	0.925	0.847	1.015	9.36	
50) T Methyl Methacr...	0.066	0.083	0.103	0.104	0.110	0.115	0.112	0.106	0.100	16.93	
51) T n-Heptane	0.253	0.254	0.268	0.250	0.257	0.248	0.238	0.223	0.249	5.39	
52) T cis-1,3-Dichlo...	0.351	0.354	0.379	0.379	0.402	0.388	0.370	0.374	5.29	6.41	
53) T 4-Methyl-2-pen...	0.190	0.198	0.219	0.223	0.218	0.230	0.218	0.206	0.213	12.77	
54) T trans-1,3-Dich...	0.265	0.263	0.325	0.325	0.355	0.345	0.333	0.314	6.04	6.04	
55) T 1,1,2-Trichlor...	0.212	0.228	0.255	0.243	0.251	0.252	0.244	0.235	0.240	0.78	
56) IR Chlorobenzene-d5	2.709	2.667	2.686	2.684	2.659	2.667	2.697	2.647	2.677	13.36	
57) S Toluene-d8 (SS2)	3.569	3.054	3.117	2.710	2.713	2.647	2.581	2.371	2.845	7.34	
58) T Toluene	1.016	0.984	1.177	1.131	1.162	1.212	1.175	1.087	1.118	10.69	
59) T 2-Hexanone	0.567	0.610	0.682	0.676	0.736	0.771	0.764	0.740	0.693	12.51	
60) T Dibromochlorom...	0.500	0.541	0.666	0.643	0.687	0.721	0.710	0.684	0.644	4.33	
61) T 1,2-Dibromoethane	1.351	1.230	1.328	1.275	1.299	1.371	1.315	1.212	1.298	10.53	
62) T n-Butyl Acetate	0.639	0.526	0.539	0.516	0.521	0.507	0.488	0.446	0.523	2.46	
63) T n-Octane	0.899	0.856	0.899	0.866	0.877	0.872	0.862	0.836	0.871	6.56	
64) T Tetrachloroethene	2.099	1.980	1.951	1.847	1.858	1.871	1.804	1.887	1.887	7.86	
65) S Chlorobenzene	3.471	3.201	3.189	2.987	3.071	3.034	2.893	2.653	3.062	11.20	
66) S Ethylbenzene	2.904	2.484	2.444	2.292	2.308	2.288	2.200	1.993	2.364	15.11	
67) S m- & p-Xylenes	0.510	0.516	0.587	0.587	0.669	0.737	0.736	0.724	0.633	6.08	
68) T Bromoform	1.735	1.728	1.842	1.740	1.943	2.016	1.956	1.824	1.848	10.14	
69) T Styrene	2.815	2.470	2.499	2.262	2.304	2.281	2.196	2.020	2.356	12.08	
70) T o-Xylene	1.296	1.341	1.217	1.136	1.168	1.110	1.035	0.910	1.152	3.80	
71) T n-Nonane	1.151	1.109	1.120	1.089	1.120	1.137	1.092	1.014	1.104	0.89	
72) T 1,1,2,2-Tetrac...	0.951	0.952	0.964	0.966	0.975	0.968	0.971	0.963	0.964	13.24	
73) S Bromofluoroben...	4.039	3.577	3.252	3.091	3.121	3.070	2.928	2.641	3.215	5.17	
74) T Cumene	1.583	1.666	1.507	1.443	1.553	1.590	1.545	1.425	1.539	11.13	
75) T alpha-Pinene	4.309	3.923	3.746	3.483	3.571	3.520	3.335	2.961	3.606	7.40	
76) T n-Propylbenzene	3.393	3.262	3.058	2.956	3.092	3.023	3.040	2.621	3.056	12.89	
77) T 3-Ethyltoluene	3.821	3.378	3.085	2.892	2.962	3.011	2.714	2.565	3.053	15.54	
78) T 4-Ethyltoluene	3.551	3.021	2.644	2.458	2.558	2.529	2.437	2.241	2.680	11.07	
79) T 1,3,5-Trimethy...	1.239	1.204	1.192	1.141	1.458	1.514	1.472	1.364	1.323	14.17	
80) T alpha-Methylst...	4.069	3.458	3.397	2.986	3.033	3.007	2.883	2.606	3.180	13.41	
81) T 1,2-Ethyltoluene	3.299	2.774	2.702	2.437	2.554	2.535	2.395	2.102	2.600	10.76	
82) T 2,2,4-Trimethy...	1.426	1.437	1.289	1.214	1.301	1.262	1.176	1.016	1.265	15.22	
83) T n-Decane	1.612	1.503	1.912	1.912	2.206	2.171	2.005	1.901	6.06	6.06	
84) T Benzyl Chloride	1.822	1.622	1.645	1.478	1.603	1.674	1.651	1.557	1.631	8.82	
85) T 1,3-Dichlorobe...	2.044	1.716	1.776	1.575	1.653	1.699	1.670	1.568	1.713	12.97	
86) T 1,4-Dichlorobe...	4.377	4.182	3.654	3.422	3.532	3.473	3.303	2.930	3.609	13.26	
87) T sec-Butylbenzene	3.963	3.909	3.479	3.234	3.401	3.315	3.076	2.574	3.369	10.79	
88) T 4-Isopropyltol...	3.027	2.856	2.585	2.417	2.580	2.563	2.431	2.117	2.572	8.38	
89) T 1,2,3-Trimethy...	1.866	1.629	1.643	1.471	1.553	1.601	1.552	1.425	1.592	9.79	
90) T 1,2-Dichlorobe...	0.921	0.995	0.823	0.795	0.987	1.001	0.938	0.799	0.907	8.91	
91) T d-Limonene	0.588	0.498	0.569	0.501	0.584	0.630	0.626	0.609	0.576	11.25	
92) T 1,2-Dibromo-3-...	1.106	1.062	1.086	1.030	1.352	1.349	1.256	1.092	1.167	14.95	
93) T n-Undecane	1.429	0.899	1.295	0.974	1.227	1.320	1.301	1.254	1.212	0.78	
94) T 1,2,4-Trichlor...											

Method Path : I:\MS13\METHODS\
 Method File : R13012319.M

Title	EPA TO-15	per SOP	VOA-TO15	(CASS TO-15/GC-MS)	2.338	3.658	2.347	3.720	4.006	3.800	3.281	3.307	20.98
95) T Naphthalene													
96) T n-Dodecane					1.159	0.696	1.263	1.298	1.178	0.952	1.091		20.92
97) T Hexachlorobuta...					1.262	0.980	0.893	0.755	0.834	0.840	0.829	0.901	17.74
98) T Cyclohexanone					1.010	0.860	0.896	0.836	0.762	0.872	0.854	0.802	8.50
99) T tert-Butylbenzene					3.242	3.109	2.704	2.543	2.601	2.548	2.391	2.067	14.23
100) T n-Butylbenzene					2.961	2.833	2.655	2.498	2.706	2.682	2.550	2.272	7.96

(#) = Out of Range

1/29/19

Primary Source Standards Concentrations (Working & Initial Calibration)

1ng/L Std. ID: S31-01221902 (2ng/L)
 4ng/L Std. ID: S31-01161906
 20ng/L Std. ID: S31-01161906
 200ng/L Std. ID: S31-01161903

Dilution Factors:

5 50 250 1000

Compounds	Source Std. mg/m ³	Primary Working Standards				Working STD Conc. (ng/L): Injection (L): ICAL Points:	4		20		200		200		200	
		200ng/L	20ng/L	4ng/L	1ng/L		0.025	0.050	0.025	0.050	0.025	0.050	0.125	25ng	50ng	100ng
Propene	1.031	206.2	20.62	4.124	1.031	0.1031	0.2062	0.5155	1.031	5.155	25.775	51.55	103.1			
Dichlorodifluoromethane	1.045	209.0	20.90	4.180	1.045	0.1045	0.2090	0.5225	1.045	5.225	26.125	52.25	104.5			
Chloromethane	1.008	201.6	20.16	4.032	1.008	0.1008	0.2016	0.5040	1.008	5.040	25.200	50.40	100.8			
Freon-114	1.028	205.6	20.56	4.112	1.028	0.1028	0.2056	0.5140	1.028	5.140	25.700	51.40	102.8			
Vinyl Chloride	1.051	210.2	21.02	4.204	1.051	0.1051	0.2102	0.5255	1.051	5.255	26.275	52.55	105.1			
1,3-Butadiene	1.049	209.8	20.98	4.196	1.049	0.1049	0.2098	0.5245	1.049	5.245	26.225	52.45	104.9			
Chloromethane	1.022	204.4	20.44	4.088	1.022	0.1022	0.2044	0.5110	1.022	5.110	25.550	51.10	102.2			
Ethanol	5.140	1028.0	102.80	20.560	5.140	0.5140	1.0280	2.5700	5.140	25.700	128.500	257.00	514.0			
Acetonitrile	1.033	206.6	20.66	4.132	1.033	0.1033	0.2066	0.5165	1.033	5.165	25.825	51.65	103.3			
Acrolein	1.028	205.6	20.56	4.112	1.028	0.1028	0.2056	0.5140	1.028	5.140	25.700	51.40	102.8			
Acetone	5.370	1074.0	107.40	21.480	5.370	0.5370	1.0740	2.6850	5.370	26.850	134.250	268.50	537.0			
Trichlorofluoromethane	1.060	212.0	21.20	4.240	1.060	0.1060	0.2120	0.5300	1.060	5.300	26.500	53.00	106.0			
Isopropanol	2.063	412.6	41.26	8.252	2.063	0.2063	0.4126	1.0315	2.063	10.315	51.575	103.15	206.3			
Acrylonitrile	1.034	206.8	20.68	4.136	1.034	0.1034	0.2068	0.5170	1.034	5.170	25.850	51.70	103.4			
1,1-Dichloroethene	1.074	214.8	21.48	4.296	1.074	0.1074	0.2148	0.5370	1.074	5.370	26.850	53.70	107.4			
tert-Butanol	2.144	428.8	42.88	8.576	2.144	0.2144	0.4288	1.0720	2.144	10.720	53.600	107.20	214.4			
Methylene Chloride	1.070	214.0	21.40	4.280	1.070	0.1070	0.2140	0.5350	1.070	5.350	26.750	53.50	107.0			
Allyl Chloride	1.067	213.4	21.34	4.268	1.067	0.1067	0.2134	0.5335	1.067	5.335	26.675	53.35	106.7			
Trichlorofluoroethane	1.065	213.0	21.30	4.260	1.065	0.1065	0.2130	0.5325	1.065	5.325	26.625	53.25	106.5			
Carbon Disulfide	1.075	215.0	21.50	4.300	1.075	0.1075	0.2150	0.5375	1.075	5.375	26.875	53.75	107.5			
trans-1,2-Dichloroethene	1.062	212.4	21.24	4.248	1.062	0.1062	0.2124	0.5310	1.062	5.310	26.550	53.10	106.2			
1,1-Dichloroethane	1.030	206.0	20.60	4.120	1.030	0.1030	0.2060	0.5150	1.030	5.150	25.750	51.50	103.0			
Methyl tert-Butyl Ether	1.089	217.8	21.78	4.356	1.089	0.1089	0.2178	0.5445	1.089	5.445	27.225	54.45	108.9			
Vinyl Acetate	5.252	1050.4	105.04	21.008	5.252	0.5252	1.0504	2.6260	5.252	26.260	131.300	262.60	525.2			
2-Butanone	1.027	205.4	20.54	4.108	1.027	0.1027	0.2054	0.5135	1.027	5.135	25.675	51.35	102.7			
cis-1,2-Dichloroethene	1.054	210.8	21.08	4.216	1.054	0.1054	0.2108	0.5270	1.054	5.270	26.350	52.70	105.4			
Diisopropyl Ether	1.081	216.2	21.62	4.324	1.081	0.1081	0.2162	0.5405	1.081	5.405	27.025	54.05	108.1			
Ethyl Acetate	2.166	433.2	43.32	8.664	2.166	0.2166	0.4332	1.0830	2.166	10.830	54.150	108.30	216.6			
n-Hexane	1.082	216.4	21.64	4.328	1.082	0.1082	0.2164	0.5410	1.082	5.410	27.050	54.10	108.2			
Chloroform	1.077	215.4	21.54	4.308	1.077	0.1077	0.2154	0.5385	1.077	5.385	26.925	53.85	107.7			
Tetrahydrofuran	1.068	213.6	21.36	4.272	1.068	0.1068	0.2136	0.5340	1.068	5.340	26.700	53.40	106.8			
Ethyl tert-Butyl Ether	1.060	212.0	21.20	4.240	1.060	0.1060	0.2120	0.5300	1.060	5.300	26.500	53.00	106.0			
1,2-Dichloroethane	1.061	212.2	21.22	4.244	1.061	0.1061	0.2122	0.5305	1.061	5.305	26.525	53.05	106.1			
1,1,1-Trichloroethane	1.081	216.2	21.62	4.324	1.081	0.1081	0.2162	0.5405	1.081	5.405	27.025	54.05	108.1			
Isopropyl Acetate	2.066	413.2	41.32	8.264	2.066	0.2066	0.4132	1.0330	2.066	10.330	51.650	103.30	206.6			
1-Butanol	2.067	413.4	41.34	8.268	2.067	0.2067	0.4134	1.0335	2.067	10.335	51.675	103.35	206.7			
Benzene	1.033	206.6	20.66	4.132	1.033	0.1033	0.2066	0.5165	1.033	5.165	25.825	51.65	103.3			
Carbon Tetrachloride	1.036	207.2	20.72	4.144	1.036	0.1036	0.2072	0.5180	1.036	5.180	25.900	51.80	103.6			
Cyclohexane	2.087	417.4	41.74	8.348	2.087	0.2087	0.4174	1.0435	2.087	10.435	52.175	104.35	208.7			
tert-Amyl Methyl Ether	1.074	214.8	21.48	4.296	1.074	0.1074	0.2148	0.5370	1.074	5.370	26.850	53.70	107.4			
1,2-Dichloropropane	1.073	214.6	21.46	4.292	1.073	0.1073	0.2146	0.5365	1.073	5.365	26.825	53.65	107.3			
Bromochloromethane	1.068	213.6	21.36	4.272	1.068	0.1068	0.2136	0.5340	1.068	5.340	26.700	53.40	106.8			
Trichloroethene	1.062	212.4	21.24	4.248	1.062	0.1062	0.2124	0.5310	1.062	5.310	26.550	53.10	106.2			
1,4-Dioxane	1.064	212.8	21.28	4.256	1.064	0.1064	0.2128	0.5320	1.064	5.320	26.600	53.20	106.4			
Isocetane	1.061	212.2	21.22	4.244	1.061	0.1061	0.2122	0.5305	1.061	5.305	26.525	53.05	106.1			
Methyl Methacrylate	2.135	427.0	42.70	8.540	2.135	0.2135	0.4270	1.0675	2.135	10.675	53.375	106.75	213.5			
n-Heptane	1.076	215.2	21.52	4.304	1.076	0.1076	0.2152	0.5380	1.076	5.380	26.900	53.80	107.6			

Primary Source Standards Concentrations (Working & Initial Calibration)

1ng/L Std. ID: S31-01221902 (2ng/L)
 4ng/L Std. ID: S31-01161906
 20ng/L Std. ID: S31-01161906
 200ng/L Std. ID: S31-01161903

Compounds	Source Std. mg/m ³	Primary Working Standards				Working STD Conc. (ng/L): Injection (L): ICAL Points:	4		20		200		200		200	
		200ng/L	20ng/L	4ng/L	1ng/L		0.025	0.050	0.025	0.050	0.025	0.050	0.125	25ng	50ng	100ng
cis-1,3-Dichloropropene	1.120	224.0	22.40	4.480	1.120	0.1120	0.2240	0.5600	1.120	5.600	28.000	26.000	56.00	112.0		
4-Methyl-2-pentanone	1.060	212.0	21.20	4.240	1.060	0.1060	0.2120	0.5300	1.060	5.300	26.500	26.500	53.00	106.0		
trans-1,3-Dichloropropene	1.055	211.0	21.10	4.220	1.055	0.1055	0.2110	0.5275	1.055	5.275	26.375	26.375	52.75	105.5		
1,1,2-Trichloroethane	1.076	215.2	21.52	4.304	1.076	0.1076	0.2152	0.5380	1.076	5.380	26.900	26.900	53.80	107.6		
Toluene	1.052	210.4	21.04	4.208	1.052	0.1052	0.2104	0.5260	1.052	5.260	26.300	26.300	52.60	105.2		
2-Hexanone	1.074	214.8	21.48	4.296	1.074	0.1074	0.2148	0.5370	1.074	5.370	26.850	26.850	53.70	107.4		
Dibromochloromethane	1.075	215.0	21.50	4.300	1.075	0.1075	0.2150	0.5375	1.075	5.375	26.875	26.875	53.75	107.5		
1,2-Dibromoethane	1.076	215.2	21.52	4.304	1.076	0.1076	0.2152	0.5380	1.076	5.380	26.900	26.900	53.80	107.6		
n-Butyl Acetate	1.085	217.0	21.70	4.340	1.085	0.1085	0.2170	0.5425	1.085	5.425	27.125	27.125	54.25	108.5		
n-Octane	1.076	215.2	21.52	4.304	1.076	0.1076	0.2152	0.5380	1.076	5.380	26.900	26.900	53.80	107.6		
Tetrachloroethene	1.058	211.6	21.16	4.232	1.058	0.1058	0.2116	0.5290	1.058	5.290	26.450	26.450	52.90	105.8		
Chlorobenzene	1.066	213.2	21.32	4.264	1.066	0.1066	0.2132	0.5330	1.066	5.330	26.650	26.650	53.30	106.6		
Ethylbenzene	1.033	206.6	20.66	4.132	1.033	0.1033	0.2066	0.5165	1.033	5.165	25.825	25.825	51.65	103.3		
m- <i>o</i> -Xylene	2.123	424.6	42.46	8.492	2.123	0.2123	0.4246	1.0615	2.123	10.615	53.075	53.075	106.15	212.3		
Bromoform	1.063	212.6	21.26	4.252	1.063	0.1063	0.2126	0.5315	1.063	5.315	26.575	26.575	53.15	106.3		
Styrene	1.060	212.0	21.20	4.240	1.060	0.1060	0.2120	0.5300	1.060	5.300	26.500	26.500	53.00	106.0		
<i>o</i> -Xylene	1.062	212.4	21.24	4.248	1.062	0.1062	0.2124	0.5310	1.062	5.310	26.550	26.550	53.10	106.2		
n-Nonane	1.071	214.2	21.42	4.284	1.071	0.1071	0.2142	0.5355	1.071	5.355	26.775	26.775	53.55	107.1		
1,1,2,2-Tetrachloroethane	1.064	212.8	21.28	4.256	1.064	0.1064	0.2128	0.5320	1.064	5.320	26.600	26.600	53.20	106.4		
Cumene	1.057	211.4	21.14	4.228	1.057	0.1057	0.2114	0.5285	1.057	5.285	26.425	26.425	52.85	105.7		
alpha-Prinene	1.035	207.0	20.70	4.140	1.035	0.1035	0.2070	0.5175	1.035	5.175	25.875	25.875	51.75	103.5		
n-Propylbenzene	1.076	215.2	21.52	4.304	1.076	0.1076	0.2152	0.5380	1.076	5.380	26.900	26.900	53.80	107.6		
3-Ethyltoluene	1.062	212.4	21.24	4.248	1.062	0.1062	0.2124	0.5310	1.062	5.310	26.550	26.550	53.10	106.2		
4-Ethyltoluene	1.061	212.2	21.22	4.244	1.061	0.1061	0.2122	0.5305	1.061	5.305	26.525	26.525	53.05	106.1		
1,3,5-Trimethylbenzene	1.057	211.4	21.14	4.228	1.057	0.1057	0.2114	0.5285	1.057	5.285	26.425	26.425	52.85	105.7		
alpha-Methylstyrene	1.058	211.6	21.16	4.232	1.058	0.1058	0.2116	0.5290	1.058	5.290	26.450	26.450	52.90	105.8		
2-Ethyltoluene	1.072	214.4	21.44	4.288	1.072	0.1072	0.2144	0.5360	1.072	5.360	26.800	26.800	53.60	107.2		
1,2,4-Trimethylbenzene	1.068	213.6	21.36	4.272	1.068	0.1068	0.2136	0.5340	1.068	5.340	26.700	26.700	53.40	106.8		
n-Decane	1.076	215.2	21.52	4.304	1.076	0.1076	0.2152	0.5380	1.076	5.380	26.900	26.900	53.80	107.6		
Benzyl Chloride	1.051	210.2	21.02	4.204	1.051	0.1051	0.2102	0.5255	1.051	5.255	26.275	26.275	52.55	105.1		
1,3-Dichlorobenzene	1.080	216.0	21.60	4.320	1.080	0.1080	0.2160	0.5400	1.080	5.400	27.000	27.000	54.00	108.0		
1,4-Dichlorobenzene	1.081	216.2	21.62	4.324	1.081	0.1081	0.2162	0.5405	1.081	5.405	27.025	27.025	54.05	108.1		
sec-Butylbenzene	1.063	212.6	21.26	4.252	1.063	0.1063	0.2126	0.5315	1.063	5.315	26.575	26.575	53.15	106.3		
p-Isopropyltoluene	1.042	208.4	20.84	4.168	1.042	0.1042	0.2084	0.5210	1.042	5.210	26.050	26.050	52.10	104.2		
1,2,3-Trimethylbenzene	1.042	208.4	20.84	4.168	1.042	0.1042	0.2084	0.5210	1.042	5.210	26.050	26.050	52.10	104.2		
1,2-Dichlorobenzene	1.089	217.8	21.78	4.356	1.089	0.1089	0.2178	0.5445	1.089	5.445	27.225	27.225	54.45	108.9		
d-Limonene	1.010	202.0	20.20	4.040	1.010	0.1010	0.2020	0.5050	1.010	5.050	25.250	25.250	50.50	101.0		
1,2-Dibromo-3-chloropropane	1.042	208.4	20.84	4.168	1.042	0.1042	0.2084	0.5210	1.042	5.210	26.050	26.050	52.10	104.2		
n-Undecane	1.057	211.4	21.14	4.228	1.057	0.1057	0.2114	0.5285	1.057	5.285	26.425	26.425	52.85	105.7		
1,2,4-Trichlorobenzene	1.064	212.8	21.28	4.256	1.064	0.1064	0.2128	0.5320	1.064	5.320	26.600	26.600	53.20	106.4		
Naphthalene	1.025	205.0	20.50	4.100	1.025	0.1025	0.2050	0.5125	1.025	5.125	25.625	25.625	51.25	102.5		
n-Dodecane	1.031	206.2	20.62	4.124	1.031	0.1031	0.2062	0.5155	1.031	5.155	25.775	25.775	51.55	103.1		
Hexachloro-1,3-butadiene	1.053	210.6	21.06	4.212	1.053	0.1053	0.2106	0.5265	1.053	5.265	26.325	26.325	52.65	105.3		
Methacrylonitrile	1.041	208.2	20.82	4.164	1.041	0.1041	0.2082	0.5205	1.041	5.205	26.025	26.025	52.05	104.1		
Cyclohexanone	0.982	196.4	19.64	3.928	0.982	0.0982	0.1964	0.4910	0.982	4.910	24.550	24.550	49.10	98.2		
tert-Butylbenzene	1.067	213.4	21.34	4.268	1.067	0.1067	0.2134	0.5335	1.067	5.335	26.675	26.675	53.35	106.7		
n-Butylbenzene	1.064	212.8	21.28	4.256	1.064	0.1064	0.2128	0.5320	1.064	5.320	26.600	26.600	53.20	106.4		

Method : I:\MS13\METHODS\R13012319.M (RTE Integrator)
 Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 Last Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration

#	ID	Conc	ISTD Conc	Path\File
1	0.1	0	13	I:\MS13\DATA\2019_01\23\01231906.D
2	0.2	0	13	I:\MS13\DATA\2019_01\23\01231907.D
3	0.5	1	13	I:\MS13\DATA\2019_01\23\01231904.D
4	1.0	1	13	I:\MS13\DATA\2019_01\23\01231905.D
5	5.0	5	13	I:\MS13\DATA\2019_01\23\01231908.D
6	25	26	13	I:\MS13\DATA\2019_01\23\01231909.D
7	50	52	13	I:\MS13\DATA\2019_01\23\01231910.D
8	100	103	13	I:\MS13\DATA\2019_01\23\01231911.D

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#	ID	Update Time				Quant Time			Acquisition Time	
1	0.1	Jan	23	11:19	2019	Jan	23	10:33	2019	23 Jan 2019 8:07
2	0.2	Jan	23	11:20	2019	Jan	23	10:34	2019	23 Jan 2019 8:40
3	0.5	Jan	23	11:20	2019	Jan	23	10:31	2019	23 Jan 2019 6:01
4	1.0	Jan	23	11:20	2019	Jan	23	10:39	2019	23 Jan 2019 6:35
5	5.0	Jan	23	11:20	2019	Jan	23	10:31	2019	23 Jan 2019 9:13
6	25	Jan	23	11:21	2019	Jan	23	10:31	2019	23 Jan 2019 9:46
7	50	Jan	23	11:21	2019	Jan	23	10:54	2019	23 Jan 2019 10:20
8	100	Jan	23	11:21	2019	Jan	23	11:18	2019	23 Jan 2019 10:53

R13012319.M

Wed Jan 23 13:58:22 2019

Data File : I:\MS13\DATA\2019_01\23\01231906.D
 Acq On : 23 Jan 2019 8:07
 Sample : 0.1ng TO-15 ICAL Std
 Misc : S31-01231908/S31-01221902 (2/20)

Vial: 14
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:33:34 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

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Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	133128	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.09	114	584644	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	230658	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.81	65	142619	12.206	ng	-0.02
Spiked Amount	12.500	Range 70 - 130	Recovery =	97.68%		
57) Toluene-d8 (SS2)	15.54	98	624855	13.133	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	105.04%		
73) Bromofluorobenzene (SS3)	18.83	174	219273	11.184	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	89.44%		

Target Compounds

						Qvalue
2) Propene	4.04	42	1193	0.095	ng	89
3) Dichlorodifluoromethan...	4.18	85	2013	0.090	ng	# 92
4) Chloromethane	4.47	50	1467	0.086	ng	80
5) 1,2-Dichloro-1,1,2,2-t...	4.70	135	1241	0.087	ng	86
6) Vinyl Chloride	4.88	62	1293	0.078	ng	69
7) 1,3-Butadiene	5.12	54	937	0.077	ng	94
8) Bromomethane	5.55	94	1047	0.091	ng	92
9) Chloroethane	0.00	64	0	N.D.	d	
10) Ethanol	6.24	45	4985	0.593	ng	79
11) Acetonitrile	6.60	41	970	0.046	ng	# 42
12) Acrolein	0.00	56	0	N.D.		
13) Acetone	6.90	58	5619	0.595	ng	# 83
14) Trichlorofluoromethane	7.09	101	1782	0.095	ng	96
15) 2-Propanol (Isopropanol)	7.44	45	5253	0.186	ng	85
16) Acrylonitrile	0.00	53	0	N.D.		
17) 1,1-Dichloroethene	8.03	96	1048	0.084	ng	99
18) 2-Methyl-2-Propanol (t...	8.32	59	4924	0.173	ng	# 77
19) Methylene Chloride	8.25	84	927	0.074	ng	100
20) 3-Chloro-1-propene (Al...	8.42	41	1307	0.082	ng	# 54
21) Trichlorotrifluoroethane	8.67	151	1225	0.093	ng	98
22) Carbon Disulfide	8.53	76	7486	0.162	ng	79
23) trans-1,2-Dichloroethene	9.53	61	902	0.061	ng	99
24) 1,1-Dichloroethane	9.76	63	1815	0.088	ng	83
25) Methyl tert-Butyl Ether	9.97	73	3229	0.091	ng	89
26) Vinyl Acetate	10.07	86	465	0.156	ng	# 1
27) 2-Butanone (MEK)	0.00	72	0	N.D.		
28) cis-1,2-Dichloroethene	10.79	61	1100	0.072	ng	79
29) Diisopropyl Ether	11.14	87	953	0.078	ng	# 51
30) Ethyl Acetate	11.17	61	189	0.047	ng	# 11
31) n-Hexane	11.08	57	1972	0.098	ng	# 100
32) Chloroform	11.13	83	1805	0.092	ng	96
34) Tetrahydrofuran (THF)	11.64	72	846	0.095	ng	# 87
35) Ethyl tert-Butyl Ether	11.74	87	1302	0.087	ng	94
36) 1,2-Dichloroethane	11.94	62	991	0.079	ng	# 42
38) 1,1,1-Trichloroethane	12.21	97	1576	0.092	ng	94
39) Isopropyl Acetate	12.69	61	1511	0.191	ng	# 68
40) 1-Butanol	12.75	56	2184	0.199	ng	87
41) Benzene	12.70	78	5663	0.107	ng	93
42) Carbon Tetrachloride	12.85	117	1323	0.085	ng	97
43) Cyclohexane	12.99	84	4162	0.200	ng	92
44) tert-Amyl Methyl Ether	13.37	73	3404	0.098	ng	89
45) 1,2-Dichloropropane	13.55	63	1158	0.095	ng	90
46) Bromodichloromethane	13.74	83	1295	0.088	ng	94
47) Trichloroethene	13.80	130	1371	0.090	ng	100
48) 1,4-Dioxane	13.84	88	865	0.081	ng	# 66
49) 2,2,4-Trimethylpentane...	13.86	57	5519	0.104	ng	87
50) Methyl Methacrylate	14.03	100	656	0.123	ng	# 55

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Data File : I:\MS13\DATA\2019_01\23\01231906.D
 Acq On : 23 Jan 2019 8:07
 Sample : 0.1ng TO-15 ICAL Std
 Misc : S31-01231908/S31-01221902 (2/20)

Vial: 14
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:33:34 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

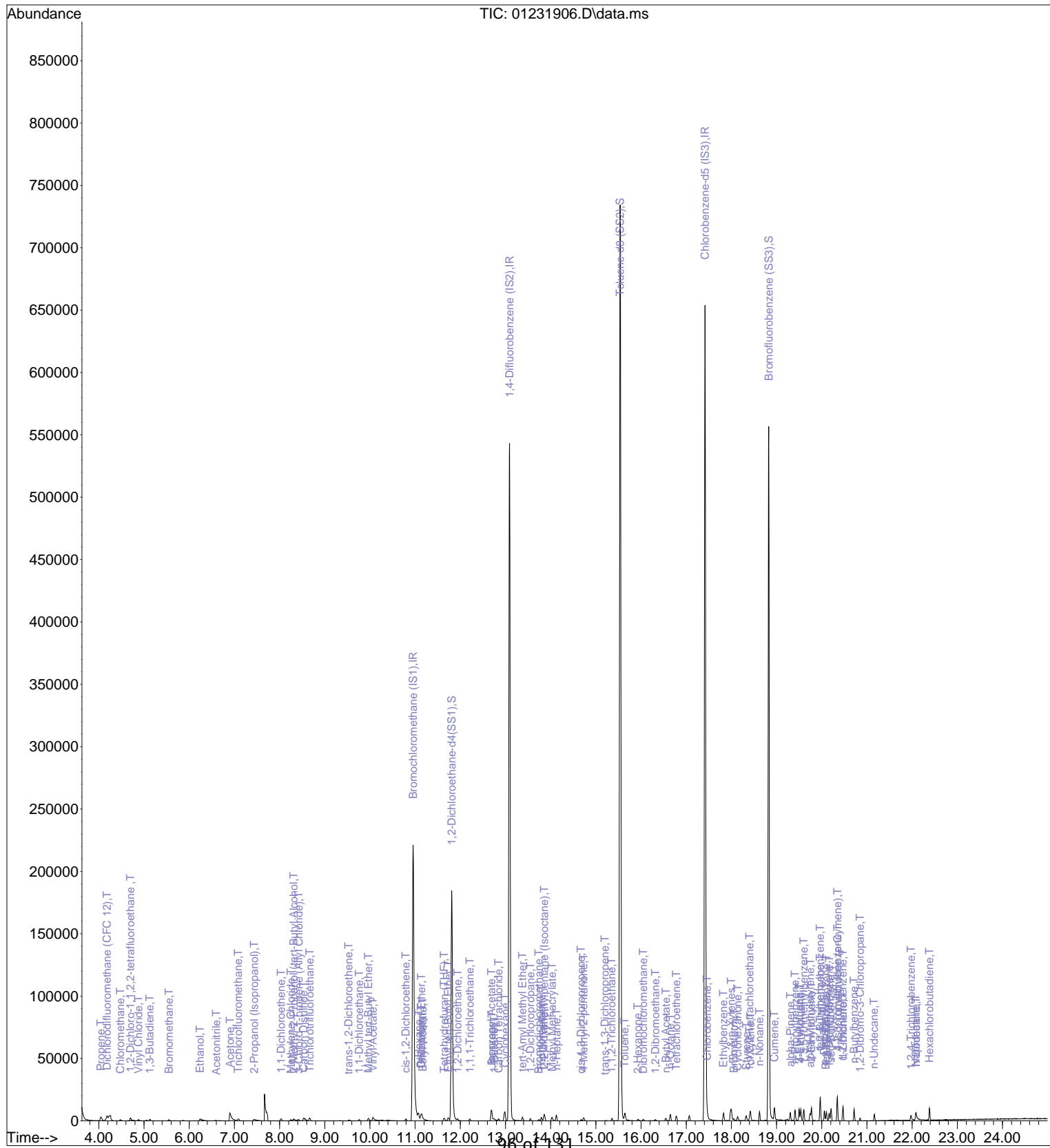
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	1274	0.098	ng	96
52) cis-1,3-Dichloropropene	14.67	75	1281	0.068	ng	76
53) 4-Methyl-2-pentanone	14.73	58	940	0.085	ng	80
54) trans-1,3-Dichloropropene	15.20	75	438	0.027	ng #	43
55) 1,1,2-Trichloroethane	15.35	97	1066	0.085	ng	98
58) Toluene	15.64	91	6928	0.129	ng	88
59) 2-Hexanone	15.92	43	2014	0.093	ng	91
60) Dibromochloromethane	16.05	129	1124	0.083	ng	94
61) 1,2-Dibromoethane	16.32	107	993	0.080	ng	97
62) n-Butyl Acetate	16.55	43	2705	0.108	ng	88
63) n-Octane	16.65	57	1269	0.128	ng #	78
64) Tetrachloroethene	16.78	166	1755	0.104	ng	96
65) Chlorobenzene	17.46	112	4129	0.114	ng	97
66) Ethylbenzene	17.83	91	6617	0.113	ng	96
67) m- & p-Xylenes	18.00	91	11376	0.254	ng	92
68) Bromoform	18.06	173	1000	0.076	ng	79
69) Styrene	18.33	104	3393	0.096	ng	96
70) o-Xylene	18.42	91	5517	0.123	ng	98
71) n-Nonane	18.62	43	2562	0.114	ng	99
72) 1,1,2,2-Tetrachloroethane	18.40	83	2260	0.105	ng	100
74) Cumene	18.95	105	7878	0.129	ng	99
75) alpha-Pinene	19.30	93	3024	0.102	ng	98
76) n-Propylbenzene	19.41	91	8556	0.123	ng	97
77) 3-Ethyltoluene	19.50	105	6650	0.111	ng	98
78) 4-Ethyltoluene	19.54	105	7480	0.129	ng	99
79) 1,3,5-Trimethylbenzene	19.60	105	6926	0.136	ng	92
80) alpha-Methylstyrene	19.74	118	2418	0.096	ng #	83
81) 2-Ethyltoluene	19.77	105	8049	0.133	ng	95
82) 1,2,4-Trimethylbenzene	19.96	105	6501	0.132	ng	97
83) n-Decane	20.06	57	2832	0.116	ng	96
84) Benzyl Chloride	20.09	91	2863	0.079	ng	80
85) 1,3-Dichlorobenzene	20.10	146	3631	0.116	ng	99
86) 1,4-Dichlorobenzene	20.16	146	4078	0.125	ng	95
87) sec-Butylbenzene	20.20	105	8586	0.124	ng	98
88) 4-Isopropyltoluene (p-...	20.35	119	7620	0.117	ng	96
89) 1,2,3-Trimethylbenzene	20.35	105	5820	0.117	ng	97
90) 1,2-Dichlorobenzene	20.47	146	3749	0.122	ng	95
91) d-Limonene	20.47	68	1717	0.098	ng	99
92) 1,2-Dibromo-3-Chloropr...	20.85	157	1130	0.103	ng	88
93) n-Undecane	21.16	57	2158	0.097	ng	94
94) 1,2,4-Trichlorobenzene	21.97	180	2805	0.131	ng #	94
95) Naphthalene	22.08	128	11176	0.191	ng	82
96) n-Dodecane	22.09	57	1465	0.076	ng	81
97) Hexachlorobutadiene	22.38	225	2453	0.148	ng	94
98) Cyclohexanone	18.14	55	1831	0.108	ng	90
99) tert-Butylbenzene	19.96	119	6384	0.125	ng	100
100) n-Butylbenzene	20.72	91	5814	0.115	ng	99

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231906.D
Acq On : 23 Jan 2019 8:07
Sample : 0.1ng TO-15 ICAL Std
Misc : S31-01231908/S31-01221902 (2/20)

Vial: 14
Operator: WA
Inst : MS13

Quant Time: Jan 23 10:33:34 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 10:31:09 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_01\23\01231907.D
 Acq On : 23 Jan 2019 8:40
 Sample : 0.2ng TO-15 ICAL Std
 Misc : S31-01231908/S31-01221902 (2/20)

Vial: 14
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:34:52 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

1/23/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	127975	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.09	114	556534	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	222857	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	137251	12.219	ng	-0.01
Spiked Amount	12.500	Range 70 - 130	Recovery =	97.76%		
57) Toluene-d8 (SS2)	15.54	98	594339	12.929	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	103.44%		
73) Bromofluorobenzene (SS3)	18.83	174	212177	11.201	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	89.60%		

Target Compounds

						Qvalue
2) Propene	4.02	42	2166	0.180	ng	89
3) Dichlorodifluoromethan...	4.17	85	4073	0.190	ng	# 96
4) Chloromethane	4.46	50	2677	0.163	ng	88
5) 1,2-Dichloro-1,1,2,2-t...	4.69	135	2540	0.184	ng	92
6) Vinyl Chloride	4.86	62	2541	0.159	ng	93
7) 1,3-Butadiene	5.12	54	1645	0.141	ng	95
8) Bromomethane	5.54	94	1852	0.168	ng	91
9) Chloroethane	5.86	64	1383	0.159	ng	87
10) Ethanol	6.22	45	7199	0.892	ng	86
11) Acetonitrile	6.52	41	2519	0.125	ng	90
12) Acrolein	0.00	56	0	N.D.	d	
13) Acetone	6.88	58	9332	1.028	ng	87
14) Trichlorofluoromethane	7.09	101	3329	0.185	ng	96
15) 2-Propanol (Isopropanol)	7.40	45	10035	0.370	ng	92
16) Acrylonitrile	0.00	53	0	N.D.		
17) 1,1-Dichloroethene	8.04	96	2122	0.176	ng	86
18) 2-Methyl-2-Propanol (t...	8.28	59	10027	0.367	ng	87
19) Methylene Chloride	8.25	84	1827	0.152	ng	91
20) 3-Chloro-1-propene (Al...	8.41	41	2691	0.176	ng	72
21) Trichlorotrifluoroethane	8.67	151	2342	0.185	ng	96
22) Carbon Disulfide	8.53	76	10196	0.229	ng	92
23) trans-1,2-Dichloroethene	9.54	61	1977	0.140	ng	85
24) 1,1-Dichloroethane	9.76	63	3636	0.184	ng	95
25) Methyl tert-Butyl Ether	9.94	73	6568	0.192	ng	96
26) Vinyl Acetate	10.06	86	1473	0.514	ng	# 1
27) 2-Butanone (MEK)	10.38	72	624	0.077	ng	# 1
28) cis-1,2-Dichloroethene	10.79	61	2471	0.169	ng	90
29) Diisopropyl Ether	11.13	87	2229	0.191	ng	94
30) Ethyl Acetate	11.14	61	994	0.256	ng	96
31) n-Hexane	11.08	57	3998	0.207	ng	# 100
32) Chloroform	11.13	83	3368	0.179	ng	95
34) Tetrahydrofuran (THF)	11.63	72	1591	0.187	ng	# 84
35) Ethyl tert-Butyl Ether	11.72	87	2584	0.180	ng	93
36) 1,2-Dichloroethane	11.94	62	2108	0.175	ng	88
38) 1,1,1-Trichloroethane	12.21	97	3203	0.197	ng	91
39) Isopropyl Acetate	12.68	61	3060	0.407	ng	95
40) 1-Butanol	12.74	56	3053	0.292	ng	99
41) Benzene	12.70	78	9863	0.196	ng	98
42) Carbon Tetrachloride	12.85	117	2651	0.178	ng	100
43) Cyclohexane	12.99	84	7799	0.394	ng	98
44) tert-Amyl Methyl Ether	13.37	73	6226	0.188	ng	97
45) 1,2-Dichloropropane	13.55	63	2196	0.189	ng	96
46) Bromodichloromethane	13.74	83	2497	0.177	ng	98
47) Trichloroethene	13.80	130	2785	0.191	ng	96
48) 1,4-Dioxane	13.82	88	1983	0.196	ng	78
49) 2,2,4-Trimethylpentane...	13.86	57	10156	0.201	ng	89
50) Methyl Methacrylate	14.02	100	1574	0.310	ng	92

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Data File : I:\MS13\DATA\2019_01\23\01231907.D
 Acq On : 23 Jan 2019 8:40
 Sample : 0.2ng TO-15 ICAL Std
 Misc : S31-01231908/S31-01221902 (2/20)

Vial: 14
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:34:52 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

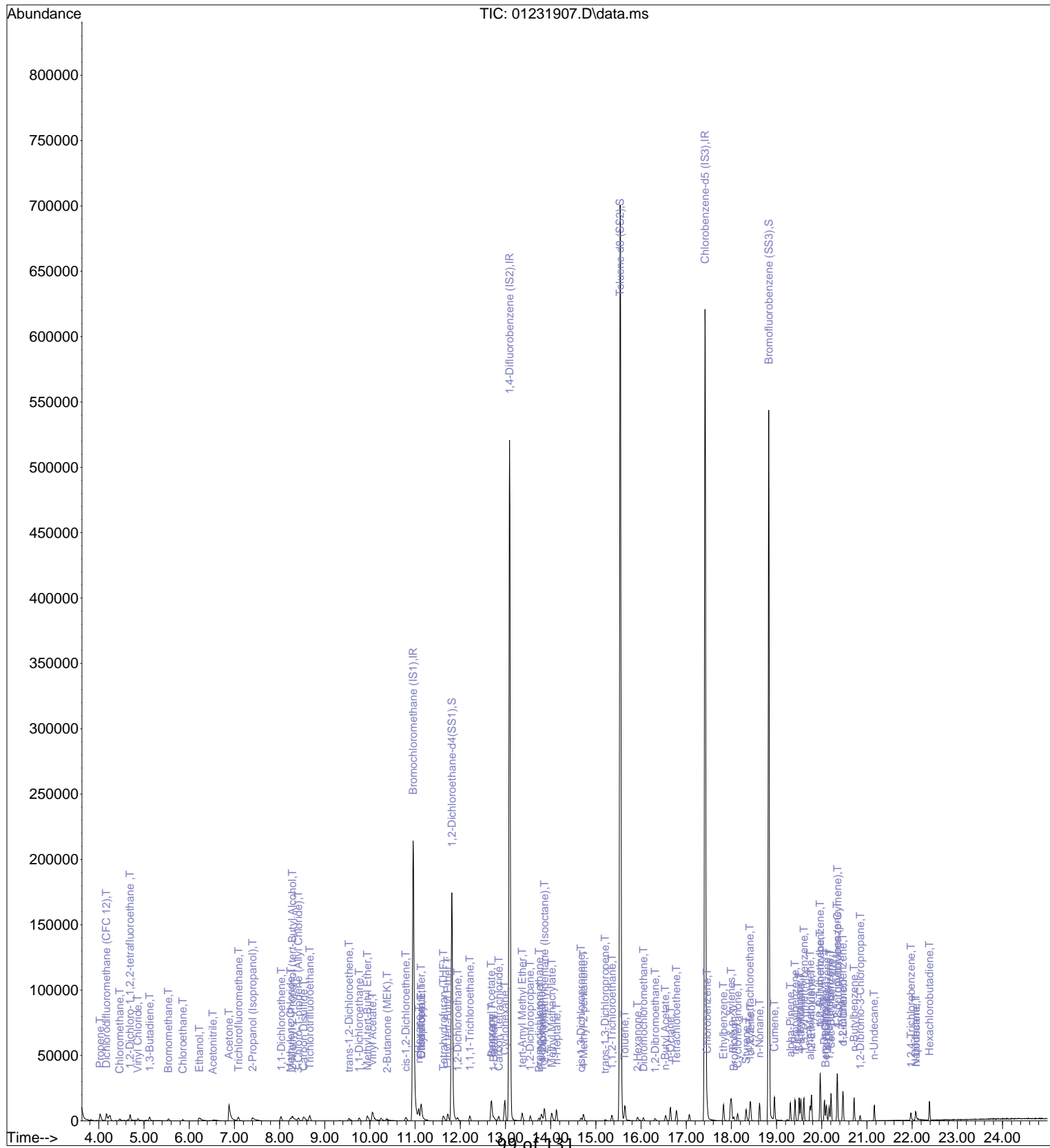
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	2430	0.197	ng	100
52) cis-1,3-Dichloropropene	14.67	75	2723	0.151	ng	95
53) 4-Methyl-2-pentanone	14.73	58	1867	0.178	ng	93
54) trans-1,3-Dichloropropene	15.21	75	1375	0.091	ng	74
55) 1,1,2-Trichloroethane	15.36	97	2184	0.182	ng	99
58) Toluene	15.64	91	11456	0.221	ng	98
59) 2-Hexanone	15.92	43	3767	0.180	ng	97
60) Dibromochloromethane	16.06	129	2339	0.179	ng	93
61) 1,2-Dibromoethane	16.31	107	2077	0.172	ng	98
62) n-Butyl Acetate	16.54	43	4757	0.197	ng	95
63) n-Octane	16.65	57	2019	0.210	ng	97
64) Tetrachloroethene	16.78	166	3229	0.198	ng	94
65) Chlorobenzene	17.46	112	7527	0.215	ng	97
66) Ethylbenzene	17.83	91	11792	0.209	ng	96
67) m- & p-Xylenes	18.00	91	18801	0.434	ng	99
68) Bromoform	18.05	173	1957	0.155	ng	88
69) Styrene	18.33	104	6533	0.192	ng	93
70) o-Xylene	18.42	91	9354	0.215	ng	98
71) n-Nonane	18.62	43	5123	0.237	ng	95
72) 1,1,2,2-Tetrachloroethane	18.41	83	4209	0.202	ng	97
74) Cumene	18.95	105	13483	0.228	ng	99
75) alpha-Pinene	19.30	93	6149	0.214	ng	98
76) n-Propylbenzene	19.41	91	15052	0.224	ng	96
77) 3-Ethyltoluene	19.50	105	12352	0.214	ng	98
78) 4-Ethyltoluene	19.54	105	12780	0.228	ng	99
79) 1,3,5-Trimethylbenzene	19.61	105	11386	0.232	ng	99
80) alpha-Methylstyrene	19.74	118	4544	0.187	ng	82
81) 2-Ethyltoluene	19.77	105	13219	0.226	ng	99
82) 1,2,4-Trimethylbenzene	19.96	105	10564	0.222	ng	96
83) n-Decane	20.06	57	5512	0.235	ng	96
84) Benzyl Chloride	20.08	91	4457	0.128	ng	93
85) 1,3-Dichlorobenzene	20.11	146	6246	0.206	ng	99
86) 1,4-Dichlorobenzene	20.16	146	6614	0.209	ng	99
87) sec-Butylbenzene	20.20	105	15852	0.236	ng	97
88) 4-Isopropyltoluene (p-...	20.35	119	14524	0.230	ng	98
89) 1,2,3-Trimethylbenzene	20.34	105	10612	0.221	ng	95
90) 1,2-Dichlorobenzene	20.47	146	6325	0.213	ng	98
91) d-Limonene	20.47	68	3584	0.213	ng	92
92) 1,2-Dibromo-3-Chloropr...	20.85	157	1852	0.174	ng	92
93) n-Undecane	21.16	57	4003	0.187	ng	97
94) 1,2,4-Trichlorobenzene	21.97	180	3412	0.164	ng	97
95) Naphthalene	22.08	128	8545	0.151	ng	85
96) n-Dodecane	22.08	57	1808	0.097	ng	85
97) Hexachlorobutadiene	22.38	225	3681	0.230	ng	99
98) Cyclohexanone	18.14	55	3011	0.183	ng	95
99) tert-Butylbenzene	19.96	119	11829	0.240	ng	98
100) n-Butylbenzene	20.72	91	10747	0.219	ng	97

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231907.D
Acq On : 23 Jan 2019 8:40
Sample : 0.2ng TO-15 ICAL Std
Misc : S31-01231908/S31-01221902 (2/20)

Vial: 14
Operator: WA
Inst : MS13

Quant Time: Jan 23 10:34:52 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 10:31:09 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



TIC: 01231907.D\data.ms

Data File : I:\MS13\DATA\2019_01\23\01231904.D
 Acq On : 23 Jan 2019 6:01
 Sample : 0.5ng R13012319 ICAL Std
 Misc : S31-01231908/S31-01161906 (2/14)

Vial: 15
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:31:37 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

WA 1/23/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	131127	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.09	114	573396	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	228390	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	141581	12.302	ng	-0.01
Spiked Amount	12.500	Range 70 - 130	Recovery =	98.40%		
57) Toluene-d8 (SS2)	15.54	98	613470	13.022	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	104.16%		
73) Bromofluorobenzene (SS3)	18.83	174	220128	11.339	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	90.72%		

Target Compounds

	R.T.	QIon	Response	Conc	Units	Qvalue
2) Propene	4.02	42	6188	0.502	ng	95
3) Dichlorodifluoromethan...	4.16	85	10871	0.494	ng	98
4) Chloromethane	4.45	50	9013	0.535	ng	93
5) 1,2-Dichloro-1,1,2,2-t...	4.68	135	6979	0.495	ng	97
6) Vinyl Chloride	4.85	62	8493	0.518	ng	94
7) 1,3-Butadiene	5.11	54	6048	0.504	ng	91
8) Bromomethane	5.53	94	5409	0.478	ng	96
9) Chloroethane	5.85	64	4318	0.486	ng	97
10) Ethanol	6.19	45	21756	2.630	ng	98
11) Acetonitrile	6.55	41	9317	0.452	ng	94
12) Acrolein	6.69	56	2575	0.364	ng	# 69
13) Acetone	6.86	58	23437	2.519	ng	85
14) Trichlorofluoromethane	7.09	101	9377	0.508	ng	98
15) 2-Propanol (Isopropanol)	7.36	45	28011	1.008	ng	100
16) Acrylonitrile	7.66	53	3291	0.250	ng	89
17) 1,1-Dichloroethene	8.03	96	6206	0.504	ng	86
18) 2-Methyl-2-Propanol (t...	8.24	59	28382	1.015	ng	93
19) Methylene Chloride	8.25	84	5865	0.478	ng	99
20) 3-Chloro-1-propene (Al...	8.41	41	7627	0.487	ng	88
21) Trichlorotrifluoroethane	8.67	151	6625	0.510	ng	97
22) Carbon Disulfide	8.53	76	25443	0.557	ng	97
23) trans-1,2-Dichloroethene	9.53	61	7120	0.491	ng	96
24) 1,1-Dichloroethane	9.76	63	9802	0.483	ng	100
25) Methyl tert-Butyl Ether	9.92	73	18137	0.519	ng	98
26) Vinyl Acetate	10.04	86	6248	2.129	ng	# 73
27) 2-Butanone (MEK)	10.33	72	3527	0.423	ng	93
28) cis-1,2-Dichloroethene	10.79	61	7706	0.514	ng	94
29) Diisopropyl Ether	11.12	87	6258	0.523	ng	# 94
30) Ethyl Acetate	11.13	61	4017	1.010	ng	99
31) n-Hexane	11.08	57	10750	0.542	ng	# 100
32) Chloroform	11.13	83	9799	0.507	ng	97
34) Tetrahydrofuran (THF)	11.60	72	4287	0.491	ng	92
35) Ethyl tert-Butyl Ether	11.71	87	7410	0.505	ng	98
36) 1,2-Dichloroethane	11.94	62	6063	0.490	ng	100
38) 1,1,1-Trichloroethane	12.21	97	8450	0.505	ng	95
39) Isopropyl Acetate	12.67	61	8208	1.061	ng	# 86
40) 1-Butanol	12.72	56	10664	0.990	ng	96
41) Benzene	12.69	78	25778	0.498	ng	98
42) Carbon Tetrachloride	12.85	117	7473	0.488	ng	99
43) Cyclohexane	12.98	84	20992	1.030	ng	96
44) tert-Amyl Methyl Ether	13.36	73	17772	0.520	ng	97
45) 1,2-Dichloropropane	13.55	63	6221	0.519	ng	97
46) Bromodichloromethane	13.74	83	7128	0.491	ng	100
47) Trichloroethene	13.80	130	7746	0.516	ng	96
48) 1,4-Dioxane	13.81	88	5254	0.503	ng	97
49) 2,2,4-Trimethylpentane...	13.86	57	27384	0.526	ng	88
50) Methyl Methacrylate	14.02	100	5039	0.962	ng	94

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Data File : I:\MS13\DATA\2019_01\23\01231904.D
 Acq On : 23 Jan 2019 6:01
 Sample : 0.5ng R13012319 ICAL Std
 Misc : S31-01231908/S31-01161906 (2/14)

Vial: 15
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:31:37 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

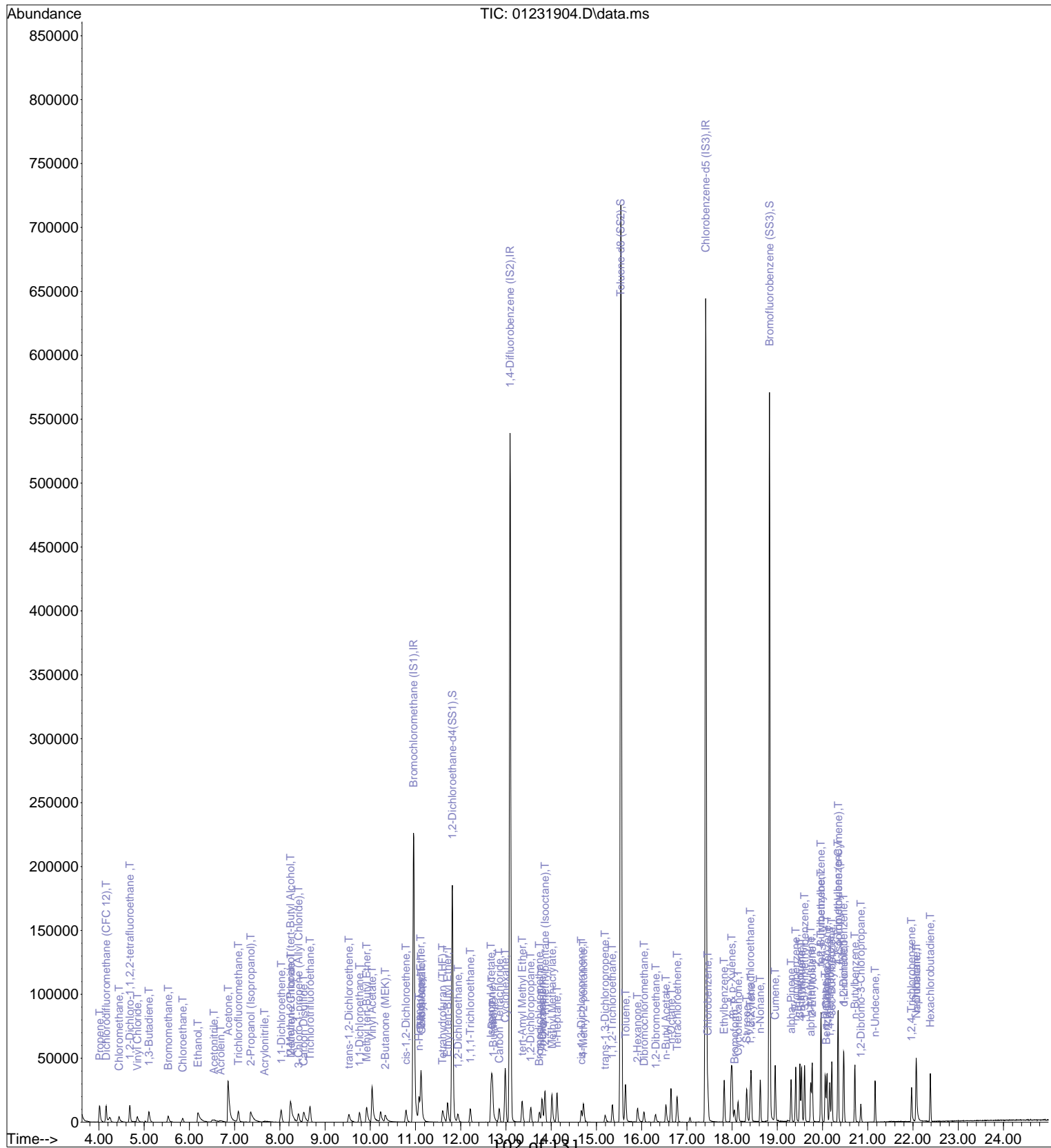
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	6608	0.519	ng	97
52) cis-1,3-Dichloropropene	14.67	75	9016	0.485	ng	98
53) 4-Methyl-2-pentanone	14.71	58	5334	0.495	ng	94
54) trans-1,3-Dichloropropene	15.19	75	6421	0.410	ng	89
55) 1,1,2-Trichloroethane	15.35	97	6286	0.510	ng	96
58) Toluene	15.65	91	29958	0.563	ng	94
59) 2-Hexanone	15.91	43	11552	0.540	ng	99
60) Dibromochloromethane	16.06	129	6694	0.500	ng	98
61) 1,2-Dibromoethane	16.31	107	6547	0.530	ng	98
62) n-Butyl Acetate	16.53	43	13162	0.533	ng	97
63) n-Octane	16.65	57	5295	0.538	ng	98
64) Tetrachloroethene	16.78	166	8693	0.520	ng	97
65) Chlorobenzene	17.46	112	18999	0.529	ng	100
66) Ethylbenzene	17.82	91	30091	0.520	ng	96
67) m- & p-Xylenes	17.99	91	47400	1.068	ng	100
68) Bromoform	18.05	173	5699	0.439	ng	96
69) Styrene	18.32	104	17838	0.510	ng	95
70) o-Xylene	18.42	91	24248	0.545	ng	100
71) n-Nonane	18.62	43	11903	0.536	ng	96
72) 1,1,2,2-Tetrachloroethane	18.40	83	10891	0.510	ng	99
74) Cumene	18.95	105	31404	0.518	ng	97
75) alpha-Pinene	19.30	93	14253	0.484	ng	96
76) n-Propylbenzene	19.41	91	36823	0.535	ng	98
77) 3-Ethyltoluene	19.50	105	29671	0.501	ng	97
78) 4-Ethyltoluene	19.54	105	29904	0.521	ng	97
79) 1,3,5-Trimethylbenzene	19.60	105	25536	0.507	ng	96
80) alpha-Methylstyrene	19.74	118	11519	0.461	ng	86
81) 2-Ethyltoluene	19.77	105	33268	0.554	ng	97
82) 1,2,4-Trimethylbenzene	19.96	105	26360	0.540	ng	98
83) n-Decane	20.06	57	12669	0.526	ng	97
84) Benzyl Chloride	20.08	91	15479	0.434	ng	92
85) 1,3-Dichlorobenzene	20.10	146	16227	0.521	ng	98
86) 1,4-Dichlorobenzene	20.16	146	17542	0.542	ng	99
87) sec-Butylbenzene	20.20	105	35482	0.516	ng	98
88) 4-Isopropyltoluene (p-...	20.35	119	33120	0.512	ng	99
89) 1,2,3-Trimethylbenzene	20.34	105	24606	0.500	ng	96
90) 1,2-Dichlorobenzene	20.47	146	16347	0.537	ng	100
91) d-Limonene	20.47	68	7596	0.440	ng	96
92) 1,2-Dibromo-3-Chloropr...	20.85	157	5415	0.497	ng	92
93) n-Undecane	21.17	57	10484	0.478	ng	97
94) 1,2,4-Trichlorobenzene	21.97	180	12589	0.592	ng	98
95) Naphthalene	22.08	128	34258	0.592	ng	96
96) n-Dodecane	22.07	57	10916	0.574	ng	94
97) Hexachlorobutadiene	22.38	225	8594	0.523	ng	99
98) Cyclohexanone	18.13	55	8038	0.477	ng	95
99) tert-Butylbenzene	19.96	119	26354	0.522	ng	99
100) n-Butylbenzene	20.72	91	25806	0.514	ng	99

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231904.D
 Acq On : 23 Jan 2019 6:01
 Sample : 0.5ng R13012319 ICAL Std
 Misc : S31-01231908/S31-01161906 (2/14)

Vial: 15
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:31:37 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_01\23\01231905.D
 Acq On : 23 Jan 2019 6:35
 Sample : 1.0ng R13012319 ICAL Std
 Misc : S31-01231908/S31-01161906 (2/14)

Vial: 15
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:39:30 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

 1/23/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.96	130	131496	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.09	114	571699	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	228613	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	141593	12.268	ng	-0.01
Spiked Amount	12.500	Range 70 - 130	Recovery =	98.16%		
57) Toluene-d8 (SS2)	15.54	98	613490	13.010	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	104.08%		
73) Bromofluorobenzene (SS3)	18.83	174	220895	11.367	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	90.96%		

Target Compounds

						Qvalue
2) Propene	4.01	42	11670	0.945	ng	93
3) Dichlorodifluoromethan...	4.16	85	21192	0.961	ng	99
4) Chloromethane	4.43	50	15705	0.929	ng	98
5) 1,2-Dichloro-1,1,2,2-t...	4.68	135	13367	0.945	ng	98
6) Vinyl Chloride	4.84	62	15750	0.958	ng	99
7) 1,3-Butadiene	5.09	54	11730	0.975	ng	95
8) Bromomethane	5.52	94	10617	0.936	ng	98
9) Chloroethane	5.85	64	8417	0.945	ng	94
10) Ethanol	6.18	45	39097	4.712	ng	99
11) Acetonitrile	6.49	41	17982	0.871	ng	98
12) Acrolein	6.66	56	6251	0.880	ng	84
13) Acetone	6.84	58	44350	4.754	ng	# 83
14) Trichlorofluoromethane	7.08	101	17946	0.969	ng	99
15) 2-Propanol (Isopropanol)	7.34	45	53344	1.914	ng	99
16) Acrylonitrile	7.63	53	10218	0.774	ng	95
17) 1,1-Dichloroethene	8.03	96	11968	0.968	ng	86
18) 2-Methyl-2-Propanol (t...	8.21	59	55574	1.981	ng	95
19) Methylene Chloride	8.24	84	11720	0.952	ng	97
20) 3-Chloro-1-propene (Al...	8.41	41	14670	0.934	ng	92
21) Trichlorotrifluoroethane	8.67	151	12451	0.955	ng	96
22) Carbon Disulfide	8.53	76	47691	1.042	ng	98
23) trans-1,2-Dichloroethene	9.52	61	14596	1.004	ng	97
24) 1,1-Dichloroethane	9.77	63	19304	0.949	ng	98
25) Methyl tert-Butyl Ether	9.91	73	34852	0.994	ng	100
26) Vinyl Acetate	10.04	86	13368	4.543	ng	# 83
27) 2-Butanone (MEK)	10.31	72	7298	0.873	ng	# 85
28) cis-1,2-Dichloroethene	10.79	61	14491	0.964	ng	94
29) Diisopropyl Ether	11.11	87	12277	1.023	ng	# 71
30) Ethyl Acetate	11.12	61	8220	2.061	ng	97
31) n-Hexane	11.08	57	20050	1.009	ng	# 100
32) Chloroform	11.13	83	18816	0.971	ng	97
34) Tetrahydrofuran (THF)	11.59	72	8419	0.961	ng	91
35) Ethyl tert-Butyl Ether	11.70	87	14886	1.011	ng	96
36) 1,2-Dichloroethane	11.93	62	11902	0.960	ng	99
38) 1,1,1-Trichloroethane	12.21	97	16351	0.980	ng	97
39) Isopropyl Acetate	12.67	61	15466	2.004	ng	# 91
40) 1-Butanol	12.70	56	19560	1.822	ng	# 63
41) Benzene	12.70	78	49801	0.965	ng	98
42) Carbon Tetrachloride	12.85	117	14697	0.962	ng	98
43) Cyclohexane	12.99	84	39673	1.952	ng	96
44) tert-Amyl Methyl Ether	13.35	73	33347	0.979	ng	98
45) 1,2-Dichloropropane	13.55	63	11849	0.992	ng	99
46) Bromodichloromethane	13.74	83	13766	0.951	ng	98
47) Trichloroethene	13.80	130	14446	0.965	ng	99
48) 1,4-Dioxane	13.80	88	10388	0.998	ng	98
49) 2,2,4-Trimethylpentane...	13.86	57	49978	0.963	ng	89
50) Methyl Methacrylate	14.01	100	10158	1.945	ng	99

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Data File : I:\MS13\DATA\2019_01\23\01231905.D
 Acq On : 23 Jan 2019 6:35
 Sample : 1.0ng R13012319 ICAL Std
 Misc : S31-01231908/S31-01161906 (2/14)

Vial: 15
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:39:30 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	12314	0.969	ng	94
52) cis-1,3-Dichloropropene	14.67	75	18150	0.979	ng	97
53) 4-Methyl-2-pentanone	14.71	58	10810	1.005	ng	95
54) trans-1,3-Dichloropropene	15.19	75	12705	0.814	ng	98
55) 1,1,2-Trichloroethane	15.36	97	11944	0.972	ng	100
58) Toluene	15.64	91	52134	0.979	ng	99
59) 2-Hexanone	15.90	43	22209	1.037	ng	97
60) Dibromochloromethane	16.05	129	13288	0.991	ng	100
61) 1,2-Dibromoethane	16.31	107	12663	1.024	ng	99
62) n-Butyl Acetate	16.53	43	25297	1.023	ng	96
63) n-Octane	16.65	57	10159	1.031	ng	96
64) Tetrachloroethene	16.79	166	16751	1.001	ng	99
65) Chlorobenzene	17.46	112	36012	1.001	ng	99
66) Ethylbenzene	17.83	91	56440	0.974	ng	96
67) m- & p-Xylenes	17.99	91	88976	2.002	ng	99
68) Bromoform	18.05	173	11408	0.879	ng	99
69) Styrene	18.32	104	33724	0.964	ng	95
70) o-Xylene	18.42	91	43939	0.986	ng	97
71) n-Nonane	18.62	43	22251	1.002	ng	98
72) 1,1,2,2-Tetrachloroethane	18.40	83	21186	0.990	ng	99
74) Cumene	18.95	105	59749	0.985	ng	99
75) alpha-Pinene	19.30	93	27308	0.925	ng	95
76) n-Propylbenzene	19.41	91	68547	0.995	ng	98
77) 3-Ethyltoluene	19.50	105	57418	0.969	ng	99
78) 4-Ethyltoluene	19.53	105	56124	0.977	ng	99
79) 1,3,5-Trimethylbenzene	19.60	105	47516	0.942	ng	97
80) alpha-Methylstyrene	19.74	118	22075	0.883	ng	87
81) 2-Ethyltoluene	19.77	105	58546	0.975	ng	99
82) 1,2,4-Trimethylbenzene	19.96	105	47609	0.975	ng	95
83) n-Decane	20.06	57	23892	0.991	ng	98
84) Benzyl Chloride	20.08	91	28882	0.809	ng	96
85) 1,3-Dichlorobenzene	20.10	146	29202	0.937	ng	100
86) 1,4-Dichlorobenzene	20.16	146	31131	0.961	ng	99
87) sec-Butylbenzene	20.20	105	66519	0.966	ng	100
88) 4-Isopropyltoluene (p-...	20.35	119	61624	0.951	ng	99
89) 1,2,3-Trimethylbenzene	20.34	105	46052	0.935	ng	96
90) 1,2-Dichlorobenzene	20.46	146	29303	0.961	ng	97
91) d-Limonene	20.47	68	14686	0.849	ng	96
92) 1,2-Dibromo-3-Chloropr...	20.85	157	9546	0.875	ng	89
93) n-Undecane	21.16	57	19919	0.908	ng	97
94) 1,2,4-Trichlorobenzene	21.97	180	18948	0.890	ng	98
95) Naphthalene	22.07	128	43994	0.759	ng	97
96) n-Dodecane	22.07	57	13115	0.689	ng	96
97) Hexachlorobutadiene	22.38	225	14537	0.884	ng	100
98) Cyclohexanone	18.13	55	15018	0.890	ng	96
99) tert-Butylbenzene	19.96	119	49631	0.981	ng	100
100) n-Butylbenzene	20.72	91	48604	0.966	ng	99

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231908.D
 Acq On : 23 Jan 2019 9:13
 Sample : 5.0ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:31:46 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

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Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.97	130	133163	12.500	ng	-0.01
37) 1,4-Difluorobenzene (IS2)	13.10	114	573427	12.500	ng	0.00
56) Chlorobenzene-d5 (IS3)	17.42	82	228793	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.82	65	141973	12.147	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	97.20%		
57) Toluene-d8 (SS2)	15.54	98	608275	12.889	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	103.12%		
73) Bromofluorobenzene (SS3)	18.83	174	223121	11.473	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	91.76%		

Target Compounds

						Qvalue
2) Propene	3.98	42	61288	4.900	ng	95
3) Dichlorodifluoromethan...	4.13	85	107398	4.808	ng	100
4) Chloromethane	4.40	50	83501	4.876	ng	98
5) 1,2-Dichloro-1,1,2,2-t...	4.66	135	66573	4.647	ng	99
6) Vinyl Chloride	4.81	62	83594	5.022	ng	98
7) 1,3-Butadiene	5.07	54	66679	5.475	ng	93
8) Bromomethane	5.50	94	56285	4.900	ng	99
9) Chloroethane	5.82	64	43477	4.818	ng	99
10) Ethanol	6.18	45	205480	24.456	ng	98
11) Acetonitrile	6.44	41	98230	4.696	ng	99
12) Acrolein	6.63	56	34941	4.860	ng	96
13) Acetone	6.82	58	222256	23.524	ng	# 82
14) Trichlorofluoromethane	7.06	101	89880	4.791	ng	99
15) 2-Propanol (Isopropanol)	7.31	45	281104	9.961	ng	96
16) Acrylonitrile	7.58	53	65893	4.932	ng	99
17) 1,1-Dichloroethene	8.01	96	62403	4.987	ng	86
18) 2-Methyl-2-Propanol (t...	8.18	59	282060	9.931	ng	95
19) Methylene Chloride	8.24	84	62044	4.977	ng	95
20) 3-Chloro-1-propene (Al...	8.40	41	81576	5.131	ng	93
21) Trichlorotrifluoroethane	8.66	151	63461	4.806	ng	94
22) Carbon Disulfide	8.50	76	221593	4.781	ng	100
23) trans-1,2-Dichloroethene	9.52	61	77142	5.241	ng	93
24) 1,1-Dichloroethane	9.77	63	99552	4.833	ng	99
25) Methyl tert-Butyl Ether	9.88	73	177677	5.003	ng	97
26) Vinyl Acetate	10.03	86	74341	24.950	ng	# 91
27) 2-Butanone (MEK)	10.29	72	39968	4.721	ng	# 90
28) cis-1,2-Dichloroethene	10.79	61	75760	4.977	ng	93
29) Diisopropyl Ether	11.10	87	62727	5.160	ng	# 72
30) Ethyl Acetate	11.10	61	44547	11.030	ng	99
31) n-Hexane	11.08	57	100179	4.977	ng	# 100
32) Chloroform	11.13	83	95915	4.890	ng	98
34) Tetrahydrofuran (THF)	11.56	72	42364	4.775	ng	97
35) Ethyl tert-Butyl Ether	11.69	87	73478	4.929	ng	98
36) 1,2-Dichloroethane	11.94	62	61320	4.884	ng	99
38) 1,1,1-Trichloroethane	12.21	97	84505	5.047	ng	97
39) Isopropyl Acetate	12.66	61	77927	10.069	ng	# 90
40) 1-Butanol	12.67	56	107106	9.945	ng	# 68
41) Benzene	12.70	78	247723	4.786	ng	98
42) Carbon Tetrachloride	12.86	117	73653	4.808	ng	99
43) Cyclohexane	12.99	84	197718	9.700	ng	97
44) tert-Amyl Methyl Ether	13.35	73	168842	4.941	ng	97
45) 1,2-Dichloropropane	13.55	63	60275	5.029	ng	100
46) Bromodichloromethane	13.74	83	73195	5.043	ng	100
47) Trichloroethene	13.80	130	74501	4.961	ng	100
48) 1,4-Dioxane	13.78	88	53416	5.117	ng	98
49) 2,2,4-Trimethylpentane...	13.86	57	250397	4.812	ng	89
50) Methyl Methacrylate	14.01	100	54065	10.321	ng	99

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Data File : I:\MS13\DATA\2019_01\23\01231908.D
 Acq On : 23 Jan 2019 9:13
 Sample : 5.0ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:31:46 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

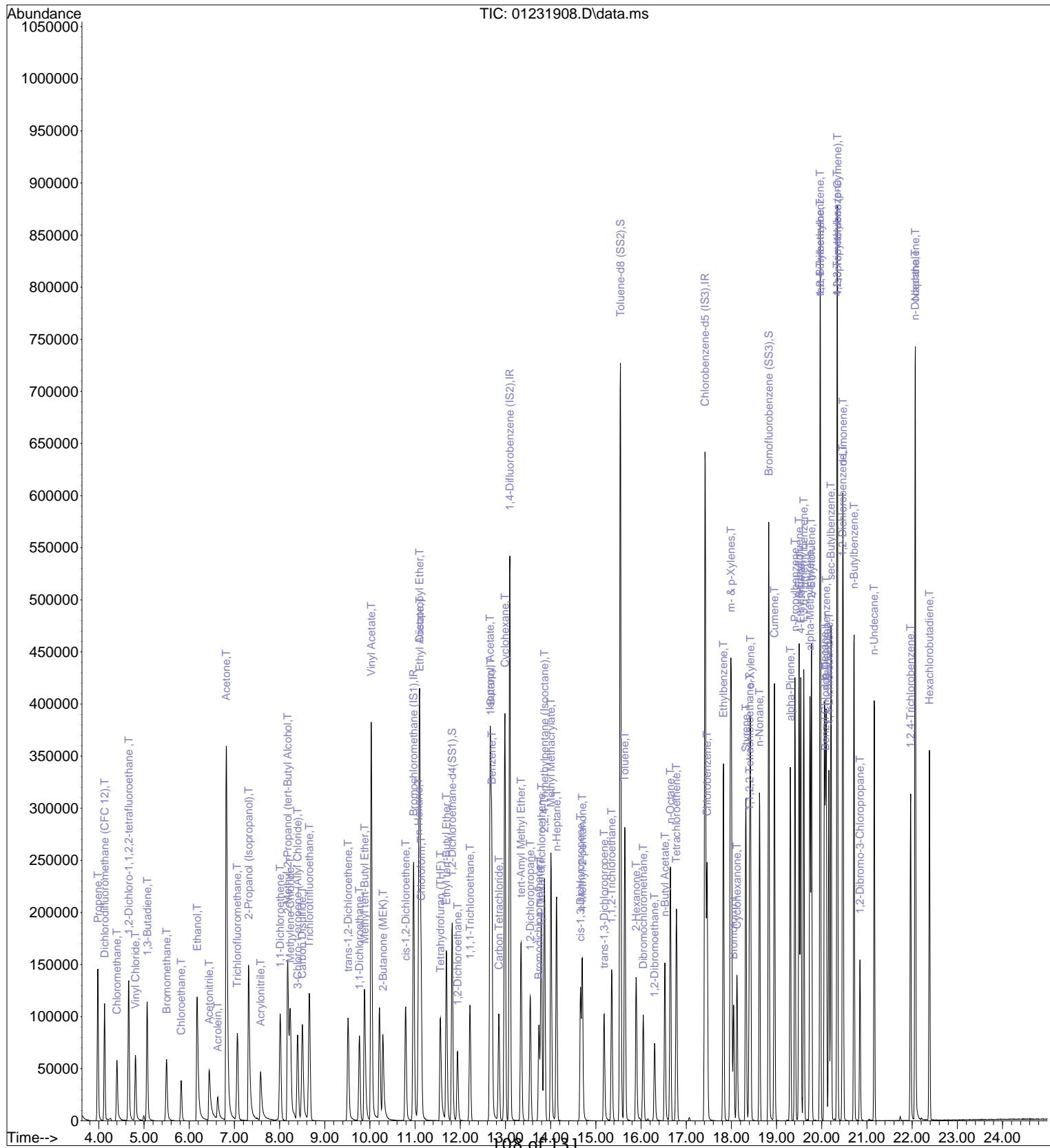
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	63371	4.974	ng	98
52) cis-1,3-Dichloropropene	14.66	75	97452	5.243	ng	100
53) 4-Methyl-2-pentanone	14.70	58	52931	4.908	ng	98
54) trans-1,3-Dichloropropene	15.18	75	78615	5.024	ng	99
55) 1,1,2-Trichloroethane	15.35	97	61981	5.026	ng	99
58) Toluene	15.64	91	261216	4.900	ng	99
59) 2-Hexanone	15.89	43	114253	5.332	ng	95
60) Dibromochloromethane	16.05	129	72456	5.401	ng	99
61) 1,2-Dibromoethane	16.30	107	67683	5.471	ng	99
62) n-Butyl Acetate	16.53	43	129007	5.215	ng	95
63) n-Octane	16.65	57	51306	5.203	ng	96
64) Tetrachloroethene	16.78	166	84886	5.067	ng	99
65) Chlorobenzene	17.46	112	181298	5.037	ng	99
66) Ethylbenzene	17.82	91	290291	5.005	ng	98
67) m- & p-Xylenes	17.99	91	448511	10.086	ng	98
68) Bromoform	18.05	173	65088	5.010	ng	99
69) Styrene	18.32	104	188503	5.384	ng	96
70) o-Xylene	18.42	91	223971	5.022	ng	97
71) n-Nonane	18.62	43	114495	5.151	ng	98
72) 1,1,2,2-Tetrachloroethane	18.40	83	109049	5.093	ng	100
74) Cumene	18.95	105	301863	4.970	ng	100
75) alpha-Pinene	19.30	93	147094	4.981	ng	95
76) n-Propylbenzene	19.41	91	351635	5.100	ng	99
77) 3-Ethyltoluene	19.50	105	300539	5.069	ng	99
78) 4-Ethyltoluene	19.54	105	287582	5.001	ng	100
79) 1,3,5-Trimethylbenzene	19.60	105	247436	4.903	ng	97
80) alpha-Methylstyrene	19.74	118	141128	5.643	ng	92
81) 2-Ethyltoluene	19.77	105	297586	4.951	ng	99
82) 1,2,4-Trimethylbenzene	19.96	105	249668	5.107	ng	96
83) n-Decane	20.06	57	128094	5.311	ng	99
84) Benzyl Chloride	20.08	91	183861	5.145	ng	100
85) 1,3-Dichlorobenzene	20.10	146	158457	5.082	ng	100
86) 1,4-Dichlorobenzene	20.16	146	163491	5.041	ng	99
87) sec-Butylbenzene	20.20	105	343634	4.985	ng	99
88) 4-Isopropyltoluene (p-...	20.35	119	324342	5.001	ng	99
89) 1,2,3-Trimethylbenzene	20.34	105	246057	4.990	ng	95
90) 1,2-Dichlorobenzene	20.46	146	154769	5.073	ng	100
91) d-Limonene	20.47	68	91243	5.271	ng	95
92) 1,2-Dibromo-3-Chloropr...	20.85	157	55738	5.106	ng	86
93) n-Undecane	21.16	57	130753	5.954	ng	97
94) 1,2,4-Trichlorobenzene	21.97	180	119437	5.606	ng	100
95) Naphthalene	22.07	128	348909	6.016	ng	99
96) n-Dodecane	22.06	57	119194	6.256	ng	98
97) Hexachlorobutadiene	22.38	225	78626	4.775	ng	100
98) Cyclohexanone	18.12	55	68504	4.059	ng	95
99) tert-Butylbenzene	19.96	119	253961	5.018	ng	100
100) n-Butylbenzene	20.72	91	263504	5.235	ng	99

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231908.D
Acq On : 23 Jan 2019 9:13
Sample : 5.0ng TO-15 ICAL Std
Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
Operator: WA
Inst : MS13

Quant Time: Jan 23 10:31:46 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 10:31:09 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_01\23\01231909.D
 Acq On : 23 Jan 2019 9:46
 Sample : 25ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:31:48 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

1/23/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.98	130	127070	12.500	ng	0.00
37) 1,4-Difluorobenzene (IS2)	13.10	114	539066	12.500	ng	0.00
56) Chlorobenzene-d5 (IS3)	17.42	82	215185	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.83	65	133196	11.943	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	95.52%		
57) Toluene-d8 (SS2)	15.54	98	573807	12.928	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	103.44%		
73) Bromofluorobenzene (SS3)	18.83	174	208213	11.383	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery =	91.04%		

Target Compounds

						Qvalue
2) Propene	3.98	42	281611	23.596	ng	96
3) Dichlorodifluoromethan...	4.13	85	489775	22.976	ng	100
4) Chloromethane	4.41	50	384457	23.528	ng	100
5) 1,2-Dichloro-1,1,2,2-t...	4.66	135	308779	22.585	ng	100
6) Vinyl Chloride	4.82	62	388805	24.478	ng	100
7) 1,3-Butadiene	5.08	54	303557	26.122	ng	93
8) Bromomethane	5.50	94	263398	24.029	ng	100
9) Chloroethane	5.83	64	204130	23.707	ng	100
10) Ethanol	6.22	45	946677	118.074	ng	97
11) Acetonitrile	6.45	41	499297	25.015	ng	97
12) Acrolein	6.64	56	176118	25.669	ng	97
13) Acetone	6.84	58	1016443	112.739	ng	# 82
14) Trichlorofluoromethane	7.07	101	406680	22.718	ng	99
15) 2-Propanol (Isopropanol)	7.34	45	1321330	49.064	ng	94
16) Acrylonitrile	7.59	53	367028	28.789	ng	97
17) 1,1-Dichloroethene	8.02	96	295186	24.719	ng	85
18) 2-Methyl-2-Propanol (t...	8.20	59	1335425	49.272	ng	96
19) Methylene Chloride	8.25	84	297023	24.969	ng	94
20) 3-Chloro-1-propene (Al...	8.41	41	388603	25.614	ng	98
21) Trichlorotrifluoroethane	8.66	151	295105	23.422	ng	94
22) Carbon Disulfide	8.51	76	1067121	24.127	ng	99
23) trans-1,2-Dichloroethene	9.53	61	376216	26.786	ng	94
24) 1,1-Dichloroethane	9.78	63	454839	23.140	ng	100
25) Methyl tert-Butyl Ether	9.88	73	825111	24.346	ng	98
26) Vinyl Acetate	10.04	86	368175	129.489	ng	96
27) 2-Butanone (MEK)	10.28	72	201439	24.934	ng	# 89
28) cis-1,2-Dichloroethene	10.80	61	358155	24.655	ng	92
29) Diisopropyl Ether	11.10	87	287847	24.816	ng	# 74
30) Ethyl Acetate	11.10	61	206917	53.689	ng	97
31) n-Hexane	11.08	57	423508	22.049	ng	# 100
32) Chloroform	11.14	83	453309	24.220	ng	99
34) Tetrahydrofuran (THF)	11.55	72	198310	23.422	ng	98
35) Ethyl tert-Butyl Ether	11.69	87	346238	24.338	ng	96
36) 1,2-Dichloroethane	11.95	62	294430	24.577	ng	100
38) 1,1,1-Trichloroethane	12.22	97	395934	25.156	ng	98
39) Isopropyl Acetate	12.66	61	355718	48.891	ng	# 92
40) 1-Butanol	12.68	56	541210	53.456	ng	# 74
41) Benzene	12.70	78	1132300	23.271	ng	99
42) Carbon Tetrachloride	12.86	117	359014	24.929	ng	100
43) Cyclohexane	12.99	84	907672	47.366	ng	98
44) tert-Amyl Methyl Ether	13.35	73	799357	24.886	ng	97
45) 1,2-Dichloropropane	13.55	63	278453	24.711	ng	100
46) Bromodichloromethane	13.74	83	354046	25.950	ng	99
47) Trichloroethene	13.80	130	352521	24.972	ng	100
48) 1,4-Dioxane	13.77	88	250996	25.577	ng	100
49) 2,2,4-Trimethylpentane...	13.86	57	1120282	22.901	ng	90
50) Methyl Methacrylate	14.01	100	264403	53.689	ng	96

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Data File : I:\MS13\DATA\2019_01\23\01231909.D
 Acq On : 23 Jan 2019 9:46
 Sample : 25ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:31:48 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

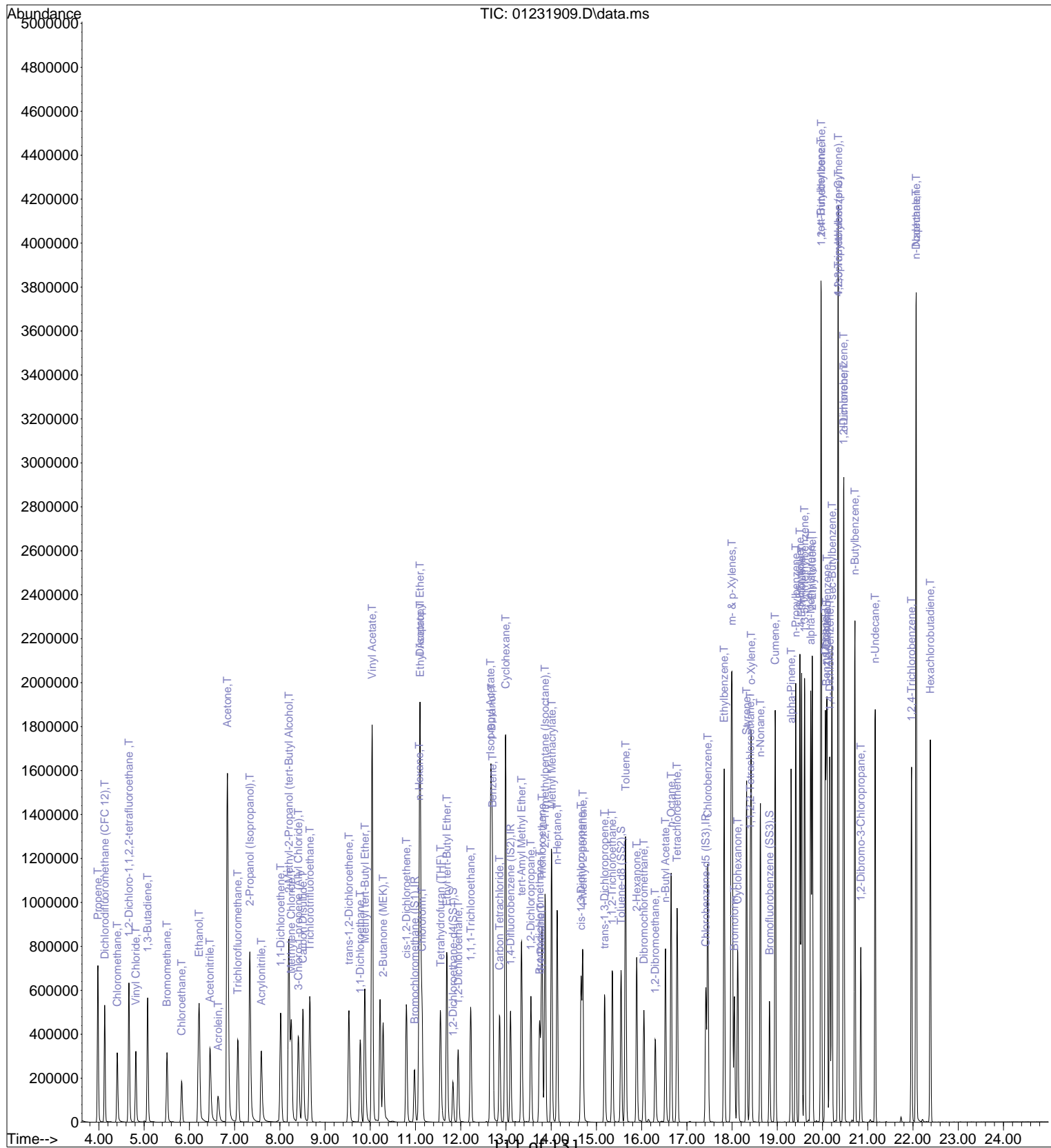
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	288030	24.050	ng	99
52) cis-1,3-Dichloropropene	14.66	75	485798	27.800	ng	100
53) 4-Methyl-2-pentanone	14.70	58	262607	25.904	ng	93
54) trans-1,3-Dichloropropene	15.18	75	404339	27.486	ng	100
55) 1,1,2-Trichloroethane	15.35	97	292085	25.197	ng	99
58) Toluene	15.65	91	1198602	23.905	ng	98
59) 2-Hexanone	15.89	43	560414	27.807	ng	93
60) Dibromochloromethane	16.05	129	356574	28.260	ng	100
61) 1,2-Dibromoethane	16.30	107	333858	28.692	ng	100
62) n-Butyl Acetate	16.53	43	640396	27.524	ng	95
63) n-Octane	16.65	57	235009	25.338	ng	97
64) Tetrachloroethene	16.78	166	396955	25.195	ng	99
65) Chlorobenzene	17.46	112	858357	25.358	ng	99
66) Ethylbenzene	17.83	91	1348824	24.725	ng	98
67) m- & p-Xylenes	18.00	91	2090753	49.990	ng	98
68) Bromoform	18.05	173	337172	27.594	ng	100
69) Styrene	18.32	104	919722	27.931	ng	96
70) o-Xylene	18.42	91	1042553	24.856	ng	96
71) n-Nonane	18.63	43	511403	24.463	ng	100
72) 1,1,2,2-Tetrachloroethane	18.40	83	520505	25.845	ng	99
74) Cumene	18.95	105	1396703	24.452	ng	99
75) alpha-Pinene	19.30	93	708141	25.497	ng	94
76) n-Propylbenzene	19.41	91	1629806	25.134	ng	98
77) 3-Ethyltoluene	19.50	105	1381613	24.774	ng	94
78) 4-Ethyltoluene	19.54	105	1374888	25.419	ng	95
79) 1,3,5-Trimethylbenzene	19.60	105	1150663	24.242	ng	96
80) alpha-Methylstyrene	19.74	118	689186	29.302	ng	92
81) 2-Ethyltoluene	19.77	105	1387145	24.536	ng	98
82) 1,2,4-Trimethylbenzene	19.97	105	1165280	25.344	ng	96
83) n-Decane	20.06	57	584605	25.770	ng	99
84) Benzyl Chloride	20.08	91	997797	29.686	ng	98
85) 1,3-Dichlorobenzene	20.10	146	777844	26.527	ng	100
86) 1,4-Dichlorobenzene	20.16	146	790407	25.911	ng	100
87) sec-Butylbenzene	20.21	105	1588997	24.509	ng	99
88) 4-Isopropyltoluene (p-...	20.35	119	1486568	24.371	ng	100
89) 1,2,3-Trimethylbenzene	20.35	105	1149185	24.778	ng	95
90) 1,2-Dichlorobenzene	20.47	146	750207	26.147	ng	100
91) d-Limonene	20.47	68	434969	26.719	ng	97
92) 1,2-Dibromo-3-Chloropr...	20.85	157	282742	27.538	ng	85
93) n-Undecane	21.17	57	613692	29.713	ng	98
94) 1,2,4-Trichlorobenzene	21.97	180	604264	30.156	ng	100
95) Naphthalene	22.07	128	1767201	32.400	ng	100
96) n-Dodecane	22.06	57	576084	32.148	ng	99
97) Hexachlorobutadiene	22.38	225	377819	24.397	ng	100
98) Cyclohexanone	18.12	55	368619	23.221	ng	94
99) tert-Butylbenzene	19.96	119	1169859	24.579	ng	99
100) n-Butylbenzene	20.72	91	1228352	25.948	ng	98

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231909.D
Acq On : 23 Jan 2019 9:46
Sample : 25ng TO-15 ICAL Std
Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
Operator: WA
Inst : MS13

Quant Time: Jan 23 10:31:48 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 10:31:09 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_01\23\01231910.D
 Acq On : 23 Jan 2019 10:20
 Sample : 50ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:54:36 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

1/23/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.98	130	129676	12.500	ng	0.00
37) 1,4-Difluorobenzene (IS2)	13.10	114	554115	12.500	ng	0.00
56) Chlorobenzene-d5 (IS3)	17.42	82	217596	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.83	65	136294	11.975	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	95.76%
57) Toluene-d8 (SS2)	15.55	98	586897	13.076	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	104.64%
73) Bromofluorobenzene (SS3)	18.83	174	211351	11.427	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	91.44%

Target Compounds

						Qvalue
2) Propene	3.98	42	550963	45.236	ng	96
3) Dichlorodifluoromethan...	4.13	85	958014	44.038	ng	99
4) Chloromethane	4.41	50	749838	44.966	ng	100
5) 1,2-Dichloro-1,1,2,2-t...	4.67	135	619578	44.407	ng	99
6) Vinyl Chloride	4.82	62	774206	47.762	ng	100
7) 1,3-Butadiene	5.08	54	615017	51.861	ng	93
8) Bromomethane	5.51	94	518382	46.340	ng	99
9) Chloroethane	5.83	64	400255	45.550	ng	100
10) Ethanol	6.24	45	1820404	222.486	ng	97
11) Acetonitrile	6.47	41	995157	48.855	ng	98
12) Acrolein	6.65	56	349588	49.928	ng	97
13) Acetone	6.85	58	1938708	210.711	ng	# 80
14) Trichlorofluoromethane	7.07	101	813055	44.506	ng	99
15) 2-Propanol (Isopropanol)	7.36	45	2536963	92.311	ng	94
16) Acrylonitrile	7.60	53	739164	56.813	ng	98
17) 1,1-Dichloroethene	8.03	96	589118	48.342	ng	# 84
18) 2-Methyl-2-Propanol (t...	8.22	59	2597364	93.906	ng	96
19) Methylene Chloride	8.26	84	587709	48.411	ng	94
20) 3-Chloro-1-propene (Al...	8.42	41	780458	50.409	ng	98
21) Trichlorotrifluoroethane	8.67	151	589650	45.859	ng	93
22) Carbon Disulfide	8.51	76	2099717	46.519	ng	99
23) trans-1,2-Dichloroethene	9.53	61	747632	52.161	ng	93
24) 1,1-Dichloroethane	9.78	63	899332	44.833	ng	100
25) Methyl tert-Butyl Ether	9.88	73	1619009	46.810	ng	98
26) Vinyl Acetate	10.05	86	716439	246.912	ng	# 84
27) 2-Butanone (MEK)	10.29	72	403390	48.927	ng	# 88
28) cis-1,2-Dichloroethene	10.80	61	705591	47.597	ng	92
29) Diisopropyl Ether	11.10	87	561552	47.440	ng	# 75
30) Ethyl Acetate	11.11	61	395905	100.662	ng	95
31) n-Hexane	11.08	57	828858	42.286	ng	# 100
32) Chloroform	11.15	83	900170	47.128	ng	100
34) Tetrahydrofuran (THF)	11.55	72	391881	45.354	ng	99
35) Ethyl tert-Butyl Ether	11.70	87	685968	47.249	ng	95
36) 1,2-Dichloroethane	11.95	62	578293	47.302	ng	99
38) 1,1,1-Trichloroethane	12.23	97	786422	48.609	ng	97
39) Isopropyl Acetate	12.66	61	684341	91.504	ng	96
40) 1-Butanol	12.69	56	1041001	100.029	ng	# 75
41) Benzene	12.70	78	2183231	43.651	ng	99
42) Carbon Tetrachloride	12.86	117	714110	48.239	ng	100
43) Cyclohexane	12.99	84	1758366	89.267	ng	99
44) tert-Amyl Methyl Ether	13.35	73	1571599	47.598	ng	97
45) 1,2-Dichloropropane	13.56	63	547191	47.242	ng	100
46) Bromodichloromethane	13.74	83	706028	50.344	ng	99
47) Trichloroethene	13.80	130	704224	48.531	ng	99
48) 1,4-Dioxane	13.78	88	499338	49.502	ng	99
49) 2,2,4-Trimethylpentane...	13.87	57	2175437	43.264	ng	90
50) Methyl Methacrylate	14.01	100	528051	104.313	ng	93

Data File : I:\MS13\DATA\2019_01\23\01231910.D
 Acq On : 23 Jan 2019 10:20
 Sample : 50ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 10:54:36 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

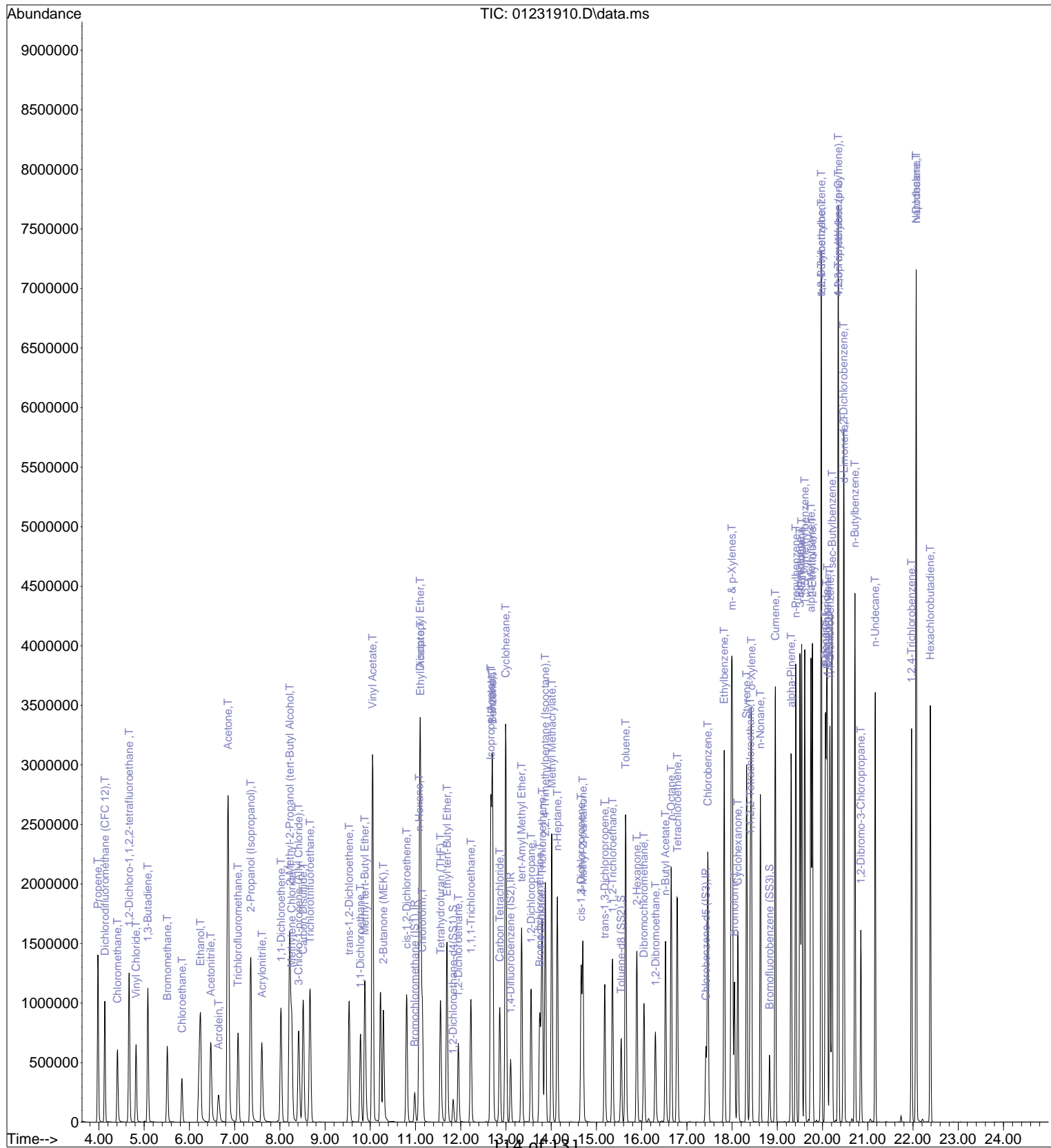
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.14	71	566594	46.024	ng	99
52) cis-1,3-Dichloropropene	14.66	75	963259	53.626	ng	100
53) 4-Methyl-2-pentanone	14.70	58	512957	49.224	ng	92
54) trans-1,3-Dichloropropene	15.18	75	805902	53.295	ng	99
55) 1,1,2-Trichloroethane	15.36	97	582332	48.870	ng	98
58) Toluene	15.65	91	2363408	46.614	ng	98
59) 2-Hexanone	15.90	43	1097984	53.877	ng	93
60) Dibromochloromethane	16.05	129	714941	56.034	ng	100
61) 1,2-Dibromoethane	16.31	107	664969	56.514	ng	100
62) n-Butyl Acetate	16.53	43	1241427	52.766	ng	94
63) n-Octane	16.65	57	456775	48.703	ng	98
64) Tetrachloroethene	16.79	166	794176	49.849	ng	99
65) Chlorobenzene	17.46	112	1673750	48.899	ng	99
66) Ethylbenzene	17.82	91	2601402	47.157	ng	99
67) m- & p-Xylenes	18.00	91	4065083	96.120	ng	98
68) Bromoform	18.05	173	680568	55.080	ng	99
69) Styrene	18.32	104	1804766	54.201	ng	97
70) o-Xylene	18.43	91	2029602	47.852	ng	96
71) n-Nonane	18.63	43	964989	45.649	ng	99
72) 1,1,2,2-Tetrachloroethane	18.40	83	1011065	49.647	ng	100
74) Cumene	18.96	105	2694136	46.644	ng	98
75) alpha-Pinene	19.30	93	1391956	49.562	ng	94
76) n-Propylbenzene	19.41	91	3123059	47.628	ng	97
77) 3-Ethyltoluene	19.50	105	2810434	49.837	ng	99
78) 4-Ethyltoluene	19.54	105	2506081	45.820	ng	98
79) 1,3,5-Trimethylbenzene	19.61	105	2241971	46.711	ng	96
80) alpha-Methylstyrene	19.74	118	1355760	57.003	ng	92
81) 2-Ethyltoluene	19.78	105	2690410	47.062	ng	97
82) 1,2,4-Trimethylbenzene	19.97	105	2226342	47.884	ng	96
83) n-Decane	20.06	57	1101364	48.012	ng	99
84) Benzyl Chloride	20.08	91	1985624	58.420	ng	96
85) 1,3-Dichlorobenzene	20.11	146	1551652	52.330	ng	99
86) 1,4-Dichlorobenzene	20.16	146	1571568	50.949	ng	100
87) sec-Butylbenzene	20.21	105	3056416	46.620	ng	98
88) 4-Isopropyltoluene (p-...	20.35	119	2789874	45.230	ng	98
89) 1,2,3-Trimethylbenzene	20.35	105	2205069	47.017	ng	96
90) 1,2-Dichlorobenzene	20.47	146	1470738	50.692	ng	99
91) d-Limonene	20.48	68	824220	50.068	ng	99
92) 1,2-Dibromo-3-Chloropr...	20.85	157	567508	54.662	ng	84
93) n-Undecane	21.17	57	1155291	55.316	ng	100
94) 1,2,4-Trichlorobenzene	21.97	180	1205152	59.477	ng	100
95) Naphthalene	22.07	128	3390515	61.472	ng	99
96) n-Dodecane	22.07	57	1057288	58.347	ng	97
97) Hexachlorobutadiene	22.38	225	770156	49.181	ng	100
98) Cyclohexanone	18.13	55	729990	45.476	ng	93
99) tert-Butylbenzene	19.97	119	2220306	46.132	ng	99
100) n-Butylbenzene	20.72	91	2361092	49.324	ng	97

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231910.D
Acq On : 23 Jan 2019 10:20
Sample : 50ng TO-15 ICAL Std
Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
Operator: WA
Inst : MS13

Quant Time: Jan 23 10:54:36 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 10:31:09 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_01\23\01231911.D
 Acq On : 23 Jan 2019 10:53
 Sample : 100ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 11:18:52 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

10A 1/23/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.99	130	130622	12.500	ng	0.01
37) 1,4-Difluorobenzene (IS2)	13.11	114	567955	12.500	ng	0.00
56) Chlorobenzene-d5 (IS3)	17.42	82	224304	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.84	65	138527	12.083	ng	0.01
Spiked Amount	12.500	Range	70 - 130	Recovery	=	96.64%
57) Toluene-d8 (SS2)	15.55	98	593656	12.831	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	102.64%
73) Bromofluorobenzene (SS3)	18.83	174	216071	11.333	ng	0.00
Spiked Amount	12.500	Range	70 - 130	Recovery	=	90.64%

Target Compounds

						Qvalue
2) Propene	3.98	42	1235949	100.742	ng	96
3) Dichlorodifluoromethan...	4.14	85	1857522	84.768	ng	99
4) Chloromethane	4.42	50	1265365	75.331	ng	99
5) 1,2-Dichloro-1,1,2,2-t...	4.67	135	1235486	87.909	ng	100
6) Vinyl Chloride	4.83	62	1531085	93.771	ng	100
7) 1,3-Butadiene	5.09	54	1199165	100.388	ng	93
8) Bromomethane	5.52	94	1034074	91.771	ng	99
9) Chloroethane	5.85	64	789553	89.203	ng	99
10) Ethanol	6.29	45	3485746	422.935	ng	97
11) Acetonitrile	6.50	41	1992641	97.116	ng	97
12) Acrolein	6.66	56	702088	99.545	ng	98
13) Acetone	6.88	58	3570661	385.272	ng	# 77
14) Trichlorofluoromethane	7.09	101	1626310	88.379	ng	99
15) 2-Propanol (Isopropanol)	7.39	45	4163038	150.381	ng	94
16) Acrylonitrile	7.62	53	1484683	113.287	ng	98
17) 1,1-Dichloroethene	8.03	96	1169842	95.299	ng	# 83
18) 2-Methyl-2-Propanol (t...	8.25	59	4887273	175.417	ng	97
19) Methylene Chloride	8.28	84	1136520	92.941	ng	89
20) 3-Chloro-1-propene (Al...	8.42	41	1535540	98.462	ng	99
21) Trichlorotrifluoroethane	8.67	151	1181859	91.252	ng	93
22) Carbon Disulfide	8.53	76	4103868	90.262	ng	99
23) trans-1,2-Dichloroethene	9.54	61	1468459	101.711	ng	92
24) 1,1-Dichloroethane	9.79	63	1755345	86.873	ng	99
25) Methyl tert-Butyl Ether	9.89	73	3168796	90.956	ng	98
26) Vinyl Acetate	10.07	86	1356871	464.242	ng	# 64
27) 2-Butanone (MEK)	10.30	72	797486	96.026	ng	# 86
28) cis-1,2-Dichloroethene	10.81	61	1387029	92.887	ng	90
29) Diisopropyl Ether	11.11	87	1076818	90.311	ng	# 79
30) Ethyl Acetate	11.13	61	717888	181.208	ng	91
31) n-Hexane	11.09	57	1530523	77.518	ng	# 100
32) Chloroform	11.16	83	1772093	92.105	ng	99
34) Tetrahydrofuran (THF)	11.55	72	775181	89.065	ng	98
35) Ethyl tert-Butyl Ether	11.70	87	1356833	92.781	ng	93
36) 1,2-Dichloroethane	11.95	62	1139631	92.541	ng	99
38) 1,1,1-Trichloroethane	12.23	97	1548462	93.379	ng	97
39) Isopropyl Acetate	12.67	61	1268000	165.413	ng	99
40) 1-Butanol	12.71	56	1948054	182.625	ng	# 76
41) Benzene	12.71	78	4015436	78.328	ng	100
42) Carbon Tetrachloride	12.87	117	1421053	93.655	ng	100
43) Cyclohexane	13.00	84	3299631	163.431	ng	98
44) tert-Amyl Methyl Ether	13.35	73	3048509	90.079	ng	97
45) 1,2-Dichloropropane	13.56	63	1063738	89.600	ng	99
46) Bromodichloromethane	13.75	83	1392144	96.849	ng	100
47) Trichloroethene	13.81	130	1389814	93.443	ng	100
48) 1,4-Dioxane	13.78	88	979074	94.696	ng	97
49) 2,2,4-Trimethylpentane...	13.88	57	4083483	79.231	ng	91
50) Methyl Methacrylate	14.02	100	1029562	198.428	ng	90

Data File : I:\MS13\DATA\2019_01\23\01231911.D
 Acq On : 23 Jan 2019 10:53
 Sample : 100ng TO-15 ICAL Std
 Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 11:18:52 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 10:31:09 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

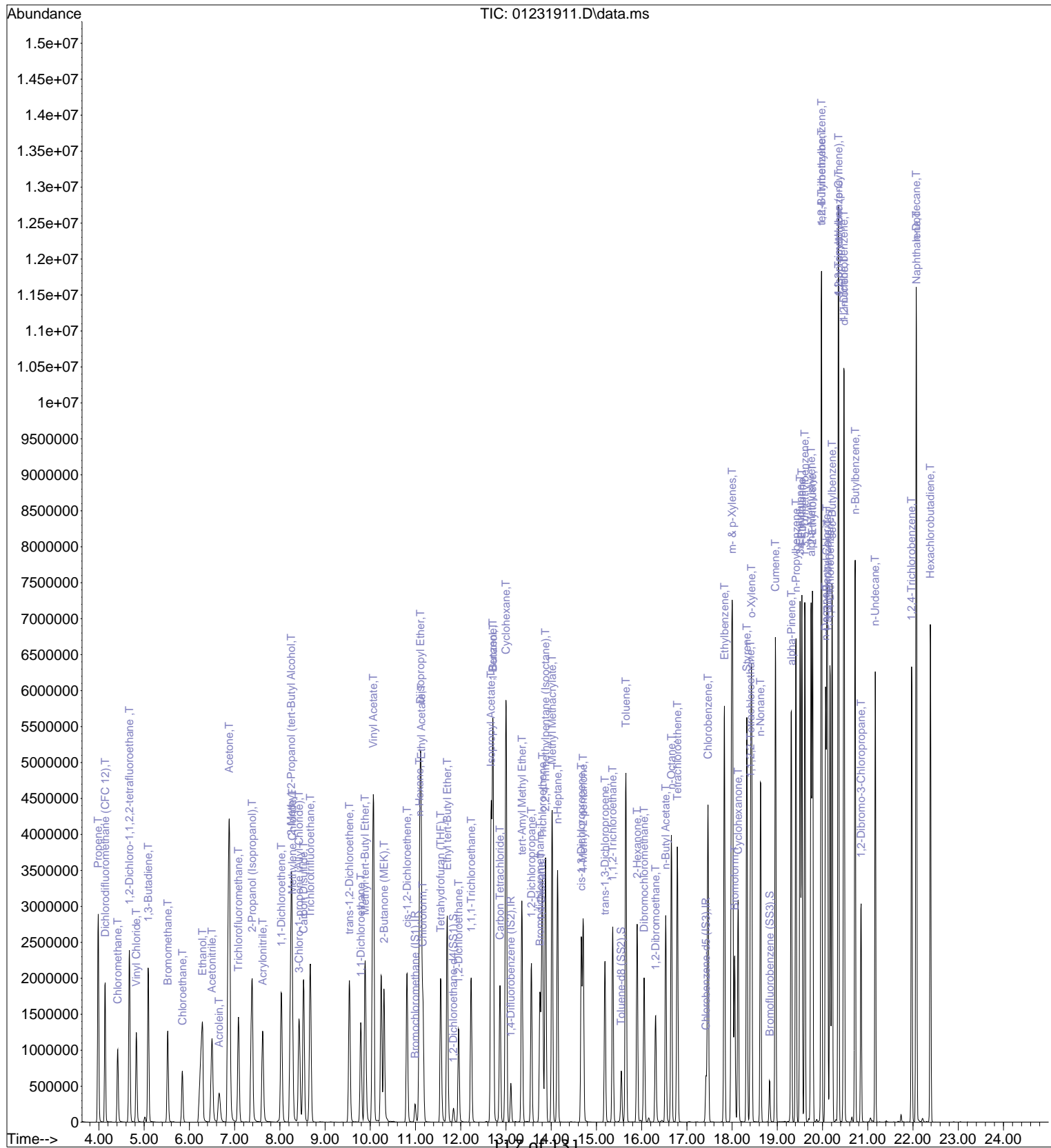
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.14	71	1090107	86.391	ng	99
52) cis-1,3-Dichloropropene	14.67	75	1884084	102.334	ng	99
53) 4-Methyl-2-pentanone	14.71	58	992068	92.881	ng	89
54) trans-1,3-Dichloropropene	15.19	75	1594371	102.868	ng	99
55) 1,1,2-Trichloroethane	15.36	97	1147869	93.984	ng	98
58) Toluene	15.65	91	4475365	85.628	ng	97
59) 2-Hexanone	15.90	43	2095409	99.745	ng	92
60) Dibromochloromethane	16.06	129	1427371	108.526	ng	100
61) 1,2-Dibromoethane	16.31	107	1320897	108.902	ng	100
62) n-Butyl Acetate	16.53	43	2358975	97.268	ng	93
63) n-Octane	16.66	57	861202	89.078	ng	99
64) Tetrachloroethene	16.79	166	1586409	96.598	ng	98
65) Chlorobenzene	17.47	112	3227442	91.470	ng	98
66) Ethylbenzene	17.83	91	4917151	86.470	ng	99
67) m- & p-Xylenes	18.00	91	7590877	174.120	ng	98
68) Bromoform	18.06	173	1381752	108.485	ng	99
69) Styrene	18.33	104	3469181	101.072	ng	97
70) o-Xylene	18.43	91	3850302	88.064	ng	95
71) n-Nonane	18.63	43	1749023	80.263	ng	96
72) 1,1,2,2-Tetrachloroethane	18.41	83	1936312	92.237	ng	100
74) Cumene	18.96	105	5008946	84.128	ng	96
75) alpha-Pinene	19.31	93	2645675	91.386	ng	95
76) n-Propylbenzene	19.41	91	5716632	84.573	ng	95
77) 3-Ethyltoluene	19.50	105	4994193	85.913	ng	91
78) 4-Ethyltoluene	19.54	105	4883267	86.612	ng	98
79) 1,3,5-Trimethylbenzene	19.61	105	4250628	85.912	ng	95
80) alpha-Methylstyrene	19.75	118	2590177	105.647	ng	90
81) 2-Ethyltoluene	19.78	105	5013066	85.068	ng	96
82) 1,2,4-Trimethylbenzene	19.98	105	4028708	84.058	ng	97
83) n-Decane	20.07	57	1961905	82.968	ng	96
84) Benzyl Chloride	20.09	91	3781785	107.939	ng	94
85) 1,3-Dichlorobenzene	20.11	146	3017833	98.733	ng	99
86) 1,4-Dichlorobenzene	20.16	146	3041985	95.669	ng	99
87) sec-Butylbenzene	20.21	105	5588775	82.698	ng	95
88) 4-Isopropyltoluene (p-...	20.35	119	4811990	75.680	ng	95
89) 1,2,3-Trimethylbenzene	20.35	105	3957709	81.864	ng	98
90) 1,2-Dichlorobenzene	20.47	146	2784546	93.105	ng	99
91) d-Limonene	20.48	68	1448812	85.378	ng	96
92) 1,2-Dibromo-3-Chloropr...	20.85	157	1138300	106.361	ng	83
93) n-Undecane	21.17	57	2071611	96.224	ng	97
94) 1,2,4-Trichlorobenzene	21.97	180	2394698	114.649	ng	99
95) Naphthalene	22.08	128	6034979	106.146	ng	98
96) n-Dodecane	22.07	57	1761991	94.328	ng	93
97) Hexachlorobutadiene	22.38	225	1567243	97.089	ng	100
98) Cyclohexanone	18.13	55	1413902	85.447	ng	92
99) tert-Butylbenzene	19.98	119	3957503	79.767	ng	97
100) n-Butylbenzene	20.72	91	4337953	87.911	ng	95

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231911.D
Acq On : 23 Jan 2019 10:53
Sample : 100ng TO-15 ICAL Std
Misc : S31-01221908/S31-01161903 (2/14)

Vial: 16
Operator: WA
Inst : MS13

Quant Time: Jan 23 11:18:52 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 10:31:09 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



Data File : I:\MS13\DATA\2019_01\23\01231912.D
 Acq On : 23 Jan 2019 11:31
 Sample : 25ng TO-15 ICV Std
 Misc : S31-01221908/S31-12271803 (1/25)

Vial: 13
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 11:53:59 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

1/23/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.97	130	131810	12.500	ng	-0.02
37) 1,4-Difluorobenzene (IS2)	13.10	114	562964	12.500	ng	-0.01
56) Chlorobenzene-d5 (IS3)	17.42	82	224891	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.83	65	137705	12.253	ng	-0.01
Spiked Amount	12.500	Range 70 - 130	Recovery	=	98.00%	
57) Toluene-d8 (SS2)	15.54	98	595617	12.368	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	98.96%	
73) Bromofluorobenzene (SS3)	18.83	174	216386	12.479	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	99.84%	

Target Compounds

						Qvalue
2) Propene	3.97	42	290238	25.307	ng	95
3) Dichlorodifluoromethan...	4.13	85	493642	25.193	ng	100
4) Chloromethane	4.41	50	413687	27.195	ng	100
5) 1,2-Dichloro-1,1,2,2-t...	4.66	135	326224	25.834	ng	100
6) Vinyl Chloride	4.82	62	414491	28.421	ng	100
7) 1,3-Butadiene	5.08	54	320393	29.205	ng	92
8) Bromomethane	5.50	94	282739	27.020	ng	99
9) Chloroethane	5.83	64	213940	26.654	ng	99
10) Ethanol	6.22	45	1011763	129.405	ng	98
11) Acetonitrile	6.45	41	513663	27.151	ng	97
12) Acrolein	6.63	56	175696	27.188	ng	98
13) Acetone	6.84	58	1042912	125.541	ng	# 81
14) Trichlorofluoromethane	7.07	101	417709	25.440	ng	99
15) 2-Propanol (Isopropanol)	7.34	45	1342639	53.071	ng	93
16) Acrylonitrile	7.59	53	378092	31.217	ng	98
17) 1,1-Dichloroethene	8.02	96	301396	27.502	ng	# 84
18) 2-Methyl-2-Propanol (t...	8.20	59	1312557	52.782	ng	96
19) Methylene Chloride	8.25	84	302373	28.711	ng	94
20) 3-Chloro-1-propene (Al...	8.41	41	415896	29.468	ng	97
21) Trichlorotrifluoroethane	8.66	151	301621	26.034	ng	93
22) Carbon Disulfide	8.51	76	1087556	25.282	ng	99
23) trans-1,2-Dichloroethene	9.52	61	379244	29.572	ng	93
24) 1,1-Dichloroethane	9.78	63	478565	26.232	ng	100
25) Methyl tert-Butyl Ether	9.87	73	825622	26.494	ng	98
26) Vinyl Acetate	10.04	86	372308	139.318	ng	96
27) 2-Butanone (MEK)	10.28	72	207861	27.296	ng	# 90
28) cis-1,2-Dichloroethene	10.80	61	359087	27.074	ng	92
29) Diisopropyl Ether	11.10	87	294686	27.567	ng	# 73
30) Ethyl Acetate	11.10	61	210958	55.937	ng	97
31) n-Hexane	11.08	57	447408	25.559	ng	# 100
32) Chloroform	11.14	83	456008	26.591	ng	99
34) Tetrahydrofuran (THF)	11.55	72	200595	25.956	ng	99
35) Ethyl tert-Butyl Ether	11.69	87	349375	26.320	ng	96
36) 1,2-Dichloroethane	11.94	62	294048	27.012	ng	99
38) 1,1,1-Trichloroethane	12.22	97	391363	26.333	ng	97
39) Isopropyl Acetate	12.66	61	359720	50.382	ng	93
40) 1-Butanol	12.68	56	538707	55.408	ng	# 73
41) Benzene	12.70	78	1163976	25.038	ng	99
42) Carbon Tetrachloride	12.86	117	362977	26.541	ng	100
43) Cyclohexane	12.99	84	917901	50.250	ng	98
44) tert-Amyl Methyl Ether	13.34	73	800206	26.373	ng	97
45) 1,2-Dichloropropane	13.55	63	280406	26.372	ng	100
46) Bromodichloromethane	13.74	83	360643	28.023	ng	99
47) Trichloroethene	13.80	130	359395	26.746	ng	100
48) 1,4-Dioxane	13.77	88	256432	27.377	ng	99
49) 2,2,4-Trimethylpentane...	13.86	57	1140295	24.936	ng	90
50) Methyl Methacrylate	14.01	100	270630	60.210	ng	95

Data File : I:\MS13\DATA\2019_01\23\01231912.D
 Acq On : 23 Jan 2019 11:31
 Sample : 25ng TO-15 ICV Std
 Misc : S31-01221908/S31-12271803 (1/25)

Vial: 13
 Operator: WA
 Inst : MS13

Quant Time: Jan 23 11:53:59 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

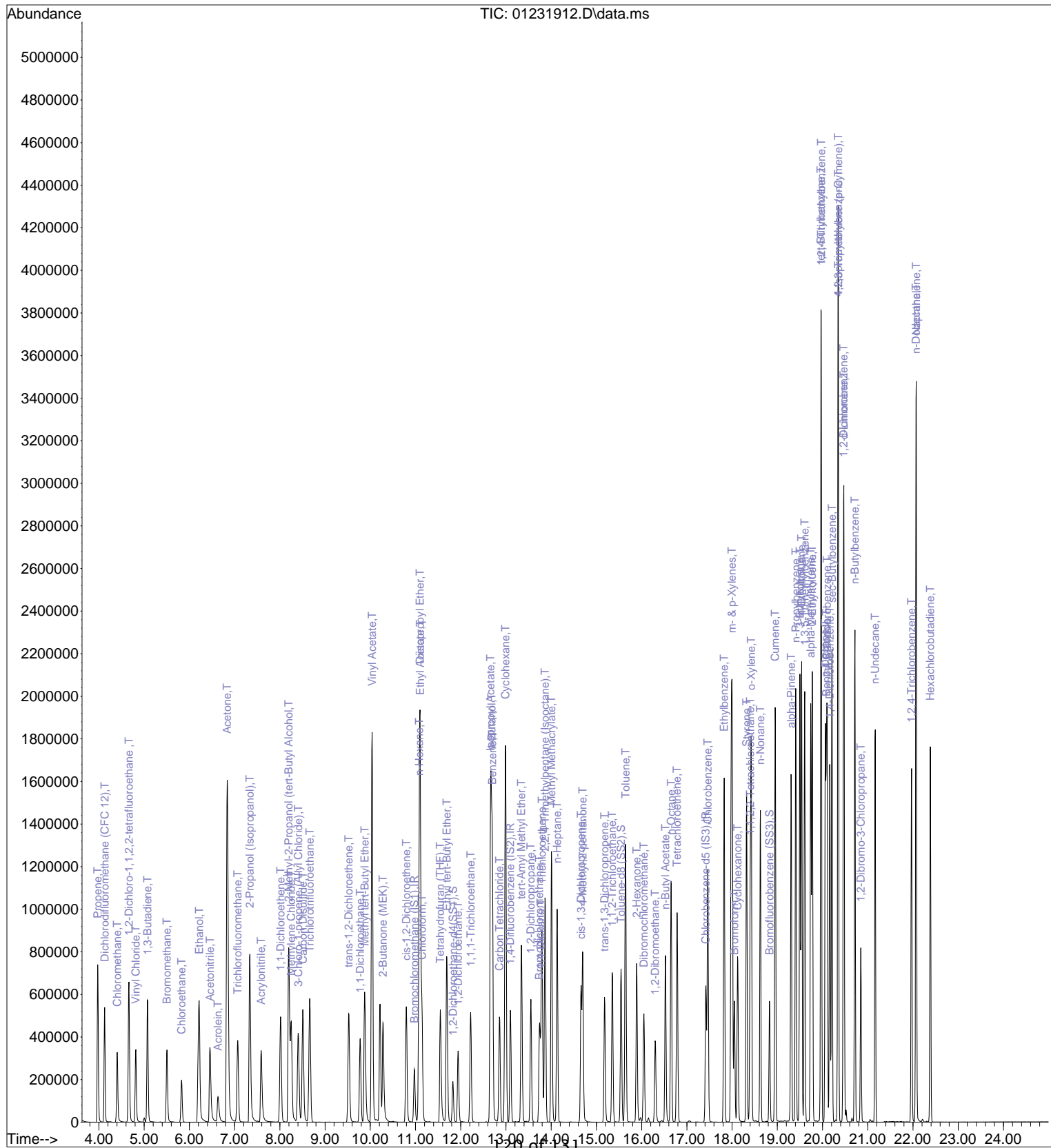
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.13	71	294825	26.312	ng	99
52) cis-1,3-Dichloropropene	14.66	75	466898	27.704	ng	100
53) 4-Methyl-2-pentanone	14.69	58	263484	27.506	ng	92
54) trans-1,3-Dichloropropene	15.18	75	409912	28.951	ng	100
55) 1,1,2-Trichloroethane	15.35	97	296327	27.429	ng	98
58) Toluene	15.65	91	1214741	23.730	ng	98
59) 2-Hexanone	15.89	43	564646	28.070	ng	93
60) Dibromochloromethane	16.05	129	360060	28.871	ng	100
61) 1,2-Dibromoethane	16.30	107	337840	29.150	ng	100
62) n-Butyl Acetate	16.53	43	638988	27.373	ng	94
63) n-Octane	16.65	57	238507	25.357	ng	97
64) Tetrachloroethene	16.78	166	403400	25.748	ng	99
65) Chlorobenzene	17.46	112	864555	25.463	ng	100
66) Ethylbenzene	17.82	91	1360808	24.698	ng	99
67) m- & p-Xylenes	18.00	91	2113341	49.688	ng	98
68) Bromoform	18.05	173	336959	29.577	ng	100
69) Styrene	18.32	104	919763	27.664	ng	96
70) o-Xylene	18.42	91	1055135	24.892	ng	96
71) n-Nonane	18.63	43	520296	25.111	ng	100
72) 1,1,2,2-Tetrachloroethane	18.40	83	519389	26.149	ng	99
74) Cumene	18.95	105	1421080	24.569	ng	99
75) alpha-Pinene	19.30	93	717581	25.916	ng	94
76) n-Propylbenzene	19.41	91	1653750	25.491	ng	98
77) 3-Ethyltoluene	19.50	105	1365282	24.834	ng	94
78) 4-Ethyltoluene	19.54	105	1440290	26.218	ng	96
79) 1,3,5-Trimethylbenzene	19.61	105	1160035	24.059	ng	97
80) alpha-Methylstyrene	19.74	118	687603	28.890	ng	92
81) 2-Ethyltoluene	19.77	105	1401382	24.495	ng	98
82) 1,2,4-Trimethylbenzene	19.97	105	1175021	25.121	ng	95
83) n-Decane	20.06	57	586875	25.783	ng	99
84) Benzyl Chloride	20.08	91	1001220	29.269	ng	98
85) 1,3-Dichlorobenzene	20.10	146	788851	26.876	ng	100
86) 1,4-Dichlorobenzene	20.16	146	804355	26.104	ng	100
87) sec-Butylbenzene	20.21	105	1609639	24.789	ng	99
88) 4-Isopropyltoluene (p-...	20.35	119	1532048	25.277	ng	100
89) 1,2,3-Trimethylbenzene	20.35	105	1204084	26.022	ng	96
90) 1,2-Dichlorobenzene	20.47	146	758450	26.474	ng	100
91) d-Limonene	20.47	68	444284	27.213	ng	96
92) 1,2-Dibromo-3-Chloropr...	20.85	157	290539	28.052	ng	85
93) n-Undecane	21.17	57	595406	28.366	ng	98
94) 1,2,4-Trichlorobenzene	21.97	180	613462	28.126	ng	100
95) Naphthalene	22.07	128	1732271	29.113	ng	100
96) n-Dodecane	22.06	57	483693	24.640	ng	99
97) Hexachlorobutadiene	22.38	225	384807	23.731	ng	100
98) Cyclohexanone	18.12	55	363308	23.435	ng	94
99) tert-Butylbenzene	19.96	119	1190104	24.957	ng	99
100) n-Butylbenzene	20.72	91	1235644	25.970	ng	98

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_01\23\01231912.D
Acq On : 23 Jan 2019 11:31
Sample : 25ng TO-15 ICV Std
Misc : S31-01221908/S31-12271803 (1/25)

Vial: 13
Operator: WA
Inst : MS13

Quant Time: Jan 23 11:53:59 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 11:21:29 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



120 of 131

Initial Calibration Verification/LABORATORY CONTROL SAMPLE CHECK SHEET

Data File Name: 01231912.D

Acq. Method File: TO15.M

IDA 1/23/19

Data File Path: I:\MS13\DATA\2019_01\23\

Sample Name: 25ng TO-15 ICV Std

Operator: WA

Misc Info: S31-01221908/S31-12271803 (

Date Acquired: 1/23/2019

11:31

Instrument Name: MS13

#	Compound Name	Ret. Time	Amt. (ng)	Spike Amt.(ng)	% Rec.	Lower Limit	Upper Limit	* OR Fail	ICV/AZ 70-130%
2)	Propene	3.97	25.3	26.43	96	53	112	*	*
3)	Dichlorodifluoromethane (CFC 12)	4.13	25.2	26.30	96	62	103	*	*
4)	Chloromethane	4.41	27.2	26.38	103	51	121	*	*
5)	1,2-Dichloro-1,1,2,2-tetrafluoroethane	4.66	25.8	26.38	98	56	111	*	*
6)	Vinyl Chloride	4.82	28.4	26.73	106	57	117	*	*
7)	1,3-Butadiene	5.08	29.2	26.28	111	53	134	*	*
8)	Bromomethane	5.50	27.0	26.48	102	65	110	*	*
9)	Chloroethane	5.83	26.7	26.75	100	64	111	*	*
10)	Ethanol	6.22	129	128.10	101	57	124	*	*
11)	Acetonitrile	6.45	27.2	25.73	106	57	126	*	*
12)	Acrolein	6.63	27.2	25.63	106	62	121	*	*
13)	Acetone	6.84	126	132.33	95	60	113	*	*
14)	Trichlorofluoromethane	7.07	25.4	26.40	96	63	104	*	*
15)	2-Propanol (Isopropanol)	7.34	53.1	51.60	103	60	124	*	*
16)	Acrylonitrile	7.59	31.2	25.88	121	66	125	*	*
17)	1,1-Dichloroethene	8.02	27.5	27.23	101	68	107	*	*
18)	2-Methyl-2-Propanol (tert-Butyl Alcohol)	8.20	52.8	54.25	97	64	114	*	*
19)	Methylene Chloride	8.25	28.7	27.08	106	66	105	FAIL	*
20)	3-Chloro-1-propene (Allyl Chloride)	8.41	29.5	27.00	109	63	127	*	*
21)	Trichlorotrifluoroethane	8.66	26.0	26.95	96	59	109	*	*
22)	Carbon Disulfide	8.51	25.3	27.20	93	67	109	*	*
23)	trans-1,2-Dichloroethene	9.52	29.6	26.73	111	70	115	*	*
24)	1,1-Dichloroethane	9.78	26.2	26.95	97	66	106	*	*
25)	Methyl tert-Butyl Ether	9.87	26.5	26.80	99	67	109	*	*
26)	Vinyl Acetate	10.04	139	133.03	104	68	136	*	*
27)	2-Butanone (MEK)	10.28	27.3	25.95	105	71	116	*	*
28)	cis-1,2-Dichloroethene	10.80	27.1	26.35	103	67	110	*	*
29)	Diisopropyl Ether	11.10	27.6	27.18	102	62	109	*	*
30)	Ethyl Acetate	11.10	55.9	54.45	103	64	127	*	*
31)	n-Hexane	11.08	25.6	26.95	95	60	115	*	*
32)	Chloroform	11.14	26.6	27.08	98	66	105	*	*
34)	Tetrahydrofuran (THF)	11.55	26.0	27.00	96	65	110	*	*
35)	Ethyl tert-Butyl Ether	11.69	26.3	26.80	98	69	109	*	*
36)	1,2-Dichloroethane	11.94	27.0	26.85	101	60	110	*	*
38)	1,1,1-Trichloroethane	12.22	26.3	26.90	98	64	108	*	*
39)	Isopropyl Acetate	12.66	50.4	51.55	98	66	119	*	*
40)	1-Butanol	12.68	55.4	51.58	107	54	143	*	*
41)	Benzene	12.70	25.0	26.38	95	67	106	*	*
42)	Carbon Tetrachloride	12.86	26.5	26.45	100	64	112	*	*
43)	Cyclohexane	12.99	50.2	52.05	96	67	110	*	*
44)	tert-Amyl Methyl Ether	13.34	26.4	27.08	98	68	112	*	*
45)	1,2-Dichloropropane	13.55	26.4	26.98	98	66	112	*	*
46)	Bromodichloromethane	13.74	28.0	26.83	104	67	113	*	*
47)	Trichloroethene	13.80	26.7	26.68	100	66	108	*	*
48)	1,4-Dioxane	13.77	27.4	26.73	103	70	116	*	*
49)	2,2,4-Trimethylpentane (Isooctane)	13.86	24.9	26.65	93	64	113	*	*

Initial Calibration Verification/LABORATORY CONTROL SAMPLE CHECK SHEET

Data File Name: 01231912.D

TO15.M

Data File Path: I:\MS13\DATA\2019_01\23\

Sample Name: 25ng TO-15 ICV Std

Operator: WA

Misc Info: S31-01221908/S31-12271803 (

Date Acquired: 1/23/2019

11:31

Instrument Name: MS13

#	Compound Name	Ret. Time	Amt. (ng)	Spike Amt.(ng)	% Rec.	Lower Limit	Upper Limit	* OR Fail	ICV/AZ 70-130%
50)	Methyl Methacrylate	14.01	60.2	53.88	112	73	118	*	*
51)	n-Heptane	14.13	26.3	26.90	98	66	110	*	*
52)	cis-1,3-Dichloropropene	14.66	27.7	26.78	103	75	120	*	*
53)	4-Methyl-2-pentanone	14.69	27.5	26.15	105	65	124	*	*
54)	trans-1,3-Dichloropropene	15.18	29.0	26.60	109	77	123	*	*
55)	1,1,2-Trichloroethane	15.35	27.4	26.85	102	68	112	*	*
58)	Toluene	15.65	23.7	26.50	89	62	111	*	*
59)	2-Hexanone	15.89	28.1	26.78	105	59	128	*	*
60)	Dibromochloromethane	16.05	28.9	26.60	109	67	123	*	*
61)	1,2-Dibromoethane	16.30	29.2	27.03	108	66	122	*	*
62)	n-Butyl Acetate	16.53	27.4	27.35	100	64	128	*	*
63)	n-Octane	16.65	25.4	27.13	94	65	114	*	*
64)	Tetrachloroethene	16.78	25.7	26.60	97	55	120	*	*
65)	Chlorobenzene	17.46	25.5	26.83	95	61	114	*	*
66)	Ethylbenzene	17.82	24.7	26.53	93	64	113	*	*
67)	m- & p-Xylenes	18.00	49.7	53.28	93	64	114	*	*
68)	Bromoform	18.05	29.6	26.68	111	65	132	*	*
69)	Styrene	18.32	27.7	26.50	105	67	124	*	*
70)	o-Xylene	18.42	24.9	26.75	93	65	114	*	*
71)	n-Nonane	18.63	25.1	26.83	94	64	117	*	*
72)	1,1,2,2-Tetrachloroethane	18.40	26.1	26.80	97	66	119	*	*
74)	Cumene	18.95	24.6	26.80	92	61	116	*	*
75)	alpha-Pinene	19.30	25.9	26.40	98	65	120	*	*
76)	n-Propylbenzene	19.41	25.5	27.23	94	63	117	*	*
77)	3-Ethyltoluene	19.50	24.8	26.83	92	60	117	*	*
78)	4-Ethyltoluene	19.54	26.2	26.80	98	63	124	*	*
79)	1,3,5-Trimethylbenzene	19.61	24.1	26.73	90	60	117	*	*
80)	alpha-Methylstyrene	19.74	28.9	26.75	108	64	131	*	*
81)	2-Ethyltoluene	19.77	24.5	27.08	90	62	116	*	*
82)	1,2,4-Trimethylbenzene	19.97	25.1	26.90	93	61	122	*	*
83)	n-Decane	20.06	25.8	26.88	96	67	120	*	*
84)	Benzyl Chloride	20.08	29.3	27.08	108	77	142	*	*
85)	1,3-Dichlorobenzene	20.10	26.9	26.98	100	61	125	*	*
86)	1,4-Dichlorobenzene	20.16	26.1	27.00	97	59	123	*	*
87)	sec-Butylbenzene	20.21	24.8	26.55	93	62	117	*	*
88)	4-Isopropyltoluene (p-Cymene)	20.35	25.3	26.95	94	58	122	*	*
89)	1,2,3-Trimethylbenzene	20.35	26.0	26.95	96	62	124	*	*
90)	1,2-Dichlorobenzene	20.47	26.5	26.95	98	61	126	*	*
91)	d-Limonene	20.47	27.2	26.38	103	66	124	*	*
92)	1,2-Dibromo-3-Chloropropane	20.85	28.1	26.15	107	67	138	*	*
93)	n-Undecane	21.17	28.4	27.03	105	68	127	*	*
94)	1,2,4-Trichlorobenzene	21.97	28.1	26.78	105	62	141	*	*
95)	Naphthalene	22.07	29.1	25.38	115	62	145	*	*
96)	n-Dodecane	22.06	24.6	25.63	96	64	152	*	*
97)	Hexachlorobutadiene	22.38	23.7	26.13	91	49	131	*	*
98)	Cyclohexanone	18.13	23.4	24.45	96	61	127	*	*
99)	tert-Butylbenzene	19.96	25.0	26.88	93	58	122	*	*
100)	n-Butylbenzene	20.72	26.0	27.00	96	64	121	*	*

Bold = 75 Compound List

Data File : I:\MS13\DATA\2019_03\11\03111901.D
 Acq On : 11 Mar 2019 6:51
 Sample : CCV R13031119_25ng
 Misc : S31-02211904/S31-02281905

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 07:15:15 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.33min
 Max. RRF Dev : 30% Max. Rel. Area : 200%

USA 3/11/19

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 IR	Bromochloromethane (IS1)	1.000	1.000	0.0	84	-0.01
2 T	Propene	1.088	1.070	1.7	84	0.00
3 T	Dichlorodifluoromethane (CF	1.858	1.763	5.1	80	0.00
4 T	Chloromethane	1.443	1.537	-6.5	86	0.00
5 T	1,2-Dichloro-1,1,2,2-tetra	1.198	1.106	7.7	79	0.00
6 T	Vinyl Chloride	1.383	1.422	-2.8	82	0.00
7 T	1,3-Butadiene	1.040	1.182	-13.7	87	0.00
8 T	Bromomethane	0.992	0.947	4.5	77	-0.01
9 T	Chloroethane	0.761	0.749	1.6	80	-0.01
10 T	Ethanol	0.741	0.750	-1.2	87	-0.06
11 T	Acetonitrile	1.794	1.940	-8.1	86	-0.03
12 T	Acrolein	0.613	0.677	-10.4	84	-0.02
13 T	Acetone	0.788	0.728	7.6	82	-0.03
14 T	Trichlorofluoromethane	1.557	1.445	7.2	80	-0.02
15 T	2-Propanol (Isopropanol)	2.399	2.509	-4.6	84	-0.05
16 T	Acrylonitrile	1.149	1.394	-21.3	84	-0.03
17 T	1,1-Dichloroethene	1.039	0.998	3.9	77	-0.01
18 T	2-Methyl-2-Propanol (tert-B	2.358	2.393	-1.5	82	-0.05
19 T	Methylene Chloride	0.999	1.021	-2.2	78	-0.02
20 T	3-Chloro-1-propene (Allyl C	1.338	1.422	-6.3	83	-0.02
21 T	Trichlorotrifluoroethane	1.099	0.991	9.8	76	-0.01
22 T	Carbon Disulfide	4.079	3.707	9.1	80	-0.02
23 T	trans-1,2-Dichloroethene	1.216	1.320	-8.6	80	-0.01
24 T	1,1-Dichloroethane	1.730	1.643	5.0	79	-0.01
25 T	Methyl tert-Butyl Ether	2.955	2.762	6.5	78	0.00
26 T	Vinyl Acetate	0.253	0.261	-3.2	79	-0.02
27 T	2-Butanone (MEK)	0.722	0.722	0.0	79	-0.02
28 T	cis-1,2-Dichloroethene	1.258	1.278	-1.6	80	-0.02
29 T	Diisopropyl Ether	1.014	0.964	4.9	77	-0.01
30 T	Ethyl Acetate	0.358	0.359	-0.3	80	-0.02
31 T	n-Hexane	1.660	1.446	12.9	79	0.00
32 T	Chloroform	1.626	1.554	4.4	79	-0.02
33 S	1,2-Dichloroethane-d4(SS1)	1.066	1.165	-9.3	93	-0.01
34 T	Tetrahydrofuran (THF)	0.733	0.777	-6.0	89	0.00
35 T	Ethyl tert-Butyl Ether	1.259	1.172	6.9	77	-0.01
36 T	1,2-Dichloroethane	1.032	1.041	-0.9	80	0.00
37 IR	1,4-Difluorobenzene (IS2)	1.000	1.000	0.0	83	0.00
38 T	1,1,1-Trichloroethane	0.330	0.320	3.0	78	-0.01
39 T	Isopropyl Acetate	0.159	0.155	2.5	81	0.00
40 T	1-Butanol	0.216	0.253	-17.1	86	-0.03
41 T	Benzene	1.032	0.949	8.0	77	0.00
42 T	Carbon Tetrachloride	0.304	0.303	0.3	78	0.00
43 T	Cyclohexane	0.406	0.377	7.1	77	0.00
44 T	tert-Amyl Methyl Ether	0.674	0.647	4.0	78	0.00
45 T	1,2-Dichloropropane	0.236	0.227	3.8	78	-0.01
46 T	Bromodichloromethane	0.286	0.292	-2.1	79	0.00
47 T	Trichloroethene	0.298	0.280	6.0	75	0.00
48 T	1,4-Dioxane	0.208	0.207	0.5	78	0.00
49 T	2,2,4-Trimethylpentane (Iso	1.015	0.927	8.7	78	-0.01
50 T	Methyl Methacrylate	0.100	0.105	-5.0	76	-0.01
51 T	n-Heptane	0.249	0.230	7.6	77	0.00
52 T	cis-1,3-Dichloropropene	0.374	0.382	-2.1	79	0.00
53 T	4-Methyl-2-pentanone	0.213	0.220	-3.3	80	0.00
54 T	trans-1,3-Dichloropropene	0.314	0.337	-7.3	79	0.00
55 T	1,1,2-Trichloroethane	0.240	0.232	3.3	77	0.00

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Data File : I:\MS13\DATA\2019_03\11\03111901.D
 Acq On : 11 Mar 2019 6:51
 Sample : CCV R13031119_25ng
 Misc : S31-02211904/S31-02281905

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 07:15:15 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.33min
 Max. RRF Dev : 30% Max. Rel. Area : 200%

Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
56 IR Chlorobenzene-d5 (IS3)	1.000	1.000	0.0	89	0.00
57 S Toluene-d8 (SS2)	2.677	2.587	3.4	87	0.00
58 T Toluene	2.845	2.271	20.2	77	0.00
59 T 2-Hexanone	1.118	1.133	-1.3	83	0.00
60 T Dibromochloromethane	0.693	0.663	4.3	77	0.00
61 T 1,2-Dibromoethane	0.644	0.622	3.4	77	0.00
62 T n-Butyl Acetate	1.298	1.276	1.7	83	0.00
63 T n-Octane	0.523	0.448	14.3	79	0.00
64 T Tetrachloroethene	0.871	0.718	17.6	74	0.00
65 T Chlorobenzene	1.887	1.571	16.7	75	0.00
66 T Ethylbenzene	3.062	2.578	15.8	76	0.00
67 T m- & p-Xylenes	2.364	1.975	16.5	77	0.00
68 T Bromoform	0.633	0.631	0.3	76	0.00
69 T Styrene	1.848	1.687	8.7	75	0.00
70 T o-Xylene	2.356	1.964	16.6	77	0.00
71 T n-Nonane	1.152	1.016	11.8	82	0.00
72 T 1,1,2,2-Tetrachloroethane	1.104	0.999	9.5	78	0.00
73 S Bromofluorobenzene (SS3)	0.964	0.955	0.9	88	0.00
74 T Cumene	3.215	2.616	18.6	76	0.00
75 T alpha-Pinene	1.539	1.349	12.3	76	0.00
76 T n-Propylbenzene	3.606	3.037	15.8	77	0.00
77 T 3-Ethyltoluene	3.056	2.639	13.6	78	0.00
78 T 4-Ethyltoluene	3.053	2.479	18.8	74	0.00
79 T 1,3,5-Trimethylbenzene	2.680	2.163	19.3	76	0.00
80 T alpha-Methylstyrene	1.323	1.258	4.9	74	0.00
81 T 2-Ethyltoluene	3.180	2.567	19.3	76	0.00
82 T 1,2,4-Trimethylbenzene	2.600	2.196	15.5	77	0.00
83 T n-Decane	1.265	1.117	11.7	79	0.00
84 T Benzyl Chloride	1.901	1.964	-3.3	80	0.00
85 T 1,3-Dichlorobenzene	1.631	1.397	14.3	75	0.00
86 T 1,4-Dichlorobenzene	1.713	1.424	16.9	75	0.00
87 T sec-Butylbenzene	3.609	2.958	18.0	76	0.00
88 T 4-Isopropyltoluene (p-Cymen)	3.369	2.851	15.4	77	0.00
89 T 1,2,3-Trimethylbenzene	2.572	2.226	13.5	78	0.00
90 T 1,2-Dichlorobenzene	1.592	1.343	15.6	75	0.00
91 T d-Limonene	0.907	0.875	3.5	78	0.00
92 T 1,2-Dibromo-3-Chloropropane	0.576	0.535	7.1	76	0.00
93 T n-Undecane	1.167	1.203	-3.1	80	0.00
94 T 1,2,4-Trichlorobenzene	1.212	1.100	9.2	74	0.00
95 T Naphthalene	3.307	3.410	-3.1	76	0.00
96 T n-Dodecane	1.091	1.165	-6.8	80	0.00
97 T Hexachlorobutadiene	0.901	0.694	23.0	74	0.01
98 T Cyclohexanone	0.862	0.849	1.5	87	0.00
99 T tert-Butylbenzene	2.651	2.177	17.9	76	0.00
100 T n-Butylbenzene	2.645	2.323	12.2	77	0.00

(#) = Out of Range

SPCC's out = 0 CCC's out = 0

Data File : I:\MS13\DATA\2019_03\11\03111901.D
 Acq On : 11 Mar 2019 6:51
 Sample : CCV R13031119_25ng
 Misc : S31-02211904/S31-02281905

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 07:15:15 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

 3/11/19

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Bromochloromethane (IS1)	10.98	130	106667	12.500	ng	-0.01
37) 1,4-Difluorobenzene (IS2)	13.10	114	447071	12.500	ng	0.00
56) Chlorobenzene-d5 (IS3)	17.42	82	192180	12.500	ng	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4(...)	11.83	65	124247	13.662	ng	-0.01
Spiked Amount	12.500	Range 70 - 130	Recovery	=	109.28%	
57) Toluene-d8 (SS2)	15.55	98	497175	12.081	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	96.64%	
73) Bromofluorobenzene (SS3)	18.83	174	183581	12.390	ng	0.00
Spiked Amount	12.500	Range 70 - 130	Recovery	=	99.12%	

Target Compounds

						Qvalue
2) Propene	3.98	42	235322	25.355	ng	96
3) Dichlorodifluoromethan...	4.13	85	392965	24.783	ng	100
4) Chloromethane	4.41	50	330438	26.843	ng	99
5) 1,2-Dichloro-1,1,2,2-t...	4.67	135	242491	23.729	ng	100
6) Vinyl Chloride	4.82	62	318899	27.021	ng	99
7) 1,3-Butadiene	5.08	54	264522	29.796	ng	95
8) Bromomethane	5.51	94	203909	24.080	ng	99
9) Chloroethane	5.83	64	163214	25.128	ng	99
10) Ethanol	6.23	45	822363	129.974	ng	97
11) Acetonitrile	6.47	41	427457	27.920	ng	98
12) Acrolein	6.64	56	148566	28.409	ng	97
13) Acetone	6.85	58	833838	124.033	ng	85
14) Trichlorofluoromethane	7.07	101	326653	24.584	ng	99
15) 2-Propanol (Isopropanol)	7.34	45	1104101	53.930	ng	94
16) Acrylonitrile	7.59	53	307571	31.381	ng	97
17) 1,1-Dichloroethene	8.02	96	228590	25.775	ng	90
18) 2-Methyl-2-Propanol (t...	8.21	59	1094314	54.379	ng	96
19) Methylene Chloride	8.26	84	233098	27.351	ng	100
20) 3-Chloro-1-propene (Al...	8.41	41	323593	28.333	ng	95
21) Trichlorotrifluoroethane	8.66	151	225186	24.018	ng	97
22) Carbon Disulfide	8.51	76	850054	24.419	ng	99
23) trans-1,2-Dichloroethene	9.53	61	299115	28.822	ng	97
24) 1,1-Dichloroethane	9.78	63	361106	24.459	ng	99
25) Methyl tert-Butyl Ether	9.88	73	641730	25.447	ng	97
26) Vinyl Acetate	10.05	86	291877	134.966	ng	# 94
27) 2-Butanone (MEK)	10.28	72	158222	25.675	ng	98
28) cis-1,2-Dichloroethene	10.80	61	287393	26.776	ng	96
29) Diisopropyl Ether	11.10	87	222396	25.708	ng	# 69
30) Ethyl Acetate	11.10	61	165812	54.330	ng	100
31) n-Hexane	11.08	57	333813	23.565	ng	# 100
32) Chloroform	11.14	83	357022	25.726	ng	99
34) Tetrahydrofuran (THF)	11.55	72	177100	28.317	ng	94
35) Ethyl tert-Butyl Ether	11.69	87	265116	24.680	ng	98
36) 1,2-Dichloroethane	11.95	62	235605	26.745	ng	100
38) 1,1,1-Trichloroethane	12.22	97	309300	26.206	ng	98
39) Isopropyl Acetate	12.66	61	287134	50.640	ng	# 87
40) 1-Butanol	12.68	56	467883	60.598	ng	# 74
41) Benzene	12.70	78	876518	23.742	ng	99
42) Carbon Tetrachloride	12.86	117	280583	25.834	ng	100
43) Cyclohexane	12.99	84	702816	48.449	ng	95
44) tert-Amyl Methyl Ether	13.35	73	620844	25.766	ng	97
45) 1,2-Dichloropropane	13.55	63	217851	25.800	ng	100
46) Bromodichloromethane	13.74	83	278511	27.251	ng	100
47) Trichloroethene	13.80	130	265611	24.891	ng	99
48) 1,4-Dioxane	13.78	88	196768	26.453	ng	97
49) 2,2,4-Trimethylpentane...	13.86	57	879320	24.214	ng	91
50) Methyl Methacrylate	14.01	100	199639	55.930	ng	97

Data File : I:\MS13\DATA\2019_03\11\03111901.D
 Acq On : 11 Mar 2019 6:51
 Sample : CCV R13031119_25ng
 Misc : S31-02211904/S31-02281905

Vial: 2
 Operator: WA
 Inst : MS13

Quant Time: Mar 11 07:15:15 2019
 Quant Method : I:\MS13\METHODS\R13012319.M
 Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 QLast Update : Wed Jan 23 11:21:29 2019
 Response via : Initial Calibration
 DataAcq Meth:TO15.M

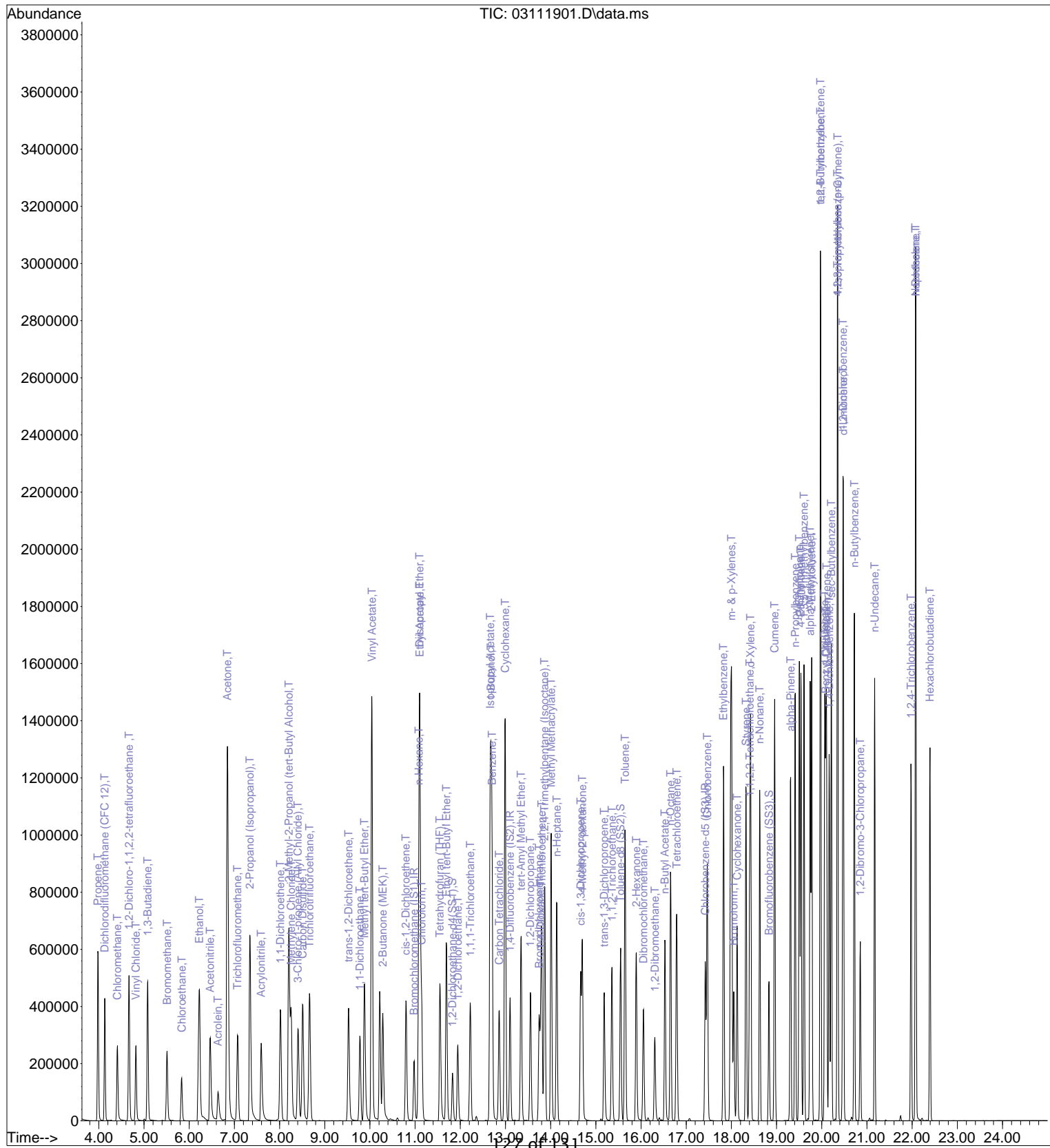
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
51) n-Heptane	14.14	71	221459	24.888	ng	98
52) cis-1,3-Dichloropropene	14.67	75	382316	28.566	ng	100
53) 4-Methyl-2-pentanone	14.70	58	208948	27.467	ng	96
54) trans-1,3-Dichloropropene	15.18	75	317511	28.238	ng	100
55) 1,1,2-Trichloroethane	15.36	97	223507	26.051	ng	100
58) Toluene	15.65	91	918137	20.989	ng	99
59) 2-Hexanone	15.90	43	467579	27.201	ng	95
60) Dibromochloromethane	16.06	129	273856	25.696	ng	100
61) 1,2-Dibromoethane	16.31	107	257132	25.963	ng	100
62) n-Butyl Acetate	16.53	43	532315	26.684	ng	96
63) n-Octane	16.65	57	185336	23.057	ng	93
64) Tetrachloroethene	16.79	166	292166	21.823	ng	99
65) Chlorobenzene	17.47	112	643493	22.178	ng	100
66) Ethylbenzene	17.82	91	1023485	21.738	ng	98
67) m- & p-Xylenes	18.00	91	1611680	44.343	ng	100
68) Bromoform	18.05	173	257887	26.489	ng	100
69) Styrene	18.33	104	687351	24.192	ng	97
70) o-Xylene	18.42	91	801502	22.127	ng	98
71) n-Nonane	18.63	43	418034	23.610	ng	96
72) 1,1,2,2-Tetrachloroethane	18.41	83	408510	24.067	ng	99
74) Cumene	18.96	105	1062690	21.500	ng	100
75) alpha-Pinene	19.31	93	536819	22.688	ng	95
76) n-Propylbenzene	19.41	91	1256120	22.658	ng	99
77) 3-Ethyltoluene	19.50	105	1077374	22.932	ng	100
78) 4-Ethyltoluene	19.54	105	1010947	21.535	ng	100
79) 1,3,5-Trimethylbenzene	19.61	105	878705	21.326	ng	98
80) alpha-Methylstyrene	19.74	118	511648	25.156	ng	91
81) 2-Ethyltoluene	19.78	105	1057528	21.631	ng	99
82) 1,2,4-Trimethylbenzene	19.97	105	901441	22.552	ng	97
83) n-Decane	20.07	57	461988	23.751	ng	99
84) Benzyl Chloride	20.08	91	793566	27.147	ng	99
85) 1,3-Dichlorobenzene	20.11	146	579892	23.119	ng	99
86) 1,4-Dichlorobenzene	20.16	146	591797	22.475	ng	100
87) sec-Butylbenzene	20.21	105	1208388	21.777	ng	99
88) 4-Isopropyltoluene (p-...	20.35	119	1141876	22.046	ng	99
89) 1,2,3-Trimethylbenzene	20.35	105	891642	22.549	ng	97
90) 1,2-Dichlorobenzene	20.47	146	562103	22.960	ng	99
91) d-Limonene	20.48	68	339751	24.353	ng	94
92) 1,2-Dibromo-3-Chloropr...	20.85	157	214319	24.215	ng	90
93) n-Undecane	21.17	57	488673	27.244	ng	97
94) 1,2,4-Trichlorobenzene	21.97	180	449693	24.127	ng	100
95) Naphthalene	22.08	128	1343621	26.425	ng	100
96) n-Dodecane	22.08	57	461720	27.524	ng	99
97) Hexachlorobutadiene	22.39	225	281020	20.280	ng	100
98) Cyclohexanone	18.13	55	320326	24.179	ng	96
99) tert-Butylbenzene	19.97	119	892611	21.904	ng	100
100) n-Butylbenzene	20.72	91	949882	23.362	ng	99

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data File : I:\MS13\DATA\2019_03\11\03111901.D
Acq On : 11 Mar 2019 6:51
Sample : CCV R13031119_25ng
Misc : S31-02211904/S31-02281905

Vial: 2
Operator: WA
Inst : MS13

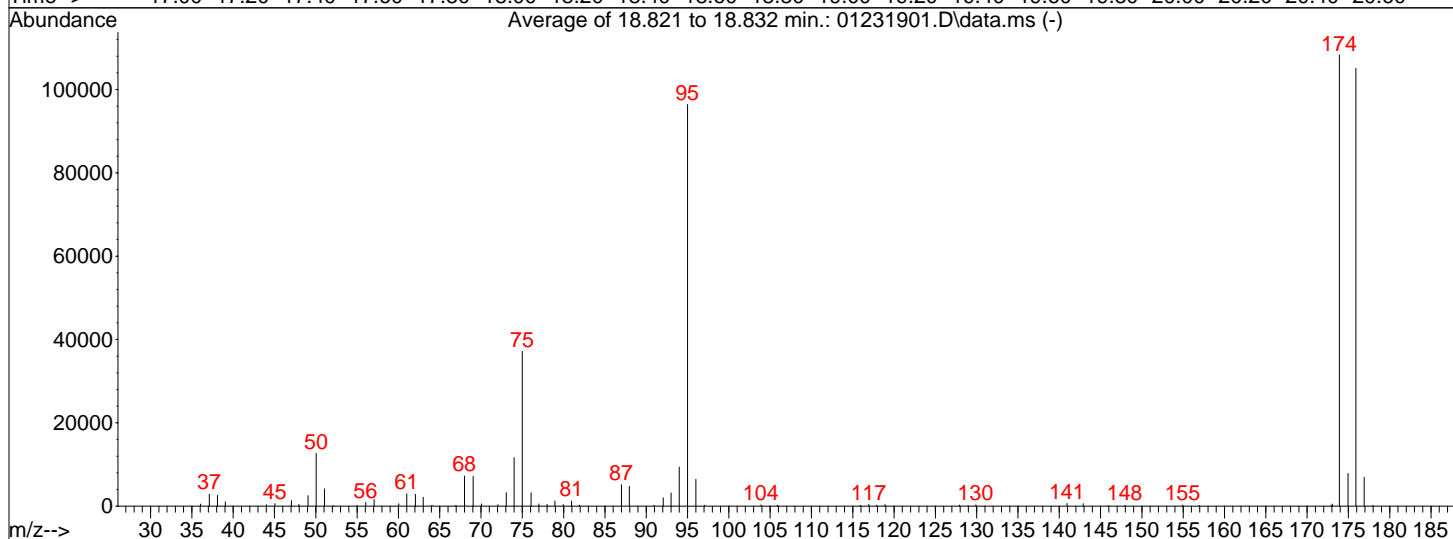
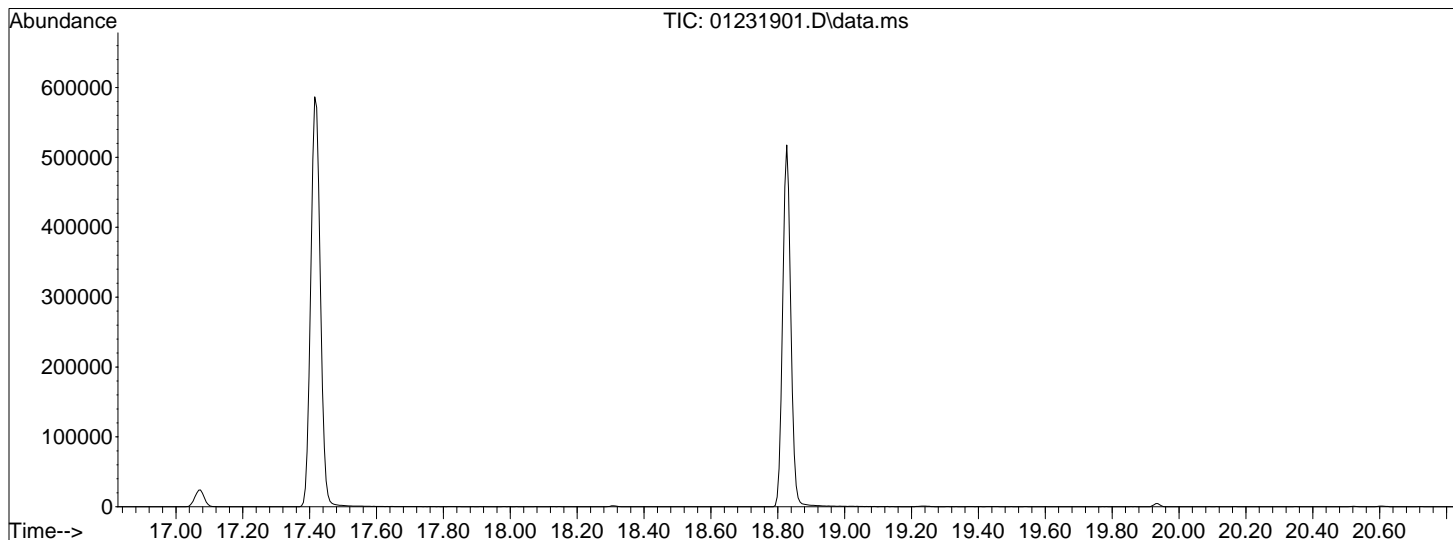
Quant Time: Mar 11 07:15:15 2019
Quant Method : I:\MS13\METHODS\R13012319.M
Quant Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
QLast Update : Wed Jan 23 11:21:29 2019
Response via : Initial Calibration
DataAcq Meth:TO15.M



Data Path : I:\MS13\DATA\2019_01\23\
 Data File : 01231901.D
 Acq On : 23 Jan 2019 4:21
 Operator : WA
 Sample : BFB Std
 Misc : S31-01221908
 ALS Vial : 2 Sample Multiplier: 1

Integration File: LSCINT.P

Method : I:\MS13\METHODS\R13012319.M
 Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 Last Update : Wed Jan 23 11:21:29 2019



AutoFind: Scans 2671, 2672, 2673; Background Corrected with Scan 2664

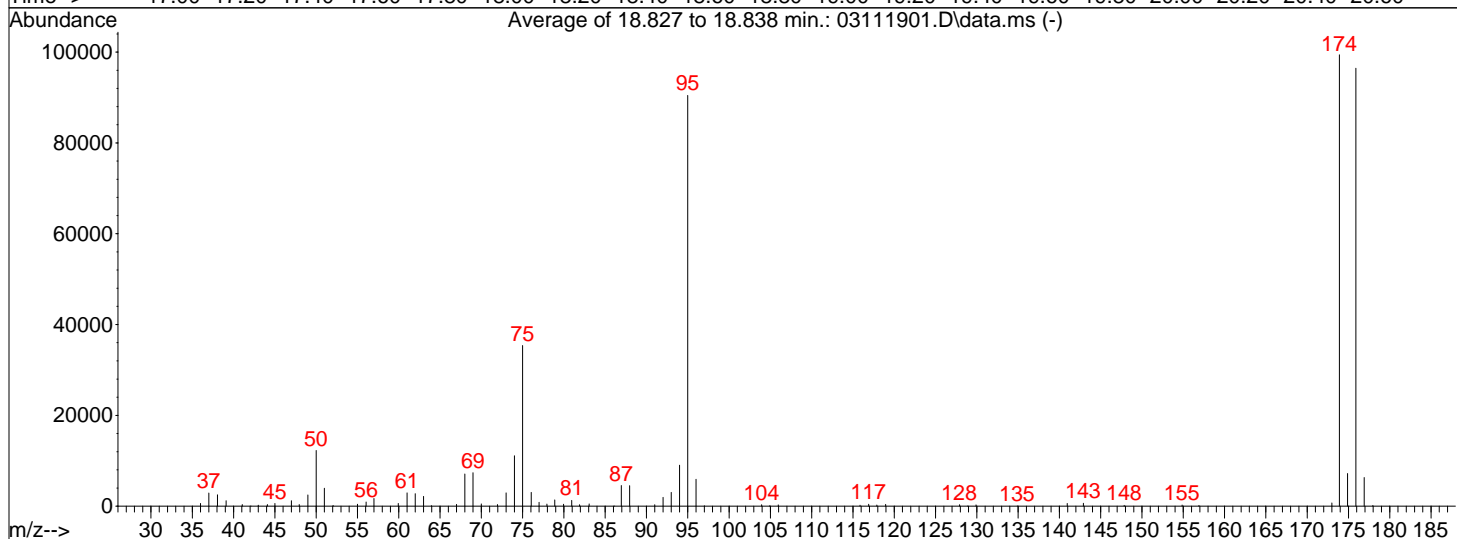
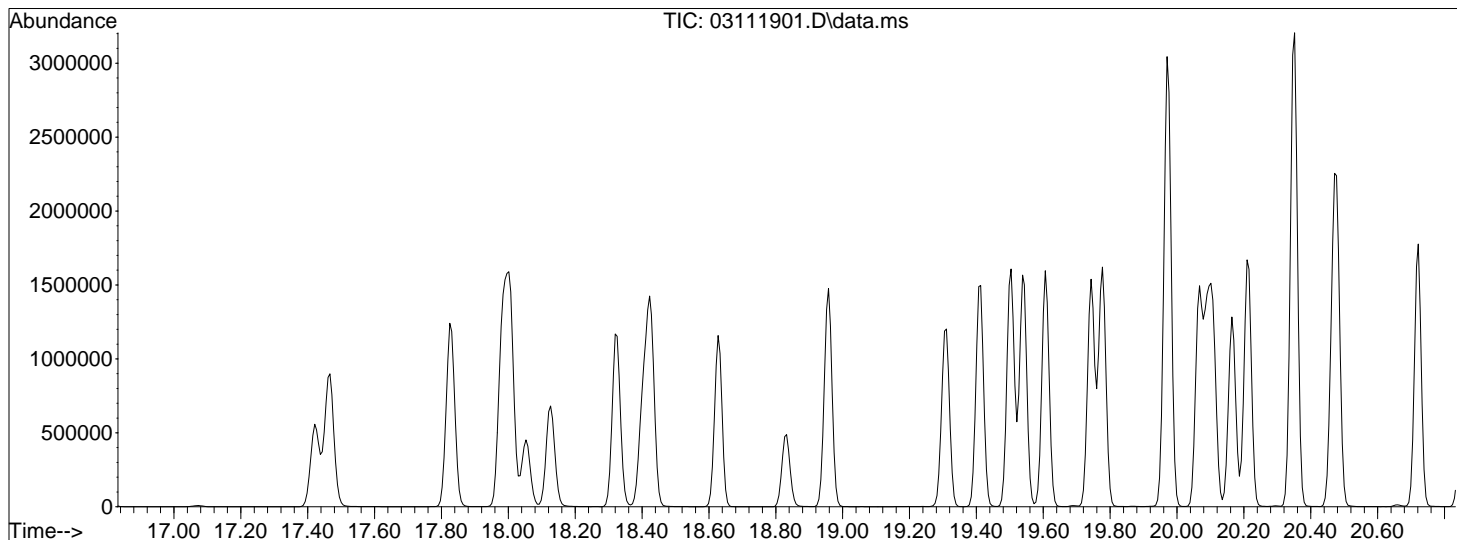
Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	8	40	13.1	12653	PASS
75	95	30	66	38.5	37123	PASS
95	95	100	100	100.0	96387	PASS
96	95	5	9	6.7	6460	PASS
173	174	0.00	2	0.5	489	PASS
174	95	50	120	112.4	108333	PASS
175	174	4	9	7.2	7833	PASS
176	174	93	101	97.0	105085	PASS
177	176	5	9	6.6	6896	PASS

WA 1/23/19

Data Path : I:\MS13\DATA\2019_03\11\
 Data File : 03111901.D
 Acq On : 11 Mar 2019 6:51
 Operator : WA
 Sample : CCV R13031119_25ng
 Misc : S31-02211904/S31-02281905
 ALS Vial : 2 Sample Multiplier: 1

Integration File: LSCINT.P

Method : I:\MS13\METHODS\R13012319.M
 Title : EPA TO-15 per SOP VOA-TO15 (CASS TO-15/GC-MS)
 Last Update : Wed Jan 23 11:21:29 2019



AutoFind: Scans 2672, 2673, 2674; Background Corrected with Scan 2665

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	8	40	13.6	12265	PASS
75	95	30	66	39.1	35389	PASS
95	95	100	100	100.0	90475	PASS
96	95	5	9	6.5	5922	PASS
173	174	0.00	2	0.7	706	PASS
174	95	50	120	109.9	99416	PASS
175	174	4	9	7.2	7181	PASS
176	174	93	101	97.0	96467	PASS
177	176	5	9	6.5	6309	PASS

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Injection Log

Directory: I:\MS13\DATA\2019_01\23\

	Date/Time	File Name	Sample ID	Misc Info	Operator	Vial	Comment	
1	1/23/19 4:21	01231901.D	BFB Std	S31-01221908	WA	2	Pass	
2	1/23/19 4:55	01231902.D	Std check	S31-1212181/S31-01181901 (2/16)	WA	2		
3	1/23/19 5:28	01231903.D	Std check	S31-1212181/S31-01181901 (2/16)	WA	2		
4	1/23/19 6:01	01231904.D	0.5ng R13012319 ICAL Std	S31-01231908/S31-01161906 (2/14)	WA	15	R13012319.M	
5	1/23/19 6:35	01231905.D	1.0ng R13012319 ICAL Std	S31-01231908/S31-01161906 (2/14)	WA	15		
6	1/23/19 8:07	01231906.D	0.1ng R13012319 ICAL Std	S31-01231908/S31-01221902 (2/20)	WA	14		
7	1/23/19 8:40	01231907.D	0.2ng R13012319 ICAL Std	S31-01231908/S31-01221902 (2/20)	WA	14		
8	1/23/19 9:13	01231908.D	5.0ng R13012319 ICAL Std	S31-01221908/S31-01161903 (2/14)	WA	16		
9	1/23/19 9:46	01231909.D	25ng R13012319 ICAL Std	S31-01221908/S31-01161903 (2/14)	WA	16		
10	1/23/19 10:20	01231910.D	50ng R13012319 ICAL Std	S31-01221908/S31-01161903 (2/14)	WA	16		
11	1/23/19 10:53	01231911.D	100ng R13012319 ICAL Std	S31-01221908/S31-01161903 (2/14)	WA	16		
12	1/23/19 11:31	01231912.D	25ng TO-15 ICV Std	S31-01221908/S31-12271803 (1/25)	WA	13		Pass all compds
R13012319.M:good for low level compounds; ranges from 0.1ng -->100ng (except: chloroethane, CS2, Naphthalene: 0.2ng --> 100ng; Acetonitrile, Acrolein, Acrylonitrile, MEK, cis/trans-1,3-DCP, Benzyl-Cl, n-Dodecane: 0.5ng --> 100ng; Ethyl Ac.: 1.0ng --> 200ng; Vinyl Ac.: 2.5ng --> 500ng								

 1/29/19

GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

APPENDIX G
PROTOCOL FOR DISCHARGING GWTP EFFLUENT

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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MEMORANDUM

DATE: August 28, 2017

PROJECT NAME: Remediation of Multiple Sites, Longhorn Army Ammunition Plant, Karnack, TX

TO: Richard Mayer Senior Project Engineer
US Environmental Protection Agency
Federal Facilities Section (6PD-F)

April Palmie Project and Grant Manager
Superfund Section, Remediation Division
Texas Commission on Environmental Quality

FROM: Rose M. Zeiler, Ph.D. Longhorn AAP Site Manager

SUBJECT: **Protocol for Discharging GWTP Effluent
Longhorn Army Ammunition Plant, Karnack, TX
(Contract: W912DY-09-D-0059, Task Order DS01)**

INTRODUCTION

The purpose of this memo is to document the protocol for discharging Longhorn Army Ammunition Plant groundwater treatment plant (GWTP) effluent to Harrison Bayou, the INF-Pond, or LHAAP-18/24.

The GWTP is designed to:

- Extract groundwater from LHAAP-18/24 and LHAAP-16 for hydraulic control;
- Remove metals by pH adjustment, polymer addition, and gravity separation;
- Remove volatile organic compounds (VOCs) by air stripping;
- Remove perchlorate in a fluidized bed reactor (FBR) and an ion exchange scavenger system; and
- Discharge the effluent continuously.

DISCHARGE CRITERIA

The discharge criteria established for discharge to Harrison Bayou are:

Parameter	Discharge Criteria (µg/L)	
	Daily Average	Daily Maximum
Volatiles		
1,1,1-Trichloroethane	3,417	7,230
1,1,2-Trichloroethane	102.5	216.9
1,1-Dichloroethane	6,633	14,032
1,1-Dichloroethene	119	253
1,2-Dichloroethane	85	181
Acetone	1,132	2,395
Benzene	85	181
Carbon Tetrachloride	85	181
Chlorobenzene	22,300	47,180
Chloroform	1,708	3,615
Ethylbenzene	26,954	57,025
Xylenes	39.5	83.6
Methylene Chloride	803	1,699
Styrene	2,829	5,987
Tetrachloroethene	85.4	180.7
Toluene	1,980	4,189
Trichloroethene	85	181
Vinyl Chloride	34	72
Anions		
Chloride	*	*
Sulfate	*	*
Perchlorate**	278	589
Metals		
Aluminum	777	1,644
Arsenic	365	772
Barium	1,000	2,000
Cadmium	1.6	3.4
Chromium, Total	355	752
Chromium, Hexavalent	58	124
Cobalt	5,433	11,495
Iron	1,132	2,395
Lead	2.2	4.6
Nickel	87	184
Manganese	7,323	15,494
Silver	1.4	3
Selenium	5.7	12
Vanadium	1,698	3,592
Zinc	146	310
Other		
Hexachlorobenzene	0.22	0.47
1,4-Dioxane		134.2
Oil and Grease		15
Chemical Oxygen Demand		200

* - Based upon flow in Harrison Bayou

** - Discharge criteria, when diverted to the INF Pond, is 17 µg/L

PROTOCOL FOR DISCHARGING GWTP EFFLUENT

In accordance with the *Sampling and Analysis Plan, Groundwater Treatment Plant and Well Fields* (SAP) Table 2-2, indicator parameters for the FBR, such as temperature, pH and oxidation reduction potential (ORP), are monitored in real time to predict FBR performance and perchlorate removal. Based upon these indicator parameters, the operator of the GWTP can make adjustments such as:

- Bring the ion exchange system online;
- Increase or decrease the addition rate of electron donor (acetic acid);
- Increase or decrease the nutrient addition rate (urea or phosphoric acid); or
- Increase or decrease the FBR recirculation rate

Samples of the GWTP effluent are collected weekly, analyzed for perchlorate, nutrients (ammonia-nitrogen and ortho-phosphate), total organic carbon (TOC), chloride, and sulfate, with the results received from the laboratory 14 days later. Other parameters (e.g. Record of Decision metals and volatiles) are collected and analyzed in GWTP effluent samples according to the frequencies listed in Table 2-1 of the SAP.

As shown in Figure 1, groundwater is continuously extracted, treated, and discharged. If Harrison Bayou is flowing and indicator parameters are within their historical optimal ranges, then the ion exchange vessels can be bypassed and the GWTP effluent sample will be collected after the FBR. If Harrison Bayou is not flowing or the indicator parameters are not within historical optimal ranges, then the ion exchange vessels will be put on line, and the GWTP effluent sample will be collected between the lead and lag ion exchange vessel. Professional judgement may also be used as to when to bring the ion exchange vessels online, such as after a power outage or during anticipated cold temperatures when the FBR has historically not performed optimally.

If a parameter is measured in the effluent at a concentration above the discharge criteria, then a confirmation sample and an effluent sample after the lag ion exchange vessel will be collected and analyzed for the parameter with a 24-hour turnaround time. Corrective measures (e.g. increased nutrient or electron donor addition rates, bring ion exchange vessels on line) will be implemented as appropriate to bring the parameter back within the discharge criteria. ***If an upset condition in the FBR leads to high concentrations of perchlorate going into the lead ion exchange vessel and breaking through at the sample location between the vessels, the lag vessel will still remove perchlorate before it is discharged to Harrison Bayou, the INF Pond, or LHAAP-18/24.*** It is estimated that the lag ion exchange vessel can remove all of the perchlorate from two weeks of typical groundwater extraction at a concentration of 920 µg/L. If the residual perchlorate concentration after the FBR and lead ion exchange vessel is only 600 µg/L, the lag ion exchange vessel could last almost 2.5 years before perchlorate would be detected in the discharged effluent.

If a parameter exceeds the discharge criteria by more than 40% (see Appendix A-2, SAP, Section 7c of Monitoring and Reporting Requirements) or reaches 920 µg/L of perchlorate, then the GWTP will be put into full recycle mode (no discharge) until the parameter is below the discharge criteria again. Appendix A-2 of the SAP requires GWTP data to be provided to TCEQ monthly including a list of noncompliance(s), if applicable.

Discharge to Harrison Bayou

As shown in Figure 1, the GWTP effluent will be discharged to Harrison Bayou as long as it has a measurable flow. The flowrate in Harrison Bayou is estimated by measuring the height of water with a staff gauge and velocity in feet/sec at intervals along the width as described in the Installation-Wide Work Plan, Standard Operating Procedures, Attachment 18 – Water Depth and Velocity Measurements (AECOM, July 2014).

The allowable flow rate of GWTP effluent that can be discharged to Harrison Bayou is given by:

$$Q_E \leq \frac{Q_S (C_C - C_A)}{(C_E - C_C)}$$

where Q_E = GWTP effluent flow Q_S = Harrison Bayou flow

C_C = Criteria concentration (100 mg/L for chloride, 50 mg/L for sulfate)

C_A = Ambient concentration = 10 mg/L

C_E = Chloride or sulfate concentration in GWTP effluent

The allowable GWTP effluent flow will be the lower of the calculated values given the measured concentrations of chloride and sulfate in the discharge stream. For each day that GWTP effluent is discharged to Harrison Bayou, the measured Harrison Bayou flow, the allowable effluent flow, and the actual effluent flow are recorded.

Discharge to INF Pond

If Harrison Bayou is not flowing, then GWTP effluent will be discharged to the Intermediate-Range Nuclear Forces (INF) Pond for temporary storage until Harrison Bayou flow resumes. Perchlorate concentration detected in the effluent must be 17 µg/L or less, when this occurs.

The INF Pond has a flexible membrane liner protected by a soil cover with a gravity discharge pipe (and valve) to Harrison Bayou. The INF Pond has a nominal capacity of 3 million gallons with a staff gage to measure the height of water stored in the pond. The GWTP operator maintains the INF Pond by visually inspecting for erosion, vegetative growth including tree growth along the anchor trench, and liner integrity and making necessary repairs. Periodically, accumulated debris must be removed from the influent and effluent piping to the INF Pond.

Prior to discharging to the INF Pond, a lead and lag ion exchange vessel will be brought online. The GWTP Operator will also confirm that the discharge valve is closed, will record the reading on the effluent totalizer, and will record the height of water using the staff gage. The GWTP Operator will then configure valves and pumps to direct GWTP effluent to the INF Pond. The height of water in the INF Pond and totalizer reading will be recorded at the beginning and end of each shift for the duration of active discharge. When the height of water in the pond reaches 3 feet below the height of the berm (freeboard), the GWTP Operator will stop discharging to the INF Pond and TCEQ will be notified. After the TCEQ acknowledges the INF Pond level, GWTP effluent may be discharged to the INF Pond again until 2 feet of freeboard is reached. The GWTP Operator will stop discharging to the INF Pond and TCEQ will be notified again. After the TCEQ acknowledges 2 feet of freeboard in the INF Pond, GWTP effluent may be discharged again until 1 foot of freeboard remains. No additional GWTP effluent can be accepted at the INF Pond until greater than 1 foot of freeboard is measured.

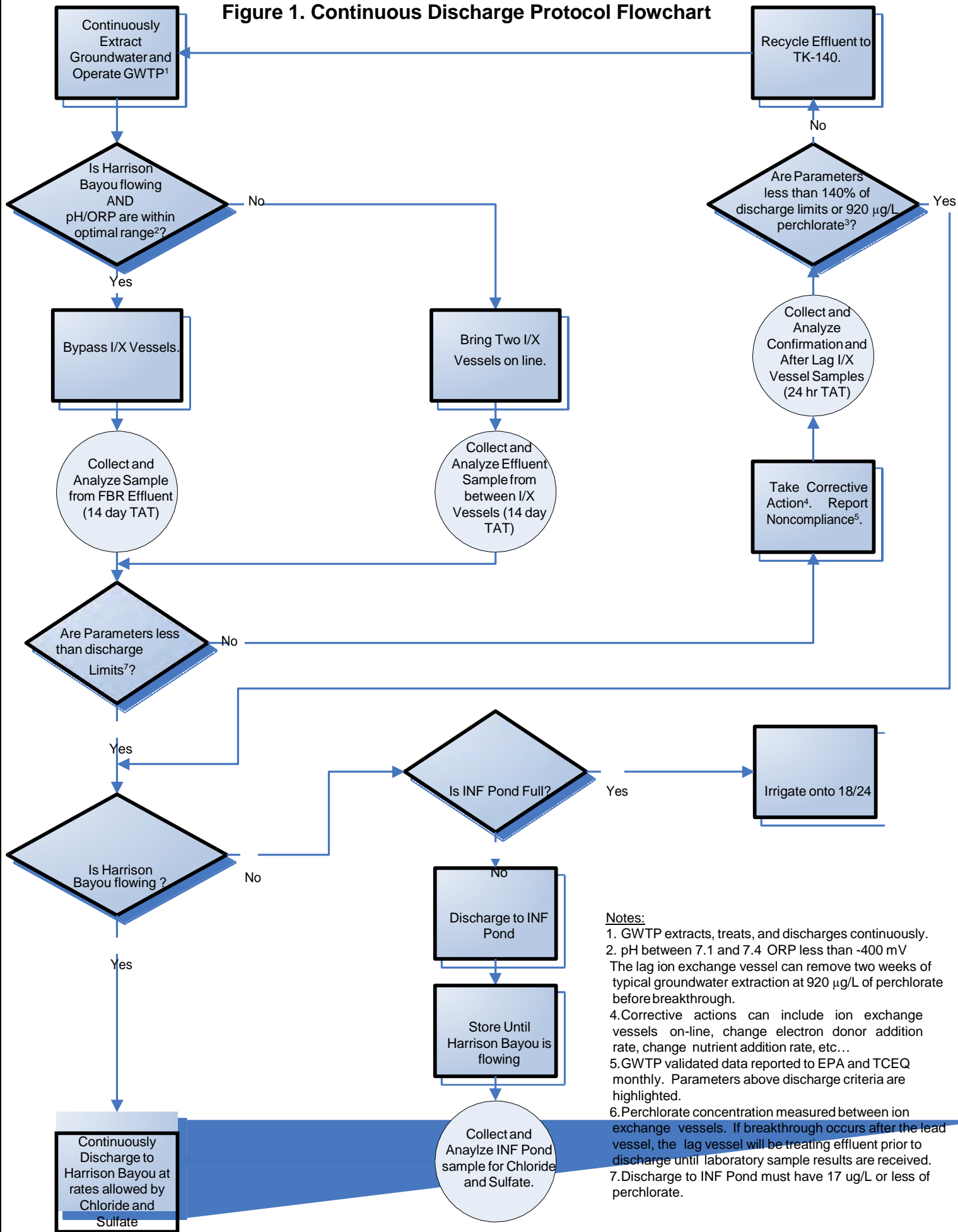
As soon as flow in Harrison Bayou returns, stored GWTP effluent from the INF Pond will be discharged. As with direct discharges from the GWTP to Harrison Bayou, the allowable flowrate of effluent from the INF Pond is calculated based upon the chloride and sulfate concentrations in the pond and the flow in Harrison Bayou. If effluent from the INF Pond and the GWTP are discharged simultaneously, total flow of both streams together should not exceed the calculated discharge level for either discharge location. For each day that INF Pond contents are discharged to Harrison Bayou, the measured Harrison Bayou flow, the allowable effluent flow, and the actual effluent flow are recorded.

Irrigation onto LHAAP-18/24

If Harrison Bayou is not flowing and the INF Pond has less than 1 foot of freeboard, then GWTP effluent will be irrigated onto LHAAP-18/24 using one of the three main sprinkler lines. To avoid pooling and runoff of irrigation water, only one line will be used for half a day at a time, with a separate line being used the second half of the day. If needed, the irrigation will occur 5 days a week for 8 hours each day (using 3 sprinklers in each line). If conditions are wet due to rain events, irrigation will not be conducted to avoid ponding and potential runoff, the GWTP will be put into recycle mode, and groundwater extraction will be interrupted if storage space is not available.

While irrigating, site inspections will be performed to ensure pooling and runoff are not occurring. During the irrigation activities, inspections will be performed twice a day, once approximately three hours and again approximately six hours into the 8-hour irrigation shift. The system will be inspected to ensure that the sprinkler heads are operating properly and not leaking large amounts of water. If ponding or runoff is observed, irrigation at that sprinkler line will cease, and irrigation at another sprinkler line will be started if possible. Volumes of GWTP effluent and twice daily inspections will be recorded daily and reported monthly until flow resumes in Harrison Bayou or greater than 1 foot of freeboard is available in the INF Pond.

Figure 1. Continuous Discharge Protocol Flowchart



- Notes:**
1. GWTP extracts, treats, and discharges continuously.
 2. pH between 7.1 and 7.4 ORP less than -400 mV The lag ion exchange vessel can remove two weeks of typical groundwater extraction at 920 µg/L of perchlorate before breakthrough.
 4. Corrective actions can include ion exchange vessels on-line, change electron donor addition rate, change nutrient addition rate, etc...
 5. GWTP validated data reported to EPA and TCEQ monthly. Parameters above discharge criteria are highlighted.
 6. Perchlorate concentration measured between ion exchange vessels. If breakthrough occurs after the lead vessel, the lag vessel will be treating effluent prior to discharge until laboratory sample results are received.
 7. Discharge to INF Pond must have 17 ug/L or less of perchlorate.

GWTP QUARTERLY EVALUATION REPORT – 1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

APPENDIX H
AIR DATA TABLES, PID READINGS, AND CALIBRATION LOGS

GWTP QUARTERLY EVALUATION REPORT –1ST QUARTER 2019
LONGHORN ARMY AMMUNITION PLANT

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**Appendix H, Table 1. Ambient Air Data - February 2019
Longhorn Army Ammunition Plant
Groundwater Treatment Plant**

Pollutant	CAS #	Short Term ESL	AMCVs	GWTP Ambient Air Concentrations (1)	Status (3)	Downwind Ambient Air Concentrations	Status (3)
		March 2012	(ST Health)			(2)	
		$\mu\text{g}/\text{m}^3$	$\mu\text{g}/\text{m}^3$	$\mu\text{g}/\text{m}^3$		$\mu\text{g}/\text{m}^3$	
1,1-Dichloroethane	75-34-3	4,000	4,047	0.67	U PASS	0.60	U PASS
1,1-Dichloroethene	75-35-4	210	714	0.69	U PASS	0.63	U PASS
1,2-Dichloroethane	107-06-2	160	162	0.68	U PASS	0.61	U PASS
Acetone	67-64-1	5,900	NA	9.0	PASS	6.3	U PASS
Benzene	71-43-2	170	575	0.80	PASS	0.69	PASS
Carbon disulfide	75-15-0	30	NA	1.4	U PASS	1.3	U PASS
Chloroform	67-66-3	100	98	0.69	U PASS	0.63	U PASS
cis-1,2-Dichloroethene	156-59-2	7,900	NA	12.0	PASS	0.64	PASS
Methylene chloride	75-09-2	3,600	12,158	2.8	PASS	1.4	PASS
Tetrachloroethene	127-18-4	2,000	6,782	0.68	U PASS	0.61	U PASS
trans-1,2-Dichloroethene	156-60-5	7,900	NA	0.68	U PASS	0.61	U PASS
Trichloroethene	79-01-6	540	537	55.0	PASS	2.1	PASS
Vinyl chloride	75-01-4	20,000	66,460	0.68	U PASS	0.61	U PASS
n-Hexane	110-54-3	5,300	6,336	1.00	PASS	0.83	PASS
Styrene	100-42-5	110	21,725	0.68	U PASS	0.61	U PASS
Toluene	108-88-3	640	15,074	1.2	PASS	0.76	PASS
Ethylbenzene	100-41-4	740	86,844	0.67	U PASS	0.60	U PASS
m,p-Xylenes	179601-23-1	180	7,382	1.4	U PASS	1.3	U PASS
o-Xylene	95-47-6	1,600	7,382	0.68	U PASS	0.61	U PASS
1,3-Dichlorobenzene	541-73-1	720	NA	0.69	U PASS	0.63	U PASS
Propene (C3 H6)	115-07-1	Asphyxiant	Asphyxiant	0.67	U NA	0.60	U NA
Dichlorodifluoromethane (CCl2F2)	75-71-8	50,000	49,452	2.5	PASS	2.6	PASS
Ethanol	64-17-5	18,800	NA	6.5	U PASS	5.9	U PASS
Trichlorofluoromethane (CCl3F)	75-69-4	28,000	56,184	1.4	PASS	1.3	PASS
Trichlorotrifluoroethane (C2Cl3F3)	76-13-1	38,000	NA	19	PASS	7.0	PASS
alpha-Pinene	80-56-8	60	3,499	0.67	U PASS	0.60	U PASS
d-Limonene	5989-27-5	1,100	NA	0.65	U PASS	0.59	U PASS

Notes:

(1) Sample collected over an 8-hour period on February 27, 2019, between 8 AM and 4 PM

(2) Sample collected over a 24-hour period beginning on February 27, 2019, at 6:30 AM and ending on February 28, 2019 at 6:30 AM

(3) Status based on comparison of air sample result to Air Monitoring Comparison Values (AMCVs). When there is no AMCV value for a chemical, the air sample concentration is compared to the short-term Effects Screening Level (ESL).

CAS # = Chemical Abstracts Service Number

GWTP = Groundwater Treatment Plant

NA - Not applicable

U = non-detect

$\mu\text{g}/\text{m}^3$ = micrograms per cubic meter

Appendix H, Table 2. Emission Stack Air Data - February 2019
Longhorn Army Ammunition Plant
Groundwater Treatment Plant

Pollutant	CAS #	Measured Air Stripper Stack Concentrations (1)		Air Stripper Emission Rates (2)		Air Stripper Emission Rates (2a)		Allowable Annual Emission (3)		Status (4)	TLV (L)	TLV Reference	Compliance section	Distance Downwind to nearest off-site Receptor (D)	(K) conversion value	Allowable Maximum Hourly Emission Limit at Nearest off-site Receptor ⁽⁶⁾⁽⁷⁾ (E) = L/K	Status (8)	
		$\mu\text{g}/\text{m}^3$	U	lb/hr	U	tpy	U	tpy	feet									lb/hr
1,1-Dichloroethane	75-34-3	67	U	5.49E-04	U	3.57E-04	U	5	PASS		405	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
1,1-Dichloroethene	75-35-4	69	U	5.66E-04	U	3.68E-04	U	5	PASS		20	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.4	PASS	
1,2-Dichloroethane	107-06-2	140	U	2.30E-03	U	1.49E-03	U	5	PASS		40	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	2.9	PASS	
Acetone	67-64-1	690	U	5.66E-03	U	3.68E-03	U	5	PASS		590	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
Benzene	71-43-2	67	U	5.49E-04	U	3.57E-04	U	5	PASS		3	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	0.21	PASS	
Carbon disulfide	75-15-0	140	U	1.15E-03	U	7.46E-04	U	5	PASS		140	31	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	2.2	PASS
Chloroform	67-66-3	69	U	5.66E-04	U	3.68E-04	U	5	PASS		10	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	0.71	PASS	
cis-1,2-Dichloroethene	156-59-2	2,800	U	4.59E-02	U	2.98E-02	U	5	PASS		793	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
Methylene chloride	75-09-2	630	U	1.03E-02	U	6.71E-03	U	5	PASS		26	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.9	PASS	
Tetrachloroethene	127-18-4	68	U	5.57E-04	U	3.62E-04	U	5	PASS		33.5	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	2.4	PASS	
trans-1,2-Dichloroethene	156-60-5	68	U	5.57E-04	U	3.62E-04	U	5	PASS		793	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
Trichloroethene	79-01-6	12,000	U	1.97E-01	U	1.28E-01	U	5	PASS		135	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	6.0	PASS	
Vinyl chloride	75-01-4	68	U	5.57E-04	U	3.62E-04	U	5	PASS		2	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	0.14	PASS	
n-Hexane	110-54-3	67	U	5.49E-04	U	3.57E-04	U	5	PASS		1,800	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
Styrene	100-42-5	68	U	5.57E-04	U	3.62E-04	U	5	PASS		21	106.262 List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.5	PASS	
Toluene	108-88-3	68	U	5.57E-04	U	3.62E-04	U	5	PASS		188	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	6.0	PASS	
Ethylbenzene	100-41-4	67	U	5.49E-04	U	3.57E-04	U	5	PASS		434	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
m,p-Xylenes	179601-23-1	140	U	1.15E-03	U	7.46E-04	U	5	PASS		434	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
o-Xylene	95-47-6	68	U	5.57E-04	U	3.62E-04	U	5	PASS		434	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
1,3-Dichlorobenzene	541-73-1	69	U	5.66E-04	U	3.68E-04	U	5	PASS		(5)	--	30 TAC 106.533(f)(1)(A)(i)	2,000	14	1.0	PASS	
Propene (C3 H6)	115-07-1	67	U	5.49E-04	U	3.57E-04	U	5	PASS		(5)	--	30 TAC 106.533(f)(1)(A)(i)	2,000	14	6.0	PASS	
Dichlorodifluoromethane (CCl2F2)	75-71-8	67	U	5.49E-04	U	3.57E-04	U	5	PASS		4,950	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
Ethanol	64-17-5	650	U	5.33E-03	U	3.46E-03	U	5	PASS		1,880	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
Trichlorofluoromethane (CCl3F)	75-69-4	68	U	5.57E-04	U	3.62E-04	U	5	PASS		5,620	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
Trichlorotrifluoroethane (C2Cl3F3)	76-13-1	1,900	U	3.12E-02	U	2.02E-02	U	5	PASS		7,670	ACGIH List	30 TAC 106.533(f)(1)(A)(ii)	2,000	14	1.0	PASS	
alpha-Pinene	80-56-8	67	U	5.49E-04	U	3.57E-04	U	5	PASS		(5)	--	30 TAC 106.533(f)(1)(A)(i)	2,000	14	1.0	PASS	
d-Limonene	5989-27-5	65	U	5.33E-04	U	3.46E-04	U	5	PASS		(5)	--	30 TAC 106.533(f)(1)(A)(i)	2,000	14	1.0	PASS	
TOTAL				0.310														

Notes:

- (1) Sample collected on February 27, 2019. The higher value of the sample or duplicate is reported.
- (2) Based on a blower flow rate of 4,390 cubic feet per minute (cfm). Note that plant operations is less than or equal to 25 hours per week. 1/2 of detection limit was used for estimating mass rate
- (2a) Based on operation of 25 hours per week, 52 weeks per year.
- (3) Per 30TAC 106.533(f)(1)(B)
- (4) Based on comparing the calculated air stripper stack sample emission rate in tons per year (tpy) to the allowable annual emission limit per chemical of 5 tpy.
- (5) No Threshold Limit Values (TLVs) for these chemicals
- (6) The maximum hourly limit allowed by 30 Texas Administrative Code (TAC) 106.262, per pollutant, is 6 pounds per hour (lb/hr) per "Figure 1: 30 TAC 106.262(a)". The E value was overridden with 6 lb/hr when the calculated E was higher.
- (7) The maximum hourly emission rate allowed by 30 TAC 106.261(a)(3) for chemicals with a limit value (L) greater than 200 mg/m³ is 1 lb/hr.
- (8) Based on comparing the calculated air stripper stack sample emission rate in lb/hr to the allowable maximum emission limit per chemical based on distance downwind to nearest off-site receptor.

CAS # = Chemical Abstracts Service Number

mg/m³ = milligrams per cubic meter $\mu\text{g}/\text{m}^3$ = micrograms per cubic meter

U = non-detect

ACGIH = American Conference of Governmental Industrial Hygienists

Appendix H, Table 3: PID Measurements for 1st Quarter 2019
Longhorn Army Ammunition Plant
Groundwater Treatment Plant

Date	Time	Location	Air Flow Rate at		Person Collecting	PID Reading	Weather Conditions
			Blower	Instrument ID			
2/13/2019	8:00	Outside GWTP Office	3980 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 39 degrees
2/13/2019	8:00	Downwind	3980 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 39 degrees
2/13/2019	8:00	Stripper	3980 ACFM	MiniRAE 3000	Kennie Moore	Max. 19.6 ppm Steady State 7.1 ppm	Clear 39 degrees
2/13/2019	14:00	Outside GWTP Office	3750 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 66 degrees
2/13/2019	14:00	Downwind	3750 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 66 degrees
2/13/2019	14:00	Stripper	3750 ACFM	MiniRAE 3000	Kennie Moore	Max. 18.9 ppm Steady State 6.4 ppm	Clear 66 degrees
2/20/2019	8:00	Outside GWTP Office	3875 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 41 degrees
2/20/2019	8:00	Downwind	3875 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 41 degrees
2/20/2019	8:00	Stripper	3875 ACFM	MiniRAE 3000	Kennie Moore	Max. 20.1 ppm Steady State 7.6 ppm	Cloudy 41 degrees
2/20/2019	14:00	Outside GWTP Office	3758 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 59 degrees
2/20/2019	14:00	Downwind	3758 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 59 degrees
2/20/2019	14:00	Stripper	3758 ACFM	MiniRAE 3000	Kennie Moore	Max. 19.0 ppm Steady State 6.1 ppm	Clear 59 degrees
2/27/2019	8:00	Outside GWTP Office	3790 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 58 degrees
2/27/2019	8:00	Downwind	3790 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 58 degrees
2/27/2019	8:00	Stripper	3790 ACFM	MiniRAE 3000	Kennie Moore	Max. 20.1 ppm Steady State 7.6 ppm	Cloudy 58 degrees
2/27/2019	14:00	Outside GWTP Office	3694 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Cloudy 67 degrees
2/27/2019	14:00	Downwind	3694 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Cloudy 67 degrees
2/27/2019	14:00	Stripper	3694 ACFM	MiniRAE 3000	Scott Beesinger	Max. 19.0 ppm Steady State 6.1 ppm	Cloudy 67 degrees
3/7/2019	8:00	Outside GWTP Office	3749 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 49 degrees
3/7/2019	8:00	Downwind	3749 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 49 degrees
3/7/2019	8:00	Stripper	3749 ACFM	MiniRAE 3000	Kennie Moore	Max. 22.9 ppm Steady State 8.3 ppm	Cloudy 49 degrees
3/7/2019	14:00	Outside GWTP Office	3771 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Cloudy 61 degrees
3/7/2019	14:00	Downwind	3771 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Cloudy 61 degrees
3/7/2019	14:00	Stripper	3771 ACFM	MiniRAE 3000	Scott Beesinger	Max. 20.4 ppm Steady State 6.9 ppm	Cloudy 61 degrees
3/13/2019	8:00	Outside GWTP Office	3701 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 67 degrees
3/13/2019	8:00	Downwind	3701 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Cloudy 67 degrees
3/13/2019	8:00	Stripper	3701 ACFM	MiniRAE 3000	Kennie Moore	Max. 23.1 ppm Steady State 7.7 ppm	Cloudy 67 degrees
3/13/2019	14:00	Outside GWTP Office	3740 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Rain 55 degrees
3/13/2019	14:00	Downwind	3740 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Rain 55 degrees
3/13/2019	14:00	Stripper	3740 ACFM	MiniRAE 3000	Scott Beesinger	Max. 21.7 ppm Steady State 5.8 ppm	Rain 55 degrees

**Appendix H, Table 3: PID Measurements for 1st Quarter 2019
Longhorn Army Ammunition Plant
Groundwater Treatment Plant**

Date	Time	Location	Air Flow Rate at		Person Collecting	PID Reading	Weather Conditions
			Blower	Instrument ID			
3/20/2019	8:00	Outside GWTP Office	3881 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 41 degrees
3/20/2019	8:00	Downwind	3881 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 41 degrees
3/20/2019	8:00	Stripper	3881 ACFM	MiniRAE 3000	Kennie Moore	Max. 21.9 ppm Steady State 5.3 ppm	Clear 41 degrees
3/20/2019	14:00	Outside GWTP Office	3688 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 74 degrees
3/20/2019	14:00	Downwind	3688 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 74 degrees
3/20/2019	14:00	Stripper	3688 ACFM	MiniRAE 3000	Scott Beesinger	Max. 23.2 ppm Steady State 6.8 ppm	Clear 74 degrees
3/28/2019	8:00	Outside GWTP Office	3784 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 44 degrees
3/28/2019	8:00	Downwind	3784 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 44 degrees
3/28/2019	8:00	Stripper	3784 ACFM	MiniRAE 3000	Kennie Moore	Max. 21.1 ppm Steady State 5.0 ppm	Clear 44 degrees
3/28/2019	14:00	Outside GWTP Office	3693 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 76 degrees
3/28/2019	14:00	Downwind	3693 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 76 degrees
3/28/2019	14:00	Stripper	3693 ACFM	MiniRAE 3000	Scott Beesinger	Max. 23.1 ppm Steady State 6.3 ppm	Clear 76 degrees
4/3/2019	8:00	Outside GWTP Office	3802 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 46 degrees
4/3/2019	8:00	Downwind	3802 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 46 degrees
4/3/2019	8:00	Stripper	3802 ACFM	MiniRAE 3000	Kennie Moore	Max. 22.7 ppm Steady State 7.4 ppm	Clear 46 degrees
4/3/2019	14:00	Outside GWTP Office	3705 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 74 degrees
4/3/2019	14:00	Downwind	3705 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 74 degrees
4/3/2019	14:00	Stripper	3705 ACFM	MiniRAE 3000	Scott Beesinger	Max. 24.3 ppm Steady State 8.2 ppm	Clear 74 degrees
4/9/2019	8:00	Outside GWTP Office	3769 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 53 degrees
4/9/2019	8:00	Downwind	3769 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 53 degrees
4/9/2019	8:00	Stripper	3769 ACFM	MiniRAE 3000	Kennie Moore	Max. 20.4 ppm Steady State 4.9 ppm	Clear 53 degrees
4/9/2019	14:00	Outside GWTP Office	3678 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 81 degrees
4/9/2019	14:00	Downwind	3678 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Clear 81 degrees
4/9/2019	14:00	Stripper	3678 ACFM	MiniRAE 3000	Scott Beesinger	Max. 22.9 ppm Steady State 6.1 ppm	Clear 81 degrees
4/15/2019	8:00	Outside GWTP Office	3847 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 49 degrees
4/15/2019	8:00	Downwind	3847 ACFM	MiniRAE 3000	Kennie Moore	0.0 ppm	Clear 49 degrees
4/15/2019	8:00	Stripper	3847 ACFM	MiniRAE 3000	Kennie Moore	Max. 19.7 ppm Steady State 4.5 ppm	Clear 49 degrees
4/15/2019	14:00	Outside GWTP Office	3678 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Rain 77 degrees
4/15/2019	14:00	Downwind	3678 ACFM	MiniRAE 3000	Scott Beesinger	0.0 ppm	Rain 77 degrees
4/15/2019	14:00	Stripper	3678 ACFM	MiniRAE 3000	Scott Beesinger	Max. 22.9 ppm Steady State 6.1 ppm	Rain 77 degrees

EQUIPMENT CALIBRATION DAILY LOG

Date: 2/13/19 Project Name: LHAAP-GWTP
 Project Number: NWD1312.0150 Recorded By: Scott Beasinger

PID	Model: <u>MiniRAE 3000</u>		Bulb: <u>11.7</u> meV		Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #: <u>PGM 7320</u>						
	Parameter	Standard	Exp. Date	Lot #	Time: <u>0630</u>	Time:	Time:
First Point Calibration	Vapor conc. (ppm)	0.0 (ambient air)	NA	NA	Initials: <u>SB</u>	Initials:	Initials:
Second Point Calibration	Vapor conc. (ppm)	<u>100ppm</u> (isobutylene)	<u>10/5/21</u>	<u>JBH-248-100-19</u>	Value: <u>To zero SB</u>	Value:	Value:

COMB. GAS/O ₂ METER	Model:				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration	O ₂ (%)				Initials:	Initials:	Initials:
	% LEL Pentane				Value:	Value:	Value:

WATER QUALITY METER	Model:				Morning Calibration/Check	Evening Check (one point only)	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration (Auto)	pH	4.00			Value:	Value:	Value:
	Conductivity (mS/cm)	4.49			Value:	Value:	Value:
	Turbidity (NTU)	0			Value:	Value:	Value:
	DO (mg/L)	8.9-9.1 (ambient air)	NA	NA	Value:	Value:	Value:
Second Point Calibration	pH	6.86			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:
Third Point Calibration	pH	9.18			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:

Additional Remarks:

EQUIPMENT CALIBRATION DAILY LOG

Date: 2/20/19	Project Name: LHAAP-GWTP
Project Number: NWD1312.0150	Recorded By: Scott Beasinger

PID	Model: Mini RAe 3000		Bulb: 11.7 10.0 meV		Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #: P&M 7320						
	Parameter	Standard	Exp. Date	Lot #	Time: 0645	Time:	Time:
First Point Calibration	Vapor conc. (ppm)	0.0 (ambient air)	NA	NA	Initials: SB	Initials:	Initials:
Second Point Calibration	Vapor conc. (ppm)	100ppm (isobutylene)	10/5/21	JBH-248-100-19	Value: TO ZERO SB	Value:	Value:

COMB. GAS/O ₂ METER	Model:				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration	O ₂ (%)				Initials:	Initials:	Initials:
	% LEL Pentane				Value:	Value:	Value:

WATER QUALITY METER	Model:				Morning Calibration/Check	Evening Check (one point only)	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration (Auto)	pH	4.00			Initials:	Initials:	Initials:
	Conductivity (mS/cm)	4.49			Value:	Value:	Value:
	Turbidity (NTU)	0			Value:	Value:	Value:
	DO (mg/L)	8.9-9.1 (ambient air)	NA	NA	Value:	Value:	Value:
Second Point Calibration	pH	6.86			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:
Third Point Calibration	pH	9.18			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:

Additional Remarks:

EQUIPMENT CALIBRATION DAILY LOG

Date: 2/27/19 Project Name: LHAAP-GWTP
 Project Number: NWD1312.0150 Recorded By: Scott Beasinger

PID	Model: <u>Mini RAE 3000</u>		Bulb: <u>11.7</u> <u>10.0 meV</u>		Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #: <u>PGM 7320</u>						
	Parameter	Standard	Exp. Date	Lot #	Time: <u>0700</u>	Time:	Time:
First Point Calibration	Vapor conc. (ppm)	0.0 (ambient air)	NA	NA	Initials: <u>SB</u>	Initials:	Initials:
Second Point Calibration	Vapor conc. (ppm)	<u>100ppm</u> (isobutylene)	<u>10/5/21</u>	<u>JBH-248-100-19</u>	Value: <u>TO ZERO</u> <u>SB</u>	Value:	Value:

COMB. GAS/O ₂ METER	Model:				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration	O ₂ (%)				Initials:	Initials:	Initials:
	% LEL Pentane				Value:	Value:	Value:

WATER QUALITY METER	Model:				Morning Calibration/Check	Evening Check (one point only)	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration (Auto)	pH	4.00			Value:	Value:	Value:
	Conductivity (mS/cm)	4.49			Value:	Value:	Value:
	Turbidity (NTU)	0			Value:	Value:	Value:
	DO (mg/L)	8.9-9.1 (ambient air)	NA	NA	Value:	Value:	Value:
Second Point Calibration	pH	6.86			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:
Third Point Calibration	pH	9.18			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:

Additional Remarks:

EQUIPMENT CALIBRATION DAILY LOG						
Date: 3/7/19	Project Name: LHAAP- GWTP					
Project Number: NW01312.0150	Recorded By: Scott Bessinger					

PID	Model: Mini Rae 3000 Bulb: 11.7 100 meV				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #: PGM 7320						
	Parameter	Standard	Exp. Date	Lot #	Time: 0640	Time:	Time:
First Point Calibration	Vapor conc. (ppm)	0.0 (ambient air)	NA	NA	Initials: SB	Initials:	Initials:
Second Point Calibration	Vapor conc. (ppm)	100ppm (isobutylene)	10/5/21	JBH-248-100-19	Value: To zero SB	Value:	Value:

COMB. GAS/O ₂ METER	Model:				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration	O ₂ (%)				Initials:	Initials:	Initials:
	% LEL Pentane				Value:	Value:	Value:

WATER QUALITY METER	Model:				Morning Calibration/Check	Evening Check (one point only)	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration (Auto)	pH	4.00			Value:	Value:	Value:
	Conductivity (mS/cm)	4.49			Value:	Value:	Value:
	Turbidity (NTU)	0			Value:	Value:	Value:
	DO (mg/L)	8.9-9.1 (ambient air)	NA	NA	Value:	Value:	Value:
Second Point Calibration	pH	6.86			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:
Third Point Calibration	pH	9.18			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:

Additional Remarks:

EQUIPMENT CALIBRATION DAILY LOG						
Date: 3/13/19			Project Name: LHAAP-GWTP			
Project Number: NWD1312.0150			Recorded By: Scott Bessinger			

PID	Model: MiniRAE 3000		Bulb: 11.7 10.6 meV		Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #: PGM 7320						
	Parameter	Standard	Exp. Date	Lot #	Time: 0650	Time:	Time:
					Initials: SB	Initials:	Initials:
First Point Calibration	Vapor conc. (ppm)	0.0 (ambient air)	NA	NA	Value: To zero SB	Value:	Value:
Second Point Calibration	Vapor conc. (ppm)	100ppm (isobutylene)	10/5/21	JBH-248-100-19	Value: SB	Value:	Value:

COMB. GAS/O ₂ METER	Model:				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
					Initials:	Initials:	Initials:
First Point Calibration	O ₂ (%)				Value:	Value:	Value:
	% LEL Pentane				Value:	Value:	Value:

WATER QUALITY METER	Model:				Morning Calibration/Check	Evening Check (one point only)	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
					Initials:	Initials:	Initials:
First Point Calibration (Auto)	pH	4.00			Value:	Value:	Value:
	Conductivity (mS/cm)	4.49			Value:	Value:	Value:
	Turbidity (NTU)	0			Value:	Value:	Value:
	DO (mg/L)	8.9-9.1 (ambient air)	NA	NA	Value:	Value:	Value:
Second Point Calibration	pH	6.86			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:
Third Point Calibration	pH	9.18			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:

Additional Remarks:

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EQUIPMENT CALIBRATION DAILY LOG	
Date: <u>3/20/19</u>	Project Name: <u>LHAAP- G WTP</u>
Project Number: <u>NWD 1312.0150</u>	Recorded By: <u>Scott Beasinger</u>

PID	Model: <u>Mini RAE 3000</u>		Bulb: <u>11.7</u> meV		Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #: <u>PGM 7320</u>						
	Parameter	Standard	Exp. Date	Lot #	Time: <u>0710</u>	Time:	Time:
					Initials: <u>SB</u>	Initials:	Initials:
First Point Calibration	Vapor conc. (ppm)	0.0 (ambient air)	NA	NA	Value: <u>To zero SB</u>	Value:	Value:
Second Point Calibration	Vapor conc. (ppm)	<u>100ppm</u> (isobutylene)	<u>10/5/21</u>	<u>JBH-248-100-19</u>	Value: <u>SB</u>	Value:	Value:

COMB. GAS/O ₂ METER	Model:				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
					Initials:	Initials:	Initials:
First Point Calibration	O ₂ (%)				Value:	Value:	Value:
	% LEL Pentane				Value:	Value:	Value:

WATER QUALITY METER	Model:				Morning Calibration/Check	Evening Check (one point only)	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
					Initials:	Initials:	Initials:
First Point Calibration (Auto)	pH	4.00			Value:	Value:	Value:
	Conductivity (mS/cm)	4.49			Value:	Value:	Value:
	Turbidity (NTU)	0			Value:	Value:	Value:
	DO (mg/L)	8.9-9.1 (ambient air)	NA	NA	Value:	Value:	Value:
Second Point Calibration	pH	6.86			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:
Third Point Calibration	pH	9.18			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:

Additional Remarks:

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EQUIPMENT CALIBRATION DAILY LOG

Date: 3/28/19 Project Name: LHAAP- G WTP
 Project Number: NWD 1312.0150 Recorded By: Scott Bessinger

PID	Model: <u>Mini Rae 3000</u>		Bulb: <u>11.7</u> 10.6 meV		Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #: <u>PGM 7320</u>						
	Parameter	Standard	Exp. Date	Lot #	Time: <u>0645</u>	Time:	Time:
First Point Calibration	Vapor conc. (ppm)	0.0 (ambient air)	NA	NA	Initials: <u>SB</u>	Initials:	Initials:
Second Point Calibration	Vapor conc. (ppm)	<u>100ppm</u> (isobutylene)	<u>10/5/21</u>	<u>JBH-248-100-19</u>	Value: <u>TO ZERO SB</u>	Value:	Value:

COMB. GAS/O ₂ METER	Model:				Morning Calibration	Evening Check	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration	O ₂ (%)				Value:	Value:	Value:
	% LEL Pentane				Value:	Value:	Value:

WATER QUALITY METER	Model:				Morning Calibration/Check	Evening Check (one point only)	Additional Calib./Check (if necessary)
	Equipment ID #:						
	Parameter	Standard	Exp. Date	Lot #	Time:	Time:	Time:
First Point Calibration (Auto)	pH	4.00			Value:	Value:	Value:
	Conductivity (mS/cm)	4.49			Value:	Value:	Value:
	Turbidity (NTU)	0			Value:	Value:	Value:
	DO (mg/L)	8.9-9.1 (ambient air)	NA	NA	Value:	Value:	Value:
Second Point Calibration	pH	6.86			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:
Third Point Calibration	pH	9.18			Value:	Value:	Value:
	Conductivity (mS/cm)	53.7			Value:	Value:	Value:
	Turbidity (NTU)	100			Value:	Value:	Value:

Additional Remarks:



**Subject: Final Minutes, Quarterly Restoration Advisory Board (RAB) Meeting
Longhorn Army Ammunition Plant (LHAAP)**

Location of Meeting: Karnack Community Center, Karnack, Texas

Date of Meeting: April 25, 2019, 5:00-5:40 PM Central Daylight Time (CDT)

Meeting Participants:

Army BRAC:	Rose M. Zeiler
USACE:	Aaron Williams and John Doran
USAEC:	Andrew Maly
USGS:	Kent Becher
Bhate:	Kim Nemmers
APTIM:	William (Bill) Foss
USEPA Region 6:	Rich Mayer, Dorelle Harrison, and Janetta Coats
TCEQ:	April Palmie
USFWS:	Eric Duerkop
RAB:	Present: Judy VanDeventer, Paul Fortune, Carol Fortune, Charles Dixon, and Richard LeTourneau Absent: Terry Britt; John Pollard, Jr.; Tom Walker; and Nigel R. Shivers
Public:	Laura-Ashley Overdyke (Executive Director of the Caddo Lake Institute [CLI]), George Rice (CLI), John Fortune (New RAB Member), Sharron McAvoy (New RAB Member), Deon Hall (New RAB Member) Brad Eskue, Robert Speight, and James Saunders

An agenda for the RAB meeting, a color copy of the Bhate Environmental Associates, Inc. (Bhate) slide presentation, and handouts (see list at end of meeting minutes) were provided for meeting attendees.

Welcome and Introduction

Mr. Paul Fortune, RAB Installation Co-Chair, called the RAB meeting to order at 5:05 pm CDT. Mr. Fortune welcomed everyone and asked if there was anyone present that had not attended before. Mr. Aaron Williams stated that Mr. John Doran was new to the RAB Meeting. Mr. Doran stated that he was with the United States Army Corps of Engineers (USACE) and was assisting Mr. Williams on the project and had been with the USACE for about 6 months.

Ms. Rose M. Zeiler welcomed everyone and explained the purpose of the RAB meeting. Ms. Zeiler explained that the meeting is held every 3 months and then presented the mission for the RAB. Ms. Zeiler stated that it is important for everyone to ask questions or provide comments and that you can just raise your hand or speak up when you need to.

Administrative

Ms. Zeiler stated that there are three RAB applicants, who were John Fortune, Sharron McAvoy, and Deon Hall. Ms. Zeiler stated that ballots should have been handed out to RAB members. Ms. Zeiler emphasized how great it was that there were three applicants. Ms. Zeiler then explained that the charter calls for a vote and requires a 2/3 majority for approval. Two of the three



applicants were present and were asked to stand up. Mr. Hall was not present. Mr. John Fortune was one of the interested RAB members and explained that he had grown up in Karnack, worked at the Army Ammunition Plant for Thiokol and had recently moved back and wanted to participate. Ms. Sharron McAvoy stated her interest in environmental protection and that she was a member of the Sierra Club. Ms. Zeiler asked for the votes to be handed to her during the meeting. **Note:** votes were tallied by Mr. Paul Fortune and Ms. Zeiler at the end of the RAB Meeting, when it was announced that each of the three applicants were approved as new RAB members.

Open Items

Ms. Zeiler noted that the RAB Meeting minutes had been sent out in November 2018. Ms. VanDeventer made a motion to accept the October 2018 RAB Meeting minutes. Mr. Fortune seconded the motion. (**Note:** no meeting was held in January 2019 due to the partial Government shutdown).

Ms. Zeiler then reminded everyone of the ongoing outreach with the website address listed in the slide packet. Ms. Zeiler stated that the meeting minutes, ongoing activities, and information by site are on the website as well as the administrative record.

Defense Environmental Restoration

Ms. Kim Nemmers reminded everyone that copies of the slides were available and that Bhaté's role was to manage most of the sites as a contractor with the exception of LHAAP-29, LHAAP-18/24, and LHAAP-47. Ms. Nemmers then pointed out the slide with the acronyms and reviewed the agenda for the meeting. Ms. Nemmers then stated that three sites will be discussed by Mr. Bill Foss.

LHAAP-03

Mr. Foss provided an update on LHAAP-03, which was discussed about a year ago. Mr. Foss stated that LHAAP-03 is a very small site that was a former waste collection pad outside of the paint shop building. Site soils are contaminated with lead and arsenic. The Record of Decision (ROD) selected a remedy of excavation. Mr. Foss explained that additional soil samples were collected in November 2018 to both narrow down what area needed to be excavated and to determine if lateral or vertical delineation could be achieved to eliminate some of the confirmation sampling following excavation. Mr. Foss stated that a Remedial Design (RD)/ Remedial Action Work Plan (RAWP) was prepared using the older data and the data collected in November 2018. The RD/RAWP is currently in Army review. Mr. Foss stated that the plan is to excavate in late summer/early fall 2019. Mr. Foss explained that the map shows the area of excavation but noted that the scale makes it look like a large area. Mr. Foss also pointed out that much of the excavation is only down to 2 to 3 feet below ground surface (bgs). Mr. Foss explained that the yellow dots were the November 2018 samples.

LHAAP-04

Mr. Foss stated that LHAAP-04 is the location of a former pilot Wastewater Treatment Plant. In 1997, the structures were demolished. Then soil excavation was completed in 2009 for mercury and perchlorate contamination. The ROD selected a remedy for perchlorate treatment. Mr. Foss



explained that the first thing completed under Bhate's contract was to install monitoring wells and collect samples. The results indicated that the plume appeared to have shifted. So, the Army provided a scope of work to complete additional sampling. Mr. Foss explained that the plume used to be centered around monitoring well 04WW04. However, 04WW05 had perchlorate detections that increased. So, this indicated that the plume may have shifted to the west. The yellow circles are the additional data collected via direct push technology to evaluate the lithology and grab a groundwater sample. Then three new wells were installed based upon the data (04WW09, 04WW10, and 04WW11). The center of the mass of the plume did shift and this information was used to develop the RD. Mr. Foss explained that the RD will use in-situ bioremediation (ISB) which uses bacteria to degrade the contaminants. Mr. Foss explained that 25 direct push points will be advanced to allow for a little over 1,400 gallons of injectate (emulsified vegetable oil [EVO], nutrients, and water) to be added. Monitoring will be completed to evaluate the injections and RD. Land use controls (LUCs) will also be monitored which basically does not allow for groundwater use except for the environmental groundwater sampling. LUCs also restrict land use to prevent residential use. Mr. Foss then showed the location of the injection points and pointed out that the target area is where the contamination is 5 times the cleanup level (hot spot) and then there are a few locations to the south and west based upon the known plume migration. Mr. Foss stated that the RD was approved so the plan is to do the work in June or July 2019. Mr. Richard LeTourneau asked about backfilling. Mr. Foss stated that the site will be backfilled with soil that is either certified clean or sampled for offsite laboratory analysis to verify that the soil does not contain any contaminants above cleanup levels. Ms. Zeiler stated that this site had a substantial excavation previously.

LHAAP-17

Mr. Foss explained that LHAAP-17 was a former burning ground and flashing ground near the groundwater treatment plant (GWTP) and site LHAAP-18/24. The site was used for burning of TNT and flashing of powder. The waste was removed in the 1980s but contamination remains. Mr. Foss explained that the contaminants are mostly explosives, perchlorate, and solvents. The ROD that was signed in 2016 included groundwater extraction for the large perchlorate plume that has concentrations that are restrictive to natural attenuation. Mr. Foss explained that soil excavation of the explosives contaminated area will also be completed. Monitoring will be completed to evaluate the remedy, and LUCs will remain. Ms. Zeiler stated that there is a contingent remedy. Mr. Foss stated that the contingent remedy will apply if the reduction of perchlorate has not occurred in 18 months and includes another active remediation ahead of monitored natural attenuation.

Mr. Foss stated that previously the presentations at the RAB included a briefing on the pre-design investigation that included an aquifer pumping test and soil and groundwater sampling. The pumping test provided the basis for the type of system to install. The soil sampling provided the extent of soil requiring excavation. Based upon that information, the RD includes approximately 5,300 cubic yards of soil that is mostly 2 to 3 feet bgs. The excavation depth will be controlled by groundwater depth because soil excavation will not extend into the groundwater, which is expected to be at 7 feet bgs. Groundwater extraction will include up to three wells and will run for 18 months. The extracted groundwater will be piped to the GWTP. The contingent remedy



will be used if the perchlorate remains above 20,000 parts per million (ppm) following the extraction for 18 months. Mr. Foss explained that monitoring of the remedy will be completed and that LUCs will be in place. Mr. Foss stated that the LUCs prohibit groundwater use and limit site usage due to the perchlorate.

Mr. Foss explained the slide with the soil excavations. Mr. Foss stated that the areas furthest out from the site boundary were being excavated due to ecological concerns. The other deeper areas are due to explosives such as TNT. Mr. Foss then pointed out the deeper areas that were determined based upon the historic data as well as the newer data collected. Mr. Foss then showed the potential wells that will be used for groundwater extraction. Mr. Foss stated that wells 17WW02 and 17WW06 will be used for extraction. Groundwater results from 17WWW01 will determine if that well is brought online for extraction also. Mr. Foss explained the purple line is the current perchlorate plume.

Overview of Sites

Ms. Zeiler explained that several remedies are being implemented this summer. Mr. Foss stated that the areas need to dry-up following all the rain this winter/spring. Ms. Zeiler stated that the website would have a schedule of the work.

Ms. Nemmers explained that field work is completed, and then a report is prepared with that information. Also, remedial action operation (RA-O) sampling is completed semi-annually at a minimum. Ms. Nemmers explained that RA-O sampling is primarily sampling of monitoring wells to evaluate the remedy once it is in place. Ms. Nemmers stated that the ion exchange vessels at the GWTP were replaced and that the ion exchange is used to ensure that there is no perchlorate in the GWTP effluent.

Ms. Nemmers stated that the ongoing documents include the RD/RAWP for LHAAP-03 that Mr. Foss discussed. Ms. Nemmers stated that even though quite a bit of sampling has occurred, the list of documents for the sites are minimal because the reports are not written until additional sampling events are completed.

For the 3 month look ahead, Ms. Nemmers presented the four sites planned for field work (LHAAP-03, LHAAP-04, LHAAP-16, and LHAAP-17) in July or August and explained soil excavation will be conducted at two of the sites (LHAAP-03 and LHAAP-17). Ms. Nemmers explained that the schedule will be dependent upon the weather and that work will be moved up if possible.

Groundwater Treatment Plant

Ms. Nemmers stated that there were handouts with the information on the slides for the GWTP as well as handouts for the RDs of LHAAP-04 and LHAAP-17. Ms. Nemmers stated that the chart had a couple of dips since that last RAB meeting, which is due to power issues to the GWTP. Currently, the GWTP is on a generator. Ms. Judy VanDeventer asked if the site used Southwestern Electric Power Company (SWEPCO) to which Ms. Nemmers stated that the GWTP does use SWEPCO. Ms. VanDeventer pointed out that there has been power loss more frequently as of late from SWEPCO. Ms. Nemmers stated that the site has not been affected recently because it is pulling its power from the generator. Ms. Nemmers then pointed out that you will



see a big jump in water processed the next month following power outage because the GWTP operators are working to fix pumps and motors while the GWTP is not operating.

Surface Water Sampling

Ms. Nemmers presented the five locations sampled for surface water and stated that results are usually non-detect and well below the action level.

LHAAP-18/24, LHAAP-29 and LHAAP-47

Mr. Williams explained that a separate contractor, HDR, is responsible for developing the Proposed Plan (PP) for the preferred remedy followed by the ROD for LHAAP-18/24, LHAAP-29, and LHAAP-47. For LHAAP-18/24, the PP was finalized in February and the public meeting is after the RAB meeting. Mr. Williams encouraged everyone to stay stating that LHAAP-18/24 is considered to be the most contaminated site at Longhorn. Mr. Williams stated that the comment period goes through May 2, 2019. Mr. Williams stated that the expectation is to have the ROD drafted by September 2019. Ms. Zeiler then stated that no comments had been received yet.

Mr. Williams explained that the LHAAP-29 PP was finalized and the public meeting was held in December 2018. The ROD has been submitted to the regulators for review.

Mr. Williams stated that the LHAAP-47 Post-Screening Investigation data had been shared at the last RAB meeting in October 2018. Mr. Williams explained that many of the shallow wells are now dry and thus the plume has shrunk. In the intermediate wells, there are somewhat higher hits but overall the remedy remains the same. Therefore, Mr. Williams stated that the remedy remains the same and so a revision to the Draft Final ROD is being prepared for submittal to the regulators in August 2019.

Mr. Williams stated that in addition to the PP that is online you can also review the Final Revised Feasibility Study (FS) that details all of the remedial alternatives evaluated for LHAAP-18/24. The FS is in the administrative record.

Mr. Williams stated that some surface water sampling needed to be completed for LHAAP-47, which was completed in March 2019. The results are either non-detect or below the cleanup levels.

Next RAB Meeting Schedule and Closing Remarks

Ms. Zeiler then discussed the next meeting with the RAB members. It was decided that the next RAB Meeting will be held on **July 25, 2019**, with the **meeting starting at 6:00 pm CDT** at the Karnack Community Center.

Adjourn

Ms. VanDeventer made the motion to adjourn, and Mr. Paul Fortune seconded the motion. The meeting adjourned at 5:41 pm CDT.



April 2019 Meeting Attachments and Handouts:

- Meeting Agenda
- Color Copy of Bhate Presentation Slides
- Groundwater Treatment Plant (GWTP) – Processed Groundwater Volumes Handout
- Surface Water Sampling Handout
- Remedial Design Handout for LHAAP-17 and LHAAP-04



LONGHORN ARMY AMMUNITION PLANT
RESTORATION ADVISORY BOARD

Karnack, Texas
(479) 635-0110

AGENDA

DATE: Thursday, April 25, 2019
TIME: 5:00 – 6:00 PM
PLACE: Karnack Community Center, Karnack, Texas

05:00 **Welcome and Introduction**

05:05 **Open Items {RMZ}**

- Purpose of the RAB Meeting
- RAB Administrative Issues
 - o RAB Applicants
 - o Minutes (October 2018 RAB Meeting)
- Ongoing Outreach/Website

05:15 **Defense Environmental Restoration Program (DERP) Update {Bhate}**

- LHAAP-03 Field Work Status Update
- LHAAP-04 Field Work Status Update
- Documents and Field Work Completed since last RAB
- Three Month Lookahead
- Groundwater Treatment Plant (GWTP) Update

05:45 **Other Defense Environmental Restoration Program (DERP) Update {RMZ}**

- LHAAP-18/24 Proposed Plan
- LHAAP-29 Record of Decision and Responsiveness Summary
- LHAAP-47 Pre-Screening Investigation (PSI) Update
- Five Year Review Update

05:55 **Next RAB Meeting Schedule and Closing Remarks {RMZ}**

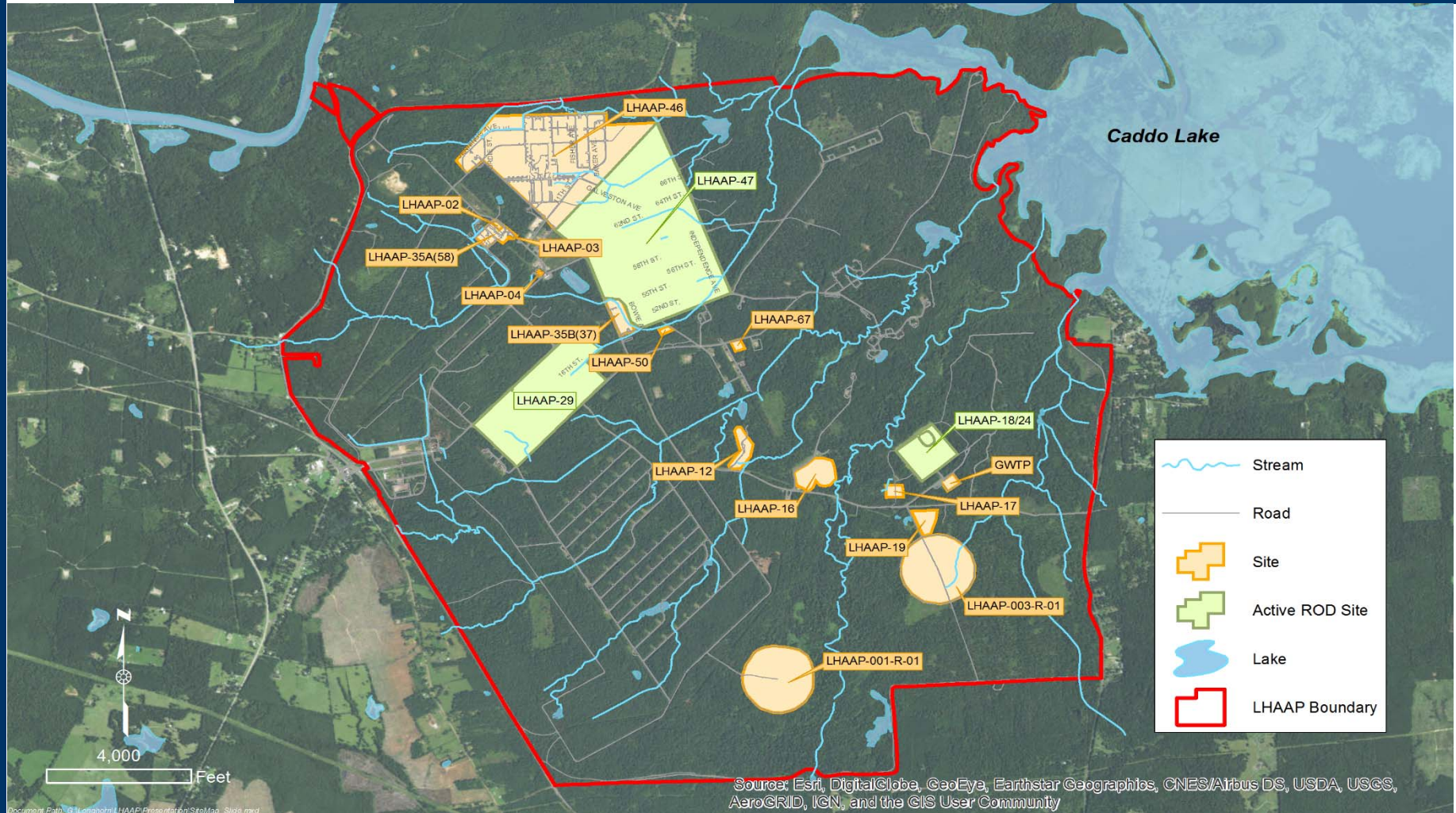
Longhorn Army Ammunition Plant Quarterly Restoration Advisory Board Meeting

Karnack Community Center
April 25, 2019
5:00 PM CDT



Restoration Advisory Board Meeting

Site Map



Restoration Advisory Board Meeting

Abbreviations and Acronyms

µg/L	Micrograms per liter	PCL	Protective Concentration Level
DERP	Defense Environmental Restoration Program	PDI	Pre-Design Investigation
ECP	Environmental Condition of Property	PSI	Pre-Screening Investigation
EISB	Enhanced In-situ Bioremediation	RAB	Restoration Advisory Board
FBR	Fluidized Bed Reactor	RA(O)	Remedial Action Operation
ft bgs	Feet below ground surface	RAWP	Remedial Action Work Plan
GWTP	Groundwater Treatment Plant	RD	Remedial Design
ISB	In-Situ Bioremediation	ROD	Record of Decision
LHAAP	Longhorn Army Ammunition Plant	TCEQ	Texas Commission on Environmental Quality
LUC	Land Use Control	TRRP	Texas Risk Reduction Program
MNA	Monitored natural attenuation	USEPA	U.S. Environmental Protection Agency
		VOCs	Volatile organic compounds

Restoration Advisory Board Meeting

Agenda

- 05:00** Welcome and Introduction
- 05:05** Open Items {RMZ}
- Purpose of the Restoration Advisory Board (RAB) Meeting
 - RAB Administrative Issues
 - RAB Applicants
 - Minutes (October 2018 RAB Meeting)
 - Ongoing Outreach/Website
- 05:15** Defense Environmental Restoration Program (DERP) Update {Bhate}
- LHAAP-03 Field Work Status Update
 - LHAAP-04 Field Work Status Update
 - LHAAP-17 Remedial Design Update
 - Documents and Field Work Completed Since Last RAB
 - Three Month Look Ahead
 - Groundwater Treatment Plant (GWTP) Update
- 05:45** Other DERP Update {AW}
- LHAAP-18/24 Proposed Plan
 - LHAAP-29 Record of Decision (ROD) and Responsiveness Summary
 - LHAAP-47 Pre-Screening Investigation (PSI) Update
 - Five Year Review Update
- 05:55** Next RAB Meeting Schedule and Closing Remarks {RMZ}

Restoration Advisory Board Meeting

Purpose of the RAB Meeting

- Held every 3 months
- The mission of the Longhorn Army Ammunition Plant (LHAAP) RAB is to promote community awareness and obtain constructive community review and comments on environmental restoration activities at the former LHAAP

Restoration Advisory Board Meeting

The Army Wants You to be Informed

- **The Army is committed to protecting human health and the environment; key to that commitment is engaging the community and increasing public participation in environmental restoration at LHAAP**
- **You are encouraged to:**
 - **Attend RAB Meetings and/or become a member of the RAB**
 - **Visit the Longhorn environmental website at www.longhornaap.com**
 - **Website is regularly updated to indicate the upcoming field events at each site including groundwater sampling, monitoring well installations, soil sampling, or remediation activities**
 - **Make suggestions for improving communication – the Army welcomes and appreciates community feedback**

Restoration Advisory Board Meeting

RAB Administrative Issues

- RAB Membership
- Discussion of October 2018 RAB Meeting minutes/motion to accept



Restoration Advisory Board Meeting

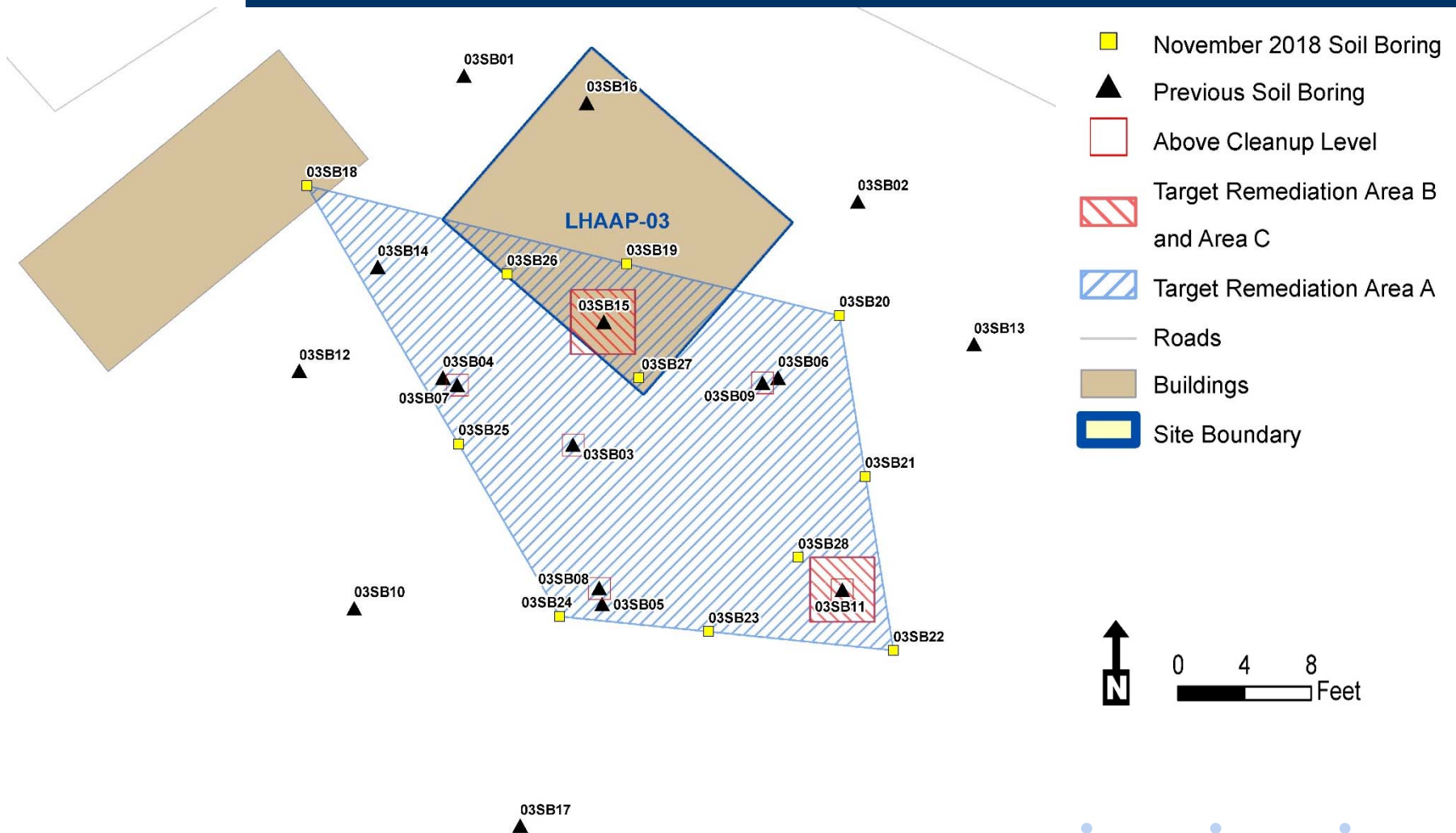
LHAAP-03 Pre-Excavation Soil Sampling

- **Site Background**
 - LHAAP-03 is the site of a former Waste Collection Pad for the Building 722-P Paint Shop
 - Building 722-P and the surrounding structures have been demolished
 - Soil is contaminated with arsenic and lead at concentrations that could be a risk to groundwater and the ROD selected excavation and offsite disposal as the remedy
 - Groundwater is being addressed as part of site LHAAP-35A(58)
- **Recent Activities**
 - Pre-excavation soil samples were collected in late November 2018 from locations surrounding the excavation area defined in the ROD
 - Samples were used to better define the area where excavation is required
 - Soil sample data will be included in the Remedial Design (RD) and Remedial Action Work Plan (RAWP)
 - RD/RAWP is currently in preparation for submittal to the U.S. Environmental Protection Agency (USEPA) and Texas Commission on Environmental Quality (TCEQ)



Restoration Advisory Board Meeting

LHAAP-03 Pre-Excavation Soil Sampling



Restoration Advisory Board Meeting

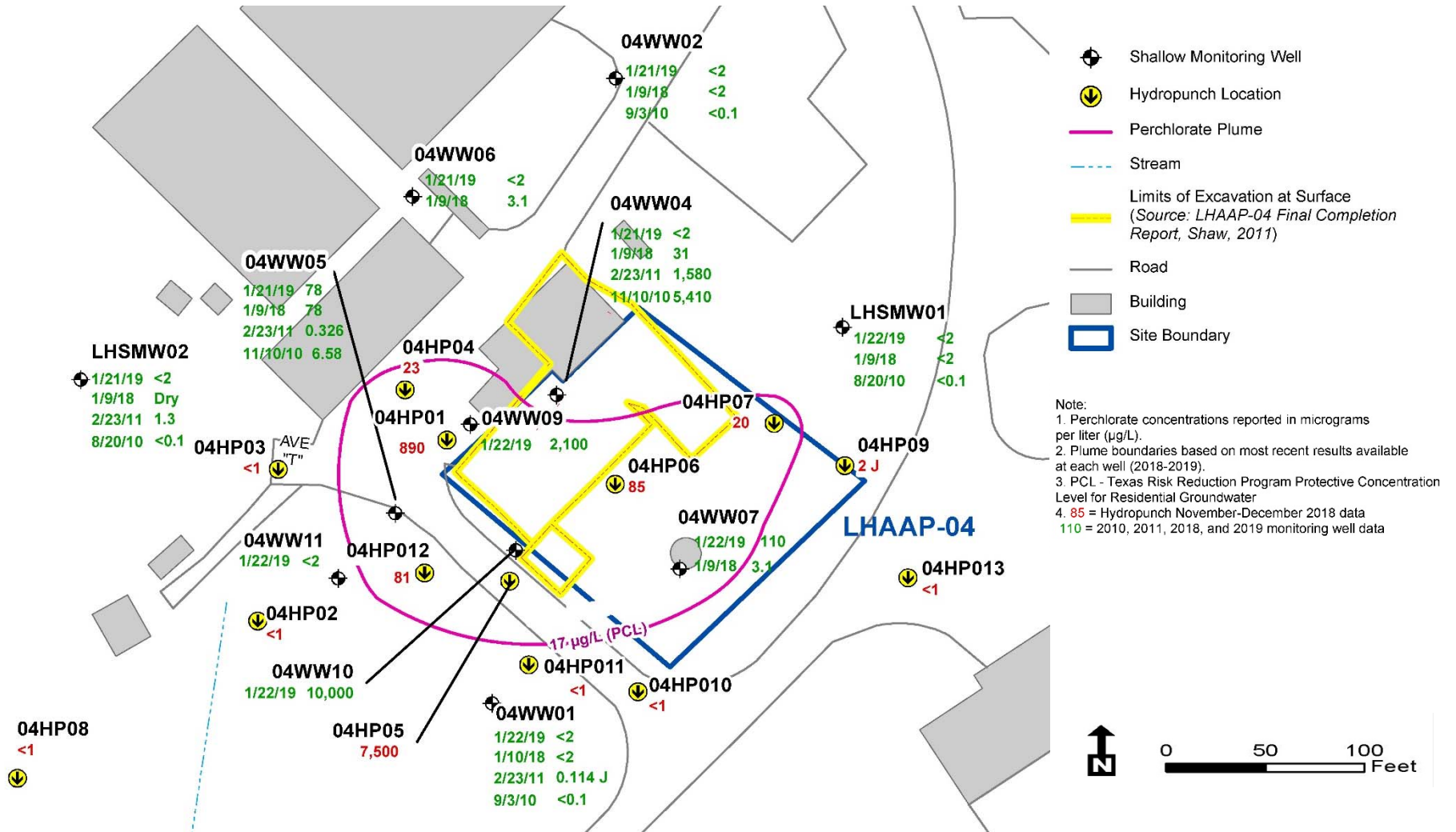
LHAAP-04 Remedial Design

- **Site Background**
 - LHAAP-04 is the site of the former Pilot Wastewater Treatment Plant, located near the former Fire Station
 - Demolition of the structures and disposal of associated wastes was conducted in 1997
 - Soil contaminated with mercury and perchlorate was excavated in 2009
 - The ROD published in October 2016 selected In-Situ Bioremediation (ISB), Long-Term Monitoring, and Land Use Controls (LUCs) as the remedy for groundwater
- **Recent Activities**
 - Groundwater sampling in January 2018 revealed that the groundwater plume may have migrated since the previous sampling in 2010-2011
 - Additional direct-push groundwater sampling was performed in November-December 2019 and additional monitoring wells were installed in January 2019
 - January 2019 sampling of the new and existing wells confirmed that the plume had migrated slightly to the southwest, but is still adequately delineated



Restoration Advisory Board Meeting

LHAAP-04 2010-2019 Perchlorate Data



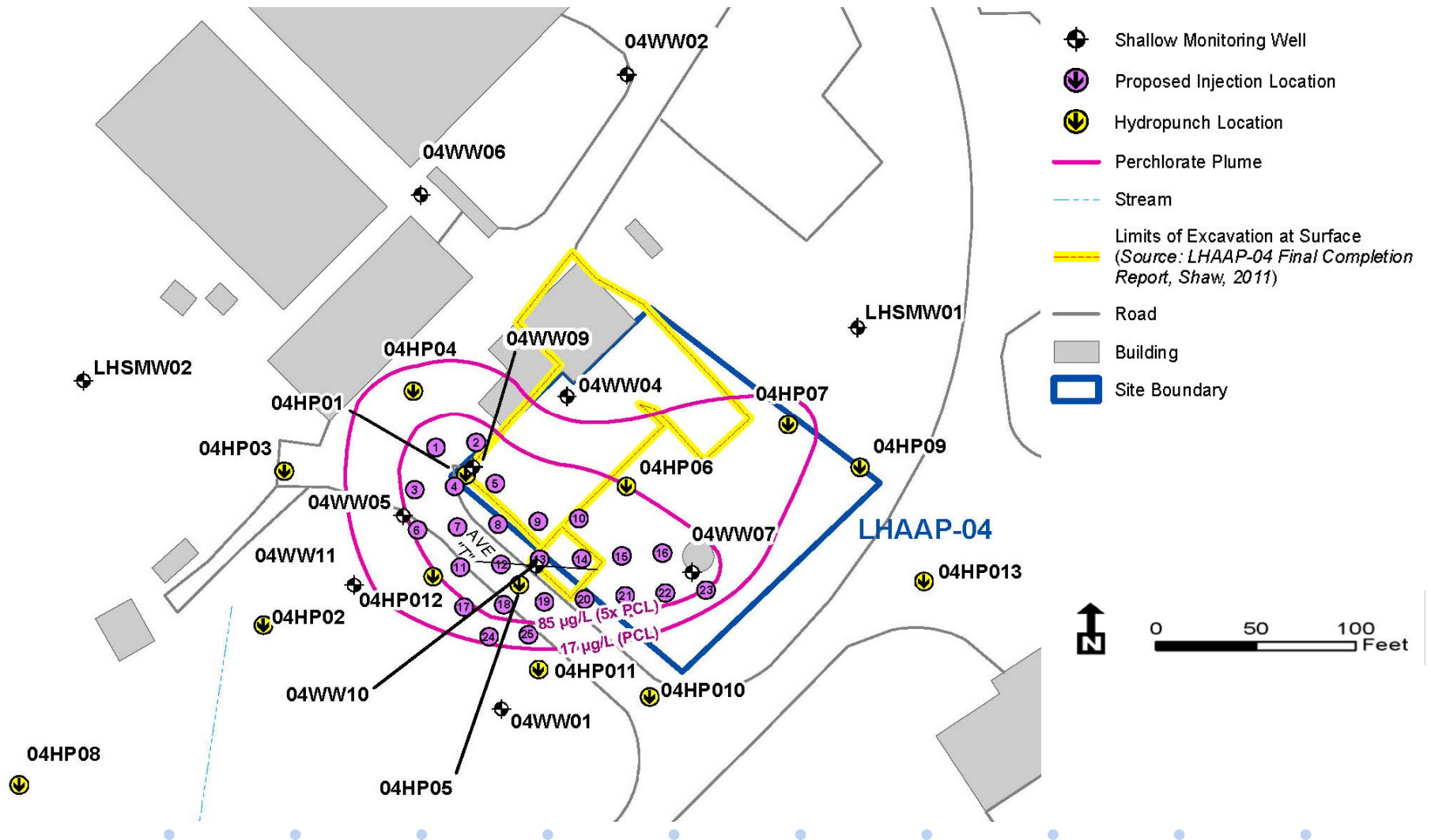
Restoration Advisory Board Meeting

LHAAP-04 Remedial Design

- **Remedial Design**
 - ISB will be implemented for the hot-spot defined as 5 times the Texas Risk Reduction Program (TRRP) Protective Concentration Level (PCL) for Residential Groundwater (17 micrograms per liter [$\mu\text{g}/\text{L}$])
 - 25 direct push injection locations will be used to inject emulsified vegetable oil (EVO)
 - Each location will receive approximately 15 gallons of EVO, 6 gallons of nutrients, and 1,463 gallons of water to treat a radius of approximately 10 feet around the location
 - Injection grid is spaced approximately 20-25 feet apart and shifted slightly to the south and west to account for future migration of the plume
- **Long-Term Monitoring and LUCs**
 - Baseline sampling of all site wells prior to injections, quarterly sampling for the first 2 years, semi-annual sampling for years 3 through 5, and annual sampling thereafter
 - LUCs include prohibition on use of groundwater (except for environmental monitoring), restriction to non-residential land use, and maintenance of remediation and monitoring systems
 - LUCs will remain in place until the concentration of perchlorate allows for unrestricted use and unlimited exposure

Restoration Advisory Board Meeting

LHAAP-04 Injection Plan



Restoration Advisory Board Meeting

LHAAP-17 Remedial Design

- **Site Background**
 - LHAAP-17 is the site of the former Burning Ground No. 2/Flashing Ground used from 1959 to 1980 for burning of bulk TNT, photo flash powder, and reject material from Universal Match Corporation
 - Waste material was reportedly removed from the burning trenches in 1984
 - Contaminants include explosives and metals in soil, and perchlorate and chlorinated solvents in groundwater
 - The ROD published in August 2016 selected Groundwater Extraction, Monitored Natural Attenuation (MNA), Soil Excavation, Long-Term Monitoring, and LUCs as the remedy
- **Recent Activities**
 - Pre-Design Investigation (PDI) (aquifer pumping test and soil and groundwater sampling) conducted in January 2018
 - Groundwater sampling to assess current plume conditions
 - Soil sampling refined the extent of the soil contamination requiring excavation
 - Aquifer pumping test provided design basis for the groundwater extraction system design

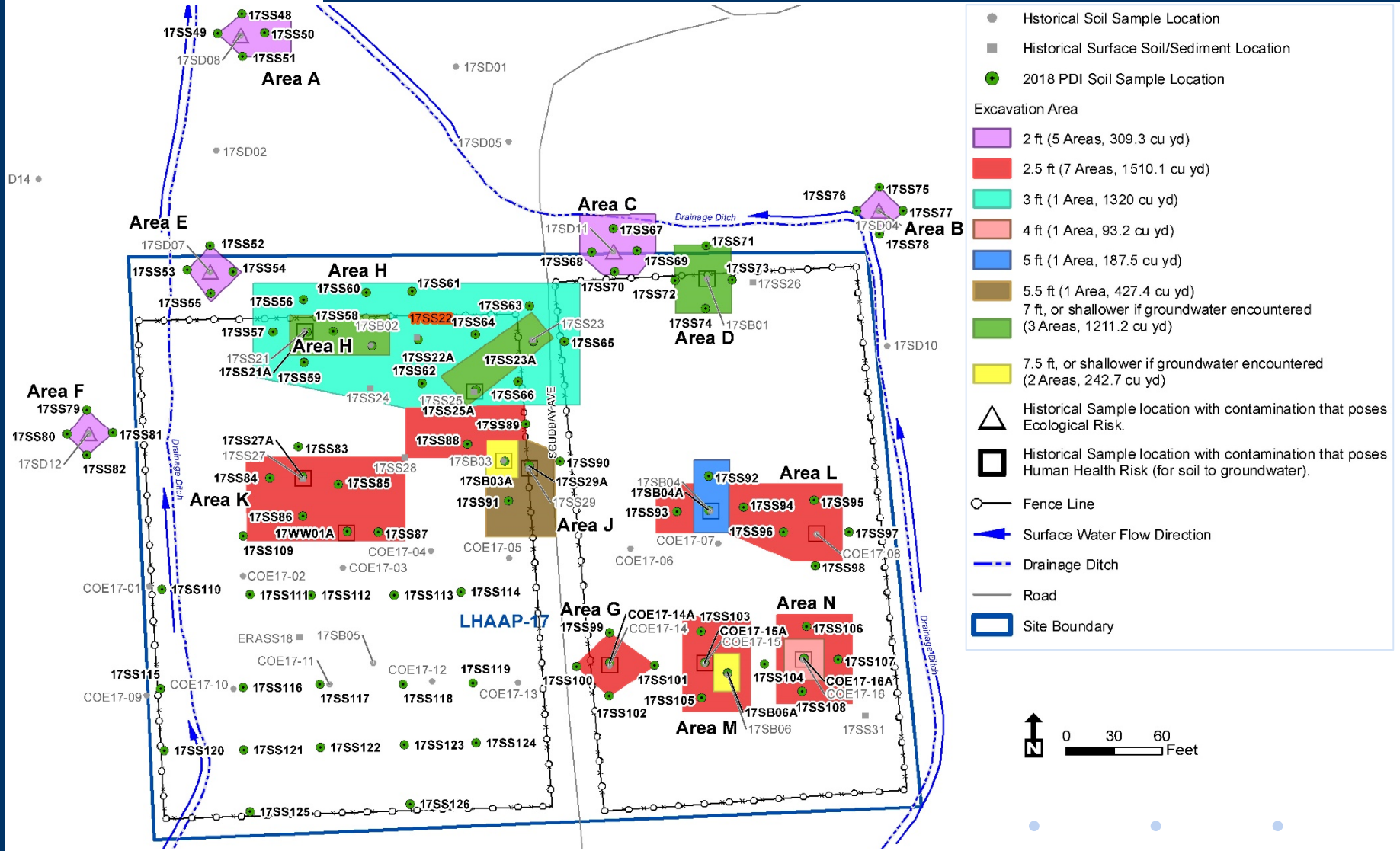
Restoration Advisory Board Meeting

LHAAP-17 Remedial Design

- **Remedial Design**
 - Approximately 5,300 in-place cubic yards of soil will be excavated based on the previous soil sampling data and transported to an offsite licensed disposal facility
 - Excavation will be backfilled with clean soil once sampling confirms contaminated soil has been removed
 - Groundwater extraction will be conducted for 18 months in up to three wells to reduce perchlorate concentrations to less than 20,000 $\mu\text{g}/\text{L}$
 - MNA will be the remedy for chlorinated solvents and perchlorate in groundwater unless perchlorate remains above 20,000 $\mu\text{g}/\text{L}$
- **Long-Term Monitoring and LUCs**
 - Extraction: Baseline sampling prior to extraction, monthly sampling for first 6 months of extraction, quarterly sampling for last 12 months of extraction
 - MNA: quarterly sampling for the first 2 years, semi-annual sampling for years 3 through 5, and annual sampling thereafter
 - LUCs include prohibiting use of groundwater (except for environmental monitoring), restricting land use to non-residential, and maintaining remediation/monitoring systems
 - LUCs will remain in place until the concentration of perchlorate allows for unrestricted use and unlimited exposure

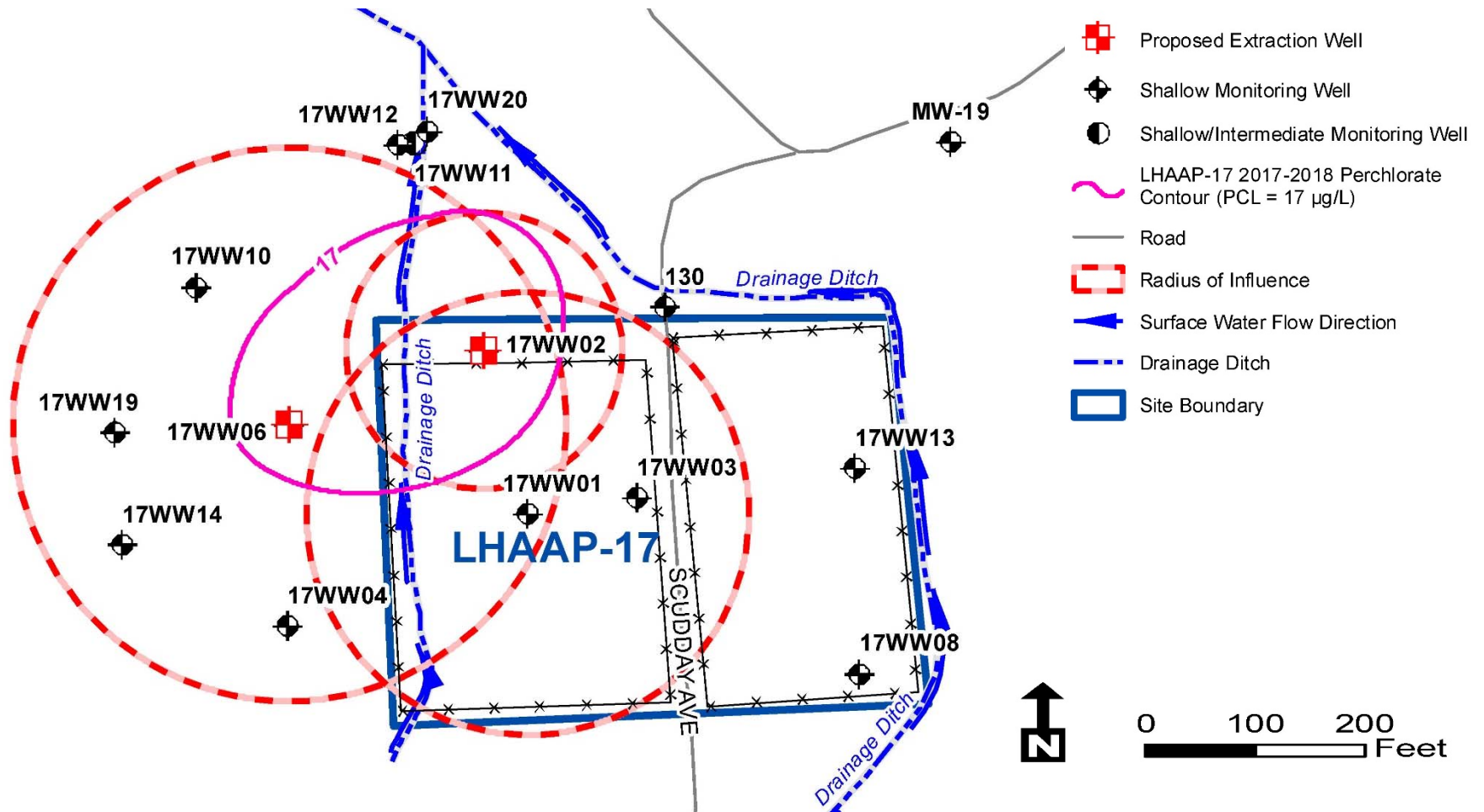
Restoration Advisory Board Meeting

LHAAP-17 Soil Excavation Areas



Restoration Advisory Board Meeting

LHAAP-17 Groundwater Extraction



Restoration Advisory Board Meeting

Completed Field Work Since Last RAB Meeting

Site	Activity
LHAAP-03	Supplemental Soil Sampling – November 2018
LHAAP-04	Supplemental Groundwater Sampling – November 2018 to January 2019
LHAAP-12	Remedial Action Operation (RA(O)) Sampling – December 2018
LHAAP-16	Annual Compliance Sampling – February 2019
LHAAP-37	RA(O) Sampling – November 2018, February 2019
LHAAP-46	RA(O) Sampling – February 2019
LHAAP-50	RA(O) Sampling – November 2018
LHAAP-58	RA(O) Sampling –December 2018, March 2019
LHAAP-67	RA(O) Sampling – October/November 2018
LHAAP-001-R	Groundwater Sampling – November 2018
LHAAP-001-R and LHAAP-003-R	Annual LUC Report-Year 1
GWTP	Replaced and disposed of ion exchange vessels (used to polish groundwater for perchlorate)
LHAAP-18/24	RA(O) Sampling – December 2018

Restoration Advisory Board Meeting

Documents in Process

Site	Document
LHAAP-03	Remedial Design and Remedial Action Work Plan
LHAAP-12	Annual RA(O) Report
GWTP	Quarterly Evaluation 4 th Quarter (October - December 2018) Quarterly Evaluation 1 st Quarter (January – March 2019)

Restoration Advisory Board Meeting

3 Month Look Ahead - Field Work

Site	Activity
LHAAP-03	Complete soil excavation
LHAAP-04	Complete ISB injections
LHAAP-16	Complete well installations and ISB injections
LHAAP-17	Complete soil excavation and extraction system installation
LHAAP-37	RA(O) Sampling – May 2019
LHAAP-50	RA(O) Sampling – May 2019
LHAAP-58	RA(O) Sampling – June 2019
LHAAP-67	RA(O) Sampling – May 2019
LHAAP-18/24	RA(O) Sampling – June 2019

Restoration Advisory Board Meeting

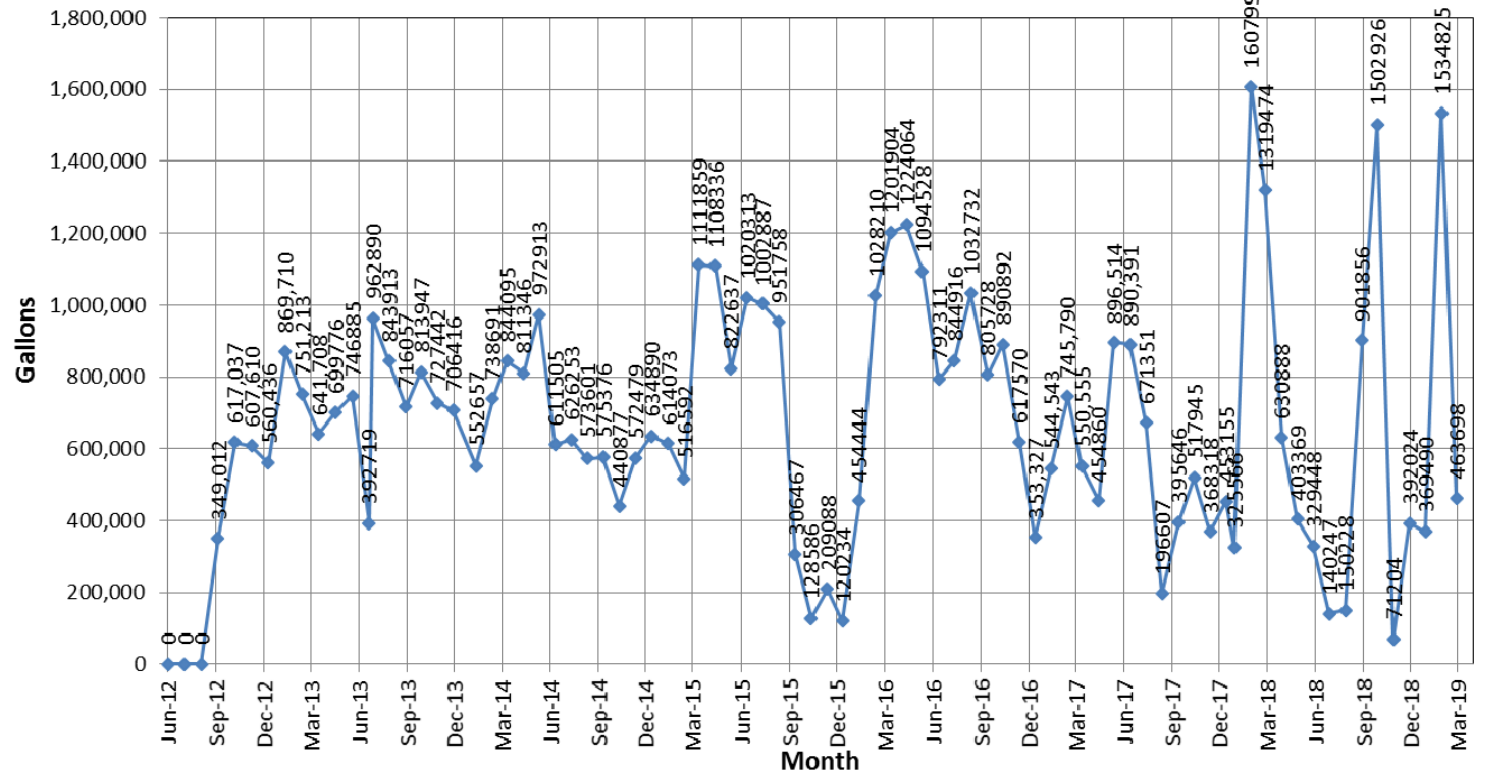
3 Month Look Ahead - Documents

Site	Document
LHAAP-03	Remedial Design and Remedial Action Work Plan
LHAAP-12	2018 RA(O) Report
GWTP, LHAAP-16, and LHAAP-18/24	Quarterly Evaluation Report: Fourth Quarter (October – December) 2018 Quarterly Evaluation Report: First Quarter (January – March 2019)

Restoration Advisory Board Meeting

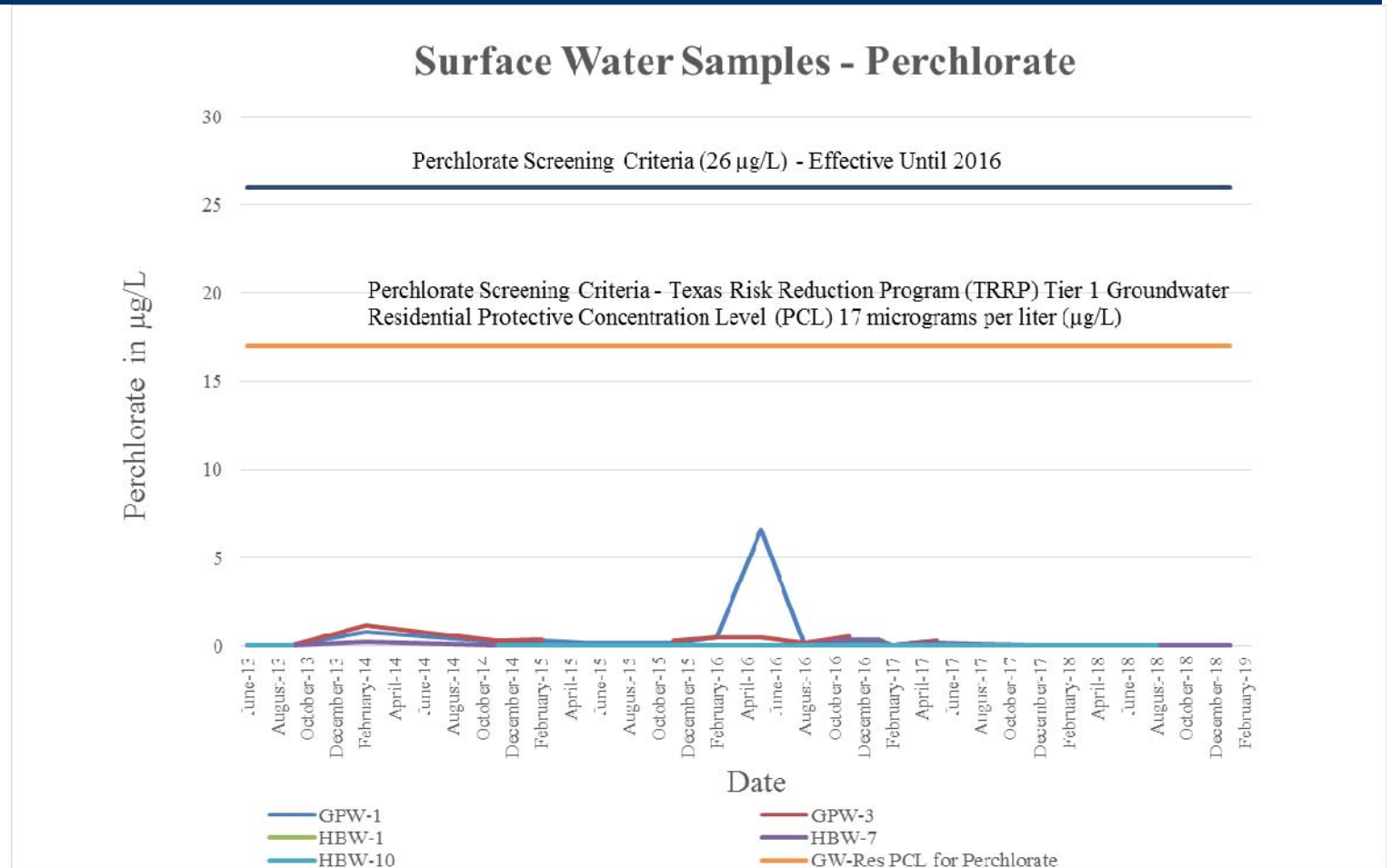
GWTP Update

Treated Groundwater Discharged Monthly from June 2012 through March 2019



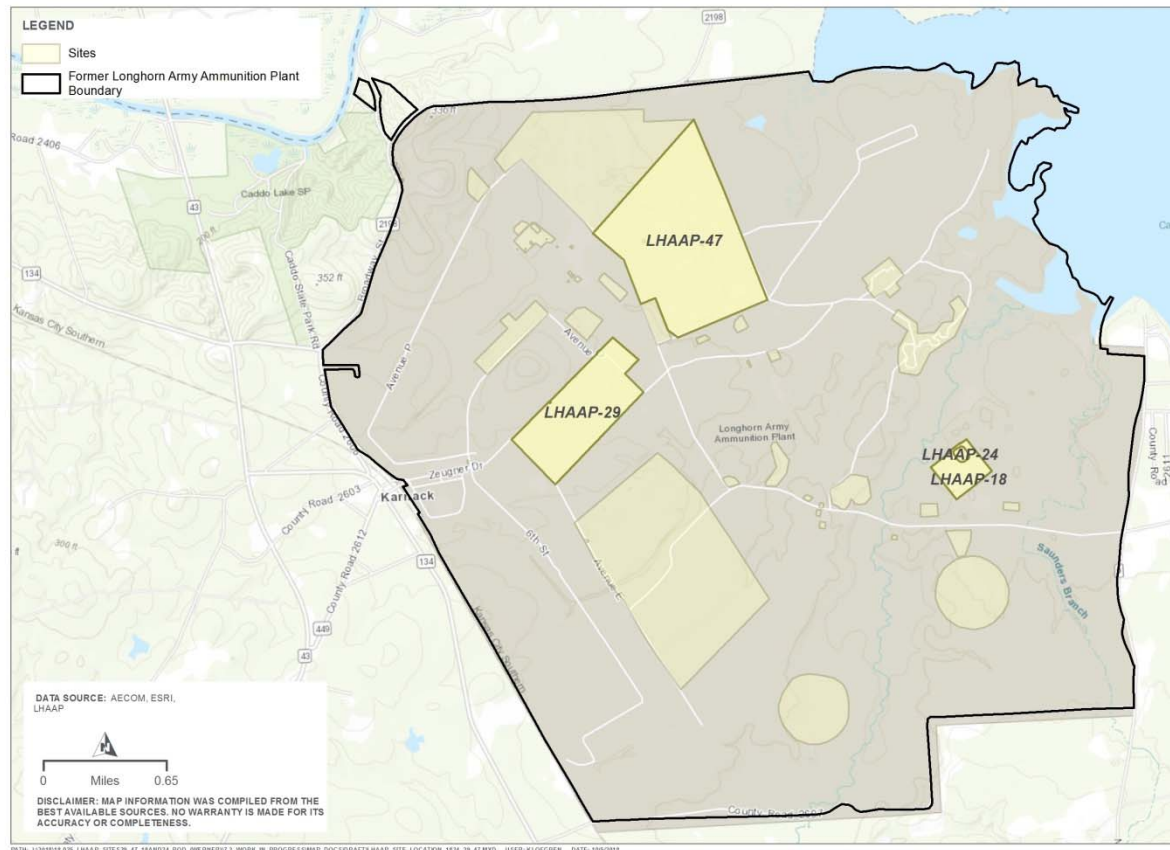
Restoration Advisory Board Meeting

Surface Water Sample Results



Restoration Advisory Board Meeting

LHAAP-18/24, 29, 47 Status Update



Restoration Advisory Board Meeting

LHAAP-18/24, 29 & 47 Document Status

- **LHAAP-18/24**
 - Proposed Plan finalized February 2019
 - Public meeting April 25, 2019, 6:00pm – 7:30pm
 - Public comment period April 2 to May 2, 2019
 - Draft Record of Decision submittal planned for September 2019
- **LHAAP-29**
 - Proposed Plan finalized November 2018 and public meeting was held December 6, 2018.
 - Draft Record of Decision submittal planned for May 2019
- **LHAAP-47**
 - Post Screening Investigation Report finalized April 2019
 - Revised Draft Final Record of Decision submittal planned for August 2019



Restoration Advisory Board Meeting

Feasibility Study for LHAAP-18/24

- LHAAP-18/24
 - Final Revised Feasibility Study located in the Administrative Record, Volume 1, 2017, Bate Stamp 00692951 - 00731961

Administrative Record located on the Longhorn environmental website at www.longhornaap.com

Restoration Advisory Board Meeting

LHAAP-47 Field Work Update

- **Work Completed**
 - Collected 4 surface water samples March 2019



Restoration Advisory Board Meeting

Next RAB Meeting Schedule & Closing Remarks

- Schedule July 2019 RAB Meeting
- Other Issues/Remarks
- Thank you for coming

Groundwater Treatment Plant - Processed Groundwater Volumes

The amount of groundwater treated is determined by measuring the number of gallons of processed water discharged.

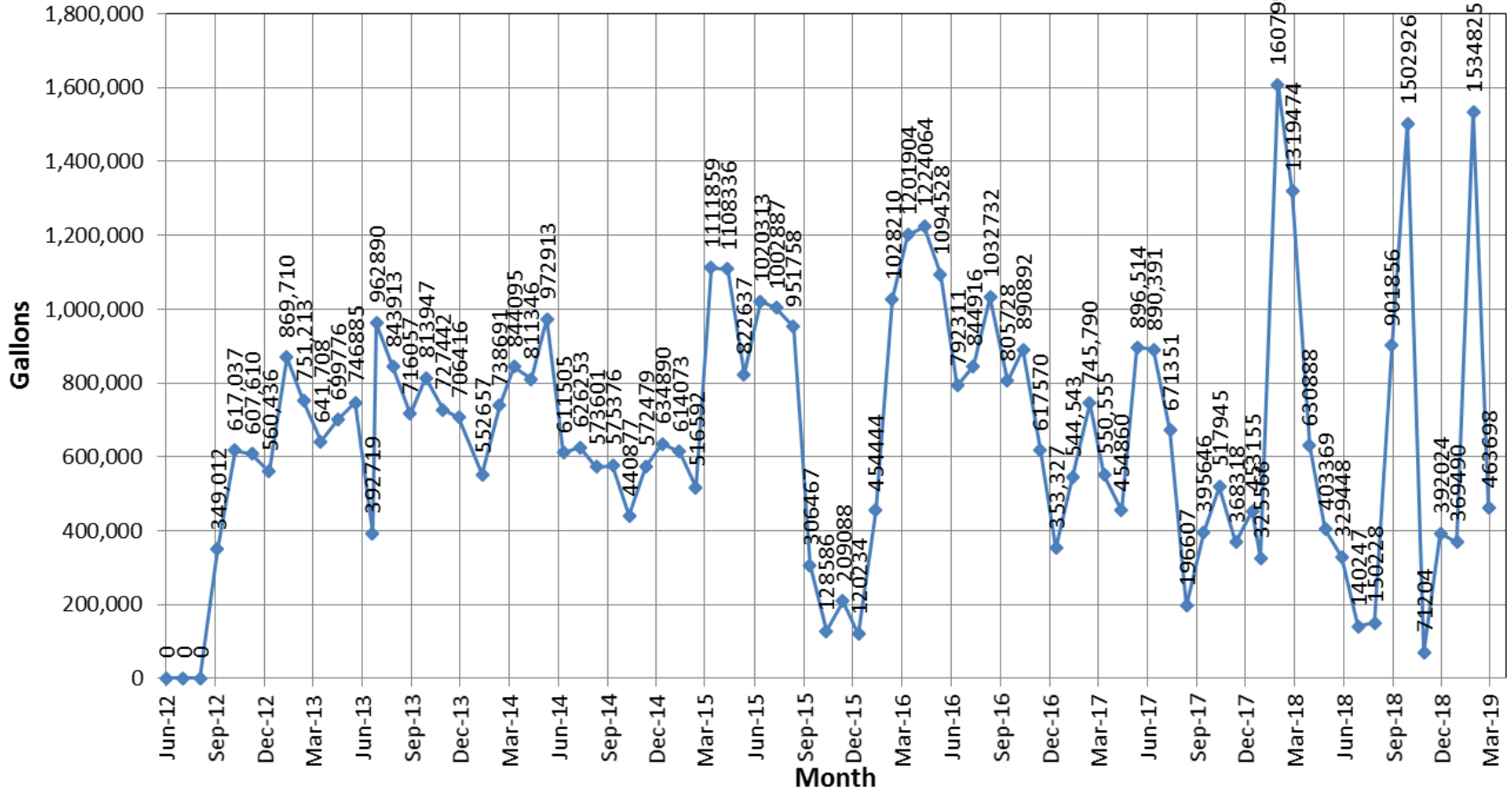
Processed Water Discharged Data

(in gallons)

Oct-07	Nov-07	Dec-07	Jan-08	Feb-08	Mar-08	Apr-08	May-08	Jun-08	Jul-08	Aug-08	Sep-08
1,041,491	848,356	804,822	792,148	665,883	818,872	791,306	568,812	776,904	748,377	690,052	617,199
Oct-08	Nov-08	Dec-08	Jan-09	Feb-09	Mar-09	Apr-09	May-09	Jun-09	Jul-09	Aug-09	Sep-09
655,059	619,274	726,118	552,299	598,144	433,800	488,807	526,958	387,644	0	414,853	735,716
Oct-09	Nov-09	Dec-09	Jan-10	Feb-10	Mar-10	Apr-10	May-10	Jun-10	Jul-10	Aug-10	Sep-10
808,322	636,306	727,492	391,898	695,343	802,656	894,731	962,121	1,257,977	1,314,924	1,041,495	1,136,547
Oct-10	Nov-10	Dec-10	Jan-11	Feb-11	Mar-11	Apr-11	May-11	Jun-11	Jul-11	Aug-11	Sep-11
956,567	705,805	849,712	811,679	668,281	1,090,348	817,325	900,338	916,552	784,369	652,524	733,456
Oct-11	Nov-11	Dec-11	Jan-12	Feb-12	Mar-12	Apr-12	May-12	Jun-12	Jul-12	Aug-12	Sep-12
748,102	658,250	684,903	865,453	725,000*	730,000*	980,000*	630,000*	0	0	0	349,012
Oct-12	Nov-12	Dec-12	Jan-13	Feb-13	Mar-13	Apr-13	May-13	Jun-13	Jul-13	Aug-13	Sep-13
617,037	607,610	560,436	869,710	751,213	641,708	699,776	746,885	392,719	962,890	843,913	716,057
Oct-13	Nov-13	Dec-13	Jan-14	Feb-14	Mar-14	Apr-14	May-14	Jun-14	Jul-14	Aug-14	Sep-14
813,974	727,442	706,416	552,657	738,691	844,095	811,346	972,913	611,505	626,253	573,601	575,376
Oct-14	Nov-14	Dec-14	Jan-15	Feb-15	Mar-15	Apr-15	May-15	Jun-15	Jul-15	Aug-15	Sep-15
440,877	572,479	634,890	614,073	516,592	1,111,859	1,108,336	822,637	1,020,313	1,002,887	951,758	306,467
Oct-15	Nov-15	Dec-15	Jan-16	Feb-16	Mar-16	Apr-16	May-16	Jun-16	Jul-16	Aug-16	Sep-16
128,586	209,088	120,234	454,444	1,028,210	1,201,904	1,224,064	1,094,528	792,311	844,916	1,032,732	805,728
Oct-16	Nov-16	Dec-16	Jan-17	Feb-17	Mar-17	Apr-17	May-17	Jun-17	Jul-17	Aug-17	Sep-17
890,892	617,570	353,327	544,543	745,790	550,555	454,860	896,514	890,391	528,538	195,198	961,324
Oct-17	Nov-17	Dec-17	Jan-18	Feb-18	Mar-18	Apr-18	May-18	Jun-18	Jul - 18	Aug-18	Sep-18
517,945	368,318	453,155	325,566	1,607,996	1,319,474	630,888	403,369	329,448	140,247	150,228	901,856
Oct-18	Nov-18	Dec-18	Jan-19	Feb-19	Mar-19						
1,502,926	71,204	392,024	369,490	1,534,825	463,698						

*Indicates Estimate

Treated Groundwater Discharged Monthly from June 2012 through March 2019



Water Discharge Location and Volume (Gallons)

Month	Total Combined to Harrison Bayou	LHAAP-18/24 Sprinklers	GWTP To INF Pond	INF Pond to Harrison Bayou	Contract Hauled Off-Site
Dec-16	0	236,688	0	0	0
Jan-17	0	0	0	0	0
Feb-17	0	0	0	0	14,355
Mar-17	127,242	0	0	0	14,400
Apr-17	113,038	0	236,821	0	0
May-17	205,665	0	534,155	0	0
Jun-17	467,830	0	294,550	490,574	0
Jul-17	0	0	528,538	0	0
Aug-17	0	0	195,197	0	0
Sep-17	0	0	309,980	651,434	0
Oct-17	0	0	517,945	0	0
Nov-17	0	0	368,318	0	0
Dec-17	0	0	453,155	560,350	0
Jan-18	325,566	0	253,177	325,566	0
Feb-18	1,607,996	0	62,017	1,430,634	0
Mar-18	1,319,474	0	0	870,816	0
Apr-18	630,888	0	0	630,888	0
May-18	403,369	0	0	403,369	0
Jun-18	193,669	0	135,779	0	0
Jul -18	0	0	140,247	0	0
Aug -18	49,409	0	100,819	0	0
Sep-18	585,397	0	316,459	524,484	0
Oct-18	1,409,106	0	93,820	1,016,285	0
Nov-18	71,204	0	0	0	0
Dec-18	392,024	0	0	0	0
Jan-19	369,490	0	0	369,490	0
Feb-19	1,534,825	0	0	1,326,485	0
Mar-19	463,698	0	0	83,250	0

Harrison Bayou and Goose Prairie Creek – Perchlorate Data

Surface water samples are collected quarterly from each location in Harrison Bayou and Goose Prairie Creek, unless the sampling location is dry.

Surface Water Sample Data (in micrograms per liter)

Quarter	3 rd	4 th	1 st	2 nd	3 rd	4 th	1 st	2 nd	3 rd	4 th	1 st
Creek Sample ID	Jul 1999	Sep 1999	Feb 2000	Apr 2000	Aug 2000	Dec 2000	Feb 2001	Apr 2001	July 2001	Oct 2001	Jan 2002
GPW-1	<1.0U	-	4	<4.0 U	<4.0 U	<4.0 U	-	2.65	<4.0 U	<4.0 U	<4.0 U
GPW-3	<1.0U	<4.0 U	17	8	<4.0 U	<4.0 U	-	2.28	<4.0 U	<4.0 U	<4.0 U
HBW-1	-	<80.0 U	310	23	-	-	<4.0 U	-	<4.0 U	<4.0 U	<4.0 U
HBW-7	-	<8.0 U	370	110	-	-	<4.0 U	-	<4.0 U	<4.0 U	<4.0 U
HBW-10	-	<8.0 U	905	650	<4.0 U	-	<4.0 U	-	<4.0 U	-	-

Quarter	2 nd	3 rd	4 th	1 st	2 nd	3 rd	3 rd	4 th	2 nd	3 rd	4 th
Creek Sample ID	June 2002	Sept 2002	Dec 2002	Feb 2003	June 2003	Aug 2003	July 2004	Dec 2006	May 2007	Aug 2007	Dec 2007
GPW-1	<4.0 U	<4.0 U	18.3	18.6	59.9	-	2.25	-	<1.0 U	<1.0 U	10.7
GPW-3	<4.0 U	<4.0 U	5.49	12.6	14.7	-	2.2	-	<1.0 U	<1.0 U	7.48
HBW-1	<4.0 U	<4.0 U	<4.0 U	-	<4.0 U	99.3	<0.2U	<1.0 U	<1.0 U	122	<1.0 U
HBW-7	<4.0 U	<4.0 U	<4.0 U	-	<4.0 U	<4.0 U	<0.2U	<1.0 U	<1.0 U	1.02	<1.0 U
HBW-10	<4.0 U	<4.0 U	<4.0 U	-	<4.0 U	-	<0.2U	<1.0 U	<1.0 U	<1.0 U	<1.0 U

Quarter	1 st	2 nd	3 rd	4 th	2 nd	3 rd	3 rd	3 rd	4 th	1 st	2 nd
Creek Sample ID	Mar 2008	Jun 2008	Sep 2008	Dec 2008	May 2009	Jul 2009	Aug 2009	Sep 2009	Dec 2009	Mar 2010	Jun 2010
GPW-1	27	<0.5U	<0.5U	<0.22U	16	<4U	NS	<1.2U	3.7	1.3J	<0.6U
GPW-3	21.9	9.42	1.1	<0.22U	8.9	<4U	NS	<0.6U	2.8	1.8J	<0.6U
HBW-1	<0.5U	<0.5U	<0.5U	<0.22U	<0.55U	<4U	NS	<1.5U	<0.275U	1.5U	<0.6U
HBW-7	<0.5U	<0.5U	<0.5U	<0.22U	<0.55U	<4U	24	<1.2U	<0.275U	1.5U	<0.6U
HBW-10	<0.5U	<0.5U	<0.5U	<0.22U	<0.55U	<4U	NS	<1.5U	<0.275U	1.2U	<0.6U

Quarter	3 rd	4 th	1 st	2 nd	3 rd	4 th	1 st	2 nd	3 rd	4 th	1 st
Creek Sample ID	Sep 2010	Dec 2010	Mar 2011	Jun 2011	Sep 2011	Dec 2011	Mar 2012	Jun 2012	Not Applicable	Jan & Feb 2013	Mar 2013
GPW-1	dry	<0.1U	8.7	dry	dry	1.76	0.163J	dry	NS	1.65	0.735
GPW-3	dry	0.199J	0.673	dry	dry	1.31	0.261	dry	NS	1.74	0.754
HBW-1	dry	<0.1U	<0.2U	dry	dry	<0.1U	0.1U	dry	NS	<0.2U	<0.2U
HBW-7	dry	<0.1U	<0.2U	dry	dry	0.171J	0.1U	dry	NS	<0.2U	<0.2U
HBW-10	dry	<0.1U	<0.2U	dry	dry	<0.1U	0.1U	dry	NS	<0.2U	<0.2U

Quarter	2 nd	3 rd	4 th	1 st	2 nd	3 rd	4 th	1 st	2 nd	3 rd	4 th
Creek Sample ID	Jun 2013	Sept 2013	Dec 2013	Feb 2014	May 2014	Aug 2014	Nov 2014	Feb 2015	May 2015	Aug 2015	Nov 2015
GPW-1	dry	<0.2 U	dry	0.766	dry	dry	0.244 J	0.311 J	0.156J	dry	0.142 J
GPW-3	dry	<0.2 U	dry	1.15	dry	dry	0.276 J	0.344 J	dry	dry	0.311 J
HBW-1	<0.2U	<0.2 U	dry	<0.2 U	dry	dry	<0.2 U	<0.2 U	dry	dry	<0.2 U
HBW-7	<0.2U	<0.2 U	dry	0.201 J	dry	dry	<0.2 U	0.124 J	dry	dry	<0.2 U
HBW-10	<0.2U	<0.2 U	dry	<0.2 U	dry	dry	<0.2 U	<0.2 U	dry	dry	<0.2 U

Quarter	1 st	2 nd	3 rd	4 th	1 st	2 nd	3 rd	4 th	1 st	2 nd	3 rd
Creek Sample ID	Feb 2016	May 2016	Aug 2016	Nov 2016	Feb 2017	May 2017	Aug 2017	Dec 2017	Mar 2018	June 2018	August 2018
GPW-1	0.447	6.59	<0.2 U	0.301 J	<1 U	0.263	dry	<4.0 U	<4.0 U	dry	<2.0 U
GPW-3	0.474	0.457	0.141	0.563	<1 U	0.274	dry	<4.0 U	<4.0 U	dry	<2.0 U
HBW-1	<0.2 U	<0.2 U	<0.2 U	<0.2 U	<1 U	<0.2 U	<0.2 U	1.1 J	<4.0 U	dry	<2.0 U
HBW-7	<0.2 U	<0.2 U	<0.2 U	0.318 J	<1 U	0.155	<0.2 U	<4.0 U	<4.0 U	dry	<2.0 U
HBW-10	<0.2 U	<0.2 U	<0.2 U	<0.2 U	<1 U	<0.2 U	0.111J	<4.0 U	<4.0 U	dry	<2.0 U

NS – not sampled

U – non-detect

J – Estimated

Dry – no surface water

Quarter	4th	1st
Creek Sample ID	Oct 2018	Jan 2019
GPW-1	<2.0 U	<2.0 U
GPW-3	<2.0 U	<2.0 U
HBW-1	<2.0 U	<2.0 U
HBW-7	<2.0 U	<2.0 U
HBW-10	<2.0 U	<2.0 U

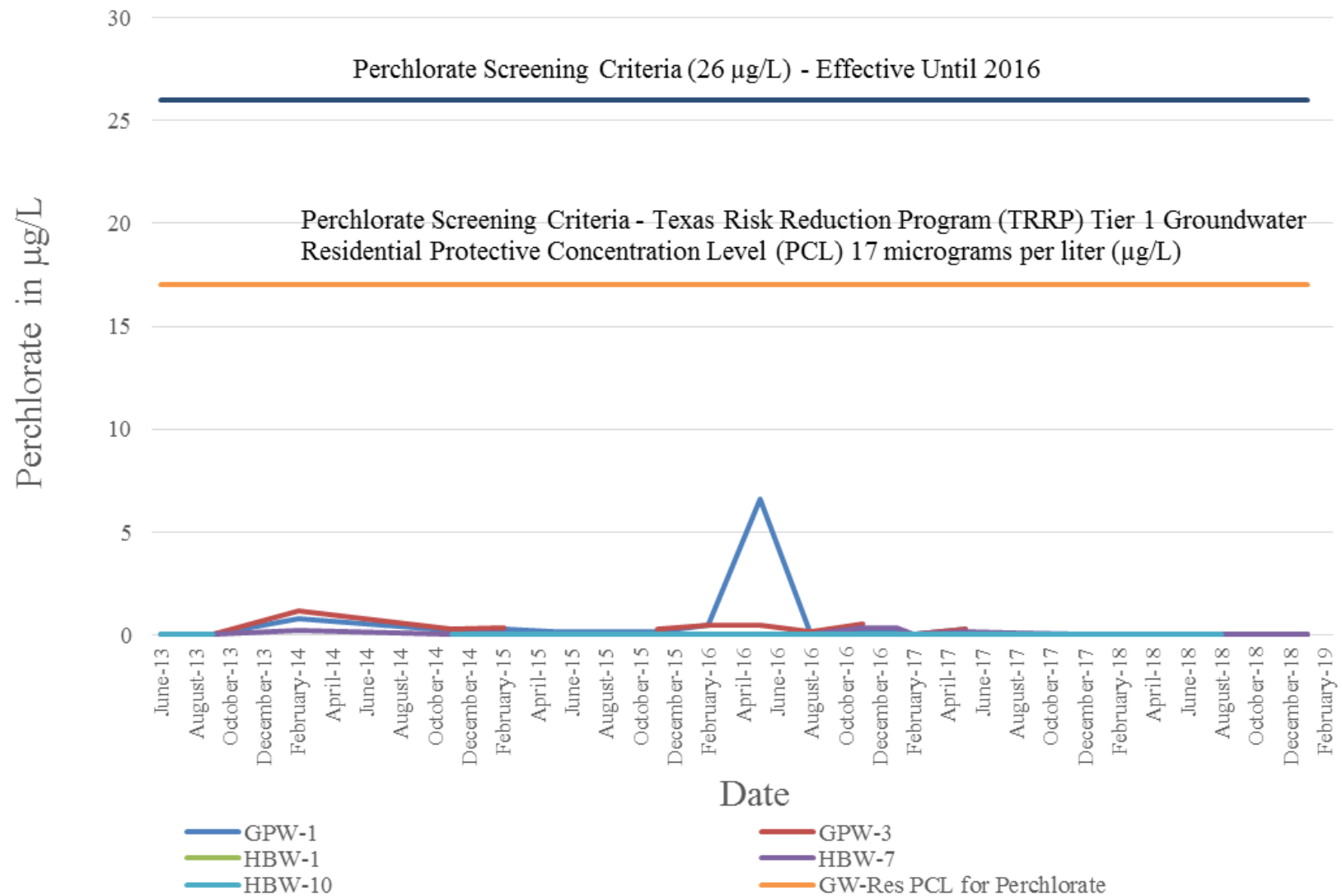
NS – not sampled

U – non-detect

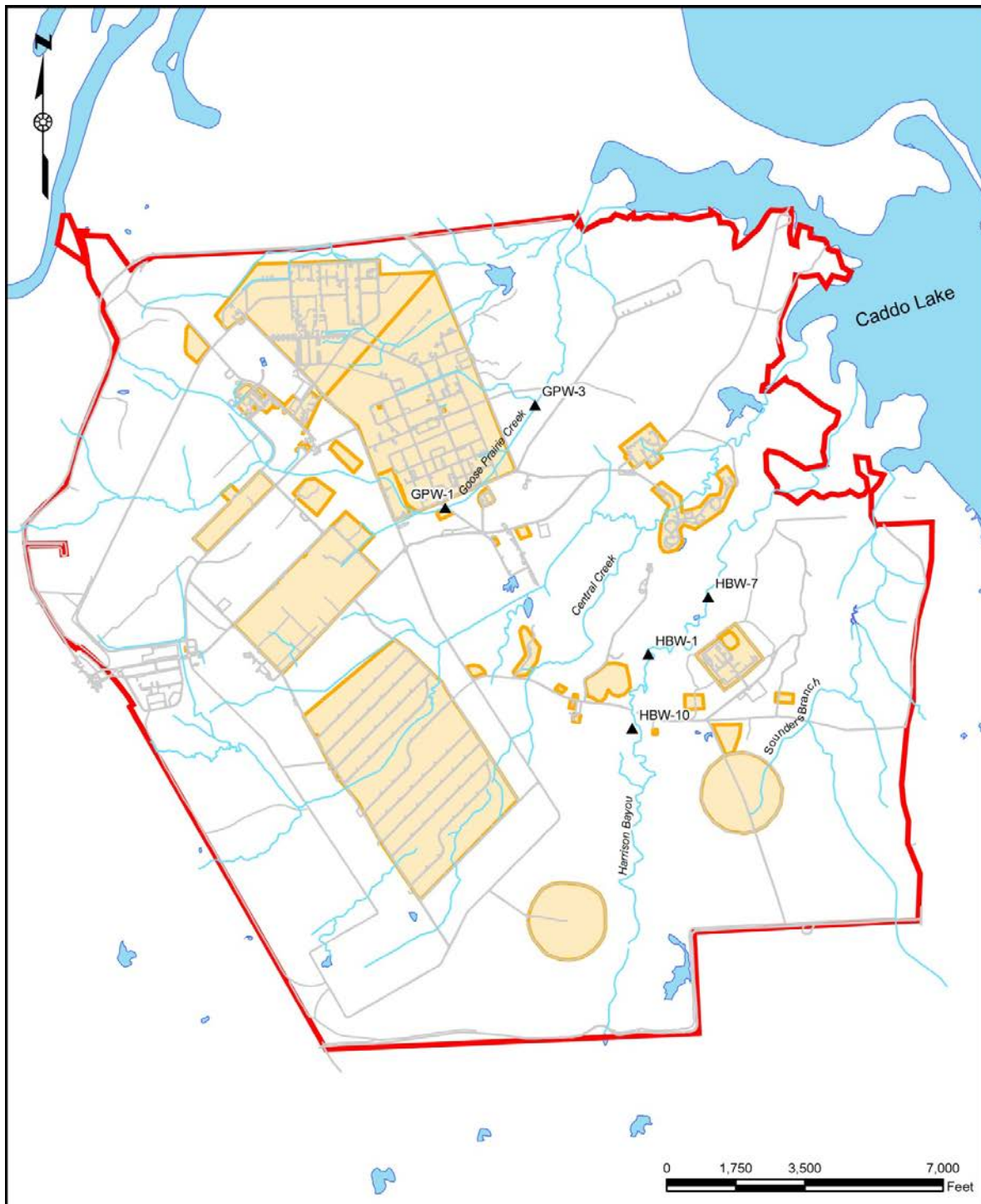
J – Estimated

Dry – no surface water

Surface Water Samples - Perchlorate



Longhorn Army Ammunition Plant Creek Sampling Locations



Legend <ul style="list-style-type: none">▲ Surface Water Sampling Location— Stream— Road■ Site■ Lake	U.S. ARMY CORPS OF ENGINEERS TULSA DISTRICT TULSA, OKLAHOMA
SURFACE WATER SAMPLING LOCATION LONGHORN ARMY AMMUNITION PLANT KARNACK, TEXAS	



DEPARTMENT OF THE ARMY
LONGHORN ARMY AMMUNITION PLANT
POST OFFICE BOX 220
RATCLIFF, AR 72951

July 30, 2019

DAIM-ODB-LO

Mr. Rich Mayer
U.S. Environmental Protection Agency, Region 6
1201 Elm Street, Suite 500
Dallas, Texas 75270-2102

Re: Draft Final Addendum Post Screening Investigation Report, LHAAP-47, Longhorn Army Ammunition Plant, Karnack, Texas, June 2019

Dear Mr. Mayer,

Two hard copies (HC) and two compact discs (CDs) of the above-referenced document are being transmitted to you for your records. The document includes revisions based upon the Environmental Protection Agency's (EPA) comments on the Draft received July 12, 2019. In accordance with the Federal Facility Agreement, this Draft Final will be considered Final after 30 days without further comment. Response to comments on the Draft version of the document are included within this Draft Final.

The document was prepared by HDR Environmental, Operations and Construction, Inc. (HDR) on behalf of the Army as part of HDR's contract for the facility. I ask that Phil Werner, HDR's Project Manager, be copied on any communications related to the project.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in cursive script that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
Longhorn AAP Site Manager

Copies furnished:

- A. Palmie, TCEQ, Austin, TX (1 hard copy, 1 CD)
- P. Bruckwicki, Caddo Lake NWR, TX (1 hard copy, 1 CD)
- A. Williams, USACE, Tulsa District, OK (1 CD)
- R. Smith, USACE, Tulsa District, OK
- A. Sherman, USAEC, San Antonio, TX (1 CD)



DEPARTMENT OF THE ARMY
LONGHORN ARMY AMMUNITION PLANT
POST OFFICE BOX 220
RATCLIFF, AR 72951

July 30, 2019

DAIM-ODB-LO

Ms. April Palmie
Texas Commission on Environmental Quality
Superfund Section, MC-136
12100 Park 35 Circle, Bldg D
Austin, TX 78753

Re: Draft Final Addendum Post Screening Investigation Report, LHAAP-47, Longhorn Army Ammunition Plant, Karnack, Texas, June 2019

Dear Ms. Palmie,

One hard copy (HC) and one compact disc (CD) of the above-referenced document are being transmitted to you for your records. The document includes revisions based upon the Environmental Protection Agency's (EPA) comments on the Draft received July 12, 2019. In accordance with the Federal Facility Agreement, this Draft Final will be considered Final after 30 days without further comment. Response to comments on the Draft version of the document are included within this Draft Final.

The document was prepared by HDR Environmental, Operations and Construction, Inc. (HDR) on behalf of the Army as part of HDR's contract for the facility. I ask that Phil Werner, HDR's Project Manager, be copied on any communications related to the project.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in cursive script that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
Longhorn AAP Site Manager

Copies furnished:

R. Mayer, USEPA Region 6, Dallas, TX (2 hard copies, 2 CDs)
P. Bruckwicki, Caddo Lake NWR, TX (1 hard copy, 1 CD)
A. Williams, USACE, Tulsa District, OK (1 CD)
R. Smith, USACE, Tulsa District, OK
A. Sherman, USAEC, San Antonio, TX (1 CD)



Comments on:

Draft Addendum PSI Report for LHAAP-47, Plant 3 Area,
Solid Rocket Motor Fuel Production, Longhorn Army Ammunition Plant, Karnack, Harrison County, Texas
Submitted: July 1, 2019

Submitted by: USACE / HDR

Responded by: Mr. Rich Mayer

U.S. Environmental Protection Agency, Region 6

Date Responded: 12 July 2019

1. Respondent concurs (C) or does not concur (D)
2. Commenter agrees (A) or does not agree (D) with response.

Number	Section/Page	Paragraph/ Line	Comment	C,D	Response	A,D
Reviewer #1:						
1	Figure 2-1		Please provide a larger sized Figure. The current Figure is difficult to see the wells, surface water sampling locations and to read the legend.	C	The paper size was increased to 11 x 17 and the figure inserted and sized accordingly.	
2	Table 2-1		The dissolved oxygen levels presented are not correct. It is impossible for the DO levels to be as high as shown, except possibly for the 14.20 mg/L result for SW4. Please correct or explain (typo errors, bad DO membrane, etc.) these high levels shown. EPA does like the info presented and formats for tables 2-1, 2-2, and 3-1.	C	The values entered into Table 2-1 were decimal point typos and have been corrected. Also, HDR appreciates the compliment.	
3	Table 3-1		The column which indicates the contaminants compliance levels for surface water should not be stated as screening levels. Please correct.	C	Text has been revised in Table 3-1 to state, "Compliance Levels".	

Draft Final
Addendum
Post-Screening
Investigation Report

for LHAAP-47, Plant 3 Area,
Solid Rocket Motor Fuel Production
Longhorn Army Ammunition Plant
Karnack, Texas

July 2019

Prepared For:



U.S. Army Corps of Engineers – Tulsa District

Prepared By:

HDR

2650 Park Tower Drive, Suite 400
Vienna, VA 22180

Contract No. W912BV-15-D-0014
Task Order No. W912BV18F0023

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Draft Final
ADDENDUM
POST-SCREENING INVESTIGATION REPORT
FOR
LHAAP-47, PLANT 3 AREA,
SOLID ROCKET MOTOR FUEL PRODUCTION
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

Prepared For:
U.S. Army Corp of Engineers Tulsa District

Prepared By:
HDR, Inc.
2650 Park Tower Drive, Suite 400
Vienna, VA 22180

Contract No. W912BV-15-D-0014
Task Order No. W912BV18F0023

July 2019

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Appendices

Appendix A – Surface Water Raw Analytical Data Package
 Appendix B – Data Validation Report

Acronyms and Abbreviations

AECOM	AECOM Technical Services, Inc.
btoc	below top of casing
ft	feet
^{GW} GW _{ing}	Tier 1 residential groundwater
GPS	Global Positioning System
IWWP	Installation Wide Work Plan
J	estimated value
LHAAP	Longhorn Army Ammunition Plant
MCL	Maximum Contaminant Level
mg/L	milligrams per liter
MSL	above mean sea level
mV	millivolt
ND	not detected
NTU	nephelometric turbidity units
PCL	Protective Concentration Level
PSI	Post Screening Investigation
ROD	Record of Decision
SDWA	Safe Drinking Water Act
TCEQ	Texas Commission on Environmental Quality
TRRP	Texas Risk Reduction Program
USEPA	U.S. Environmental Protection Agency
ug/L	micrograms per liter
μS/cm	microsiemens per centimeter
VOC	volatile organic compound
VRS	Virtual Reference Station
WP	Work Plan
°C	degrees Celsius



1 Introduction

This addendum to the Post-Screening Investigation (PSI) Report has been prepared for the former Longhorn Army Ammunition Plant (LHAAP) site LHAAP-47 (Plant 3 Area), located in Karnack, Texas. The purpose of this addendum is to summarize the results of the surface water sample collection and analysis performed at LHAAP-47 in March and April 2019. Due to drought conditions and the lack of surface water in Goose Prairie Creek and its tributaries during the summer of 2018 when the PSI field investigation was conducted, surface water sampling could not be performed. The decision was subsequently made to perform this activity when surface water was present and could be attributed to groundwater recharge and not overland flow from precipitation events.

This document describes the PSI activities completed at site LHAAP-47. The PSI effort was performed in accordance with the Installation Wide Work Plan (IWWP) (AECOM, 2014) and PSI Work Plan (WP) (AECOM, 2016). The IWWP and PSI WP were reviewed by and concurrence was obtained from the Texas Commission on Environmental Quality (TCEQ) and the U.S. Environmental Protection Agency (USEPA).

The objective of the surface water sampling is to re-assess and update the LHAAP-47 groundwater contribution to surface water in Goose Prairie Creek. The addendum PSI data will be used to support revision of the Draft Final Record of Decision (ROD), as necessary.

2 Investigation Activities

This section presents information on the activities completed at LHAAP-47 to re-assess the current surface water conditions. The following provides a description of the activities completed to collect and analyze surface water samples from Goose Prairie Creek.

Surface water samples were collected on 7 March 2019, as part of the PSI surface water re-assessment. Four samples were collected at various locations in Goose Prairie Creek and its tributaries, as shown on **Figure 2-1**. The surface water samples were analyzed for volatile organic compounds (VOCs) and perchlorate to assess whether contaminated groundwater was impacting surface water. Additionally, water quality parameters were collected at each sampling location using an YSI 556 water quality meter and a Hanna Turbidimeter. **Table 2-1** presents the water quality parameters collected at each surface water location.

To evaluate whether groundwater was contributing to surface water, water levels were gauged in nearby wells, while creek bottom elevations were collected at each sample location (as shown in **Table 2-2**). Water level measurements were collected in accordance with the procedures identified in the IWWP and PSI WPs from the up gradient wells.

Creek bottom elevations were collected using a Trimble R2 Global Positioning System (GPS) receiver receiving real-time corrections from a Virtual Reference Station (VRS) subscription. At each sample location, the R2 receiver, which was mounted on a 2-meter survey pole, was placed in the center of the creek. The survey pole was set up using a tripod and leveled properly. The field team then collected several points at each location, those with the highest accuracy metrics were used for the final creek bottom elevations. A second mobilization was conducted 25 April 2019 to revisit creek bottom elevations at sample location SW2 due to abnormal readings collected in March. It was determined that elevation readings were collected while the receiver was in roving mode, which did not fix an accurate elevation. The coordinate data collected from the other three surface water sample locations were confirmed they were collected in fixed mode and did not need reassessment.

Prior to collecting GPS data, the system was verified to be working properly by collecting a coordinate on an established survey monument (monument C-22) near the sampling areas. The coordinate data comparison for this verification is shown in **Table 2-2**. The location of the C-22 monument is shown on **Figure 2-1**.



Table 2-1. Water Quality Parameters

Sample Location	Turbidity (NTUs)	Temperature (°C)	Conductivity (µS/cm)	pH	Dissolved Oxygen (mg/L)	Oxidation-Reduction Potential (mV)	Water Clarity	Estimated Channel Depth (feet)	Estimated Channel Width (feet)	Estimated Water Depth (feet)	Cut Bank
SW1	15.5	7.85	257	7.95	8.340	97.2	cloudy	6	10	1	no
SW2	14.3	8.47	86	7.67	4.532	96.9	cloudy	5	8	3	no
SW3	14.3	9.32	82	6.61	2.230	137.9	clear	8	12	0.3	no
SW4	14.6	8.92	81	7.35	1.420	90.1	cloudy	5	8	2	yes

NTUs - nephelometric turbidity units
 °C - degrees Celsius
 µS/cm - microsiemens per centimeter
 mg/L - milligrams per liter
 mV - millivolt



Table 2-2. Sample and Well Survey Location Data

Location ID	Associated Sample Area	Creek Bottom Elevation (ft MSL) ²	Groundwater Elevation (ft MSL) ²	Groundwater Elevation Minus Creek Bottom Elevation ³	Easting ¹	Northing ¹	Top of Casing Elevation (ft MSL) ²	Depth to Groundwater (ft btoc) ²
Monument Location Data								
C-22 ¹	-	213.89	-	-	3307823.95	6955892.44	-	-
Collection Date: 3/07/2019	Surface Water Sample Location Data							
SW1	-	184.97	-	-	3310243.92	6957976.18	-	-
SW2	-	180.35*	-	-	3311050.55	6958996.49	-	-
SW3	-	179.67	-	-	3311800.52	6960219.95	-	-
SW4	-	175.57	-	-	3312417.66	6962387.87	-	-
Collection Date: 3/07/2019	Monitoring Well Location Data							
08WW01	SW1	-	188.92	3.95	3310498.19	6958043.02	199.31	10.39
LHSMW53	SW1	-	190.71	5.74	6958096.73	3309903.12	197.61	6.90
47WW42	SW2	-	175.32	-15.97	6958850.75	3310575.55	195.22	19.90
LHSMW55R	SW2	-	175.09	-16.20	6959029.23	3310180.19	198.76	23.67
47WW21	SW3	-	172.64	-7.03	6959832.47	3311452.33	187.59	14.95
47WW33	SW4	-	170.73	-4.84	6961743.00	3312137.00	181.58	10.85

1 - NAD_1983_StatePlane_Texas_North_Central_FIPS_4202_Feet

2 - North American Vertical Datum of 1988

FIPS: Federal Information Processing Standard

ft - feet

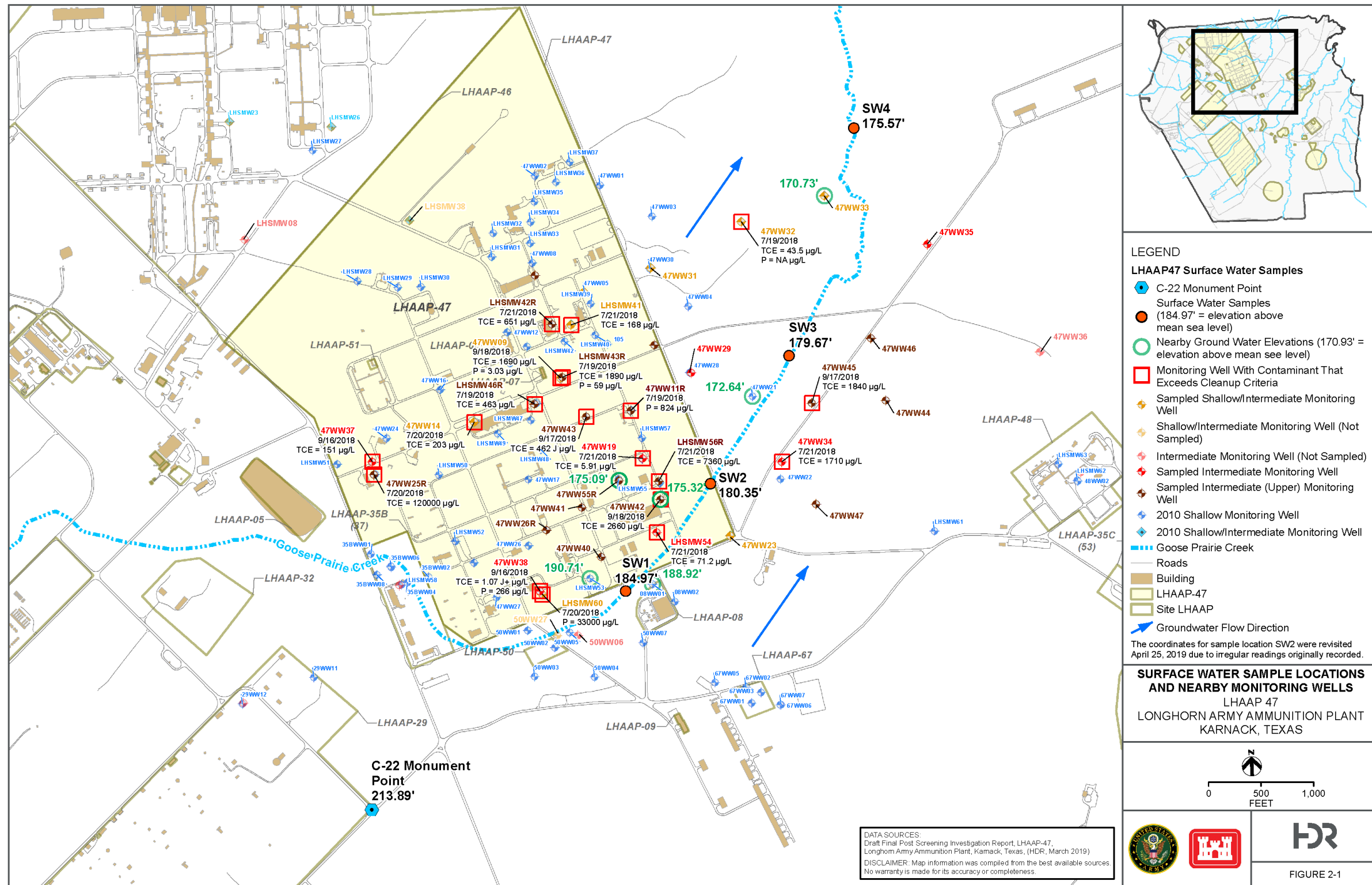
MSL - mean sea level

btoc - below top of casing

* - Elevation height recorded during 25 April 2019



Figure 2-1. Surface Water Sample Locations and Nearby Monitoring Wells



PATH: Z:\2018\18-025_LHAAP_SITE\29_47_18AND24_ROD_WERNER\7_2_WORK_IN_PROGRESS\MAP_DOCS\DRAT\LHAAP47\LHAAP47_PSI_TCE_PERCH_MM_2018RESULTS_ELEVATIONS.MXD - USER: KJLGREN - DATE: 6/28/2019



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3 Investigation Findings

This section presents sample results from the PSI surface water field investigation. The surface water results presented in this section have been compared to Safe Drinking Water Act (SDWA) Maximum Contaminant Levels (MCLs), where they exist. Where an MCL has not been promulgated, the Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Levels (PCL) (^{GW}GW_{ing}) (TRRP Groundwater PCL) comparison levels were used. Surface water results are discussed in relation to these screening levels. The results of the sampling and analyses are discussed below.

General observations of the surface water conditions and creek morphology were recorded during sampling. Field activities were completed under sunny skies; the previous rain event had occurred three days prior to sample collection. In general, surface water was found to be cloudy with low laminar flow at locations with deeper water. In areas with shallow water, the water appeared clear with slightly more turbulent flow. Since sampling was performed post-winter, tree foliage was minimal and all sample locations were surrounded by heavy woods and dense branch cover. Creek bottoms were very sandy at all locations except SW2, which was very rocky. Samples were collected in one day; weather conditions were overcast, damp, and cool with no measurable precipitation.

Four surface water and one duplicate sample were collected and submitted to MicroBac Laboratories for perchlorate and VOC analyses. Perchlorate was detected at each surface water location at low concentrations, well below the TRRP Groundwater PCL value of 17 micrograms per liter ($\mu\text{g/L}$). Acetone was detected in three of the four samples collected, plus the duplicate sample. These values were qualified as estimated, with possible high bias (J+), due to acetone detections in the associated trip blanks. The detections, however, were well below the TRRP Groundwater PCL value of 22,000 $\mu\text{g/L}$. Trichloroethene was detected at one surface water sample location at an estimated value of 0.373 $\mu\text{g/L}$ below the MCL. No other VOCs were detected in any of the surface water samples. A summary of the sample results are provided in **Table 3-1** presented below.

A comparison of the creek bottom elevations to the groundwater elevations identified that groundwater near surface water location SW1 was higher than the creek bottom and was likely contributing to surface water flow. Groundwater elevations at all other surface water locations were found to be lower than the creek bottom elevations.



Table 3-1. Surface Water Results

Chemical Abstracts Service Number	Chemical Name	Sample ID	SW1	DUP1	SW2	SW3	SW4
		Collection Date	03/07/2019				
		Compliance Levels (µg/L)					
SW6850-Perchlorate (µg/L)							
14797-73-0	Perchlorate	17 ⁽²⁾	0.542 J-	0.548	0.530	0.486	0.430
SW8260B-VOCs (µg/L)							
67-64-1	Acetone	22,000 ⁽²⁾	3.56 J+	3.76 J+	3.16 J+	3.65 J+	ND
75-35-4	1,1-Dichloroethene	7 ⁽¹⁾	ND	ND	ND	ND	ND
107-06-2	1,2-Dichloroethane	5 ⁽¹⁾	ND	ND	ND	ND	ND
67-66-3	Chloroform	80 ⁽¹⁾	ND	ND	ND	ND	ND
156-59-2	cis-1,2-Dichloroethene	70 ⁽¹⁾	ND	ND	ND	ND	ND
127-18-4	Tetrachloroethene	5 ⁽¹⁾	ND	ND	ND	ND	ND
156-60-5	trans-1,2-Dichloroethene	100 ⁽¹⁾	ND	ND	ND	ND	ND
79-01-6	Trichloroethene	5 ⁽¹⁾	ND	ND	ND	0.373 J	ND
75-01-4	Vinyl chloride	2 ⁽¹⁾	ND	ND	ND	ND	ND

ND = not detected

(1) = EPA MCL

(2) = TRRP Groundwater PCL

ug/L = micrograms per liter

J = estimated value

J+ = estimated value bias high

J- = estimated value bias low



4 References

AECOM, 2014, Final Installation-Wide Work Plan for Longhorn Army Ammunition Plant, Karnack, Texas, July.

AECOM, 2016, Draft Final Post-Screening Investigation Work Plan, LHAAP-47, Plant Area 3, Longhorn Army Ammunition Plant, Karnack, Texas, December.



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A

Surface Water Raw
Analytical Data Package



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Laboratory Report Number: L19030638

Lynn Lutz
HDR Environmental
HDR 9781 S. Meridian Blvd
Englweood, CO 80112

Please find enclosed the analytical results for the samples you submitted to Microbac Laboratories. Review and compilation of your report was completed by Microbac's Ohio Valley Division (OVD). If you have any questions, comments, or require further assistance regarding this report, please contact your service representative listed below.

Laboratory Contact:
Stephanie Mossburg – Team Chemist/Data Specialist
(740) 373-4071
Stephanie.Mossburg@microbac.com

I certify that all test results meet all of the requirements of the DoD QSM and other applicable contract terms and conditions. Any exceptions are attached to this cover page or addressed in the method narratives presented in the report. All results for soil samples are reported on a 'dry-weight' basis unless specified otherwise. Analytical results for water and wastes are reported on a 'as received' basis unless specified otherwise. A statement of uncertainty for each analysis is available upon request. This laboratory report shall not be reproduced, except in full, without the written approval of Microbac Laboratories, DoD ELAP certification number 2936.01. The reported results are related only to the samples analyzed as received.

This report was certified on March 25 2019



Leslie Bucina – Laboratory Manager

State of Origin: TX
Accrediting Authority: Texas Commission on Environmental Quality ID:T104704252-07-TX
QAPP: DOD Ver 5.0



Microbac Laboratories * Ohio Valley Division
158 Starlite Drive, Marietta, OH 45750 * T: (740) 373-4071 F: (740) 373-4835 * www.microbac.com

Lab Report #: L19030638

Lab Project #: 2886.016

Project Name: Longhorn Army Ammunition DOD

Lab Contact: Stephanie Mossburg

Record of Sample Receipt and Inspection

Comments/Discrepancies

This is the record of the shipment conditions and the inspection records for the samples received and reported as a sample delivery group (SDG). All of the samples were inspected and observed to conform to our receipt policies, except as noted below.

There were no discrepancies.

Discrepancy	Resolution

Coolers

Cooler #	Temperature Gun	Temperature	COC #	Airbill #	Temp Required?
00114989	I	3.0		1001915721210004575000785913246353	X

Inspection Checklist

#	Question	Result
1	Were shipping coolers sealed?	Yes
2	Were custody seals intact?	Yes
3	Were cooler temperatures in range of 0-6?	Yes
4	Was ice present?	Yes
5	Were COC's received/information complete/signed and dated?	Yes
6	Were sample containers intact and match COC?	Yes
7	Were sample labels intact and match COC?	Yes
8	Were the correct containers and volumes received?	Yes
9	Were samples received within EPA hold times?	Yes
10	Were correct preservatives used? (water only)	Yes
11	Were pH ranges acceptable? (voa's excluded)	Yes
12	Were VOA samples free of headspace (less than 6mm)?	Yes



Lab Report #: L19030638

Lab Project #: 2886.016

Project Name: Longhorn Army Ammunition DOD

Lab Contact: Stephanie Mossburg

Samples Received

Client ID	Laboratory ID	Date Collected	Date Received
SW1	L19030638-01	03/07/2019 11:30	03/09/2019 09:22
SW1 MS	L19030638-02	03/07/2019 11:30	03/09/2019 09:22
SW1 MSD	L19030638-03	03/07/2019 11:30	03/09/2019 09:22
SW2	L19030638-04	03/07/2019 12:30	03/09/2019 09:22
SW3	L19030638-05	03/07/2019 13:00	03/09/2019 09:22
SW4	L19030638-06	03/07/2019 14:45	03/09/2019 09:22
DUP01	L19030638-07	03/07/2019 10:00	03/09/2019 09:22
TB01	L19030638-08	03/07/2019 08:00	03/09/2019 09:22

Microbac REPORT L19030638
PREPARED FOR HDR Environmental
WORK ID:

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1.0 Summary Data

1.1 Narratives



Login Number: L19030638
Department: General Chromatography
Analyst: John W. Richards Jr.

METHOD

Analysis SW-846 6850

HOLDING TIMES

Sample Preparation: All holding times were met.

Sample Analysis: All holding times were met.

PREPARATION

Sample preparation proceeded normally.

CALIBRATION

Initial Calibration: For all compounds that yielded a %RSD greater than 15%, linear or higher order equations were applied. All acceptance criteria were met.

Alternate Source Standards: All acceptance criteria were met.

Continuing Calibration and Tune: All acceptance criteria were met.

BATCH QA/QC

Method Blank: All acceptance criteria were met.

Laboratory Control Sample: All acceptance criteria were met.

Matrix Spikes: Recovery out of range was observed for the following analyte: Perchlorate. Please see the applicable QC report for a detailed presentation of the failures.

Sample #	Analyte	Date	Result	Lower	Upper	Type
L19030638-03	Perchlorate	2019-03-18 20:48:40	80.5	84	119	Recovery

SAMPLES

Samples: All acceptance criteria were met.

Internal Standards: All acceptance criteria were met.

Manual Integration Reason Codes

Reason #1: Data System Fails to Select Correct Peak In some cases the chromatography system selects and integrates the 'wrong peak'. In this case the analyst must correct the selection and force the system to integrate the proper peak. Other times the system may miss the peak completely.

Reason #2: Data System Splits the Peak Incorrectly or Integrates a False Peak as a Rider Peak This phenomena is common at low concentrations where the signal:noise ratio is low. A single compound (peak) is incorrectly split into multiple peaks or integrated as a main peak with one or more rider peaks resulting in low area counts for the target compound.

Reason #3: Improperly Integrated Isomers and/or coeluting compounds. This system often fails to distinguish coeluting compounds and or isomers. The integration areas and concentrations are wrong, and they must be corrected by manual integration. Prime examples are benzo(k)fluoranthene and benzo(b)fluoranthene which are often unresolved and integrated improperly when both are present at low concentrations in standards or samples.

Reason #4: System Establishes Incorrect Baseline There are numerous situations in chromatography where the system establishes the baseline incorrectly. Some baseline errors will be obvious to the analyst and should be corrected via manual procedures.

Reason #5: Miscellaneous Other situations involving integration errors may require in-depth review and technical judgment. These cases should be brought to the attention of the laboratory management. If the form of manual integration is not clearly covered by these four cases, then review and approval by the Laboratory Director or the QA/QC Supervisor will be required.

I certify that this data package is in compliance with the terms and conditions agreed to by the client and Microbac Laboratories Inc., both technically and for completeness, except for the conditions noted above. Release of the data contained in this hard copy data package has been authorized by the Laboratory Manager or designated person, as verified by the following signature.

Narrative ID: 147535

Approved By: Eric Lawson





Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	8260
Prep Batch Number(s):		Reviewer Name:	Anthony Canter
LRC Date:	2019-03-25 00:00:00		

Laboratory Data Package Cover Page

R1	Field chain-of-custody documentation;
R2	Sample identification cross-reference;
R3	Test reports (analytical data sheets) for each environmental sample that includes: (a) Items consistent with NELAC Chapter 5, (b) dilution factors, (c) preparation methods, (d) cleanup methods, and (e) a.if required for the project, tentatively identified compounds (TICs).
R4	Surrogate recovery data including: (a) Calculated recovery (%R), and (b) the laboratory's surrogate QC limits.
R5	Test reports/summary forms for blank samples;
R6	Test reports/summary forms for laboratory control samples (LCSs) including: (a) LCS spiking amounts, (b) calculated %R for each analyte, and (c) the laboratory's LCS QC limits.
R7	Test reports for project matrix spike/matrix spike duplicates (MS/MSDs) including: (a) samples associated with the MS/MSD clearly identified, (b) MS/MSD spiking compounds, (c) concentration of each MS/MSD analyte measured in the parent and spiked samples, (d) calculated %Rs and relative percent differences (RPDs), and (e) the laboratory's MS/MSD QC limits.
R8	Laboratory analytical duplicate (if applicable) recovery and precision: (a) the amount of analyte measured in the duplicate, (b) the calculated RPD, and (c) the laboratory's QC limits for analytical duplicates.
R9	List of method quantitation limits (MQLs) and detectability check sample results for each analyte for each method and matrix.
R10	Other problems or anomalies.

Name (Printed)	Signature	Official Title (Printed)	Date
Anthony Canter		Volatiles Lab Supervisor	2019-03-25 13:47:36



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	8260
Prep Batch Number(s):		Reviewer Name:	Anthony Canter
LRC Date:	2019-03-25 00:00:00		

Description	Yes	No	NA	NR	ER#
Chain-of-custody (C-O-C)					
Did samples meet the laboratory's standard conditions of sample acceptability upon receipt?	X				
Were all departures from standard conditions described in an exception report?	X				
Sample and quality control (QC) identification	X				
Are all field sample ID numbers cross-referenced to the laboratory ID numbers?	X				
Are all laboratory ID numbers cross-referenced to the corresponding QC data?	X				
Test reports					
Were all samples prepared and analyzed within holding times?	X				
Other than those results < MQL, were all other raw values bracketed by calibration standards?	X				
Were calculations checked by a peer or supervisor?	X				
Were all analyte identifications checked by a peer or supervisor?	X				
Were sample detection limits reported for all analytes not detected?	X				
Were all results for soil and sediment samples reported on a dry weight basis?	X				
Were % moisture (or solids) reported for all soil and sediment samples?	X				
Were bulk soils/solids samples for volatile analysis extracted with methanol per SW846 Method 5035?			X		
If required for the project, are TICs reported?			X		
Surrogate recovery data					
Were surrogates added prior to extraction?	X				
Were surrogate percent recoveries in all samples within the laboratory QC limits?	X				
Test reports/summary forms for blank samples	X				
Were appropriate type(s) of blanks analyzed?	X				
Were blanks analyzed at the appropriate frequency?	X				
Were method blanks taken through the entire analytical process, including preparation and, if applicable, cleanup procedures?	X				
Were blank concentrations < MQL?	X				
Laboratory control samples (LCS):					
Were all COCs included in the LCS?	X				



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	8260
Prep Batch Number(s):		Reviewer Name:	Anthony Canter
LRC Date:	2019-03-25 00:00:00		

Was each LCS taken through the entire analytical procedure, including prep and cleanup steps?	X				
Were LCSs analyzed at the required frequency?	X				
Were LCS (and LCSD, if applicable) %Rs within the laboratory QC limits?	X				
Does the detectability check sample data document the laboratory's capability to detect the COCs at the MDL used to calculate the SDLs?	X				
Was the LCSD RPD within QC limits?	X				
Matrix spike (MS) and matrix spike duplicate (MSD) data					
Were the project/method specified analytes included in the MS and MSD?			X		
Were MS/MSD analyzed at the appropriate frequency?			X		
Were MS (and MSD, if applicable) %Rs within the laboratory QC limits?			X		
Were MS/MSD RPDs within laboratory QC limits?			X		
Analytical duplicate data					
Were appropriate analytical duplicates analyzed for each matrix?			X		
Were analytical duplicates analyzed at the appropriate frequency?			X		
Were RPDs or relative standard deviations within the laboratory QC limits?			X		
Method quantitation limits (MQLs):					
Are the MQLs for each method analyte included in the laboratory data package?	X				
Do the MQLs correspond to the concentration of the lowest non-zero calibration standard?	X				
Are unadjusted MQLs and DCSs included in the laboratory data package?	X				
Other problems/anomalies					
Are all known problems/anomalies/special conditions noted in this LRC and ER?	X				
Was applicable and available technology used to lower the SDL to minimize the matrix interference effects on the sample results?	X				
Is the laboratory NELAC-accredited under the Texas Laboratory Accreditation Program for the analytes, matrices and methods associated with this laboratory data package?	X				
Initial calibration (ICAL)					
Were response factors and/or relative response factors for each analyte within QC limits?	X				
Were percent RSDs or correlation coefficient criteria met?	X				



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	8260
Prep Batch Number(s):		Reviewer Name:	Anthony Canter
LRC Date:	2019-03-25 00:00:00		

Was the number of standards recommended in the method used for all analytes?	X				
Were all points generated between the lowest and highest standard used to calculate the curve?	X				
Are ICAL data available for all instruments used?	X				
Has the initial calibration curve been verified using an appropriate second source standard?	X				
Initial and continuing calibration verification (ICCV and CCV) and continuing calibration blank (CCB):					
Was the CCV analyzed at the method-required frequency?	X				
Were percent differences for each analyte within the method-required QC limits?	X				
Was the ICAL curve verified for each analyte?	X				
Was the absolute value of the analyte concentration in the inorganic CCB < MDL?			X		
Mass spectral tuning					
Was the appropriate compound for the method used for tuning?	X				
Were ion abundance data within the method-required QC limits?	X				
Internal standards (IS)					
Were IS area counts and retention times within the method-required QC limits?	X				
Raw data (NELAC Section 5.5.10)					
Were the raw data (for example, chromatograms, spectral data) reviewed by an analyst?	X				
Were data associated with manual integrations flagged on the raw data?	X				
Dual column confirmation					
Did dual column confirmation results meet the method-required QC?			X		
Tentatively identified compounds (TICs)					
If TICs were requested, were the mass spectra and TIC data subject to appropriate checks?			X		
Interference Check Sample (ICS) results					
Were percent recoveries within method QC limits?			X		
Serial dilutions, post digestion spikes, and method of standard additions					
Were percent differences, recoveries, and the linearity within the QC limits specified in the method?			X		
Method detection limit (MDL) studies					



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	8260
Prep Batch Number(s):		Reviewer Name:	Anthony Canter
LRC Date:	2019-03-25 00:00:00		

Was a MDL study performed for each reported analyte?	X				
Is the MDL either adjusted or supported by the analysis of DCSs?	X				
Proficiency test reports					
Was the laboratory's performance acceptable on the applicable proficiency tests or evaluation studies?	X				
Standards documentation					
Are all standards used in the analyses NIST-traceable or obtained from other appropriate sources?	X				
Compound/analyte identification procedures					
Are the procedures for compound/analyte identification documented?	X				
Demonstration of analyst competency (DOC)					
Was DOC conducted consistent with NELAC Chapter 5?	X				
Is documentation of the analyst's competency up-to-date and on file?	X				
Verification/validation documentation for methods (NELAC Chapter 5)					
Are all the methods used to generate the data documented, verified, and validated, where applicable?	X				
Laboratory standard operating procedures (SOPs)					
Are laboratory SOPs current and on file for each method performed	X				

1. Items identified by the letter "R" must be included in the laboratory data package submitted in the TRRP-required report(s). Items identified by the letter "S" should be retained and made available upon request for the appropriate retention period;
2. O = organic analyses; I = inorganic analyses (and general chemistry, when applicable);
3. NA = Not applicable;
4. NR = Not reviewed;
5. ER# = Exception Report identification number (an Exception Report should be completed for an item if "NR" or "No" is checked).

The Exception Report for each "No" or "Not Reviewed (NR)" item in Laboratory Review Checklist and for each analyte, matrix, and method for which the laboratory does not hold NELAC accreditation under the Texas Laboratory Accreditation Program.

Release Statement: I am responsible for the release of this laboratory data package. This laboratory is NELAC accredited under the Texas Laboratory Accreditation Program for all the methods, analytes, and matrices reported in this data package except as noted in the Exception Reports. The data have been reviewed and are technically compliant with



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	8260
Prep Batch Number(s):		Reviewer Name:	Anthony Canter
LRC Date:	2019-03-25 00:00:00		

the requirements of the methods used, except where noted by the laboratory in the Exception Reports. By my signature below, I affirm to the best of my knowledge all problems/anomalies observed by the laboratory have been identified in the Laboratory Review Checklist, and no information affecting the quality of the data has been knowingly withheld.

Check, if applicable: This laboratory meets an exception under 30 TAC §25.6 and was last inspection by TCEQ or _____ on **(enter date of last inspection)**. Any findings affecting the data in this laboratory data package are noted in the Exception Reports herein. The official signing the cover page of the report in which these data are used is responsible for releasing this data package and is by signature affirming the above release statement is true.

Exceptions Report

There are no exceptions.



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	6850
Prep Batch Number(s):	WG699914	Reviewer Name:	Eric Lawson
LRC Date:	2019-03-20 00:00:00		

Laboratory Data Package Cover Page

R1	Field chain-of-custody documentation;
R2	Sample identification cross-reference;
R3	Test reports (analytical data sheets) for each environmental sample that includes: (a) Items consistent with NELAC Chapter 5, (b) dilution factors, (c) preparation methods, (d) cleanup methods, and (e) a.if required for the project, tentatively identified compounds (TICs).
R4	Surrogate recovery data including: (a) Calculated recovery (%R), and (b) the laboratory's surrogate QC limits.
R5	Test reports/summary forms for blank samples;
R6	Test reports/summary forms for laboratory control samples (LCSs) including: (a) LCS spiking amounts, (b) calculated %R for each analyte, and (c) the laboratory's LCS QC limits.
R7	Test reports for project matrix spike/matrix spike duplicates (MS/MSDs) including: (a) samples associated with the MS/MSD clearly identified, (b) MS/MSD spiking compounds, (c) concentration of each MS/MSD analyte measured in the parent and spiked samples, (d) calculated %Rs and relative percent differences (RPDs), and (e) the laboratory's MS/MSD QC limits.
R8	Laboratory analytical duplicate (if applicable) recovery and precision: (a) the amount of analyte measured in the duplicate, (b) the calculated RPD, and (c) the laboratory's QC limits for analytical duplicates.
R9	List of method quantitation limits (MQLs) and detectability check sample results for each analyte for each method and matrix.
R10	Other problems or anomalies.

Name (Printed)	Signature	Official Title (Printed)	Date
Eric Lawson		Chemist III	2019-03-20 20:04:37



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	6850
Prep Batch Number(s):	WG699914	Reviewer Name:	Eric Lawson
LRC Date:	2019-03-20 00:00:00		

Description	Yes	No	NA	NR	ER#
Chain-of-custody (C-O-C)					
Did samples meet the laboratory's standard conditions of sample acceptability upon receipt?	X				
Were all departures from standard conditions described in an exception report?	X				
Sample and quality control (QC) identification	X				
Are all field sample ID numbers cross-referenced to the laboratory ID numbers?	X				
Are all laboratory ID numbers cross-referenced to the corresponding QC data?	X				
Test reports					
Were all samples prepared and analyzed within holding times?	X				
Other than those results < MQL, were all other raw values bracketed by calibration standards?	X				
Were calculations checked by a peer or supervisor?	X				
Were all analyte identifications checked by a peer or supervisor?	X				
Were sample detection limits reported for all analytes not detected?	X				
Were all results for soil and sediment samples reported on a dry weight basis?			X		
Were % moisture (or solids) reported for all soil and sediment samples?			X		
Were bulk soils/solids samples for volatile analysis extracted with methanol per SW846 Method 5035?			X		
If required for the project, are TICs reported?			X		
Surrogate recovery data					
Were surrogates added prior to extraction?			X		
Were surrogate percent recoveries in all samples within the laboratory QC limits?			X		
Test reports/summary forms for blank samples	X				
Were appropriate type(s) of blanks analyzed?	X				
Were blanks analyzed at the appropriate frequency?	X				
Were method blanks taken through the entire analytical process, including preparation and, if applicable, cleanup procedures?	X				
Were blank concentrations < MQL?	X				
Laboratory control samples (LCS):					
Were all COCs included in the LCS?	X				



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	6850
Prep Batch Number(s):	WG699914	Reviewer Name:	Eric Lawson
LRC Date:	2019-03-20 00:00:00		

Was each LCS taken through the entire analytical procedure, including prep and cleanup steps?	X				
Were LCSs analyzed at the required frequency?	X				
Were LCS (and LCSD, if applicable) %Rs within the laboratory QC limits?	X				
Does the detectability check sample data document the laboratory's capability to detect the COCs at the MDL used to calculate the SDLs?	X				
Was the LCSD RPD within QC limits?			X		
Matrix spike (MS) and matrix spike duplicate (MSD) data					
Were the project/method specified analytes included in the MS and MSD?	X				
Were MS/MSD analyzed at the appropriate frequency?	X				
Were MS (and MSD, if applicable) %Rs within the laboratory QC limits?		X			1
Were MS/MSD RPDs within laboratory QC limits?	X				
Analytical duplicate data					
Were appropriate analytical duplicates analyzed for each matrix?			X		
Were analytical duplicates analyzed at the appropriate frequency?			X		
Were RPDs or relative standard deviations within the laboratory QC limits?			X		
Method quantitation limits (MQLs):					
Are the MQLs for each method analyte included in the laboratory data package?	X				
Do the MQLs correspond to the concentration of the lowest non-zero calibration standard?	X				
Are unadjusted MQLs and DCSs included in the laboratory data package?	X				
Other problems/anomalies					
Are all known problems/anomalies/special conditions noted in this LRC and ER?	X				
Was applicable and available technology used to lower the SDL to minimize the matrix interference effects on the sample results?	X				
Is the laboratory NELAC-accredited under the Texas Laboratory Accreditation Program for the analytes, matrices and methods associated with this laboratory data package?	X				
Initial calibration (ICAL)					
Were response factors and/or relative response factors for each analyte within QC limits?	X				
Were percent RSDs or correlation coefficient criteria met?	X				



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	6850
Prep Batch Number(s):	WG699914	Reviewer Name:	Eric Lawson
LRC Date:	2019-03-20 00:00:00		

Was the number of standards recommended in the method used for all analytes?	X				
Were all points generated between the lowest and highest standard used to calculate the curve?	X				
Are ICAL data available for all instruments used?	X				
Has the initial calibration curve been verified using an appropriate second source standard?	X				
Initial and continuing calibration verification (ICCV and CCV) and continuing calibration blank (CCB):					
Was the CCV analyzed at the method-required frequency?	X				
Were percent differences for each analyte within the method-required QC limits?	X				
Was the ICAL curve verified for each analyte?	X				
Was the absolute value of the analyte concentration in the inorganic CCB < MDL?			X		
Mass spectral tuning					
Was the appropriate compound for the method used for tuning?	X				
Were ion abundance data within the method-required QC limits?	X				
Internal standards (IS)					
Were IS area counts and retention times within the method-required QC limits?	X				
Raw data (NELAC Section 5.5.10)					
Were the raw data (for example, chromatograms, spectral data) reviewed by an analyst?	X				
Were data associated with manual integrations flagged on the raw data?	X				
Dual column confirmation					
Did dual column confirmation results meet the method-required QC?			X		
Tentatively identified compounds (TICs)					
If TICs were requested, were the mass spectra and TIC data subject to appropriate checks?			X		
Interference Check Sample (ICS) results					
Were percent recoveries within method QC limits?			X		
Serial dilutions, post digestion spikes, and method of standard additions					
Were percent differences, recoveries, and the linearity within the QC limits specified in the method?			X		
Method detection limit (MDL) studies					



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	6850
Prep Batch Number(s):	WG699914	Reviewer Name:	Eric Lawson
LRC Date:	2019-03-20 00:00:00		

Was a MDL study performed for each reported analyte?	X				
Is the MDL either adjusted or supported by the analysis of DCSs?	X				
Proficiency test reports					
Was the laboratory's performance acceptable on the applicable proficiency tests or evaluation studies?	X				
Standards documentation					
Are all standards used in the analyses NIST-traceable or obtained from other appropriate sources?	X				
Compound/analyte identification procedures					
Are the procedures for compound/analyte identification documented?	X				
Demonstration of analyst competency (DOC)					
Was DOC conducted consistent with NELAC Chapter 5?	X				
Is documentation of the analyst's competency up-to-date and on file?	X				
Verification/validation documentation for methods (NELAC Chapter 5)					
Are all the methods used to generate the data documented, verified, and validated, where applicable?	X				
Laboratory standard operating procedures (SOPs)					
Are laboratory SOPs current and on file for each method performed	X				

1. Items identified by the letter "R" must be included in the laboratory data package submitted in the TRRP-required report(s). Items identified by the letter "S" should be retained and made available upon request for the appropriate retention period;
2. O = organic analyses; I = inorganic analyses (and general chemistry, when applicable);
3. NA = Not applicable;
4. NR = Not reviewed;
5. ER# = Exception Report identification number (an Exception Report should be completed for an item if "NR" or "No" is checked).

The Exception Report for each "No" or "Not Reviewed (NR)" item in Laboratory Review Checklist and for each analyte, matrix, and method for which the laboratory does not hold NELAC accreditation under the Texas Laboratory Accreditation Program.

Release Statement: I am responsible for the release of this laboratory data package. This laboratory is NELAC accredited under the Texas Laboratory Accreditation Program for all the methods, analytes, and matrices reported in this data package except as noted in the Exception Reports. The data have been reviewed and are technically compliant with the requirements of the methods used, except where noted by the laboratory in the Exception Reports. By my signature



Texas Risk Reduction Program (TRRP) Checklist

Laboratory Name:	Microbac OVD	Laboratory Log Number:	L19030638
Project Name:		Method:	6850
Prep Batch Number(s):	WG699914	Reviewer Name:	Eric Lawson
LRC Date:	2019-03-20 00:00:00		

below, I affirm to the best of my knowledge all problems/anomalies observed by the laboratory have been identified in the Laboratory Review Checklist, and no information affecting the quality of the data has been knowingly withheld.

Check, if applicable: This laboratory meets an exception under 30 TAC §25.6 and was last inspection by TCEQ or _____ on **(enter date of last inspection)**. Any findings affecting the data in this laboratory data package are noted in the Exception Reports herein. The official signing the cover page of the report in which these data are used is responsible for releasing this data package and is by signature affirming the above release statement is true.

Exceptions Report

1. Sample 03 MSD yielded a recovery for perchlorate that was below the acceptance limit.

1.2 Certificate of Analysis

Lab Report #: L19030638
 Lab Project #: 2886.016
 Project Name: Longhorn Army Ammunition DOD
 Lab Contact: Stephanie Mossburg

Certificate of Analysis

Sample #: L19030638-01	PrePrep Method: N/A	Instrument: HPMS11
Client ID: SW1	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/05/2019 21:06
Workgroup #: WG699219	Analyst: KFR	Run Date: 03/12/2019 15:30
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 11M29837
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.56	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	96.8	81	118			
4-Bromofluorobenzene	99.7	85	114			
Dibromofluoromethane	96.0	80	119			
Toluene-d8	99.9	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-01	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW1	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 20:22
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 1LM.LM45229
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.542		0.400	0.200	0.100

Lab Report #: L19030638
 Lab Project #: 2886.016
 Project Name: Longhorn Army Ammunition DOD
 Lab Contact: Stephanie Mossburg

Certificate of Analysis

Sample #: L19030638-02	PrePrep Method: N/A	Instrument: HPMS11
Client ID: SW1 MS	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/05/2019 21:06
Workgroup #: WG699219	Analyst: KFR	Run Date: 03/12/2019 13:30
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 11M29833
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	18.8		2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	20.0		1.00	0.500	0.250
Acetone	67-64-1	18.1		10.0	5.00	2.50
Chloroform	67-66-3	19.9		0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	20.6		1.00	0.500	0.250
Tetrachloroethene	127-18-4	19.5		1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	20.0		1.00	0.500	0.250
Trichloroethene	79-01-6	19.4		1.00	0.500	0.250
Vinyl chloride	75-01-4	14.4		1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	96.5	81	118			
4-Bromofluorobenzene	101	85	114			
Dibromofluoromethane	101	80	119			
Toluene-d8	100	89	112			

Sample #: L19030638-02	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW1 MS	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 20:35
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 1LM.LM45230
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.735		0.400	0.200	0.100

Lab Report #: L19030638

Lab Project #: 2886.016

Project Name: Longhorn Army Ammunition DOD

Lab Contact: Stephanie Mossburg

Certificate of Analysis

Sample #: L19030638-03	PrePrep Method: N/A	Instrument: HPMS11
Client ID: SW1 MSD	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/05/2019 21:06
Workgroup #: WG699219	Analyst: KFR	Run Date: 03/12/2019 14:00
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 11M29834
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	18.2		2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	20.1		1.00	0.500	0.250
Acetone	67-64-1	18.4		10.0	5.00	2.50
Chloroform	67-66-3	19.4		0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	20.5		1.00	0.500	0.250
Tetrachloroethene	127-18-4	19.4		1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	19.7		1.00	0.500	0.250
Trichloroethene	79-01-6	19.4		1.00	0.500	0.250
Vinyl chloride	75-01-4	14.2		1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	99.3	81	118			
4-Bromofluorobenzene	106	85	114			
Dibromofluoromethane	102	80	119			
Toluene-d8	102	89	112			

Sample #: L19030638-03	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW1 MSD	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 20:48
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 1LM.LM45231
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.703		0.400	0.200	0.100

Certificate of Analysis

Sample #: L19030638-04	PrePrep Method: N/A	Instrument: HPMS8
Client ID: SW2	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/04/2019 17:07
Workgroup #: WG699028	Analyst: EEA	Run Date: 03/11/2019 17:01
Collect Date: 03/07/2019 12:30	Dilution: 1	File ID: 8M429076
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.16	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	108	81	118			
4-Bromofluorobenzene	112	85	114			
Dibromofluoromethane	106	80	119			
Toluene-d8	105	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-04	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW2	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:01
Collect Date: 03/07/2019 12:30	Dilution: 1	File ID: 1LM.LM45232
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.530		0.400	0.200	0.100

Certificate of Analysis

Sample #: L19030638-05	PrePrep Method: N/A	Instrument: HPMS8
Client ID: SW3	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/04/2019 17:07
Workgroup #: WG699028	Analyst: EEA	Run Date: 03/11/2019 17:30
Collect Date: 03/07/2019 13:00	Dilution: 1	File ID: 8M429077
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.65	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.373	J	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	106	81	118			
4-Bromofluorobenzene	106	85	114			
Dibromofluoromethane	104	80	119			
Toluene-d8	103	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-05	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW3	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:14
Collect Date: 03/07/2019 13:00	Dilution: 1	File ID: 1LM.LM45233
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.486		0.400	0.200	0.100

Lab Report #: L19030638

Lab Project #: 2886.016

Project Name: Longhorn Army Ammunition DOD

Lab Contact: Stephanie Mossburg

Certificate of Analysis

Sample #: L19030638-06	PrePrep Method: N/A	Instrument: HPMS8
Client ID: SW4	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/04/2019 17:07
Workgroup #: WG699028	Analyst: EEA	Run Date: 03/11/2019 17:59
Collect Date: 03/07/2019 14:45	Dilution: 1	File ID: 8M429078
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	5.00	U	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	108	81	118			
4-Bromofluorobenzene	111	85	114			
Dibromofluoromethane	104	80	119			
Toluene-d8	103	89	112			
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-06	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW4	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:27
Collect Date: 03/07/2019 14:45	Dilution: 1	File ID: 1LM.LM45234
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.430		0.400	0.200	0.100

Certificate of Analysis

Sample #: L19030638-07	PrePrep Method: N/A	Instrument: HPMS8
Client ID: DUP01	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/04/2019 17:07
Workgroup #: WG699028	Analyst: EEA	Run Date: 03/11/2019 18:28
Collect Date: 03/07/2019 10:00	Dilution: 1	File ID: 8M429079
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.76	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	111	81	118			
4-Bromofluorobenzene	114	85	114			
Dibromofluoromethane	106	80	119			
Toluene-d8	106	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-07	PrePrep Method: N/A	Instrument: LCMS1
Client ID: DUP01	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:40
Collect Date: 03/07/2019 10:00	Dilution: 1	File ID: 1LM.LM45235
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.548		0.400	0.200	0.100

Lab Report #: L19030638

Lab Project #: 2886.016

Project Name: Longhorn Army Ammunition DOD

Lab Contact: Stephanie Mossburg

Certificate of Analysis

Sample #: L19030638-08	PrePrep Method: N/A	Instrument: HPMS11
Client ID: TB01	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/05/2019 21:06
Workgroup #: WG699219	Analyst: KFR	Run Date: 03/12/2019 14:59
Collect Date: 03/07/2019 08:00	Dilution: 1	File ID: 11M29836
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	2.67	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	98.6	81	118			
4-Bromofluorobenzene	98.2	85	114			
Dibromofluoromethane	99.4	80	119			
Toluene-d8	99.6	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Certificate of Analysis

2.0 Full Sample Data Package

2.1 Volatiles Data

2.1.1 Volatiles GCMS Data (8260)

2.1.1.1 Summary Data

Certificate of Analysis

Sample #: L19030638-01	PrePrep Method: N/A	Instrument: HPMS11
Client ID: SW1	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/05/2019 21:06
Workgroup #: WG699219	Analyst: KFR	Run Date: 03/12/2019 15:30
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 11M29837
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.56	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	96.8	81	118			
4-Bromofluorobenzene	99.7	85	114			
Dibromofluoromethane	96.0	80	119			
Toluene-d8	99.9	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-02	PrePrep Method: N/A	Instrument: HPMS11
Client ID: SW1 MS	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/05/2019 21:06
Workgroup #: WG699219	Analyst: KFR	Run Date: 03/12/2019 13:30
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 11M29833
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	18.8		2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	20.0		1.00	0.500	0.250
Acetone	67-64-1	18.1		10.0	5.00	2.50
Chloroform	67-66-3	19.9		0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	20.6		1.00	0.500	0.250
Tetrachloroethene	127-18-4	19.5		1.00	0.500	0.250

Certificate of Analysis

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
trans-1,2-Dichloroethene	156-60-5	20.0		1.00	0.500	0.250
Trichloroethene	79-01-6	19.4		1.00	0.500	0.250
Vinyl chloride	75-01-4	14.4		1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	96.5	81	118			
4-Bromofluorobenzene	101	85	114			
Dibromofluoromethane	101	80	119			
Toluene-d8	100	89	112			

Sample #: L19030638-03

PrePrep Method: N/A

Instrument: HPMS11

Client ID: SW1 MSD

Prep Method: 5030B/5030C/5035A

Prep Date: N/A

Matrix: Water

Analytical Method: 8260B

Cal Date: 03/05/2019 21:06

Workgroup #: WG699219

Analyst: KFR

Run Date: 03/12/2019 14:00

Collect Date: 03/07/2019 11:30

Dilution: 1

File ID: 11M29834

Sample Tag: 01

Units: ug/L

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	18.2		2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	20.1		1.00	0.500	0.250
Acetone	67-64-1	18.4		10.0	5.00	2.50
Chloroform	67-66-3	19.4		0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	20.5		1.00	0.500	0.250
Tetrachloroethene	127-18-4	19.4		1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	19.7		1.00	0.500	0.250
Trichloroethene	79-01-6	19.4		1.00	0.500	0.250
Vinyl chloride	75-01-4	14.2		1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	99.3	81	118			
4-Bromofluorobenzene	106	85	114			
Dibromofluoromethane	102	80	119			
Toluene-d8	102	89	112			

Certificate of Analysis

Sample #: L19030638-04	PrePrep Method: N/A	Instrument: HPMS8
Client ID: SW2	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/04/2019 17:07
Workgroup #: WG699028	Analyst: EEA	Run Date: 03/11/2019 17:01
Collect Date: 03/07/2019 12:30	Dilution: 1	File ID: 8M429076
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.16	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	108	81	118			
4-Bromofluorobenzene	112	85	114			
Dibromofluoromethane	106	80	119			
Toluene-d8	105	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-05	PrePrep Method: N/A	Instrument: HPMS8
Client ID: SW3	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/04/2019 17:07
Workgroup #: WG699028	Analyst: EEA	Run Date: 03/11/2019 17:30
Collect Date: 03/07/2019 13:00	Dilution: 1	File ID: 8M429077
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.65	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250

Certificate of Analysis

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.373	J	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	106	81	118			
4-Bromofluorobenzene	106	85	114			
Dibromofluoromethane	104	80	119			
Toluene-d8	103	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-06

PrePrep Method: N/A

Instrument: HPMS8

Client ID: SW4

Prep Method: 5030B/5030C/5035A

Prep Date: N/A

Matrix: Water

Analytical Method: 8260B

Cal Date: 03/04/2019 17:07

Workgroup #: WG699028

Analyst: EEA

Run Date: 03/11/2019 17:59

Collect Date: 03/07/2019 14:45

Dilution: 1

File ID: 8M429078

Sample Tag: 01

Units: ug/L

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	5.00	U	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	108	81	118			
4-Bromofluorobenzene	111	85	114			
Dibromofluoromethane	104	80	119			
Toluene-d8	103	89	112			
U	Analyte was not detected. The concentration is below the reported LOD.					

Certificate of Analysis

Sample #: L19030638-07	PrePrep Method: N/A	Instrument: HPMS8
Client ID: DUP01	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/04/2019 17:07
Workgroup #: WG699028	Analyst: EEA	Run Date: 03/11/2019 18:28
Collect Date: 03/07/2019 10:00	Dilution: 1	File ID: 8M429079
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	3.76	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	111	81	118			
4-Bromofluorobenzene	114	85	114			
Dibromofluoromethane	106	80	119			
Toluene-d8	106	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Sample #: L19030638-08	PrePrep Method: N/A	Instrument: HPMS11
Client ID: TB01	Prep Method: 5030B/5030C/5035A	Prep Date: N/A
Matrix: Water	Analytical Method: 8260B	Cal Date: 03/05/2019 21:06
Workgroup #: WG699219	Analyst: KFR	Run Date: 03/12/2019 14:59
Collect Date: 03/07/2019 08:00	Dilution: 1	File ID: 11M29836
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
1,1-Dichloroethene	75-35-4	1.00	U	2.00	1.00	0.500
1,2-Dichloroethane	107-06-2	0.500	U	1.00	0.500	0.250
Acetone	67-64-1	2.67	J	10.0	5.00	2.50
Chloroform	67-66-3	0.250	U	0.500	0.250	0.125
cis-1,2-Dichloroethene	156-59-2	0.500	U	1.00	0.500	0.250
Tetrachloroethene	127-18-4	0.500	U	1.00	0.500	0.250

Certificate of Analysis

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
trans-1,2-Dichloroethene	156-60-5	0.500	U	1.00	0.500	0.250
Trichloroethene	79-01-6	0.500	U	1.00	0.500	0.250
Vinyl chloride	75-01-4	0.500	U	1.00	0.500	0.250
Surrogate	Recovery	Lower Limit	Upper Limit	Q		
1,2-Dichloroethane-d4	98.6	81	118			
4-Bromofluorobenzene	98.2	85	114			
Dibromofluoromethane	99.4	80	119			
Toluene-d8	99.6	89	112			
J	Estimated value ; the analyte concentration was less than the LOQ.					
U	Analyte was not detected. The concentration is below the reported LOD.					

Certificate of Analysis

2.1.1.2 QC Summary Data

Example 8260 Calculations

1.0 Calculating the Response Factor (RF) from the initial calibration (ICAL) data:

$$RF = [(Ax) (Cis)] / [(Ais) (Cx)]$$

where:

Ax = Area of the characteristic ion for the compound being measured:	3399156
Cis = Concentration of the specific internal standard (ug/mL)	25
Ais = Area of the characteristic ion of the specific internal standard	846471
Cx = Concentration of the compound in the standard being measured (ug/mL)	100
RF = Calculated Response Factor	1.0039

Example

2.0 Calculating the concentration (C) of a compound in water using the average RF: *

$$Cx = [(Ax) (Cis) (Vn)(D)] / [(Ais) (RF) (Vs)]$$

where:

Ax = Area of the characteristic ion for the compound being measured	3122498
Cis = Concentration of the specific internal standard (ug/L)	25
D = Dilution factor for sample as a multiplier (10x = 10)	1
Ais = Area of the characteristic ion of the specific internal standard	611048
RF = Average RF from the ICAL	1.004
Vs = Purge volume of sample (mL)	10
Vn = Nominal purge volume of sample (mL) (10.0 mL)	10
Cx = Concentration of the compound in the sample being measured (ug/L)	127.2428

Example

3.0 Calculating the concentration (C) of a compound in soil using the average RF: *

$$Cx = [(Ax) (Cis) (Wn)(D)] / [(Ais) (RF) (Ws)]$$

where:

Ax = Area of the characteristic ion for the compound being measured	3122498
Cis = Concentration of the specific internal standard (ug/L)	25
D = Dilution factor for sample as a multiplier (10x = 10)	1
Ais = Area of the characteristic ion of the specific internal standard	611048
RF = Average RF from the ICAL	1.004
Ws = Weight of sample purged (g)	5
Wn = Nominal purge weight (g) (5.0 g)	5
Cx = Concentration of the compound in the sample being measured (ug/L)	127.2428

Example

Dry weight correction:

Percent solids (PCT_S)	50
Cd = (Cx) (100)/PCT_S	254.4856

* Concentrations appearing on the instrument quantitation reports are on-column results and do not take into account initial volume, final volume, and the dilution factor.

4.0 Concentration from Linear Regression

Step 1: Retrieve Curve Data From Plot, $y = mx + b$

y = response ratio = response of analyte / response of IS = Ax/Ais

x = amount ratio = concentration analyte/concentration internal standard = Cx / Cis

m = slope from curve = 0.213

b = intercept from curve = - 0.00642

Step 2: Calculate y from Quantitation Report

$$y = 86550/593147 = 0.1459$$

Step 3: Solve for x

$$x = (y - b)/m = [(0.1459 - (-0.00642))/0.213] = 0.7152$$

Step 4: Solve for analyte concentration Cx

$$Cx = Cis (x) = (25.0)(0.7152) = 17.88$$

Example Spreadsheet Calculation:

Slope from curve, m:	0.213
Intercept from curve, b:	-0.00642
Area of analyte, Ax:	86550
Area of Internal Standard, Ais:	593147
Concentration of IS, Cis	25.00
Response Ratio:	0.145917
Amount Ratio:	0.715195
Concentration:	17.87988
Units of Internal Standard:	ug/L

5.0 Concentration from Quadratic Regression**Step 1 - Retrieve Curve Data from Plot, $y = Ax^2 + Bx + C$**

Where:

$$Ax^2 + Bx + (C - y) = 0$$

A, B, C = constants from the ICAL quadratic regression

y = Response ratio = Area of analyte/Area of internal standard (IS)

x = Amount ratio = Concentration of analyte/concentration of IS

Step 2: Calculate y from Quantitation Report

$$y = Ax/Ais$$

Step 3: Solve for x using the quadratic formula

$$Ax^2 + Bx + C - y = 0$$

$$x = \frac{b \pm \sqrt{(b^2 - 4a(c - y))}}{2a} \quad (\text{Two possible solutions})$$

Step 4: Solve for analyte concentration Cx

$$Cx = (Cis)(\text{Amount ratio})$$

Example Spreadsheet Calculation:

Value of A from plot:	-0.00629
Value of B from plot:	0.511
Value of C from plot:	-0.0276
Area of unknown from quantitation report:	293821
Area of IS from quantitation report:	784848
Response ratio, y:	0.374367
C - y:	-0.40197
Root 1 - Computed amount ratio, X1:	80.44567
Root 2 - Computed amount ratio, X2:	0.794396 use this solution
Concentration of IS, Cis:	25.00
Concentration of analyte, Cx:	19.86 ug/L

Instrument Run Log

Instrument: HPMS8 Dataset: 111218
 Analyst1: EEA Analyst2: NA
 Method: 8260B SOP: MSV01 Rev: 19
 Method: 5030/5035 SOP: PAT01 Rev: 16

Maintenance Log ID: _____

Internal Standard: STD90422 Surrogate Standard: STD90202
 CCV: STD90360 LCS: STD90635 MS/MSD: STD90635

Column 1 ID: RTX502.2 Column 2 ID: NA
 Workgroups: WG684281 ICAL A9

Comments:

File ID	Sample Information	pH	Mat	Dil	Reference	Date/Time
8M427929	WG684281-01 50ng/ BFB STD	NA	1	1	STD90475	11/12/18 13:12
8M427930	WG684281-02 5 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 13:37
8M427931	WG684281-03 20 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 14:06
8M427932	WG684281-04 50 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 14:34
8M427933	WG684281-05 100 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 15:03
8M427934	WG684281-06 200 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 15:31
8M427935	WG684281-07 300 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 15:59
8M427936	WG684281-08 400 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 16:28
8M427937	WG684281-09 500 ug/L ICAL A9 8260	NA	1	1	STD90626	11/12/18 16:57
8M427938	RINSE	NA	1	1	STD90626	11/12/18 17:25
8M427939	WG684281-10 100 ug/L ICV/ALT A9 8260	NA	1	1	STD90627	11/12/18 17:54
8M427940	WG684351-01 BLANK 8260	NA	1	1		11/12/18 18:22
8M427941	WG684351-02 P+A 100 ug	NA	1	1		11/12/18 18:52
8M427942	WG684351-03 P+A 100 ug/L 8260	NA	1	1	STD90627	11/12/18 19:21
8M427943	WG684351-04 P+A 100 ug/L 8260	NA	1	1	STD90627	11/12/18 19:49
8M427944	WG684351-05 P+A 100 ug/L 8260	NA	1	1	STD90627	11/12/18 20:17

Approved: November 15, 2018

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Instrument Run Log

Instrument: HPMS11 Dataset: 021919
 Analyst1: KFR Analyst2: NA
 Method: 8260B SOP: MSV01, OVAP MSV01 Rev: 25.0
 Method: 5035A/5030B/5030C SOP: PATO1, OVAP PATO1 Rev: 19,1

Maintenance Log ID: _____

Internal Standard: STD91917 Surrogate Standard: STD91918
 CCV: STD92151 LCS: STD92076 MS/MSD: NA
 Column 1 ID: RTX502.2 Column 2 ID: NA
 Workgroups: WG696624

Comments:

File ID	Sample Information	pH	Mat	Dil	Reference	Date/Time
11M29623	WG696624-01 50ng BFB 8260	NA	1	1	STD91899	02/19/19 10:43
11M29624	WG696624-02 CCV 50ug/L 8260	NA	1	1	STD91899	02/19/19 11:12
11M29625	WG696624-02 CCV 50ug/L 8260	NA	1	1	STD91899	02/19/19 11:48
11M29626	WG696507-12 ICV 50ug/L 8260	NA	1	1	STD91879	02/19/19 12:18
11M29628	WG696624-03 5ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 13:17
11M29629	WG696624-04 20ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 13:47
11M29630	WG696624-05 50ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 14:16
11M29631	WG696624-06 100ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 14:45
11M29632	WG696624-07 200ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 15:15
11M29633	WG696624-08 300ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 15:44
11M29634	WG696624-09 400ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 16:14
11M29635	WG696624-10 500ug/L ICAL A9 8260	NA	1	1	STD92151	02/19/19 16:44
11M29636	RINSE	NA	1	1		02/19/19 17:13
11M29637	WG696624-11 ICV/ALT 100ug/L ICAL A9 82	NA	1	1	STD92076	02/19/19 17:43

Approved: February 20, 2019

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Instrument Run Log

Instrument: HPMS8 Dataset: 030419
 Analyst1: EEA Analyst2: NA
 Method: 8260B SOP: MSV01 Rev: 26
 Method: 5030/5035 SOP: PAT01 Rev: 18

Maintenance Log ID: _____

Internal Standard: STD92348 Surrogate Standard: STD92349
 CCV: STD92326 LCS: STD92354 MS/MSD: NA
 Column 1 ID: RTX502.2 Column 2 ID: NA
 Workgroups: WG698192 ICAL

Comments:

File ID	Sample Information	pH	Mat	Dil	Reference	Date/Time
8M428954	WG698192-01 50ng BFB 8260	NA	1	1	STD91948	03/04/19 11:54
8M428956	WG698192-02 0.3 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 12:45
8M428957	WG698192-03 0.4 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 13:15
8M428958	WG698192-04 1 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 13:43
8M428959	WG698192-05 2 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 14:12
8M428960	WG698192-06 5 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 14:42
8M428961	WG698192-07 20 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 15:11
8M428962	WG698192-08 50 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 15:40
8M428963	WG698192-09 100 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 16:09
8M428964	WG698192-10 200 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 16:38
8M428965	WG698192-11 300 ug/L ICAL 8260	NA	1	1	STD92326	03/04/19 17:07
8M428967	WG698192-12 ALT 50 ug/L ICAL 8260	NA	1	1	STD92354	03/04/19 18:06

Approved: March 05, 2019

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Instrument Run Log

Instrument: HPMS11 Dataset: 030519
 Analyst1: KFR Analyst2: NA
 Method: 8260B SOP: MSV01, OVAP MSV01 Rev: 25.0
 Method: 5035A/5030B/5030C SOP: PATO1, OVAP PATO1 Rev: 19,1

Maintenance Log ID: _____

Internal Standard: STD92347 Surrogate Standard: STD92346
 CCV: STD92320 LCS: STD92316 MS/MSD: NA
 Column 1 ID: RTX502.2 Column 2 ID: NA
 Workgroups: WG698387, WG698731

Comments:

File ID	Sample Information	pH	Mat	Dil	Reference	Date/Time
11M29730	WG698387-02 CCV 50ug/kg 8260	NA	1	1	STD92189	03/05/19 12:58
11M29735	WG698387-01 50ng BFB 8260	NA	1	1	STD91899	03/05/19 15:39
11M29736	RINSE	NA	1	1		03/05/19 16:10
11M29737	WG698387-02 STD 0.3ug/L 8260	NA	1	1	STD92320	03/05/19 16:39
11M29738	WG698387-03 STD 0.4ug/L 8260	NA	1	1	STD92320	03/05/19 17:09
11M29739	WG698387-04 STD 1ug/L 8260	NA	1	1	STD92320	03/05/19 17:38
11M29740	WG698387-05 STD 2ug/L 8260	NA	1	1	STD92320	03/05/19 18:08
11M29741	WG698387-06 STD 5ug/L 8260	NA	1	1	STD92320	03/05/19 18:38
11M29742	WG698387-07 STD 20ug/L 8260	NA	1	1	STD92320	03/05/19 19:08
11M29743	WG698387-08 STD 50ug/L 8260	NA	1	1	STD92320	03/05/19 19:38
11M29744	WG698387-09 STD 100ug/L 8260	NA	1	1	STD92320	03/05/19 20:08
11M29745	WG698387-10 STD 200ug/L 8260	NA	1	1	STD92320	03/05/19 20:37
11M29746	WG698387-11 STD 300ug/L 8260	NA	1	1	STD92320	03/05/19 21:06
11M29747	RINSE	NA	1	1		03/05/19 21:35
11M29748	WG698387-12 ICV 50ug/L 8260	NA	1	1	STD92316	03/05/19 22:04
11M29749	RINSE	NA	1	1		03/05/19 22:33
11M29750	WG698731-01 BLANK 8260	NA	1	1		03/05/19 23:02
11M29751	L19010001-01 A LOQ 8260	NA	1	1	STD92184	03/05/19 23:30
11M29752	L19010001-01 B LOQ 8260	NA	1	1	STD92185	03/06/19 00:00
11M29753	RINSE	NA	1	1		03/06/19 00:29
11M29754	RINSE	NA	1	1		03/06/19 00:58

Approved: March 07, 2019

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Instrument Run Log

Instrument: HPMS8 Dataset: 031119
 Analyst1: EEA Analyst2: NA
 Method: 8260B SOP: MSV01 Rev: 26
 Method: 5030/5035 SOP: PAT01 Rev: 18

Maintenance Log ID: _____

Internal Standard: STD92348 Surrogate Standard: STD92349
 CCV: STD92326 LCS: STD92354 MS/MSD: NA
 Column 1 ID: RTX502.2 Column 2 ID: NA
 Workgroups: WG699028

Comments:

File ID	Sample Information	pH	Mat	Dil	Reference	Date/Time
8M429060	WG699027-01 50ng BFB 8260	NA	1	1	STD92421	03/11/19 08:56
8M429061	WG699027-02 CCV 50 ug/L 8260	NA	1	1	STD92326	03/11/19 09:19
8M429062	WG000000-01 A9 100 ug/L 8260	NA	1	1	STD00000	03/11/19 10:12
8M429063	WG699028-01 BLANK 8260	NA	1	1		03/11/19 10:41
8M429064	WG699028-02 LCS 20 ug/L 8260	NA	1	1	STD92354	03/11/19 11:09
8M429065	WG699028-03 LCS DUP 20 ug/L 8260	NA	1	1	STD92354	03/11/19 11:38
8M429066	L19030582-02 A 826-BETX	<2	1	1		03/11/19 12:07
8M429067	L19030584-02 A 826-BETX	<2	1	1		03/11/19 12:36
8M429068	L19030586-02 A 826-BETX	<2	1	1		03/11/19 13:05
8M429069	L19030582-01 A 826-BETX	<2	1	1		03/11/19 13:35
8M429070	L19030583-01 A 826-BETX	<2	1	1		03/11/19 14:04
8M429071	L19030584-01 A 826-BETX	<2	1	1		03/11/19 14:34
8M429072	L19030585-01 A 826-BETX	<2	1	1		03/11/19 15:03
8M429073	L19030586-01 A 826-BETX	<2	1	1		03/11/19 15:32
8M429074	L19030587-01 A 826-BETX	<2	1	1		03/11/19 16:02
8M429075	L19030239-01 A 10X 826-TC-SPE	NA	17	10		03/11/19 16:31
8M429076	L19030638-04 A 826-SPE	<2	1	1		03/11/19 17:01
8M429077	L19030638-05 A 826-SPE	<2	1	1		03/11/19 17:30
8M429078	L19030638-06 A 826-SPE	<2	1	1		03/11/19 17:59
8M429079	L19030638-07 A 826-SPE	<2	1	1		03/11/19 18:28
8M429080	L19030631-01 B 826-SPE1	NA	12	100		03/11/19 18:57
8M429081	RINSE	NA	1	1		03/11/19 19:26
8M429082	RINSE	NA	1	1		03/11/19 19:54
8M429083	WG699028-04 CCV 50 ug/L 8260	NA	1	1	STD92326	03/11/19 20:23
8M429084	RINSE	NA	1	1		03/11/19 20:51
8M429085	WG698729-01 A 10X 826-TC-SPE	NA	17	10		03/11/19 21:20

Approved: March 12, 2019

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Instrument Run Log

Instrument: HPMS11 Dataset: 031219
 Analyst1: KFR Analyst2: NA
 Method: 8260B SOP: MSV01, OVAP MSV01 Rev: 26.0
 Method: 5035A/5030B/5030C SOP: PATO1, OVAP PATO1 Rev: 18,1

Maintenance Log ID: _____

Internal Standard: STD92347 Surrogate Standard: STD92346
 CCV: STD92189 STD92320 LCS: STD92316 MS/MSD: NA
 Column 1 ID: RTX502.2 Column 2 ID: NA
 Workgroups: WG699219

Comments:

File ID	Sample Information	pH	Mat	Dil	Reference	Date/Time
11M29828	WG699218-01 50ng BFB 8260	NA	1	1	STD92290	03/12/19 10:56
11M29829	WG699218-02 CCV 50ug/kg 8260	NA	1	1	STD92189	03/12/19 11:21
11M29830	WG000000-01 A9 50ug/L 8260	NA	1	1	STD92151	03/12/19 12:02
11M29831	WG699219-01 BLANK 8260	NA	1	1		03/12/19 12:31
11M29832	WG699219-02 LCS 20ug/L 8260	NA	1	1	STD92316	03/12/19 13:00
11M29833	L19030638-02 A MS 826-SPE	<2	1	1	STD92316	03/12/19 13:30
11M29834	L19030638-03 A MSD 826-SPE	<2	1	1	STD92316	03/12/19 14:00
11M29835	RINSE	NA	1	1		03/12/19 14:30
11M29836	L19030638-08 A TBLK 826-SPE	<2	1	1		03/12/19 14:59
11M29837	L19030638-01 A REF 826-SPE	<2	1	1		03/12/19 15:30
11M29838	L19030660-01 A 826-SPE	<2	1	1		03/12/19 16:00
11M29839	L19030660-02 A 826-SPE	<2	1	1		03/12/19 16:37
11M29840	L19030660-05 A 826-SPE	<2	1	1		03/12/19 17:06
11M29841	L19030660-06 A 826-SPE	<2	1	1		03/12/19 17:37
11M29842	L19030660-08 A 826-SPE	<2	1	1		03/12/19 18:13
11M29843	L19030660-03 2X A 826-SPE	<2	1	2	SEDIMENT	03/12/19 18:43
11M29844	L19030660-04 2X A 826-SPE	3	1	2	SEDIMENT	03/12/19 19:13
11M29845	L19030660-09 2X A 826-SPE	<2	1	2	SEDIMENT	03/12/19 19:43
11M29846	L19030660-10 2X A 826-SPE	<2	1	2	SEDIMENT	03/12/19 20:13
11M29847	L19030660-07 2X A 826-SPE	4	1	2	SEDIMENT	03/12/19 20:42
11M29848	RINSE	NA	1	1		03/12/19 21:12
11M29849	L19030687-03 A 826-BETX	<2	1	1		03/12/19 21:41
11M29850	L19030687-01 A 826-BETX	<2	1	1		03/12/19 22:10
11M29851	WG699219-03 QCMRL 50ug/L 8260	NA	1	1	STD92320	03/12/19 22:39
11M29852	RINSE	NA	1	1		03/12/19 23:09
11M29853	RINSE	NA	1	1		03/12/19 23:39

Approved: March 18, 2019

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



Data Checklist

Date: 12-NOV-2018
 Analyst: EEA
 Analyst: NA
 Method: 8260B
 Instrument: HPMS8
 Curve Workgroup: WG684281
 Runlog ID: 93326
 Analytical Workgroups: _____

System Performance Check	NA
BFB	X
Initial Calibration	X
Average RF	X
Linear Reg or Higher Order Curve	X
Second Source standard % Difference	X
Continuing Calibration /Check Standards	X
Project/Client Specific Requirements	X
Special Standards	NA
Blanks	X
TCL's	X
Surrogates	X
LCS (Laboratory Control Sample)	X
Recoveries	X
Surrogates	X
MS/MSD/Duplicates	NA
Samples	X
TCL Hits	X
Spectra of TCL Hits	EEA
Surrogates	X
Internal Standards Criteria	X
Library Searches	NA
Calculations & Correct Factors	X
Dilutions Run	NA
Reruns	NA
Manual Integrations	NA
Case Narrative	X
Results Reporting/Data Qualifiers	X
KOBRA Workgroup Data	X
Check for Completeness	X
Primary Reviewer	EEA
Secondary Reviewer	ADC
Check for compliance with method and project specific requirements	X
Check the completeness of reported information	X
Check the information for the report narrative	X
Check the reasonableness of the results	X

Primary Reviewer:
14-NOV-2018



Secondary Reviewer:
15-NOV-2018




Data Checklist

Date: 19-FEB-2019
 Analyst: KFR
 Analyst: NA
 Method: 8260B
 Instrument: HPMS11
 Curve Workgroup: NA
 Runlog ID: 95301
 Analytical Workgroups: WG696624

System Performance Check	NA
BFB	X
Initial Calibration	X
Average RF	X
Linear Reg or Higher Order Curve	X
Second Source standard % Difference	X
Continuing Calibration /Check Standards	X
Project/Client Specific Requirements	X
Special Standards	NA
Blanks	X
TCL's	X
Surrogates	X
LCS (Laboratory Control Sample)	X
Recoveries	X
Surrogates	X
MS/MSD/Duplicates	NA
Samples	NA
TCL Hits	X
Spectra of TCL Hits	KFR
Surrogates	X
Internal Standards Criteria	X
Library Searches	NA
Calculations & Correct Factors	X
Dilutions Run	NA
Reruns	X
Manual Integrations	NA
Case Narrative	X
Results Reporting/Data Qualifiers	X
KOBRA Workgroup Data	X
Check for Completeness	X
Primary Reviewer	KFR
Secondary Reviewer	ADC
Check for compliance with method and project specific requirements	X
Check the completeness of reported information	X
Check the information for the report narrative	X
Check the reasonableness of the results	X

Primary Reviewer:
20-FEB-2019

Secondary Reviewer:
20-FEB-2019

Karissa Reynolds

ADC



Data Checklist

Date: 04-MAR-2019
 Analyst: EEA
 Analyst: NA
 Method: 8260B
 Instrument: HPMS8
 Curve Workgroup: WG698192
 Runlog ID: 95571
 Analytical Workgroups: _____

System Performance Check	NA
BFB	X
Initial Calibration	X
Average RF	X
Linear Reg or Higher Order Curve	X
Second Source standard % Difference	X
Continuing Calibration /Check Standards	X
Project/Client Specific Requirements	X
Special Standards	NA
Blanks	X
TCL's	X
Surrogates	X
LCS (Laboratory Control Sample)	X
Recoveries	X
Surrogates	X
MS/MSD/Duplicates	NA
Samples	NA
TCL Hits	X
Spectra of TCL Hits	EEA
Surrogates	X
Internal Standards Criteria	X
Library Searches	NA
Calculations & Correct Factors	X
Dilutions Run	NA
Reruns	NA
Manual Integrations	NA
Case Narrative	X
Results Reporting/Data Qualifiers	X
KOBRA Workgroup Data	X
Check for Completeness	X
Primary Reviewer	EEA
Secondary Reviewer	ADC
Check for compliance with method and project specific requirements	X
Check the completeness of reported information	X
Check the information for the report narrative	X
Check the reasonableness of the results	X

Primary Reviewer:
05-MAR-2019



Secondary Reviewer:
05-MAR-2019




Data Checklist

Date: 05-MAR-2019
 Analyst: KFR
 Analyst: NA
 Method: 8260B
 Instrument: HPMS11
 Curve Workgroup: NA
 Runlog ID: 95609
 Analytical Workgroups: WG698387

System Performance Check	NA
BFB	X
Initial Calibration	X
Average RF	X
Linear Reg or Higher Order Curve	X
Second Source standard % Difference	X
Continuing Calibration /Check Standards	X
Project/Client Specific Requirements	X
Special Standards	NA
Blanks	X
TCL's	X
Surrogates	X
LCS (Laboratory Control Sample)	X
Recoveries	X
Surrogates	X
MS/MSD/Duplicates	NA
Samples	X
TCL Hits	X
Spectra of TCL Hits	KFR
Surrogates	X
Internal Standards Criteria	X
Library Searches	NA
Calculations & Correct Factors	X
Dilutions Run	NA
Reruns	NA
Manual Integrations	NA
Case Narrative	X
Results Reporting/Data Qualifiers	X
KOBRA Workgroup Data	X
Check for Completeness	X
Primary Reviewer	KFR
Secondary Reviewer	ADC
Check for compliance with method and project specific requirements	X
Check the completeness of reported information	X
Check the information for the report narrative	X
Check the reasonableness of the results	X

Primary Reviewer:
07-MAR-2019

Karissa Reynolds

Secondary Reviewer:
07-MAR-2019

Adrian Carter




Data Checklist

Date: 11-MAR-2019
 Analyst: EEA
 Analyst: NA
 Method: 8260B
 Instrument: HPMS8
 Curve Workgroup: NA
 Runlog ID: 95685
 Analytical Workgroups: WG699028

System Performance Check	NA
BFB	X
Initial Calibration	X
Average RF	X
Linear Reg or Higher Order Curve	X
Second Source standard % Difference	X
Continuing Calibration /Check Standards	X
Project/Client Specific Requirements	X
Special Standards	NA
Blanks	X
TCL's	X
Surrogates	X
LCS (Laboratory Control Sample)	X
Recoveries	X
Surrogates	X
MS/MSD/Duplicates	X
Samples	X
TCL Hits	X
Spectra of TCL Hits	EEA
Surrogates	X
Internal Standards Criteria	X
Library Searches	NA
Calculations & Correct Factors	X
Dilutions Run	X
Reruns	NA
Manual Integrations	NA
Case Narrative	X
Results Reporting/Data Qualifiers	X
KOBRA Workgroup Data	X
Check for Completeness	X
Primary Reviewer	EEA
Secondary Reviewer	ADC
Check for compliance with method and project specific requirements	X
Check the completeness of reported information	X
Check the information for the report narrative	X
Check the reasonableness of the results	X

Primary Reviewer:
12-MAR-2019



Secondary Reviewer:
12-MAR-2019




Data Checklist

Date: 12-MAR-2019
 Analyst: KFR
 Analyst: NA
 Method: 8260B
 Instrument: HPMS11
 Curve Workgroup: NA
 Runlog ID: 95794
 Analytical Workgroups: WG699219

System Performance Check	NA
BFB	X
Initial Calibration	X
Average RF	X
Linear Reg or Higher Order Curve	X
Second Source standard % Difference	X
Continuing Calibration /Check Standards	X
Project/Client Specific Requirements	X
Special Standards	NA
Blanks	X
TCL's	X
Surrogates	X
LCS (Laboratory Control Sample)	X
Recoveries	X
Surrogates	X
MS/MSD/Duplicates	X
Samples	X
TCL Hits	X
Spectra of TCL Hits	KFR
Surrogates	X
Internal Standards Criteria	X
Library Searches	NA
Calculations & Correct Factors	X
Dilutions Run	X
Reruns	NA
Manual Integrations	NA
Case Narrative	X
Results Reporting/Data Qualifiers	X
KOBRA Workgroup Data	X
Check for Completeness	X
Primary Reviewer	KFR
Secondary Reviewer	EEA
Check for compliance with method and project specific requirements	X
Check the completeness of reported information	X
Check the information for the report narrative	X
Check the reasonableness of the results	X

Primary Reviewer:
15-MAR-2019

Secondary Reviewer:
18-MAR-2019

Karissa Reynolds

E. E. A.



Analytical Method:8260B
Login Number:L19030638

AAB#:WG699028

Client ID	ID	Date Collected	TCLP Date	Time Held	Max Hold	Q	Extract Date	Time Held	Max Hold	Q	Run Date	Time Held	Max Hold	Q
SW2	04	03/07/19					03/11/2019	4.2	14		03/11/19	4.2	14	
SW3	05	03/07/19					03/11/2019	4.2	14		03/11/19	4.2	14	
SW4	06	03/07/19					03/11/2019	4.1	14		03/11/19	4.1	14	
DUP01	07	03/07/19					03/11/2019	4.4	14		03/11/19	4.4	14	

* = SEE PROJECT QAPP REQUIREMENTS

HOLD_TIMES - Modified 03/06/2008
PDF File ID: 6344147
Report generated 03/16/2019 10:31



Analytical Method:8260B
Login Number:L19030638

AAB#:WG699219

Client ID	ID	Date Collected	TCLP Date	Time Held	Max Hold	Q	Extract Date	Time Held	Max Hold	Q	Run Date	Time Held	Max Hold	Q
SW1	01	03/07/19					03/12/2019	5.2	14		03/12/19	5.2	14	
SW1 MS	02	03/07/19					03/12/2019	5.1	14		03/12/19	5.1	14	
SW1 MSD	03	03/07/19					03/12/2019	5.1	14		03/12/19	5.1	14	
TB01	08	03/07/19					03/12/2019	5.3	14		03/12/19	5.3	14	

* = SEE PROJECT QAPP REQUIREMENTS

HOLD_TIMES - Modified 03/06/2008
PDF File ID: 6344147
Report generated 03/16/2019 10:31



Login Number: L19030638
 Instrument Id: HPMS8
 Workgroup (AAB#): WG699028

Method: 8260
 CAL ID: HPMS8-04-MAR-19
 Matrix: Water

Sample Number	Dilution	Tag	1	2	3	4
L19030638-04	1.00	01	108	106	112	105
L19030638-05	1.00	01	106	104	106	103
L19030638-06	1.00	01	108	104	111	103
L19030638-07	1.00	01	111	106	114	106
WG699028-01	1.00	01	103	103	109	104
WG699028-02	1.00	01	105	104	104	102
WG699028-03	1.00	01	106	106	102	102
WG699028-04	1.00	01	104	103	101	101

Surrogates	Surrogate Limits		
1 - 1,2-Dichloroethane-d4	81	-	118
2 - Dibromofluoromethane	80	-	119
3 - 4-Bromofluorobenzene	85	-	114
4 - Toluene-d8	89	-	112

Underline = Result out of surrogate limits

DL = surrogate diluted out

ND = surrogate not detected



Login Number: L19030638
 Instrument Id: HPMS11
 Workgroup (AAB#): WG699219

Method: 8260
 CAL ID: HPMS11-05-MAR-19
 Matrix: Water

Sample Number	Dilution	Tag	1	2	3	4
L19030638-01	1.00	01	96.8	96.0	99.7	99.9
L19030638-02	1.00	01	96.5	101	101	100
L19030638-03	1.00	01	99.3	102	106	102
L19030638-08	1.00	01	98.6	99.4	98.2	99.6
WG699219-01	1.00	01	95.7	98.9	106	102
WG699219-02	1.00	01	98.5	101	100	99.8
WG699219-03	1.00	01	100	105	98.7	101

Surrogates	Surrogate Limits		
1 - 1,2-Dichloroethane-d4	81	-	118
2 - Dibromofluoromethane	80	-	119
3 - 4-Bromofluorobenzene	85	-	114
4 - Toluene-d8	89	-	112

Underline = Result out of surrogate limits

DL = surrogate diluted out

ND = surrogate not detected



METHOD BLANK SUMMARY

Login Number: L19030638 Work Group: WG699219
 Blank File ID: 11M29831 Blank Sample ID: WG699219-01
 Prep Date: 03/12/19 12:31 Instrument ID: HPMS11
 Analyzed Date: 03/12/19 12:31 Method: 8260B
 Analyst: KFR

This Method Blank Applies To The Following Samples:

Client ID	Lab Sample ID	Lab File ID	Time Analyzed	TAG
LCS	WG699219-02	11M29832	03/12/19 13:00	01
SW1 MS	L19030638-02	11M29833	03/12/19 13:30	01
SW1 MSD	L19030638-03	11M29834	03/12/19 14:00	01
TB01	L19030638-08	11M29836	03/12/19 14:59	01
SW1	L19030638-01	11M29837	03/12/19 15:30	01
QCMRL	WG699219-03	11M29851	03/12/19 22:39	01

Report Name: BLANK_SUMMARY
 PDF File ID: 6344148
 Report generated 03/16/2019 10:31



METHOD BLANK SUMMARY

Login Number: L19030638 Work Group: WG699028
 Blank File ID: 8M429063 Blank Sample ID: WG699028-01
 Prep Date: 03/11/19 10:41 Instrument ID: HPMS8
 Analyzed Date: 03/11/19 10:41 Method: 8260B
 Analyst: EEA

This Method Blank Applies To The Following Samples:

Client ID	Lab Sample ID	Lab File ID	Time Analyzed	TAG
LCS	WG699028-02	8M429064	03/11/19 11:09	01
LCS2	WG699028-03	8M429065	03/11/19 11:38	01
SW2	L19030638-04	8M429076	03/11/19 17:01	01
SW3	L19030638-05	8M429077	03/11/19 17:30	01
SW4	L19030638-06	8M429078	03/11/19 17:59	01
DUP01	L19030638-07	8M429079	03/11/19 18:28	01
QCMRL	WG699028-04	8M429083	03/11/19 20:23	01

Report Name: BLANK_SUMMARY
 PDF File ID: 6344148
 Report generated 03/16/2019 10:31



Login Number: L19030638 Prep Date: 03/12/19 12:31 Sample ID: WG699219-01
 Instrument ID: HPMS11 Run Date: 03/12/19 12:31 Prep Method: 5030B/5030C/503
 File ID: 11M29831 Analyst: KFR Method: 8260B
 Workgroup (AAB#): WG699219 Matrix: Water Units: ug/L
 Contract #: _____ Cal ID: HPMS11-05-MAR-19

Analytes	DL	LOQ	Concentration	Dilution	Qualifier
1,1-Dichloroethene	0.500	2.00	0.500	1	U
1,2-Dichloroethane	0.250	1.00	0.250	1	U
Acetone	2.50	10.0	2.50	1	U
Chloroform	0.125	0.500	0.125	1	U
cis-1,2-Dichloroethene	0.250	1.00	0.250	1	U
Tetrachloroethene	0.250	1.00	0.250	1	U
trans-1,2-Dichloroethene	0.250	1.00	0.250	1	U
Trichloroethene	0.250	1.00	0.250	1	U
Vinyl chloride	0.250	1.00	0.250	1	U

Surrogates	% Recovery	Surrogate Limits	Qualifier
1,2-Dichloroethane-d4	95.7	81 - 118	PASS
4-Bromofluorobenzene	106	85 - 114	PASS
Dibromofluoromethane	98.9	80 - 119	PASS
Toluene-d8	102	89 - 112	PASS

DL Method Detection Limit
 LOQ Reporting/Practical Quantitation Limit
 ND Analyte Not detected at or above reporting limit
 * |Analyte concentration| > 1/2 RL

Report Name: BLANK
 PDF ID: 6344149
 16-MAR-2019 10:31



Login Number: L19030638 Prep Date: 03/11/19 10:41 Sample ID: WG699028-01
 Instrument ID: HPMS8 Run Date: 03/11/19 10:41 Prep Method: 5030B/5030C/503
 File ID: 8M429063 Analyst: EEA Method: 8260B
 Workgroup (AAB#): WG699028 Matrix: Water Units: ug/L
 Contract #: _____ Cal ID: HPMS8-04-MAR-19

Analytes	DL	LOQ	Concentration	Dilution	Qualifier
1,1-Dichloroethene	0.500	2.00	0.500	1	U
1,2-Dichloroethane	0.250	1.00	0.250	1	U
Acetone	2.50	10.0	2.50	1	U
Chloroform	0.125	0.500	0.125	1	U
cis-1,2-Dichloroethene	0.250	1.00	0.250	1	U
Tetrachloroethene	0.250	1.00	0.250	1	U
trans-1,2-Dichloroethene	0.250	1.00	0.250	1	U
Trichloroethene	0.250	1.00	0.250	1	U
Vinyl chloride	0.250	1.00	0.250	1	U

Surrogates	% Recovery	Surrogate Limits	Qualifier
1,2-Dichloroethane-d4	103	81 - 118	PASS
4-Bromofluorobenzene	109	85 - 114	PASS
Dibromofluoromethane	103	80 - 119	PASS
Toluene-d8	104	89 - 112	PASS

DL Method Detection Limit
 LOQ Reporting/Practical Quantitation Limit
 ND Analyte Not detected at or above reporting limit
 * |Analyte concentration| > 1/2 RL

Report Name: BLANK
 PDF ID: 6344149
 16-MAR-2019 10:31



Login Number: L19030638 Run Date: 03/12/2019 Sample ID: WG699219-02
 Instrument ID: HPMS11 Run Time: 13:00 Prep Method: 5030B/5030C/503
 File ID: 11M29832 Analyst: KFR Method: 8260B
 Workgroup (AAB#): WG699219 Matrix: Water Units: ug/L
 QC Key: DOD5 Lot#: STD92316 Cal ID: HPMS11-05-MAR-19

Analytes	Expected	Found	% Rec	LCS Limits	Q
1,1-Dichloroethene	20.0	19.3	96.3	71 - 131	
1,2-Dichloroethane	20.0	20.2	101	73 - 128	
Acetone	20.0	14.7	73.7	39 - 160	
Chloroform	20.0	20.0	99.9	79 - 124	
cis-1,2-Dichloroethene	20.0	21.2	106	78 - 123	
Tetrachloroethene	20.0	20.0	99.9	74 - 129	
trans-1,2-Dichloroethene	20.0	20.9	105	75 - 124	
Trichloroethene	20.0	20.1	100	79 - 123	
Vinyl chloride	20.0	15.3	76.3	58 - 137	

Surrogates	% Recovery	Surrogate Limits	Qualifier
1,2-Dichloroethane-d4	98.5	81 - 118	PASS
4-Bromofluorobenzene	100	85 - 114	PASS
Dibromofluoromethane	101	80 - 119	PASS
Toluene-d8	99.8	89 - 112	PASS

* EXCEEDS %REC LIMIT

LCS - Modified 03/06/2008
 PDF File ID: 6351178
 Report generated: 03/25/2019 09:45



Login Number: L19030638 Analyst: EEA Prep Method: 5030B/5030C/503
 Instrument ID: HPMS8 Matrix: Water Method: 8260B
 Workgroup (AAB#): WG699028 Units: ug/L
 QC Key: DOD5 Lot #: STD92354
 Sample ID: WG699028-02 LCS File ID: 8M429064 Run Date: 03/11/2019 11:09
 Sample ID: WG699028-03 LCS2 File ID: 8M429065 Run Date: 03/11/2019 11:38

Analytes	LCS			LCS2			%RPD	%Rec Limits	RPD Lmt	Q
	Known	Found	% REC	Known	Found	% REC				
1,1-Dichloroethene	20.0	20.1	100	20.0	19.1	95.5	4.97	71 - 131	20	
1,2-Dichloroethane	20.0	20.8	104	20.0	20.3	102	2.16	73 - 128	20	
Acetone	20.0	14.7	73.3	20.0	14.9	74.4	1.53	39 - 160	20	
Chloroform	20.0	20.0	100	20.0	19.5	97.4	2.79	79 - 124	20	
cis-1,2-Dichloroethene	20.0	20.7	104	20.0	20.2	101	2.51	78 - 123	20	
Tetrachloroethene	20.0	20.7	103	20.0	19.6	97.8	5.60	74 - 129	20	
trans-1,2-Dichloroethene	20.0	20.5	102	20.0	19.9	99.4	3.01	75 - 124	20	
Trichloroethene	20.0	20.9	105	20.0	20.5	102	2.23	79 - 123	20	
Vinyl chloride	20.0	18.7	93.3	20.0	17.9	89.6	4.11	58 - 137	20	

Surogates	LCS	LCS2	Surrogate Limits	Qualifier
	% Recovery	% Recovery		
1,2-Dichloroethane-d4	105	106	81 - 118	PASS
Dibromofluoromethane	104	106	80 - 119	PASS
4-Bromofluorobenzene	104	102	85 - 114	PASS
Toluene-d8	102	102	89 - 112	PASS

* EXCEEDS %REC LIMIT
EXCEEDS RPD LIMIT



Loginum: L19030638 Cal ID: HPMS11- 05-MAR-19
 Instrument ID: HPMS11 Contract #: _____
 Parent ID: L19030638-01 File ID: 11M29837 Dil: 1
 Sample ID: L19030638-02 MS File ID: 11M29833 Dil: 1
 Sample ID: L19030638-03 MSD File ID: 11M29834 Dil: 1

Worknum: WG699219
 Prep Method: 5030B/5030C/
 Method: 5035A
 Matrix: 8260B
 Units: Water
ug/L

Analyte	Parent	MS Spiked	MS Found	MS %Rec	MSD Spiked	MSD Found	MSD %Rec	%RPD	%Rec Limits	RPD Limit	Q
1,1-Dichloroethene	U	20.0	18.8	93.8	20.0	18.2	91.2	2.76	71 - 131	20	
1,2-Dichloroethane	U	20.0	20.0	100	20.0	20.1	100	0.224	73 - 128	20	
Acetone	3.56	20.0	18.1	72.6	20.0	18.4	74	1.47	39 - 160	20	
Chloroform	U	20.0	19.9	99.3	20.0	19.4	97	2.42	79 - 124	20	
cis-1,2-Dichloroethene	U	20.0	20.6	103	20.0	20.5	102	0.391	78 - 123	20	
Tetrachloroethene	U	20.0	19.5	97.5	20.0	19.4	97	0.543	74 - 129	20	
trans-1,2-Dichloroethene	U	20.0	20.0	99.9	20.0	19.7	98.5	1.42	75 - 124	20	
Trichloroethene	U	20.0	19.4	96.9	20.0	19.4	96.9	0.0217	79 - 123	20	
Vinyl chloride	U	20.0	14.4	72.2	20.0	14.2	70.9	1.78	58 - 137	20	

* FAILS %REC LIMIT

FAILS RPD LIMIT

BFB

Login Number: L19030638
Instrument: HPMS11
Analyst: KFR
Workgroup: WG698264

Tune ID: WG698264-01
Run Date: 02/19/2019
Run Time: 10:43
File ID: 11M29623
Cal ID: HPMS11-18-FEB-19

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50.0	95.0	15.0	40.0	24.8	6454	PASS
75.0	95.0	30.0	60.0	51.0	13278	PASS
95.0	95.0	100	100	100	26034	PASS
96.0	95.0	5.00	9.00	7.76	2021	PASS
173	174	0	2.00	0	0	PASS
174	95.0	50.0	100	83.6	21765	PASS
175	174	5.00	9.00	7.68	1671	PASS
176	174	95.0	101	99.3	21610	PASS
177	176	5.00	9.00	6.27	1355	PASS

This check relates to the following samples:

Lab ID	Client ID	Tag	Date Analyzed	Q
WG696624-03	STD	01	02/19/2019 13:17	
WG696624-04	STD	01	02/19/2019 13:47	
WG696624-05	STD	01	02/19/2019 14:16	
WG696624-06	STD-CCV	01	02/19/2019 14:45	
WG696624-07	STD	01	02/19/2019 15:15	
WG696624-08	STD	01	02/19/2019 15:44	
WG696624-09	STD	01	02/19/2019 16:14	
WG696624-10	STD	01	02/19/2019 16:44	
WG696624-11	SSCV	01	02/19/2019 17:43	

* Sample past 12 hour tune limit



BFB

Login Number: L19030638
Instrument: HPMS11
Analyst: KFR
Workgroup: WG698387

Tune ID: WG698387-01
Run Date: 03/05/2019
Run Time: 15:39
File ID: 11M29735

Cal ID: HPMS11-05-MAR-19

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50.0	95.0	15.0	40.0	28.5	5208	PASS
75.0	95.0	30.0	60.0	55.0	10035	PASS
95.0	95.0	100	100	100	18251	PASS
96.0	95.0	5.00	9.00	5.67	1034	PASS
173	174	0	2.00	0	0	PASS
174	95.0	50.0	100	74.8	13653	PASS
175	174	5.00	9.00	8.31	1135	PASS
176	174	95.0	101	98.9	13499	PASS
177	176	5.00	9.00	6.20	837	PASS

This check relates to the following samples:

Lab ID	Client ID	Tag	Date Analyzed	Q
WG698387-02	STD	01	03/05/2019 16:39	
WG698387-03	STD	01	03/05/2019 17:09	
WG698387-04	STD	01	03/05/2019 17:38	
WG698387-05	STD	01	03/05/2019 18:08	
WG698387-06	STD	01	03/05/2019 18:38	
WG698387-07	STD	01	03/05/2019 19:08	
WG698387-08	STD-CCV	01	03/05/2019 19:38	
WG698387-09	STD	01	03/05/2019 20:08	
WG698387-10	STD	01	03/05/2019 20:37	
WG698387-11	STD	01	03/05/2019 21:06	
WG698387-12	SSCV	01	03/05/2019 22:04	

* Sample past 12 hour tune limit



BFB

Login Number: L19030638 Tune ID: WG699218-01
 Instrument: HPMS11 Run Date: 03/12/2019
 Analyst: KFR Run Time: 10:56
 Workgroup: WG699218 File ID: 11M29828
 Cal ID: HPMS11-05-MAR-19

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50.0	95.0	15.0	40.0	23.7	7906	PASS
75.0	95.0	30.0	60.0	52.2	17441	PASS
95.0	95.0	100	100	100	33392	PASS
96.0	95.0	5.00	9.00	6.51	2173	PASS
173	174	0	2.00	0	0	PASS
174	95.0	50.0	100	83.3	27800	PASS
175	174	5.00	9.00	7.67	2131	PASS
176	174	95.0	101	96.6	26850	PASS
177	176	5.00	9.00	6.71	1801	PASS

This check relates to the following samples:

Lab ID	Client ID	Tag	Date Analyzed	Q
WG699218-02	CCV	01	03/12/2019 11:21	
WG699219-01	BLANK	01	03/12/2019 12:31	
WG699219-02	LCS	01	03/12/2019 13:00	
L19030638-02	SW1 MS	01	03/12/2019 13:30	
L19030638-03	SW1 MSD	01	03/12/2019 14:00	
L19030638-08	TB01	01	03/12/2019 14:59	
L19030638-01	SW1	01	03/12/2019 15:30	
WG699219-03	QCMRL	01	03/12/2019 22:39	

* Sample past 12 hour tune limit



BFB

Login Number: L19030638 Tune ID: WG684281-01
 Instrument: HPMS8 Run Date: 11/12/2018
 Analyst: EEA Run Time: 13:12
 Workgroup: WG684281 File ID: 8M427929
 Cal ID: HPMS8-12-NOV-18

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50.0	95.0	15.0	40.0	22.0	18596	PASS
75.0	95.0	30.0	60.0	48.9	41416	PASS
95.0	95.0	100	100	100	84642	PASS
96.0	95.0	5.00	9.00	6.98	5906	PASS
173	174	0	2.00	0.256	193	PASS
174	95.0	50.0	100	89.2	75469	PASS
175	174	5.00	9.00	7.05	5319	PASS
176	174	95.0	101	99.7	75216	PASS
177	176	5.00	9.00	6.87	5164	PASS

This check relates to the following samples:

Lab ID	Client ID	Tag	Date Analyzed	Q
WG684281-02	STD	01	11/12/2018 13:37	
WG684281-03	STD	01	11/12/2018 14:06	
WG684281-04	STD	01	11/12/2018 14:34	
WG684281-05	STD-CCV	01	11/12/2018 15:03	
WG684281-06	STD	01	11/12/2018 15:31	
WG684281-07	STD	01	11/12/2018 15:59	
WG684281-08	STD	01	11/12/2018 16:28	
WG684281-09	STD	01	11/12/2018 16:57	
WG684281-10	SSCV	01	11/12/2018 17:54	

* Sample past 12 hour tune limit



BFB

Login Number: L19030638 Tune ID: WG698192-01
 Instrument: HPMS8 Run Date: 03/04/2019
 Analyst: EEA Run Time: 11:54
 Workgroup: WG698192 File ID: 8M428954
 Cal ID: HPMS8-

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50.0	95.0	15.0	40.0	20.6	13778	PASS
75.0	95.0	30.0	60.0	45.4	30314	PASS
95.0	95.0	100	100	100	66746	PASS
96.0	95.0	5.00	9.00	6.45	4307	PASS
173	174	0	2.00	0.231	137	PASS
174	95.0	50.0	100	88.7	59210	PASS
175	174	5.00	9.00	7.60	4497	PASS
176	174	95.0	101	98.1	58093	PASS
177	176	5.00	9.00	6.48	3763	PASS

This check relates to the following samples:

Lab ID	Client ID	Tag	Date Analyzed	Q
WG698192-02	STD	01	03/04/2019 12:45	
WG698192-03	STD	01	03/04/2019 13:15	
WG698192-04	STD	01	03/04/2019 13:43	
WG698192-05	STD	01	03/04/2019 14:12	
WG698192-06	STD	01	03/04/2019 14:42	
WG698192-07	STD	01	03/04/2019 15:11	
WG698192-08	STD-CCV	01	03/04/2019 15:40	
WG698192-09	STD	01	03/04/2019 16:09	
WG698192-10	STD	01	03/04/2019 16:38	
WG698192-11	STD	01	03/04/2019 17:07	
WG698192-12	SSCV	01	03/04/2019 18:06	

* Sample past 12 hour tune limit



BFB

Login Number: L19030638
Instrument: HPMS8
Analyst: EEA
Workgroup: WG699027

Tune ID: WG699027-01
Run Date: 03/11/2019
Run Time: 08:56
File ID: 8M429060
Cal ID: HPMS8-04-MAR-19

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50.0	95.0	15.0	40.0	21.3	11408	PASS
75.0	95.0	30.0	60.0	45.9	24533	PASS
95.0	95.0	100	100	100	53437	PASS
96.0	95.0	5.00	9.00	7.71	4118	PASS
173	174	0	2.00	0.392	184	PASS
174	95.0	50.0	100	87.8	46904	PASS
175	174	5.00	9.00	7.10	3330	PASS
176	174	95.0	101	95.1	44586	PASS
177	176	5.00	9.00	7.15	3188	PASS

This check relates to the following samples:

Lab ID	Client ID	Tag	Date Analyzed	Q
WG699027-02	CCV	01	03/11/2019 09:19	
WG699028-01	BLANK	01	03/11/2019 10:41	
WG699028-02	LCS	01	03/11/2019 11:09	
WG699028-03	LCS2	01	03/11/2019 11:38	
L19030638-04	SW2	01	03/11/2019 17:01	
L19030638-05	SW3	01	03/11/2019 17:30	
L19030638-06	SW4	01	03/11/2019 17:59	
L19030638-07	DUP01	01	03/11/2019 18:28	
WG699028-04	QCMRL	01	03/11/2019 20:23	
WG698729-01	FBLK1	DL01	03/11/2019 21:20	*

* Sample past 12 hour tune limit



Calibration Table Report
 Method: A9FOOWT.M
 Title: Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Calibration: Wed Feb 20 09:41:04 2019
 Curve: WG696624
 Calibration Files

Compound	R ²								Avg	%RSD	Linear	Quadratic
	5	20	50	100	200	300	400	500				
11M29628.D	11M29629.D	11M29630.D	11M29631.D	11M29632.D	11M29633.D	11M29634.D	11M29635.D					
Fluorobenzene	ISTD											
Acetonitrile	0.037	0.043	0.039	0.036	0.034	0.036	0.035	0.036	0.037	8.060		
3-Chloro-1-propene	0.526	0.498	0.507	0.514	0.466	0.445	0.414	0.391	0.470	10.567		
2-Chloro-1,3-butadiene	0.594	0.553	0.561	0.595	0.533	0.508	0.469	0.439	0.531	10.635		
Methacrylonitrile	0.245	0.218	0.221	0.216	0.198	0.216	0.204	0.202	0.215	6.918		
Isobutyl Alcohol			0.015	0.014	0.014	0.016	0.015		0.015	6.329		
1-Butanol	0.006	0.008	0.008	0.009	0.008	0.009	0.009	0.009	0.008	11.670		
Cyclohexanone		0.023	0.021	0.021	0.020	0.021	0.020	0.021	0.021	5.842		
2-Nitropropane	0.100	0.090	0.095	0.096	0.093	0.104	0.104	0.103	0.098	5.484		
Ethyl Acetate	0.302	0.301	0.304	0.298	0.272	0.290	0.272	0.261	0.288	5.855		
Methyl methacrylate	0.307	0.285	0.293	0.287	0.262	0.273	0.256	0.245	0.276	7.559		
Chlorobenzene-d5	ISTD											
1,4-Dichlorobenzene-d4	ISTD											

Wed Feb 20 09:42:09 2019

Calibration Table Report
 Method: 8260WT.M
 Title: 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Calibration: Wed Mar 06 14:13:24 2019
 Curve: WG698387
 Calibration Files

Compound	Concentration (ppm)											Avg	%RSD	Linear R ²	Quadratic R ²
	0.3	0.4	1	2	5	20	50	100	200	300					
Fluorobenzene	ISTD														
Dichlorodifluoromethane			0.348	0.350	0.330	0.401	0.404	0.381	0.371			0.369	7.615		
Chloromethane	0.901	0.771	0.748	0.686	0.718	0.679	0.620	0.550				0.709	14.773		
Vinyl Chloride	0.596	0.529	0.558	0.526	0.565	0.543	0.490	0.430				0.530	9.622		
1,3-Butadiene		0.545	0.497	0.305	0.255	0.202	0.182	0.174	0.163	0.290	51.794		0.992		
Bromomethane		0.230	0.218	0.197	0.197	0.186	0.189	0.202				0.203	7.837		
Chloroethane		0.205	0.193	0.190	0.213	0.213	0.207	0.212				0.205	4.665		
Trichlorofluoromethane	0.464	0.480	0.513	0.488	0.516	0.527	0.512	0.509				0.501	4.218		
Diethyl ether		0.265	0.269	0.266	0.247	0.240	0.269		0.279	0.262	5.252				
Isoprene		0.365	0.366	0.363	0.403	0.410	0.416	0.412		0.423	0.395	6.470			
Acrolein			0.045	0.046	0.044	0.045	0.046		0.047	0.045	2.821				
1,1,2-Trichloro-1,2,2-Trifluoroet		0.224	0.236	0.229	0.256	0.263	0.254	0.260		0.246	6.393				
Acetone			0.100	0.099	0.093	0.085	0.088	0.083	0.083	0.091	7.863				
1,1-Dichloroethene	0.417	0.497	0.453	0.474	0.503	0.503	0.495	0.485		0.478	6.261				
Tert-Butyl Alcohol			0.023	0.024	0.026	0.025	0.027		0.029	0.025	8.004				
Dimethyl Sulfide	0.298	0.285	0.306	0.328	0.329	0.326	0.330	0.328	0.316	5.493					
Iodomethane		0.168	0.184	0.283	0.335	0.325	0.344		0.273	28.636	0.998				
Methyl acetate		0.162	0.189	0.200	0.204	0.203	0.209	0.213	0.197	8.753					
Methylene Chloride		0.254	0.255	0.260	0.274	0.274	0.272	0.277		0.267	3.702				
Carbon Disulfide		0.710	0.723	0.725	0.800	0.794	0.776	0.719	0.632	0.735	7.476				
Acrylonitrile		0.083	0.100	0.101	0.102	0.110	0.110		0.099	0.101	8.767				
Methyl Tert Butyl Ether	0.766	0.740	0.738	0.731	0.779	0.784	0.766	0.743		0.756	2.681				
trans-1,2-Dichloroethene	0.217	0.239	0.253	0.252	0.274	0.279	0.277	0.283		0.259	8.917				
n-Hexane			0.378	0.369	0.400	0.413	0.414	0.420	0.400	0.399	4.760				
Diisopropyl ether	1.021	1.041	1.034	0.961	0.910	0.910	0.974		0.878	0.974	6.485				
Vinyl Acetate			0.219	0.326	0.348	0.339	0.394	0.343	0.328	17.719	0.995				
1,1-Dichloroethane	0.562	0.557	0.540	0.542	0.588	0.589	0.583	0.564		0.566	3.441				
Ethyl-Tert-Butyl ether		1.021	1.051	1.034	0.978	0.938	0.984		0.887	0.985	5.842				
2-Butanone			0.106	0.108	0.125	0.122	0.117	0.117	0.120	0.116	5.983				
Propionitrile	0.029	0.036	0.035	0.037	0.037	0.037	0.035		0.028	0.034	11.563				
2,2-Dichloropropane		0.342	0.389	0.358	0.407	0.412	0.418	0.412		0.391	7.647				
cis-1,2-Dichloroethene	0.234	0.284	0.288	0.284	0.316	0.315	0.316	0.317		0.294	9.793				
Chloroform	0.581	0.482	0.492	0.494	0.523	0.521	0.515	0.498		0.513	6.027				
1-Bromopropane			0.055	0.065	0.065	0.065	0.066	0.066	0.067	0.064	6.952				
Bromochloromethane	0.123	0.161	0.172	0.185	0.189	0.185	0.191			0.173	14.001				
Tetrahydrofuran	0.092	0.074	0.072	0.072	0.073	0.073			0.073	0.075	9.560				
Dibromofluoromethane			0.289	0.269	0.301	0.298	0.286			0.289	4.386				
1,1,1-Trichloroethane	0.425	0.412	0.444	0.439	0.482	0.498	0.509	0.510		0.465	8.485				
Cyclohexane		0.541	0.537	0.515	0.557	0.575	0.572	0.571	0.533	0.550	3.968				
1,1-Dichloropropene		0.307	0.346	0.330	0.356	0.370	0.370	0.367		0.349	6.367				
Carbon Tetrachloride		0.304	0.355	0.355	0.404	0.439	0.460	0.449		0.395	14.848				
Tert-Amyl-Methyl ether		0.73	0.776	0.761	0.725	0.708	0.751		0.69	0.7344	4.1499				
1,2-Dichloroethane-d4			0.363	0.323	0.36	0.351	0.335			0.3463	4.8933				
1,2-Dichloroethane	0.406	0.433	0.427	0.451	0.454	0.466	0.471	0.461	0.445	0.4461	4.6706				
Benzene		1.09	1.001	1.093	1.015	1.085	1.088	1.04	0.899	1.0386	6.4759				
Trichloroethene	0.287	0.258	0.279	0.302	0.284	0.304	0.312	0.314	0.311	0.2946	6.4663				
Methylcyclohexane		0.385	0.389	0.388	0.426	0.433	0.435	0.437	0.427	0.415	5.6404				
1,2-Dichloropropane	0.27	0.297	0.32	0.291	0.323	0.321	0.322	0.324		0.3085	6.541				
1,4-Dioxane			0.002	0.002	0.002	0.002	0.002		0.002	0.0021	5.7317				
Bromodichloromethane	0.319	0.315	0.323	0.349	0.392	0.408	0.407	0.404		0.3646	11.576				
Dibromomethane		0.15	0.165	0.177	0.183	0.188	0.184	0.187		0.1762	7.9773				
2-Chloroethyl Vinyl Ether			0.114	0.142	0.153	0.154	0.163	0.169	0.169	0.1493	13.144				
4-Methyl-2-Pentanone			0.095	0.11	0.111	0.104	0.106	0.111	0.1064	5.7105					
cis-1,3-Dichloropropene	0.394	0.355	0.372	0.39	0.437	0.451	0.448	0.44		0.4109	9.1114				
Dimethyl Disulfide			0.17	0.244	0.267	0.28	0.292			0.2509	19.253	0.999			
Chlorobenzene-d5	ISTD														
Toluene-d8				1.314	1.167	1.296	1.268	1.176		1.2442	5.4772				
Toluene	1.556	1.487	1.45	1.547	1.589	1.573	1.474	1.172		1.481	9.0786				
Ethyl Methacrylate		0.335	0.364	0.389	0.451	0.456	0.451	0.444	0.456	0.4183	11.541				
trans-1,3-Dichloropropene		0.409	0.432	0.451	0.507	0.536	0.538	0.522		0.4849	10.97				
1,1,2-Trichloroethane	0.364	0.291	0.293	0.299	0.324	0.325	0.314	0.313		0.3153	7.472				
2-Hexanone			0.189	0.23	0.251	0.252	0.238	0.237	0.243	0.2343	9.1319				
1,3-Dichloropropane	0.447	0.504	0.535	0.505	0.501	0.536	0.534	0.519	0.497	0.5087	5.4558				
Tetrachloroethene		0.304	0.287	0.295	0.321	0.327	0.328	0.337		0.3141	6.0162				
Dibromochloromethane		0.299	0.304	0.365	0.395	0.393	0.397			0.3589	12.849				
1,2-Dibromoethane		0.287	0.281	0.286	0.312	0.323	0.314	0.312		0.3022	5.638				
1-Chlorohexane		0.442	0.445	0.455	0.502	0.52	0.518	0.52	0.51	0.4889	7.1805				
Chlorobenzene	1.088	1.097	0.984	0.957	0.986	1.061	1.092	1.065	0.95	1.0311	5.8969				
1,1,1,2-Tetrachloroethane			0.319	0.333	0.398	0.439	0.459	0.484		0.4053	16.74	0.992			
Ethylbenzene	0.456	0.461	0.497	0.484	0.507	0.54	0.576	0.592	0.607	0.5245	10.808				
m-,p-Xylene	0.581	0.584	0.607	0.601	0.601	0.661	0.685	0.652	0.727	0.6332	7.9869				
o-Xylene	0.524	0.6	0.565	0.617	0.595	0.649	0.673	0.659	0.648	0.6144	7.9247				
Styrene		0.947	0.918	0.982	1.001	1.096	1.144	1.088	0.954	1.0163	8.0974				
Bromoform			0.148	0.172	0.166	0.225	0.26	0.257	0.267	0.2137	23.656	0.998			
Isopropylbenzene	1.521	1.621	1.535	1.52	1.56	1.676	1.696	1.514	1.2	1.5381	9.3868				
1,4-Dichlorobenzene-d4	ISTD														
1,1,2,2-Tetrachloroethane	0.479	0.626	0.633	0.665	0.713	0.756	0.667	0.679		0.6522	12.485				
p-Bromofluorobenzene				0.987	0.875	1.037	0.965	0.919		0.9567	6.5367				
1,2,3-Trichloropropane		0.17	0.227	0.215	0.233	0.241	0.221	0.217		0.2178	10.565				
trans-1,4-Dichloro-2-Butene		0.232	0.208	0.264	0.286	0.306	0.296	0.307	0.298	0.2746	13.421				

n-Propylbenzene	3.825	3.77	3.713	3.979	4.105	4.125	3.44	2.537	3.6868	13.987	
Bromobenzene	0.869	0.845	0.84	0.914	0.917	0.985	0.926	0.907	0.9004	5.3071	
1,3,5-Trimethylbenzene	2.444	2.76	2.567	2.602	2.729	2.816	2.939	2.595	2.102	2.6173	9.3004
2-Chlorotoluene	2.791	2.593	2.652	2.771	2.747	2.852	2.531	2.051	2.6234	9.7295	
4-Chlorotoluene	2.209	2.257	2.181	2.217	2.259	2.376	2.079	1.671	2.1561	9.8709	
a-Methylstyrene		1.286	1.405	1.435	1.633	1.665	1.606	1.48	1.412	1.4903	8.8698
tert-Butylbenzene		0.621	0.602	0.64	0.651	0.678	0.649	0.671	0.6446	4.1475	
1,2,4-Trimethylbenzene	2.622	2.831	2.707	2.685	2.733	2.931	3.026	2.648	2.129	2.7013	9.3786
sec-Butylbenzene	3.222	3.208	3.421	3.342	3.58	3.665	3.13	2.427	3.2493	11.711	
p-Isopropyltoluene	2.717	2.754	2.893	2.8	2.944	3.105	3.213	2.78	2.222	2.8254	9.9368
1,3-Dichlorobenzene	1.652	1.678	1.536	1.654	1.654	1.715	1.834	1.687	1.538	1.6609	5.4265
1,4-Dichlorobenzene	1.616	1.673	1.609	1.601	1.643	1.724	1.845	1.687	1.529	1.6586	5.4208
n-Butylbenzene	2.507	2.421	2.436	2.552	2.777	2.967	2.604	2.106	2.5463	10.032	
1,2-Dichlorobenzene	1.577	1.517	1.558	1.554	1.603	1.656	1.769	1.608	1.485	1.5919	5.2505
1,2-Dibromo-3-Chloropropane				0.11	0.128	0.151	0.138	0.146	0.1347	11.997	
1,2,4-Trichlorobenzene		0.87	0.93	0.954	1.053	1.153	1.113	1.136	1.0298	10.885	
Hexachlorobutadiene		0.39	0.472	0.512	0.533	0.57	0.554	0.601	0.519	13.554	
Naphthalene	2.268	2.064	2.103	2.131	2.326	2.549	2.282	1.916	2.205	8.7724	
1,2,3-Trichlorobenzene	0.882	0.914	0.878	0.875	0.969	1.089	1.055	1.078	0.9675	9.6848	

Wed Mar 06 14:13:53 2019

Calibration Table Report
 Method: A9FOOWT.M
 Title: A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Calibration: Tue Nov 13 11:28:33 2018
 Curve: WG684281
 Calibration Files

Compound	5 20 50 100 200 300 400 500									Avg	%RSD
	8M427930.D	8M427931.D	8M427932.D	8M427933.D	8M427934.D	8M427935.D	8M427936.D	8M427937.D			
Fluorobenzene	ISTD										
Acetonitrile		0.021	0.023	0.021	0.025	0.025	0.023	0.025	0.023	7.012	
3-Chloro-1-propene	0.440	0.458	0.460	0.474	0.478	0.476	0.447	0.426	0.457	4.070	
2-Chloro-1,3-butadiene	0.415	0.459	0.466	0.479	0.489	0.487	0.460	0.438	0.462	5.466	
Ethyl Acetate		0.183	0.202	0.208	0.222	0.227	0.212	0.216	0.210	6.943	
Methacrylonitrile	0.076	0.086	0.094	0.097	0.102	0.105	0.100	0.103	0.095	10.375	
Isobutyl Alcohol			0.007	0.006	0.007	0.007	0.007	0.008	0.007	10.078	
1-Butanol									0.000	0.000	
Methyl methacrylate	0.150	0.178	0.196	0.209	0.225	0.232	0.219	0.222	0.204	13.626	
2-Nitropropane		0.052	0.062	0.067	0.076	0.079	0.076	0.079	0.070	14.773	
Chlorobenzene-d5	ISTD										
1,4-Dichlorobenzene-d4	ISTD										
Cyclohexanone			0.023	0.022	0.027	0.028	0.025	0.028	0.026	9.715	

Tue Nov 13 12:22:27 2018

Calibration Table Report
 Method: 8260WTR.M
 Title: Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Calibration: Mon Mar 04 17:44:12 2019
 Curve: WG698192
 Calibration Files

Compound	Concentration (ppm)										Avg	%RSD	r ²	
	0.3	0.4	1	2	5	20	50	100	200	300			Linear	Quadratic
Fluorobenzene	ISTD													
Dichlorodifluoromethane			0.379	0.399	0.396	0.501	0.488	0.438	0.443		0.435	10.795		
Chloromethane			0.693	0.702	0.643	0.663	0.611	0.535	0.531		0.625	11.194		
Vinyl Chloride	0.464	0.482	0.521	0.507	0.559	0.543	0.492				0.510	6.668		
1,3-Butadiene				0.442	0.400	0.325	0.272			0.263	0.329	22.926	0.995	
Bromomethane			0.311	0.357	0.342	0.324	0.282	0.259	0.272		0.307	12.069		
Chloroethane			0.235	0.269	0.275	0.292	0.290	0.270	0.277		0.273	7.015		
Trichlorofluoromethane	0.471	0.479	0.520	0.518	0.571	0.559	0.507	0.514			0.517	6.710		
Diethyl ether			0.241	0.288	0.281	0.293	0.283	0.267		0.279	0.276	6.331		
Isoprene				0.499	0.508	0.535	0.544	0.502	0.514		0.519	5.17	3.251	
Acrolein				0.030	0.029	0.032	0.033	0.032		0.034	0.032	5.791		
1,1,2-Trichloro-1,2,2-Trifluoroet		0.264	0.288	0.300	0.324	0.319	0.291	0.300			0.298	6.728		
Acetone				0.069	0.083	0.082	0.076	0.077	0.074		0.077	6.846		
1,1-Dichloroethene	0.511	0.504	0.542	0.545	0.571	0.559	0.512	0.521			0.533	4.600		
Tert-Butyl Alcohol			0.027	0.024	0.026	0.023	0.024			0.025	0.025	6.414		
Dimethyl Sulfide				0.371	0.366	0.400	0.404	0.375	0.384	0.388	0.384	3.701		
Iodomethane	0.054	0.067	0.110	0.284	0.414	0.403	0.404				0.248	67.275	0.993	
Methyl acetate				0.171	0.201	0.205	0.198	0.211	0.211		0.199	7.458		
Methylene Chloride			0.299	0.345	0.344	0.355	0.345	0.318	0.326		0.333	5.907		
Carbon Disulfide			0.973	1.026	1.011	1.069	1.064	0.970	0.954	0.909	0.997	5.575		
Acrylonitrile				0.087	0.089	0.103	0.102	0.101		0.099	0.097	7.358		
Methyl Tert Butyl Ether			0.725	0.817	0.793	0.858	0.819	0.763	0.777		0.793	5.465		
trans-1,2-Dichloroethene	0.479	0.481	0.515	0.518	0.546	0.538	0.496	0.503			0.509	4.824		
n-Hexane			0.475	0.500	0.495	0.526	0.525	0.482	0.506	0.490	0.500	3.736		
Diisopropyl ether		1.159	1.346	1.305	1.351	1.311	1.185			1.172	1.261	6.787		
Vinyl Acetate				0.436	0.496	0.480	0.470	0.439		0.487	0.468	5.352		
1,1-Dichloroethane	0.624	0.624	0.681	0.667	0.708	0.691	0.637	0.637			0.659	4.919		
Ethyl-Tert-Butyl ether		0.997	1.151	1.116	1.156	1.118	1.020				1.093	6.231		
2-Butanone				0.096	0.118	0.119	0.111	0.119		0.117	0.113	7.905		
Propionitrile			0.006	0.027	0.031	0.036	0.035	0.034		0.037	0.029	37.074	0.999	
2,2-Dichloropropane			0.456	0.498	0.490	0.511	0.496	0.446	0.448		0.478	5.593		
cis-1,2-Dichloroethene		0.314	0.340	0.369	0.375	0.399	0.388	0.359	0.365		0.364	7.419		
Chloroform	0.654	0.582	0.564	0.585	0.577	0.616	0.592	0.545	0.546		0.585	5.881		
1-Bromopropane				0.069	0.075	0.082	0.083	0.077	0.080	0.080	0.078	6.256		
Bromochloromethane			0.172	0.204	0.210	0.227	0.221	0.206	0.212		0.207	8.443		
Tetrahydrofuran			0.073	0.081	0.074	0.082	0.077	0.073		0.075	0.076	5.141		
Dibromofluoromethane				0.244	0.240	0.261	0.259	0.242			0.249	4.013		
1,1,1-Trichloroethane		0.452	0.510	0.522	0.546	0.539	0.490	0.495			0.508	6.302		
Cyclohexane			0.627	0.674	0.691	0.710	0.717	0.654	0.673	0.664	0.676	4.374		
1,1-Dichloropropene	0.354	0.404	0.440	0.446	0.465	0.456	0.421	0.423			0.426	8.233		
Tert-Amyl-Methyl ether			0.745	0.878	0.853	0.885	0.853	0.783			0.833	6.736		
Carbon Tetrachloride		0.42	0.455	0.464	0.497	0.49	0.45	0.453			0.46134	5.67645		
1,2-Dichloroethane-d4				0.237	0.243	0.258	0.253	0.239			0.24601	3.71655		
Heptane											0	0		
1,2-Dichloroethane	0.339	0.385	0.409	0.409	0.442	0.426	0.393	0.395			0.3998	7.73245		
Benzene		1.295	1.259	1.398	1.393	1.462	1.406	1.268	1.202		1.33547	6.82851		
Trichloroethene	0.375	0.316	0.34	0.361	0.377	0.392	0.388	0.357	0.368		0.36384	6.57282		
Methylcyclohexane			0.525	0.567	0.587	0.598	0.591	0.536	0.558	0.546	0.56348	4.76329		
1,2-Dichloropropane		0.318	0.325	0.381	0.367	0.393	0.391	0.358	0.367		0.36249	7.75569		
Bromodichloromethane	0.377	0.364	0.43	0.428	0.462	0.452	0.418	0.425			0.41972	8.03818		
1,4-Dioxane				0.002	0.002	0.002	0.002	0.002		0.002	0.00204	13.6522		
Dibromomethane		0.142	0.165	0.169	0.183	0.174	0.164	0.17			0.16685	7.43399		
2-Chloroethyl Vinyl Ether				0.128	0.154	0.164	0.159	0.173		0.173	0.15839	10.5996		
4-Methyl-2-Pentanone				0.079	0.1	0.103	0.101	0.109		0.106	0.09965	10.9254		
cis-1,3-Dichloropropene	0.432	0.454	0.485	0.498	0.557	0.547	0.507	0.514			0.49934	8.55427		
Dimethyl Disulfide				0.272	0.313	0.324	0.303	0.315		0.312	0.30664	5.9179		
Chlorobenzene-d5	ISTD													
Toluene-d8				1.352	1.252	1.319	1.282	1.162			1.27345	5.72054		
Toluene	1.823	1.799	1.984	2.022	2.058	1.94	1.733	1.575			1.8666	8.80527		
Ethyl Methacrylate		0.264	0.384	0.413	0.492	0.493	0.468	0.489		0.492	0.43707	18.547	1.000	
Paraldehyde											0	0		
trans-1,3-Dichloropropene		0.458	0.536	0.577	0.626	0.603	0.563	0.564			0.56117	9.63234		
1,1,2-Trichloroethane	0.276	0.299	0.339	0.343	0.363	0.346	0.319	0.325			0.32628	8.57325		
2-Hexanone				0.072	0.122	0.129	0.128	0.142		0.141	0.12236	20.9232	0.999	
1,3-Dichloropropane	0.475	0.517	0.587	0.591	0.64	0.588	0.549	0.55			0.56193	9.06418		
Tetrachloroethene	0.364	0.371	0.408	0.418	0.419	0.408	0.377	0.382			0.39336	5.65135		
Dibromochloromethane		0.35	0.403	0.415	0.45	0.442	0.416	0.423			0.4141	7.87795		
1,2-Dibromoethane	0.249	0.28	0.321	0.336	0.358	0.341	0.321	0.329			0.31683	11.1504		
1-Chlorohexane			0.609	0.625	0.676	0.68	0.679	0.625	0.638	0.642	0.64683	4.313		
Chlorobenzene	1.418	1.188	1.208	1.309	1.322	1.352	1.287	1.176	1.128		1.26541	7.53272		
1,1,1,2-Tetrachloroethane			0.399	0.48	0.464	0.502	0.489	0.461	0.461		0.46523	7.07824		
Ethylbenzene	0.627	0.643	0.639	0.718	0.715	0.744	0.728	0.677	0.67		0.68465	6.31414		
m-,p-Xylene	0.862	0.806	0.799	0.857	0.886	0.91	0.872	0.792	0.725		0.83421	6.93508		
o-Xylene	0.773	0.727	0.726	0.835	0.842	0.894	0.868	0.815	0.806		0.80945	7.25366		
Styrene		1.161	1.309	1.361	1.481	1.443	1.34	1.286			1.34009	7.85616		
Bromoform			0.202	0.227	0.244	0.283	0.281	0.271	0.29		0.25676	12.9003		
Isopropylbenzene		1.89	1.969	2.151	2.205	2.264	2.176	1.945	1.756		2.04451	8.7673		
1,4-Dichlorobenzene-d4	ISTD													
1,1,2,2-Tetrachloroethane	0.641	0.626	0.772	0.74	0.784	0.719	0.684	0.677			0.7053	8.24185		
p-Bromofluorobenzene				0.924	0.875	0.939	0.932	0.841			0.90228	4.71386		

1,2,3-Trichloropropane		0.168	0.221	0.221	0.233	0.221	0.205	0.208		0.21099	10.1005	
trans-1,4-Dichloro-2-Butene				0.205	0.233	0.249	0.249	0.239	0.247	0.246	0.23843	6.59335
n-Propylbenzene	4.572	4.432	4.971	4.973	5.031	4.724	4.094	3.397			4.5243	12.2898
Bromobenzene	0.878	0.953	1.126	1.128	1.128	1.079	1.005	0.984			1.03519	9.11264
1,3,5-Trimethylbenzene	2.984	3.19	3.573	3.577	3.552	3.435	3.067	2.714			3.26143	9.90939
2-Chlorotoluene	2.658	3.069	2.941	3.458	3.365	3.144	2.784	2.513			2.99144	11.102
4-Chlorotoluene	2.485	2.528	3.162	2.759	2.813	2.651	2.397	2.116			2.61382	11.9401
a-Methylstyrene			1.804	1.927	1.941	2.011	1.872	1.782	1.763		1.87148	4.95057
tert-Butylbenzene		0.679	0.778	0.809	0.802	0.77	0.72	0.71			0.75242	6.61504
1,2,4-Trimethylbenzene	3.122	3.099	3.555	3.641	3.648	3.501	3.118	2.732			3.30188	10.0521
sec-Butylbenzene	4.009	4.079	4.513	4.615	4.629	4.39	3.879	3.277			4.17398	11.0555
p-Isopropyltoluene	3.396	3.4	3.854	3.91	3.981	3.804	3.373	2.906			3.57808	10.338
1,3-Dichlorobenzene	1.944	1.916	2.132	2.189	2.161	2.075	1.924	1.803			2.01804	6.93274
1,4-Dichlorobenzene	2.16	1.959	2.124	2.155	2.173	2.08	1.908	1.791			2.04364	6.91852
n-Butylbenzene	3.285	3.043	3.372	3.518	3.601	3.472	3.098	2.693			3.26014	9.23843
1,2-Dichlorobenzene	1.863	1.846	1.826	1.938	1.986	2.008	1.926	1.784	1.677		1.87269	5.58313
1,2-Dibromo-3-Chloropropane				0.114	0.122	0.14	0.132	0.126	0.129		0.12712	6.88681
1,2,4-Trichlorobenzene		1.35	1.238	1.374	1.374	1.466	1.427	1.334	1.306		1.35883	5.18072
Hexachlorobutadiene	0.601	0.569	0.667	0.662	0.687	0.677	0.634	0.624			0.64028	6.38782
Naphthalene	2.132	2.329	2.596	2.61	2.803	2.656	2.439	2.253			2.47706	9.19022
1,2,3-Trichlorobenzene	1.176	1.168	1.314	1.276	1.347	1.302	1.208	1.195			1.24827	5.5729

Tue Mar 05 10:59:07 2019

Login Number: L19030638 Run Date: 03/05/2019 Sample ID: WG698387-12
 Instrument ID: HPMS11 Run Time: 22:04 Method: 8260B
 File ID: 11M29748 Analyst: KFR QC Key: DOD5
 ICal Workgroup: WG698387 Cal ID: HPMS11 - 05-MAR-19

Analyte		Expected	Found	Units	RF	%D	UCL	Q
1,1-Dichloroethene	CCC	50.0	51.2	ug/L	0.490	2.40	20	
Chloroform	CCC	50.0	51.1	ug/L	0.524	2.10	20	
Vinyl Chloride	CCC	50.0	45.2	ug/L	0.478	9.70	20	
Bromoform	SPCC	50.0	46.7	ug/L	0.241	6.70	20	
Chloromethane	SPCC	50.0	43.4	ug/L	0.615	13.3	20	
1,1-Dichloroethane	SPCC	50.0	51.4	ug/L	0.581	2.80	20	
Chlorobenzene	SPCC	50.0	52.6	ug/L	1.08	5.20	20	
1,1,2,2-Tetrachloroethane	SPCC	50.0	49.9	ug/L	0.651	0.200	20	
1,2-Dichloroethane		50.0	52.1	ug/L	0.465	4.10	20	
Acetone		50.0	44.2	ug/L	0.0808	11.6	20	
cis-1,2-Dichloroethene		50.0	53.2	ug/L	0.313	6.30	20	
Tetrachloroethene		50.0	51.7	ug/L	0.325	3.40	20	
trans-1,2-Dichloroethene		50.0	52.6	ug/L	0.273	5.30	20	
Trichloroethene		50.0	54.3	ug/L	0.320	8.60	20	

* Exceeds %D Limit

CCC Calibration Check Compounds
 SPCC System Performance Check Compounds



Login Number: L19030638 Run Date: 03/04/2019 Sample ID: WG698192-12
 Instrument ID: HPMS8 Run Time: 18:06 Method: 8260B
 File ID: 8M428967 Analyst: EEA QC Key: DOD5
 ICal Workgroup: WG698192 Cal ID: HPMS8 - 04-MAR-19

Analyte		Expected	Found	Units	RF	%D	UCL	Q
1,1-Dichloroethene	CCC	50.0	51.6	ug/L	0.550	3.10	20	
Chloroform	CCC	50.0	51.7	ug/L	0.604	3.40	20	
Vinyl Chloride	CCC	50.0	44.8	ug/L	0.457	10.4	20	
1,1,2,2-Tetrachloroethane	SPCC	50.0	47.6	ug/L	0.671	4.80	20	
Bromoform	SPCC	50.0	55.7	ug/L	0.286	11.4	20	
Chloromethane	SPCC	50.0	49.0	ug/L	0.613	2.00	20	
1,1-Dichloroethane	SPCC	50.0	53.0	ug/L	0.698	5.90	20	
Chlorobenzene	SPCC	50.0	53.8	ug/L	1.36	7.50	20	
1,2-Dichloroethane		50.0	53.5	ug/L	0.428	7.00	20	
Acetone		50.0	46.4	ug/L	0.0714	7.30	20	
cis-1,2-Dichloroethene		50.0	54.6	ug/L	0.397	9.20	20	
Tetrachloroethene		50.0	53.7	ug/L	0.423	7.40	20	
trans-1,2-Dichloroethene		50.0	53.3	ug/L	0.543	6.60	20	
Trichloroethene		50.0	57.0	ug/L	0.415	13.9	20	

* Exceeds %D Limit

CCC Calibration Check Compounds
 SPCC System Performance Check Compounds



Login Number: L19030638 Run Date: 03/11/2019 Sample ID: WG699027-02
Instrument ID: HPMS8 Run Time: 09:19 Method: 8260B
File ID: 8M429061 Analyst: EEA QC Key: DOD5
Workgroup (AAB#): WG699028 Cal ID: HPMS8 - 04-MAR-19
Matrix: WATER

Analyte		Expected	Found	UNITS	RF	%D	UCL	Q
1,2-Dichloropropane	CCC	50.0	48.0	ug/L	0.348	4.04	20	
Ethylbenzene	CCC	50.0	47.7	ug/L	0.653	4.58	20	
Toluene	CCC	50.0	46.1	ug/L	1.72	7.73	20	
1,1-Dichloroethene	CCC	50.0	46.6	ug/L	0.497	6.82	20	
Chloroform	CCC	50.0	45.3	ug/L	0.530	9.31	20	
Vinyl Chloride	CCC	50.0	46.6	ug/L	0.475	6.85	20	
1,1,2,2-Tetrachloroethane	SPCC	50.0	44.7	ug/L	0.630	10.6	20	
1,1-Dichloroethane	SPCC	50.0	46.6	ug/L	0.613	6.88	20	
Bromoform	SPCC	50.0	47.3	ug/L	0.243	5.34	20	
Chlorobenzene	SPCC	50.0	45.1	ug/L	1.14	9.71	20	
Chloromethane	SPCC	50.0	42.1	ug/L	0.526	15.8	20	
1,2-Dichloroethane		50.0	46.8	ug/L	0.374	6.44	20	
Acetone		50.0	40.0	ug/L	0.0616	20.0	20	
cis-1,2-Dichloroethene		50.0	47.6	ug/L	0.346	4.90	20	
Tetrachloroethene		50.0	46.3	ug/L	0.364	7.36	20	
trans-1,2-Dichloroethene		50.0	47.2	ug/L	0.481	5.64	20	
Trichloroethene		50.0	45.7	ug/L	0.333	8.55	20	

* Exceeds %D Criteria

CCC Calibration Check Compounds
SPCC System Performance Check Compounds

CCV - Modified 03/05/2008
PDF File ID: 6344154
Report generated 03/16/2019 10:29



Login Number: L19030638 Run Date: 03/12/2019 Sample ID: WG699218-02
 Instrument ID: HPMS11 Run Time: 11:21 Method: 8260B
 File ID: 11M29829 Analyst: KFR QC Key: DOD5
 Workgroup (AAB#): WG699219 Cal ID: HPMS11 - 05-MAR-19
 Matrix: WATER

Analyte		Expected	Found	UNITS	RF	%D	UCL	Q
1,2-Dichloropropane	CCC	50.0	48.0	ug/L	0.296	4.09	20	
Ethylbenzene	CCC	50.0	52.0	ug/L	0.546	4.01	20	
Toluene	CCC	50.0	50.0	ug/L	1.48	0.00600	20	
1,1-Dichloroethene	CCC	50.0	49.9	ug/L	0.477	0.225	20	
Chloroform	CCC	50.0	47.9	ug/L	0.492	4.18	20	
Vinyl Chloride	CCC	50.0	41.3	ug/L	0.437	17.5	20	
1,1,2,2-Tetrachloroethane	SPCC	50.0	46.9	ug/L	0.612	6.19	20	
1,1-Dichloroethane	SPCC	50.0	48.1	ug/L	0.545	3.72	20	
Bromoform	SPCC	50.0	46.5	ug/L	0.240	6.98	20	
Chlorobenzene	SPCC	50.0	49.6	ug/L	1.02	0.842	20	
Chloromethane	SPCC	50.0	34.5	ug/L	0.490	30.9	20	*
1,2-Dichloroethane		50.0	46.4	ug/L	0.414	7.30	20	
Acetone		50.0	40.6	ug/L	0.0743	18.7	20	
cis-1,2-Dichloroethene		50.0	50.7	ug/L	0.298	1.40	20	
Tetrachloroethene		50.0	52.1	ug/L	0.327	4.20	20	
trans-1,2-Dichloroethene		50.0	51.8	ug/L	0.269	3.65	20	
Trichloroethene		50.0	49.5	ug/L	0.291	1.08	20	

* Exceeds %D Criteria

CCC Calibration Check Compounds
 SPCC System Performance Check Compounds

CCV - Modified 03/05/2008
 PDF File ID: 6344154
 Report generated 03/16/2019 10:29



Login Number: L19030638 Run Date: 03/12/2019 Sample ID: WG699219-03
 Instrument ID: HPMS11 Run Time: 22:39 Prep Method: 5030B/5030C/503
 File ID: 11M29851 Analyst: KFR Method: 8260B
 Workgroup (AAB#): WG699219 Matrix: Water Units: ug/L
 Contract #: _____ Cal ID: HPMS11-05-MAR-19

Analytes	Expected	Found	% Rec	Limits	Q
1,1-Dichloroethene	50.0	47.2	94.5	50 - 150	
1,2-Dichloroethane	50.0	49.5	99.0	50 - 150	
Acetone	50.0	44.5	89.0	50 - 150	
Chloroform	50.0	47.7	95.4	50 - 150	
cis-1,2-Dichloroethene	50.0	50.9	102	50 - 150	
Tetrachloroethene	50.0	48.5	96.9	50 - 150	
trans-1,2-Dichloroethene	50.0	49.7	99.5	50 - 150	
Trichloroethene	50.0	57.2	114	50 - 150	
Vinyl chloride	50.0	43.4	86.8	50 - 150	

Surrogates	% Recovery	Surrogate Limits	Qualifier
1,2-Dichloroethane-d4	100	81 - 118	PASS
4-Bromofluorobenzene	98.7	85 - 114	PASS
Dibromofluoromethane	105	80 - 119	PASS
Toluene-d8	101	89 - 112	PASS

* FAILS %REC LIMIT

Login Number: L19030638 Run Date: 03/11/2019 Sample ID: WG699028-04
 Instrument ID: HPMS8 Run Time: 20:23 Prep Method: 5030B/5030C/503
 File ID: 8M429083 Analyst: EEA Method: 8260B
 Workgroup (AAB#): WG699028 Matrix: Water Units: ug/L
 Contract #: _____ Cal ID: HPMS8-04-MAR-19

Analytes	Expected	Found	% Rec	Limits	Q
1,1-Dichloroethene	50.0	50.7	101	50 - 150	
1,2-Dichloroethane	50.0	52.0	104	50 - 150	
Acetone	50.0	45.3	90.5	50 - 150	
Chloroform	50.0	49.7	99.5	50 - 150	
cis-1,2-Dichloroethene	50.0	51.6	103	50 - 150	
Tetrachloroethene	50.0	48.0	96.0	50 - 150	
trans-1,2-Dichloroethene	50.0	51.5	103	50 - 150	
Trichloroethene	50.0	54.1	108	50 - 150	
Vinyl chloride	50.0	50.7	101	50 - 150	

Surrogates	% Recovery	Surrogate Limits	Qualifier
1,2-Dichloroethane-d4	104	81 - 118	PASS
4-Bromofluorobenzene	101	85 - 114	PASS
Dibromofluoromethane	103	80 - 119	PASS
Toluene-d8	101	89 - 112	PASS

* FAILS %REC LIMIT

Login Number: L19030638
Instrument ID: HPMS8
Workgroup (AAB#): WG699028

ICAL CCV Number: WG698192-08
CAL ID: HPMS8-04-MAR-19
Matrix: WATER

Sample Number	Dilution	Tag	IS-1	IS-2	IS-3
WG698192-08	NA	NA	217564	400675	533134
Upper Limit	NA	NA	435128	801350	1066268
Lower Limit	NA	NA	108782	200338	266567
<u>L19030638-04</u>	1.00	01	173005	343409	454746
L19030638-05	1.00	01	182078	350977	465976
L19030638-06	1.00	01	174153	341702	457216
L19030638-07	1.00	01	165281	327300	434603
WG699028-01	1.00	01	196164	381131	508811
WG699028-02	1.00	01	215609	399992	523789
WG699028-03	1.00	01	217474	402726	524940

IS-1 - 1,4-Dichlorobenzene-d4
IS-2 - Chlorobenzene-d5
IS-3 - Fluorobenzene

Underline = Response outside limits



Login Number: L19030638
Instrument ID: HPMS11
Workgroup (AAB#): WG699219

ICAL CCV Number: WG698387-08
CAL ID: HPMS11-05-MAR-19
Matrix: WATER

Sample Number	Dilution	Tag	IS-1	IS-2	IS-3
WG698387-08	NA	NA	148978	299517	395124
Upper Limit	NA	NA	297956	599034	790248
Lower Limit	NA	NA	74489	149759	197562
<u>L19030638-01</u>	1.00	01	<u>140428</u>	<u>281096</u>	<u>379422</u>
L19030638-02	1.00	01	148384	291460	384567
L19030638-03	1.00	01	153036	305010	401795
L19030638-08	1.00	01	140720	282943	375235
WG699219-01	1.00	01	149164	314358	428800
WG699219-02	1.00	01	156356	309185	402710

IS-1 - 1,4-Dichlorobenzene-d4
IS-2 - Chlorobenzene-d5
IS-3 - Fluorobenzene

Underline = Response outside limits



Microbac Laboratories Inc.
INTERNAL STANDARD RETENTION TIME SUMMARY
(COMPARED TO MIDPOINT OF ICAL)

00936528

Login Number: L19030638
Instrument ID: HPMS8
Workgroup (AAB#): WG699028

ICAL CCV Number: WG698192-08
CAL ID: HPMS8-04-MAR-19
Matrix: WATER

Sample Number	Dilution	Tag	IS-1	IS-2	IS-3
WG698192-08	NA	NA	17.62	14.6	10.74
Upper Limit	NA	NA	18.12	15.1	11.24
Lower Limit	NA	NA	17.12	14.1	10.24
<u>L19030638-04</u>	1.00	01	17.62	14.6	10.74
L19030638-05	1.00	01	17.61	14.59	10.74
L19030638-06	1.00	01	17.62	14.6	10.73
L19030638-07	1.00	01	17.61	14.59	10.74
WG699028-01	1.00	01	17.61	14.59	10.74
WG699028-02	1.00	01	17.61	14.6	10.73
WG699028-03	1.00	01	17.62	14.6	10.73

IS-1 - 1,4-Dichlorobenzene-d4
IS-2 - Chlorobenzene-d5
IS-3 - Fluorobenzene

Underline = Response outside limits



Microbac Laboratories Inc.
INTERNAL STANDARD RETENTION TIME SUMMARY
(COMPARED TO MIDPOINT OF ICAL)

00936529

Login Number: L19030638
Instrument ID: HPMS11
Workgroup (AAB#): WG699219

ICAL CCV Number: WG698387-08
CAL ID: HPMS11-05-MAR-19
Matrix: WATER

Sample Number	Dilution	Tag	IS-1	IS-2	IS-3
WG698387-08	NA	NA	16.83	14.02	10.39
Upper Limit	NA	NA	17.33	14.52	10.89
Lower Limit	NA	NA	16.33	13.52	9.89
<u>L19030638-01</u>	1.00	01	<u>16.833</u>	<u>14.0206</u>	<u>10.3914</u>
L19030638-02	1.00	01	16.8329	14.0205	10.3913
L19030638-03	1.00	01	16.8329	14.0205	10.381
L19030638-08	1.00	01	16.833	14.0207	10.3915
WG699219-01	1.00	01	16.8223	14.0203	10.3911
WG699219-02	1.00	01	16.8227	14.0207	10.3812

IS-1 - 1,4-Dichlorobenzene-d4
IS-2 - Chlorobenzene-d5
IS-3 - Fluorobenzene

Underline = Response outside limits



2.1.1.3 Sample Data

Data File : C:\MSDCHEM\1\DATA\031219\11M29837.D Vial: 9
 Acq On : 12 Mar 2019 15:30 Operator: KFR
 Sample : L19030638-01 A REF 826-SPE Inst : hpms11
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 16 10:12:53 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3914	96	379422	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0206	117	281096	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	140428	25.0000	ug/L	0.0000
System Monitoring Compounds						
37) Dibromofluoromethane	9.3988	111	105079	23.9918	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	95.9672%	
43) 1,2-Dichloroethane-d4	10.0088	65	127216	24.2084	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	96.8336%	
57) Toluene-d8	12.2525	98	349412	24.9775	ug/L	0.0000
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	99.9100%	
78) p-Bromofluorobenzene	15.4164	95	133942	24.9245	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	99.6980%	
Target Compounds						
3) Chloromethane	3.5466	50	1702	0.1581	ug/L #	75
9) Diethyl ether	5.8213	59	1149	0.2890	ug/L #	1
13) Acetone	6.1108	43	4940	3.5613	ug/L	96

(#) = qualifier out of range (m) = manual integration
 11M29837.D 8260WT.M Sat Mar 16 10:12:53 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\031219\11M29837.D

Vial: 9

Acq On : 12 Mar 2019 15:30

Operator: KFR

Sample : L19030638-01 A REF 826-SPE

Inst : hpms11

Misc : 1,1

Multiplr: 1.00

MS Integration Params: rteint.p

Quant Time: Mar 16 10:12 2019

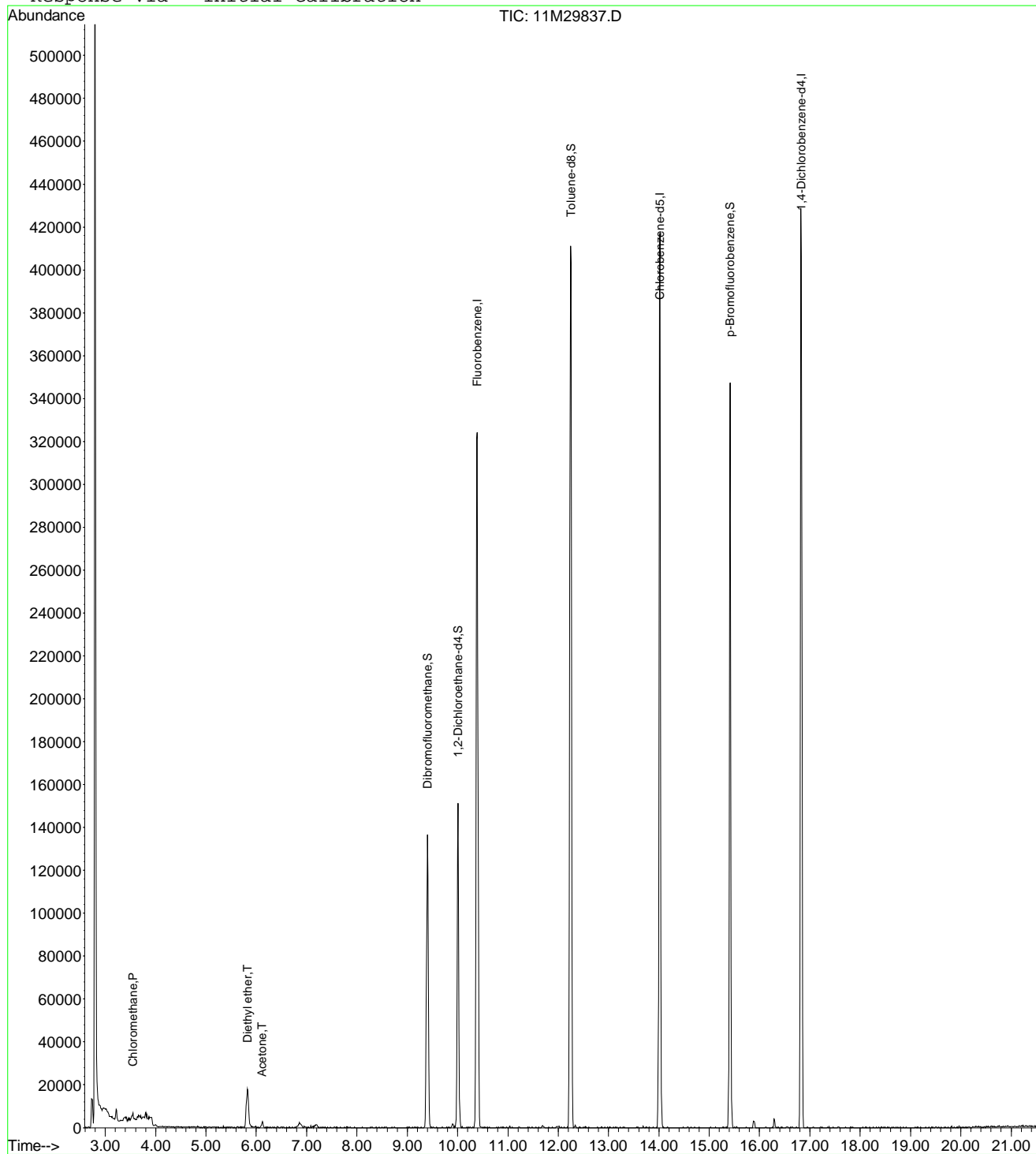
Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)

Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11

Last Update : Wed Mar 06 14:13:24 2019

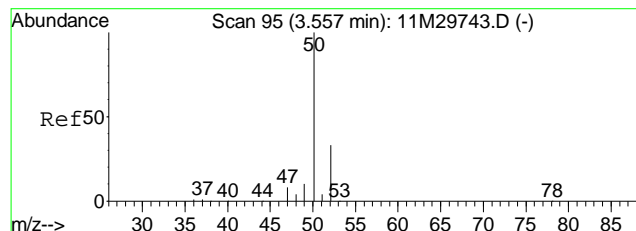
Response via : Initial Calibration



11M29837.D 8260WT.M

Sat Mar 16 10:12:54 2019

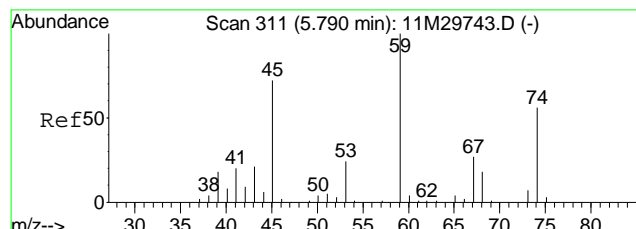
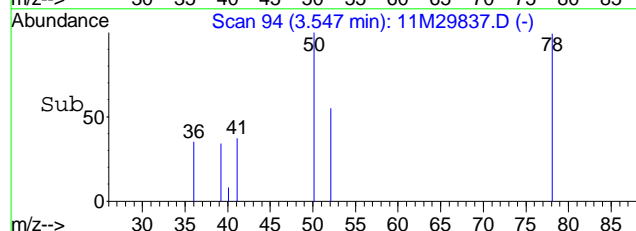
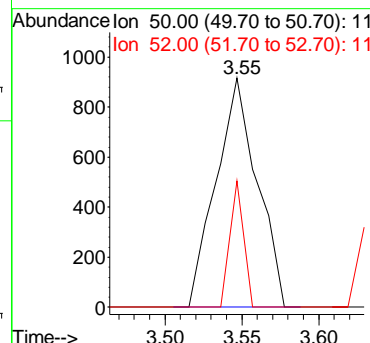
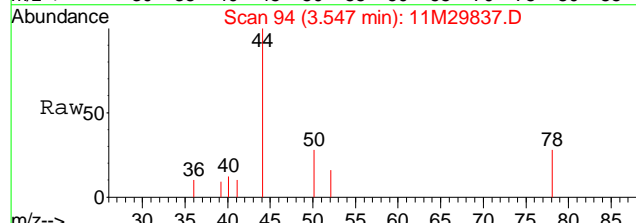
Page 2



#3
 Chloromethane
 Concen: 0.1581 ug/L
 RT: 3.55 min Scan# 94
 Delta R.T. -0.01 min
 Lab File: 11M29837.D
 Acq: 12 Mar 2019 15:30

Tgt Ion: 50 Resp: 1702

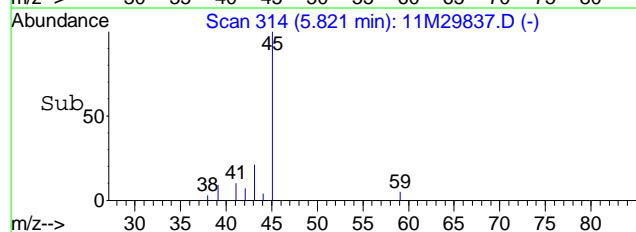
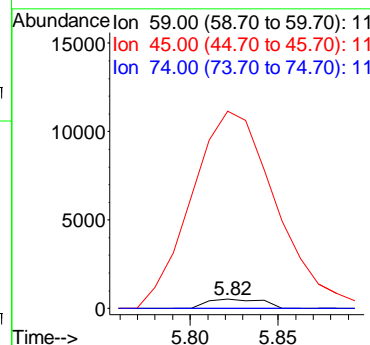
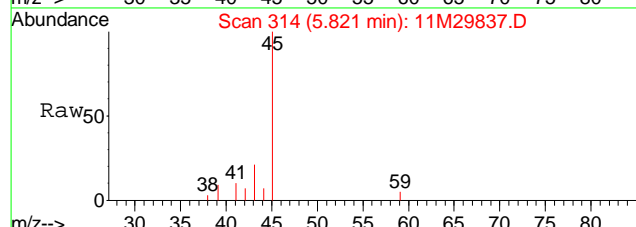
Ion	Ratio	Lower	Upper
50	100		
52	18.4	19.6	45.6#

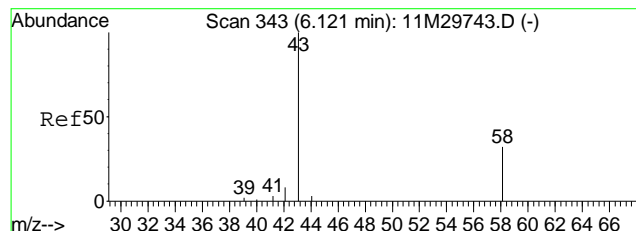


#9
 Diethyl ether
 Concen: 0.2890 ug/L
 RT: 5.82 min Scan# 314
 Delta R.T. 0.03 min
 Lab File: 11M29837.D
 Acq: 12 Mar 2019 15:30

Tgt Ion: 59 Resp: 1149

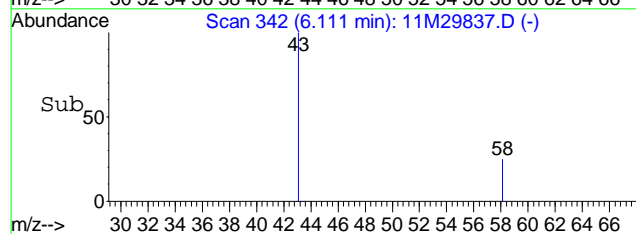
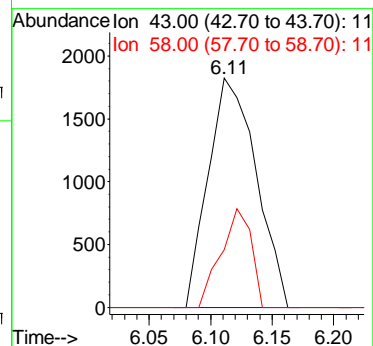
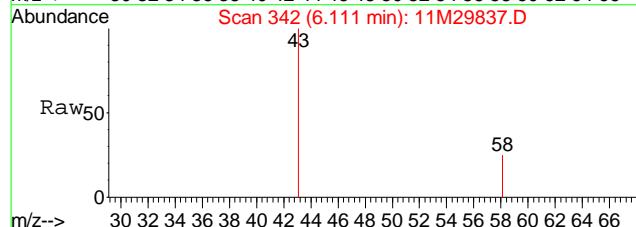
Ion	Ratio	Lower	Upper
59	100		
45	3269.5	43.1	100.5#
74	0.0	34.4	80.2#





#13
 Acetone
 Concen: 3.5613 ug/L
 RT: 6.11 min Scan# 342
 Delta R.T. -0.01 min
 Lab File: 11M29837.D
 Acq: 12 Mar 2019 15:30

Tgt Ion	Ratio	Lower	Upper
43	100		
58	27.2	17.8	41.4



Data File : C:\MSDCHEM\1\DATA\031219\11M29833.D Vial: 5
 Acq On : 12 Mar 2019 13:30 Operator: KFR
 Sample : L19030638-02 A MS 826-SPE Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 15:48:15 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3913	96	384567	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0205	117	291460	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8329	152	148384	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3987	111	112107	25.2540	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	= 101.0160%		
43) 1,2-Dichloroethane-d4	10.0088	65	128552	24.1354	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	= 96.5416%		
57) Toluene-d8	12.2421	98	363045	25.0292	ug/L	-0.0105
Spiked Amount	25.0000	Range 88 - 110	Recovery	= 100.1168%		
78) p-Bromofluorobenzene	15.4164	95	143624	25.2932	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	= 101.1728%		

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1226	85	80317	14.1375	ug/L	98
3) Chloromethane	3.5569	50	129369	11.8562	ug/L	99
4) Vinyl Chloride	3.7740	62	117559	14.4322	ug/L	99
5) 1,3-Butadiene	3.8154	54	43352	13.3665	ug/L	98
6) Bromomethane	4.6528	94	66389	21.2731	ug/L	97
7) Chloroethane	4.7976	64	57741	18.3359	ug/L	99
8) Trichlorofluoromethane	5.2732	101	129392	16.7861	ug/L	99
9) Diethyl ether	5.7902	59	386259	95.8462	ug/L	96
10) Isoprene	5.8212	67	106672	17.5698	ug/L	99
11) Acrolein	6.0177	56	63038	90.0764	ug/L	97
12) 1,1,2-Trichloro-1,2,2-Trif	6.0384	101	72940	19.2900	ug/L	100
13) Acetone	6.1314	43	25433	18.0896	ug/L	99
14) 1,1-Dichloroethene	6.3382	61	138053	18.7592	ug/L	100
15) Tert-Butyl Alcohol	6.4416	59	69257	176.6046	ug/L	99
16) Dimethyl Sulfide	6.5864	62	100022	20.5596	ug/L	98
17) Iodomethane	6.8345	142	91374	18.9965	ug/L	100
18) Methyl acetate	6.8552	43	53504	17.6427	ug/L #	88
19) Methylene Chloride	7.0930	84	84413	20.5833	ug/L	91
20) Carbon Disulfide	7.1344	76	188811	16.7042	ug/L	99
21) Acrylonitrile	7.2688	53	29793	19.2224	ug/L	95
22) Methyl Tert Butyl Ether	7.2998	73	219636	18.8868	ug/L	100
23) trans-1,2-Dichloroethene	7.5273	96	79628	19.9746	ug/L	99
24) n-Hexane	7.5996	57	99128	16.1442	ug/L	97
25) Diisopropyl ether	7.9305	45	1483348	98.9823	ug/L	99
26) Vinyl Acetate	8.0959	43	147917	28.4864	ug/L	99
27) 1,1-Dichloroethane	8.1270	63	170585	19.6099	ug/L	100
28) Ethyl-Tert-Butyl ether	8.4785	59	1548861	102.2538	ug/L	99
29) 2-Butanone	8.6543	43	30958	17.2866	ug/L	98
30) Propionitrile	8.7577	54	51668	98.7468	ug/L	98
31) 2,2-Dichloropropane	8.8611	77	133409	22.1807	ug/L	99
32) cis-1,2-Dichloroethene	8.9335	96	93129	20.5775	ug/L	99
33) Chloroform	9.1299	83	156881	19.8695	ug/L	98
34) 1-Bromopropane	9.2540	122	19355	19.7444	ug/L	95
35) Bromochloromethane	9.3470	130	55718	20.9965	ug/L	96
36) Tetrahydrofuran	9.3677	42	105670	91.0754	ug/L	97
38) 1,1,1-Trichloroethane	9.6365	97	150906	21.1081	ug/L	97
39) Cyclohexane	9.6572	56	125463	14.8228	ug/L	98
40) 1,1-Dichloropropene	9.8123	75	109170	20.3096	ug/L	99
41) Carbon Tetrachloride	9.9571	117	133653	21.9845	ug/L	99
42) Tert-Amyl-Methyl ether	9.9157	73	1179267	104.3820	ug/L	98

(#) = qualifier out of range (m) = manual integration
 11M29833.D 8260WT.M Tue Mar 12 15:48:16 2019

Data File : C:\MSDCHEM\1\DATA\031219\11M29833.D Vial: 5
 Acq On : 12 Mar 2019 13:30 Operator: KFR
 Sample : L19030638-02 A MS 826-SPE Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 15:48:15 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1225	62	137489	20.0351	ug/L	98
45) Benzene	10.1535	78	323838	20.2690	ug/L	99
46) Trichloroethene	10.8669	130	87832	19.3832	ug/L	99
47) Methylcyclohexane	10.9393	83	109903	17.2166	ug/L	98
48) 1,2-Dichloropropane	11.0634	63	91785	19.3395	ug/L	98
49) 1,4-Dioxane	11.3426	88	4204	129.0965	ug/L	95
50) Bromodichloromethane	11.3529	83	114675	20.4454	ug/L	99
51) Dibromomethane	11.4356	93	55041	20.3028	ug/L	97
53) 4-Methyl-2-Pentanone	11.6527	58	30325	18.5331	ug/L	97
54) cis-1,3-Dichloropropene	11.9423	75	137501	21.7519	ug/L	99
55) Dimethyl Disulfide	12.1904	79	75627	19.3160	ug/L	95
58) Toluene	12.3455	91	356314	20.6372	ug/L	99
59) Ethyl Methacrylate	12.4282	69	100139	20.5345	ug/L	95
60) trans-1,3-Dichloropropene	12.5006	75	117448	20.7762	ug/L	100
61) 1,1,2-Trichloroethane	12.7074	97	73443	19.9814	ug/L	100
62) 2-Hexanone	12.6454	43	49499	18.1183	ug/L #	72
63) 1,3-Dichloropropane	12.9969	76	123021	20.7445	ug/L	96
64) Tetrachloroethene	13.1106	164	71408	19.4983	ug/L	98
65) Dibromochloromethane	13.3588	129	89145	21.3055	ug/L	99
66) 1,2-Dibromoethane	13.5966	107	71097	20.1830	ug/L	99
67) 1-Chlorohexane	13.6690	91	106626	18.7056	ug/L	96
68) Chlorobenzene	14.0619	112	242793	20.1983	ug/L	99
69) 1,1,1,2-Tetrachloroethane	14.0929	131	96570	19.3196	ug/L	99
70) Ethylbenzene	14.0825	106	121025	19.7914	ug/L	94
71) m-,p-Xylene	14.1653	106	307571	41.6625	ug/L	95
72) o-Xylene	14.6926	106	148617	20.7483	ug/L	97
73) Styrene	14.7340	104	254520	21.4821	ug/L	99
74) Bromoform	15.1889	173	57475	19.5067	ug/L	98
75) Isopropylbenzene	15.0855	105	367862	20.5151	ug/L	99
77) 1,1,2,2-Tetrachloroethane	15.2923	83	85758	22.1546	ug/L	98
79) 1,2,3-Trichloropropane	15.4681	110	26632	20.6052	ug/L	77
80) trans-1,4-Dichloro-2-Buten	15.5094	53	30635	18.7987	ug/L	88
81) n-Propylbenzene	15.5611	91	472472	21.5916	ug/L	99
82) Bromobenzene	15.6852	156	109154	20.4255	ug/L	95
83) 1,3,5-Trimethylbenzene	15.7369	105	326778	21.0358	ug/L	99
84) 2-Chlorotoluene	15.8196	91	318728	20.4696	ug/L	100
85) 4-Chlorotoluene	15.8610	91	255627	19.9748	ug/L	99
86) a-Methylstyrene	16.1091	118	185178	20.9348	ug/L	97
87) tert-Butylbenzene	16.1608	134	75502	19.7334	ug/L	98
88) 1,2,4-Trimethylbenzene	16.2125	105	334743	20.8785	ug/L	100
89) sec-Butylbenzene	16.4193	105	406703	21.0883	ug/L	98
90) p-Isopropyltoluene	16.5641	119	356283	21.2457	ug/L	98
91) 1,3-Dichlorobenzene	16.7502	146	203188	20.6109	ug/L	98
92) 1,4-Dichlorobenzene	16.8639	146	205943	20.9199	ug/L	99
93) n-Butylbenzene	17.0500	91	318673	21.0859	ug/L	99
94) 1,2-Dichlorobenzene	17.3292	146	199286	21.0924	ug/L	97
95) 1,2-Dibromo-3-Chloropropan	18.2494	75	16540	20.6838	ug/L	97
96) 1,2,4-Trichlorobenzene	19.3144	180	131500	21.5152	ug/L	99
97) Hexachlorobutadiene	19.4488	225	64656	20.9885	ug/L	98
98) Naphthalene	19.6556	128	292138	22.3216	ug/L	99
99) 1,2,3-Trichlorobenzene	19.9451	180	124914	21.7534	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29833.D 8260WT.M Tue Mar 12 15:48:16 2019

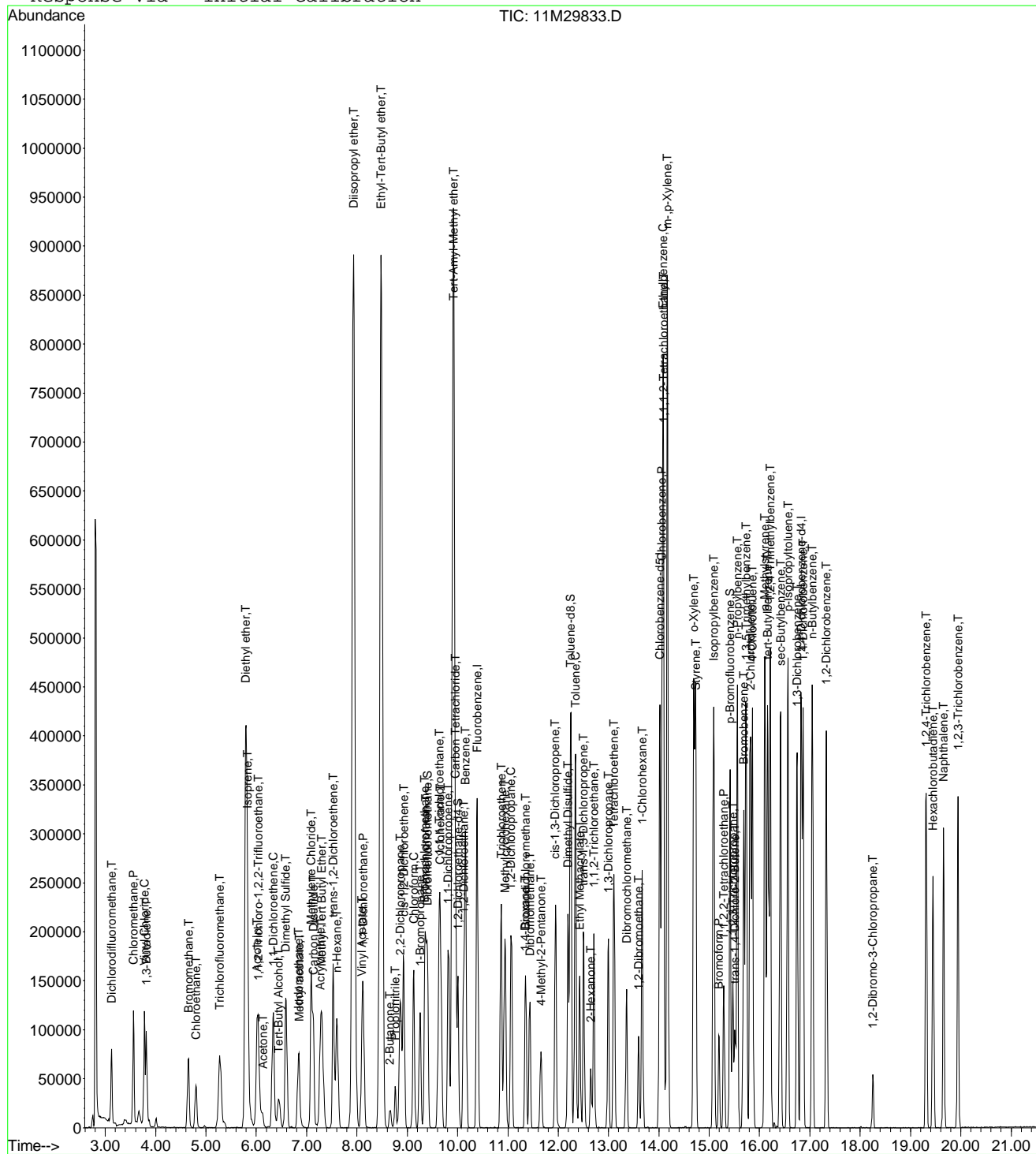
Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29833.D
Acq On : 12 Mar 2019 13:30
Sample : L19030638-02 A MS 826-SPE
Misc : 1,1 STD92316
MS Integration Params: rteint.p
Quant Time: Mar 12 15:48 2019

Vial: 5
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:13:24 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\031219\11M29834.D Vial: 6
 Acq On : 12 Mar 2019 14:00 Operator: KFR
 Sample : L19030638-03 A MSD 826-SPE Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 15:48:17 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3810	96	401795	25.0000	ug/L	-0.0105
56) Chlorobenzene-d5	14.0205	117	305010	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8329	152	153036	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3987	111	118196	25.4840	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	= 101.9360%		
43) 1,2-Dichloroethane-d4	10.0088	65	138169	24.8287	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	= 99.3148%		
57) Toluene-d8	12.2421	98	385857	25.4201	ug/L	-0.0105
Spiked Amount	25.0000	Range 88 - 110	Recovery	= 101.6804%		
78) p-Bromofluorobenzene	15.4164	95	155418	26.5382	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	= 106.1528%		

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1020	85	80150	13.5032	ug/L	97
3) Chloromethane	3.5465	50	137095	12.0255	ug/L	99
4) Vinyl Chloride	3.7637	62	120653	14.1769	ug/L	99
5) 1,3-Butadiene	3.8050	54	46237	13.7099	ug/L	97
6) Bromomethane	4.6322	94	66753	20.4726	ug/L	100
7) Chloroethane	4.7873	64	57755	17.5540	ug/L	99
8) Trichlorofluoromethane	5.2629	101	132120	16.4050	ug/L	99
9) Diethyl ether	5.7799	59	400099	95.0235	ug/L	95
10) Isoprene	5.8109	67	107918	17.0129	ug/L	98
11) Acrolein	6.0074	56	64880	88.7334	ug/L	98
12) 1,1,2-Trichloro-1,2,2-Trif	6.0281	101	75497	19.1101	ug/L	99
13) Acetone	6.1108	43	26966	18.3575	ug/L	97
14) 1,1-Dichloroethene	6.3279	61	140305	18.2477	ug/L	98
15) Tert-Butyl Alcohol	6.4416	59	73072	178.3434	ug/L	98
16) Dimethyl Sulfide	6.5761	62	101894	20.0464	ug/L	99
17) Iodomethane	6.8242	142	97304	19.3332	ug/L	98
18) Methyl acetate	6.8449	43	53606	16.9184	ug/L #	82
19) Methylene Chloride	7.0827	84	84341	19.6839	ug/L	94
20) Carbon Disulfide	7.1240	76	194241	16.4478	ug/L	98
21) Acrylonitrile	7.2688	53	30283	18.7008	ug/L	99
22) Methyl Tert Butyl Ether	7.2895	73	225429	18.5538	ug/L	100
23) trans-1,2-Dichloroethene	7.5273	96	82023	19.6932	ug/L	97
24) n-Hexane	7.5893	57	100831	15.7174	ug/L	99
25) Diisopropyl ether	7.9305	45	1538618	98.2681	ug/L	99
26) Vinyl Acetate	8.0960	43	151765	28.0099	ug/L	98
27) 1,1-Dichloroethane	8.1166	63	172792	19.0119	ug/L	100
28) Ethyl-Tert-Butyl ether	8.4785	59	1598878	101.0299	ug/L	98
29) 2-Butanone	8.6543	43	32953	17.6116	ug/L	99
30) Propionitrile	8.7577	54	54194	99.1334	ug/L	100
31) 2,2-Dichloropropane	8.8611	77	132488	21.0831	ug/L	99
32) cis-1,2-Dichloroethene	8.9231	96	96922	20.4973	ug/L	96
33) Chloroform	9.1196	83	159988	19.3942	ug/L	99
34) 1-Bromopropane	9.2540	122	19435	18.9759	ug/L	95
35) Bromochloromethane	9.3470	130	57636	20.7880	ug/L	95
36) Tetrahydrofuran	9.3677	42	110026	90.7637	ug/L	96
38) 1,1,1-Trichloroethane	9.6262	97	153558	20.5581	ug/L	98
39) Cyclohexane	9.6572	56	125596	14.2023	ug/L	96
40) 1,1-Dichloropropene	9.8123	75	110716	19.7141	ug/L	98
41) Carbon Tetrachloride	9.9467	117	136517	21.4927	ug/L	98
42) Tert-Amyl-Methyl ether	9.9157	73	1222188	103.5426	ug/L	98

(#) = qualifier out of range (m) = manual integration
 11M29834.D 8260WT.M Tue Mar 12 15:48:18 2019

Data File : C:\MSDCHEM\1\DATA\031219\11M29834.D Vial: 6
 Acq On : 12 Mar 2019 14:00 Operator: KFR
 Sample : L19030638-03 A MSD 826-SPE Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 15:48:17 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1122	62	143971	20.0801	ug/L	98
45) Benzene	10.1535	78	331414	19.8538	ug/L	99
46) Trichloroethene	10.8670	130	91787	19.3874	ug/L	98
47) Methylcyclohexane	10.9393	83	111003	16.6433	ug/L	97
48) 1,2-Dichloropropane	11.0634	63	94993	19.1572	ug/L	100
49) 1,4-Dioxane	11.3426	88	4376	128.6164	ug/L	96
50) Bromodichloromethane	11.3529	83	118778	20.2689	ug/L	100
51) Dibromomethane	11.4357	93	55827	19.7098	ug/L	96
53) 4-Methyl-2-Pentanone	11.6528	58	31090	18.1859	ug/L	97
54) cis-1,3-Dichloropropene	11.9423	75	138778	21.0126	ug/L	100
55) Dimethyl Disulfide	12.1904	79	75810	18.6301	ug/L	89
58) Toluene	12.3352	91	363444	20.1150	ug/L	98
59) Ethyl Methacrylate	12.4283	69	101283	19.8464	ug/L	96
60) trans-1,3-Dichloropropene	12.5006	75	119544	20.2076	ug/L	100
61) 1,1,2-Trichloroethane	12.7074	97	74073	19.2575	ug/L	98
62) 2-Hexanone	12.6454	43	52725	18.4417	ug/L #	75
63) 1,3-Dichloropropane	12.9969	76	124959	20.1352	ug/L	98
64) Tetrachloroethene	13.1107	164	74323	19.3927	ug/L	98
65) Dibromochloromethane	13.3588	129	90562	20.6827	ug/L	100
66) 1,2-Dibromoethane	13.5966	107	73699	19.9922	ug/L	98
67) 1-Chlorohexane	13.6690	91	107453	18.0133	ug/L	96
68) Chlorobenzene	14.0619	112	249114	19.8035	ug/L	99
69) 1,1,1,2-Tetrachloroethane	14.0929	131	99276	18.9904	ug/L	100
70) Ethylbenzene	14.0826	106	126550	19.7755	ug/L	96
71) m-,p-Xylene	14.1653	106	309475	40.0581	ug/L	93
72) o-Xylene	14.6926	106	156195	20.8375	ug/L	99
73) Styrene	14.7340	104	258441	20.8440	ug/L	99
74) Bromoform	15.1992	173	57703	18.7437	ug/L	99
75) Isopropylbenzene	15.0855	105	377185	20.1005	ug/L	99
77) 1,1,2,2-Tetrachloroethane	15.2923	83	87397	21.8917	ug/L	98
79) 1,2,3-Trichloropropane	15.4681	110	28306	21.2346	ug/L	95
80) trans-1,4-Dichloro-2-Buten	15.5094	53	31342	18.6479	ug/L	89
81) n-Propylbenzene	15.5611	91	479217	21.2341	ug/L	100
82) Bromobenzene	15.6852	156	112466	20.4056	ug/L	94
83) 1,3,5-Trimethylbenzene	15.7369	105	335778	20.9581	ug/L	99
84) 2-Chlorotoluene	15.8196	91	325026	20.2396	ug/L	100
85) 4-Chlorotoluene	15.8610	91	260729	19.7542	ug/L	99
86) a-Methylstyrene	16.1091	118	187499	20.5528	ug/L	98
87) tert-Butylbenzene	16.1608	134	75262	19.0727	ug/L	96
88) 1,2,4-Trimethylbenzene	16.2125	105	340472	20.5903	ug/L	99
89) sec-Butylbenzene	16.4193	105	412349	20.7311	ug/L	99
90) p-Isopropyltoluene	16.5641	119	362103	20.9364	ug/L	98
91) 1,3-Dichlorobenzene	16.7502	146	206075	20.2683	ug/L	99
92) 1,4-Dichlorobenzene	16.8639	146	208212	20.5074	ug/L	99
93) n-Butylbenzene	17.0500	91	322284	20.6766	ug/L	99
94) 1,2-Dichlorobenzene	17.3292	146	200079	20.5326	ug/L	99
95) 1,2-Dibromo-3-Chloropropan	18.2494	75	17130	20.7704	ug/L	98
96) 1,2,4-Trichlorobenzene	19.3144	180	133856	21.2349	ug/L	98
97) Hexachlorobutadiene	19.4488	225	66894	21.0549	ug/L	98
98) Naphthalene	19.6556	128	293786	21.7652	ug/L	99
99) 1,2,3-Trichlorobenzene	19.9451	180	126194	21.3082	ug/L	99

(#) = qualifier out of range (m) = manual integration
 11M29834.D 8260WT.M Tue Mar 12 15:48:18 2019

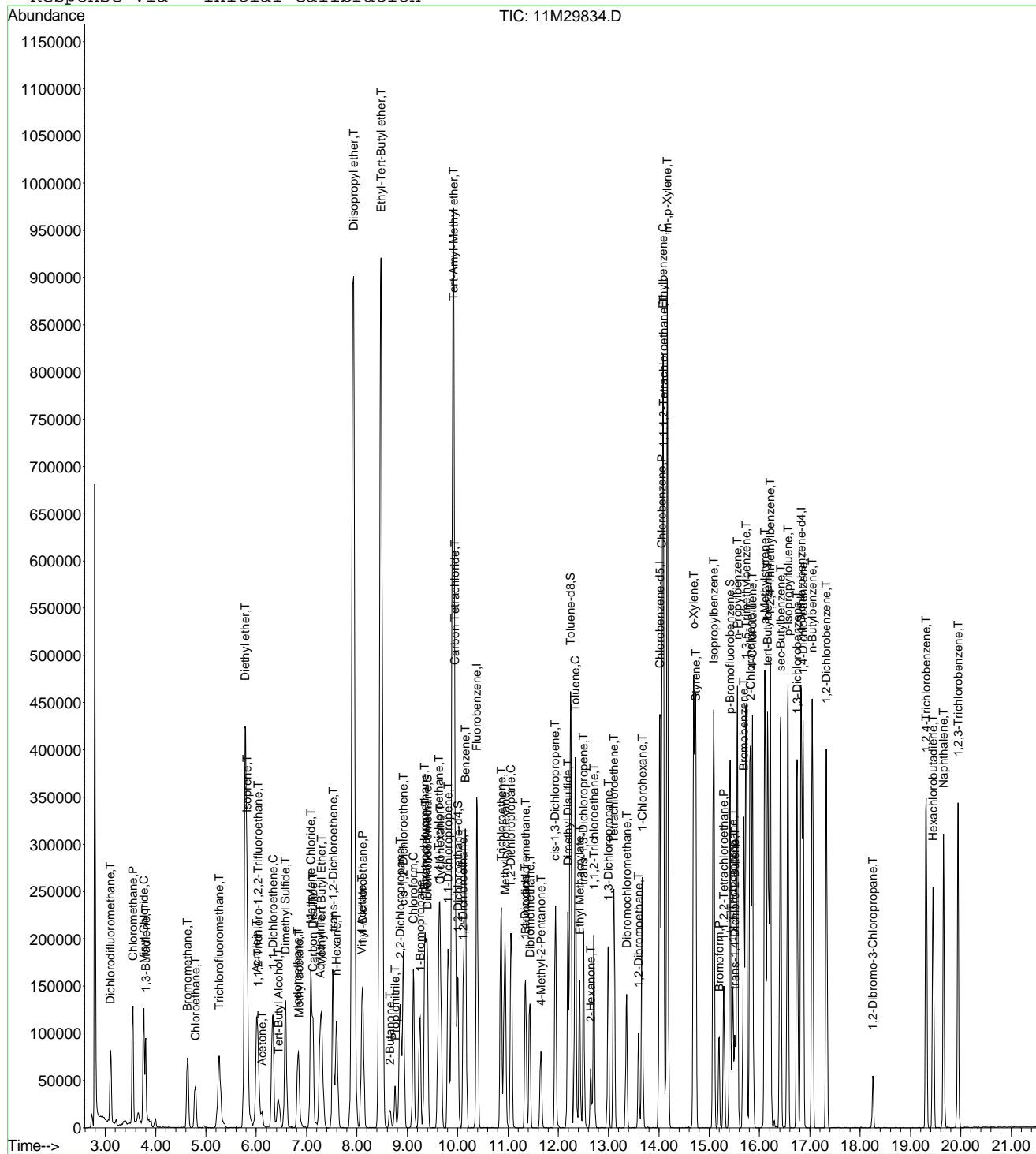
Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29834.D
 Acq On : 12 Mar 2019 14:00
 Sample : L19030638-03 A MSD 826-SPE
 Misc : 1,1 STD92316
 MS Integration Params: rteint.p
 Quant Time: Mar 12 15:48 2019

Vial: 6
 Operator: KFR
 Inst : hpms11
 Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429076.D Vial: 16
 Acq On : 11 Mar 2019 17:01 Operator: EEA
 Sample : L19030638-04 A 826-SPE Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 09:00:34 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

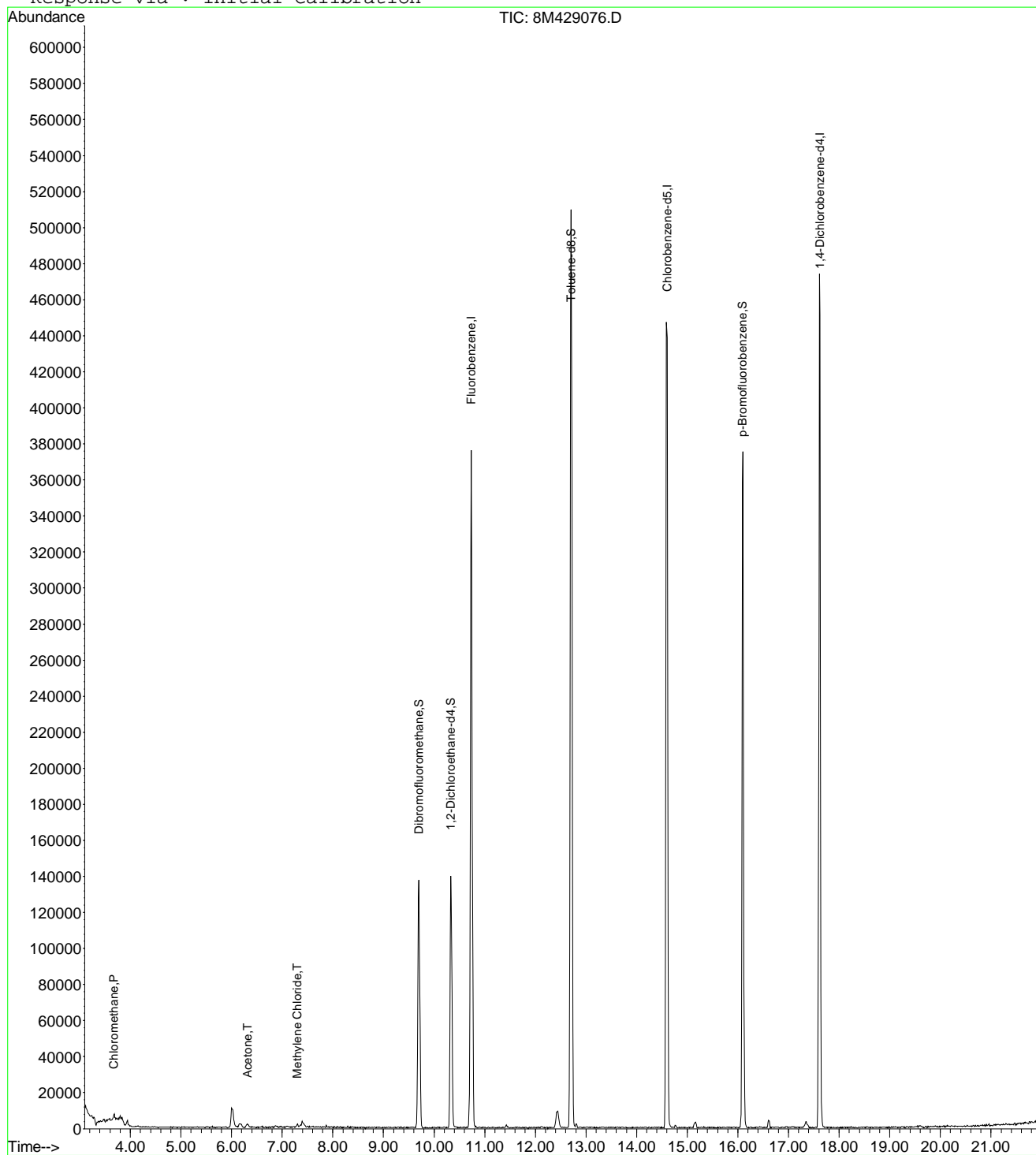
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	454746	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	343409	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	173005	25.00	ug/L	0.00
System Monitoring Compounds						
37) Dibromofluoromethane	9.70	111	120294	26.5379	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	106.16%	
43) 1,2-Dichloroethane-d4	10.33	65	121109	27.0638	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	108.24%	
58) Toluene-d8	12.71	98	459517	26.2692	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	105.08%	
80) p-Bromofluorobenzene	16.09	95	174326	27.9193	ug/L	-0.01
Spiked Amount	25.000	Range 86 - 115	Recovery	=	111.68%	
Target Compounds						
						Qvalue
3) Chloromethane	3.67	50	4517	0.3971	ug/L	94
13) Acetone	6.31	43	4429	3.1617	ug/L	83
19) Methylene Chloride	7.29	84	1034	0.1706	ug/L	73

(#) = qualifier out of range (m) = manual integration
 8M429076.D 8260WTR.M Tue Mar 12 09:00:35 2019

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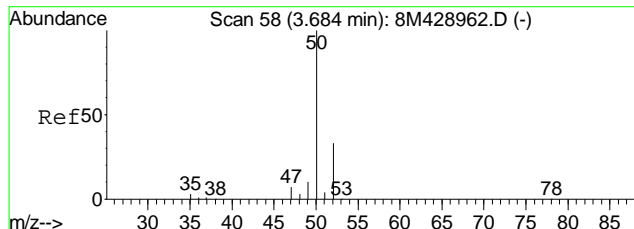
Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429076.D Vial: 16
Acq On : 11 Mar 2019 17:01 Operator: EEA
Sample : L19030638-04 A 826-SPE Inst : HPMS8
Misc : 1,1 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 12 9:00 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



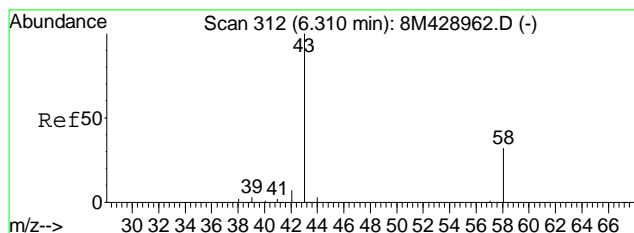
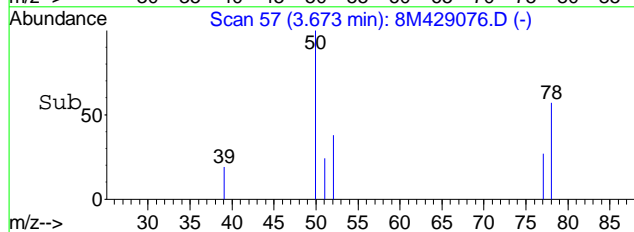
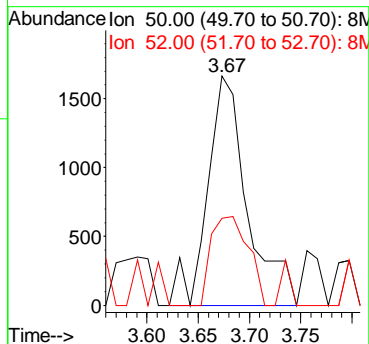
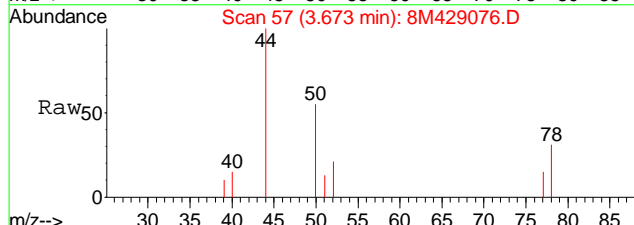
8M429076.D 8260WTR.M Tue Mar 12 09:00:35 2019

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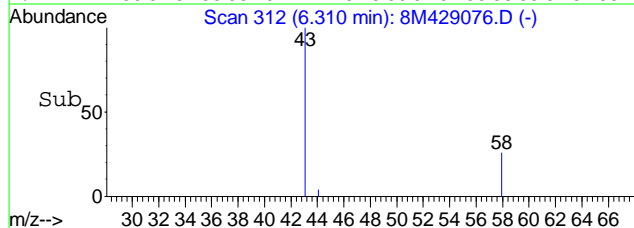
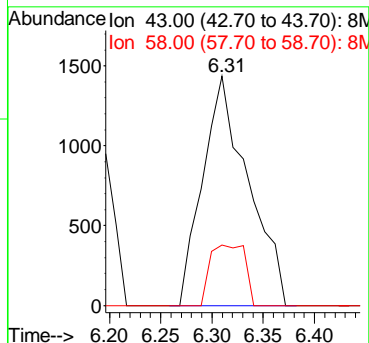
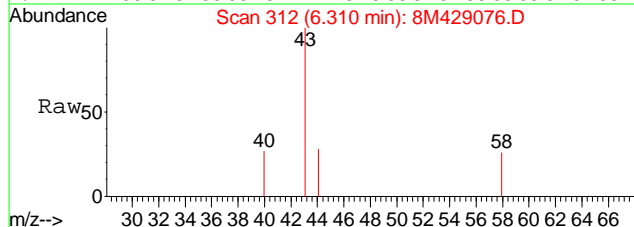
#3
 Chloromethane
 Concen: 0.40 ug/L
 RT: 3.67 min Scan# 57
 Delta R.T. -0.01 min
 Lab File: 8M429076.D
 Acq: 11 Mar 2019 17:01

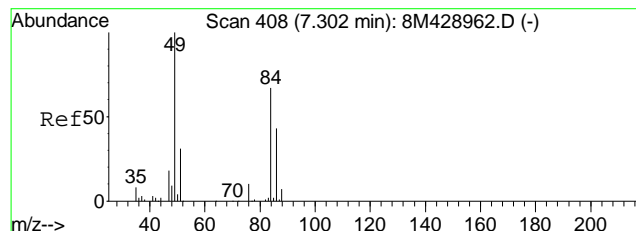
Tgt Ion	Resp	Lower	Upper
50	4517		
50	100		
52	36.3	19.7	46.1



#13
 Acetone
 Concen: 3.16 ug/L
 RT: 6.31 min Scan# 312
 Delta R.T. 0.00 min
 Lab File: 8M429076.D
 Acq: 11 Mar 2019 17:01

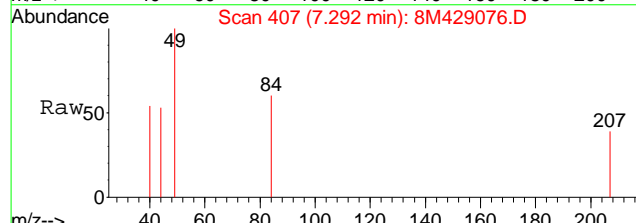
Tgt Ion	Resp	Lower	Upper
43	4429		
43	100		
58	20.3	17.6	41.2



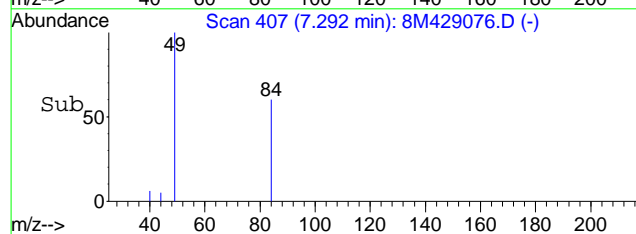
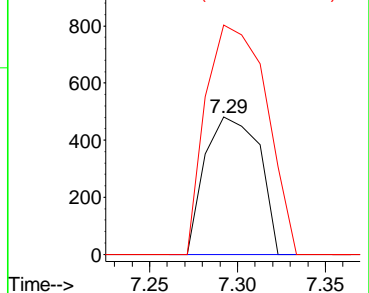


#19
 Methylene Chloride
 Concen: 0.17 ug/L
 RT: 7.29 min Scan# 407
 Delta R.T. -0.01 min
 Lab File: 8M429076.D
 Acq: 11 Mar 2019 17:01

Tgt Ion	Ratio	Lower	Upper
84	100		
49	186.0	90.7	211.7



Abundance Ion 84.00 (83.70 to 84.70): 8N
 Ion 49.00 (48.70 to 49.70): 8N



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429077.D Vial: 17
 Acq On : 11 Mar 2019 17:30 Operator: EEA
 Sample : L19030638-05 A 826-SPE Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 09:00:37 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

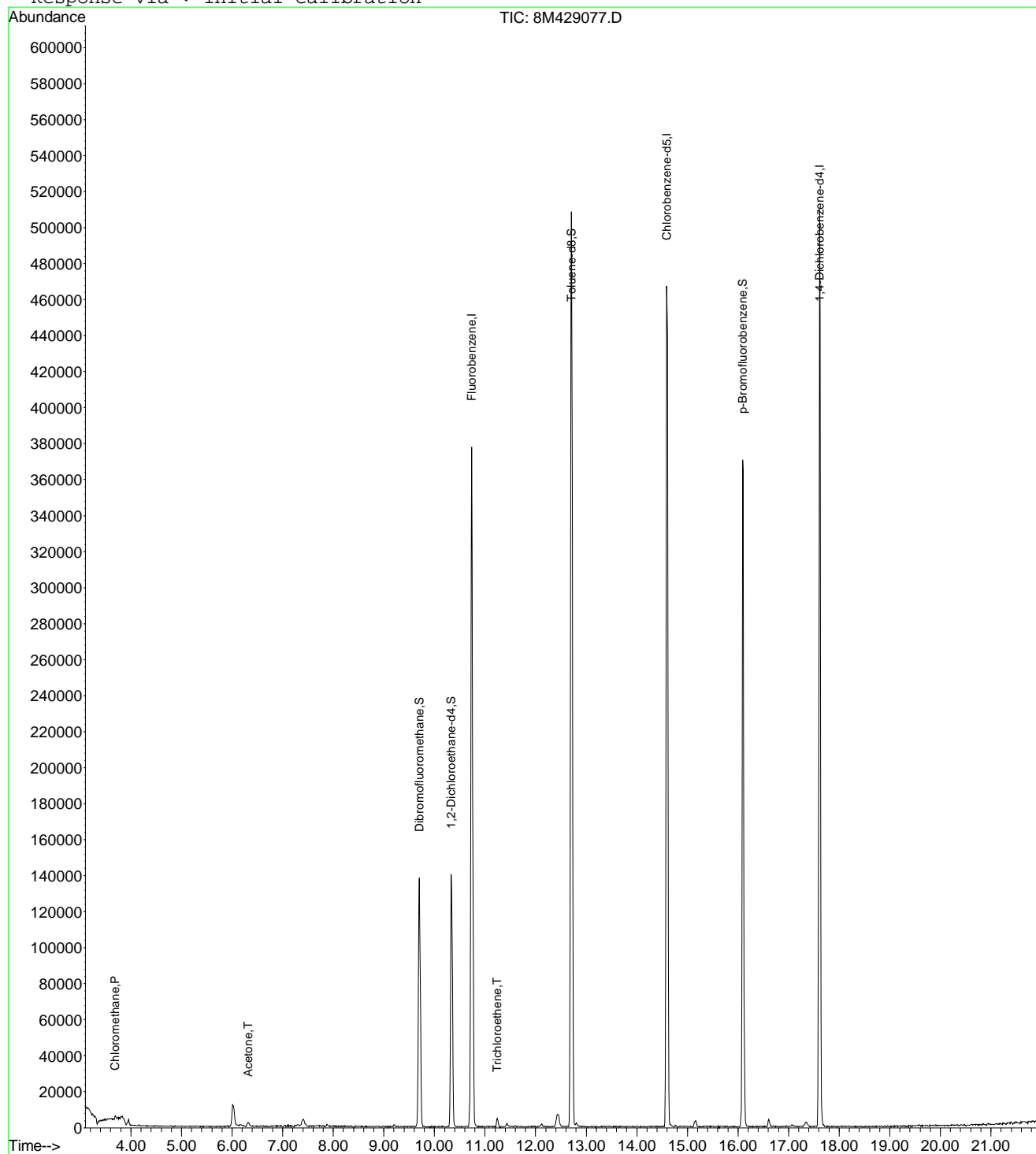
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	465976	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.59	117	350977	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.61	152	182078	25.00	ug/L	0.00
System Monitoring Compounds						
37) Dibromofluoromethane	9.70	111	120553	25.9541	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	103.80%	
43) 1,2-Dichloroethane-d4	10.33	65	121077	26.4045	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	105.60%	
58) Toluene-d8	12.71	98	460535	25.7597	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	103.04%	
80) p-Bromofluorobenzene	16.09	95	174740	26.5910	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	106.36%	
Target Compounds						
						Qvalue
3) Chloromethane	3.69	50	2748	0.2358	ug/L	# 64
13) Acetone	6.31	43	5246	3.6547	ug/L	83
47) Trichloroethene	11.24	130	2529	0.3729	ug/L	92

(#) = qualifier out of range (m) = manual integration
 8M429077.D 8260WTR.M Tue Mar 12 09:00:37 2019

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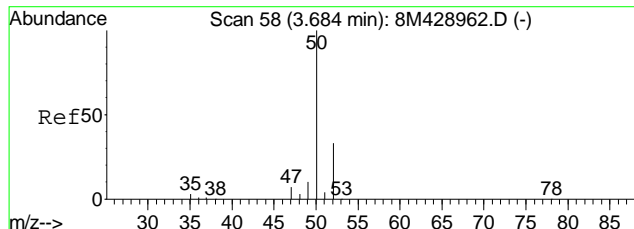
Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429077.D Vial: 17
Acq On : 11 Mar 2019 17:30 Operator: EEA
Sample : L19030638-05 A 826-SPE Inst : HPMS8
Misc : 1,1 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 12 9:00 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



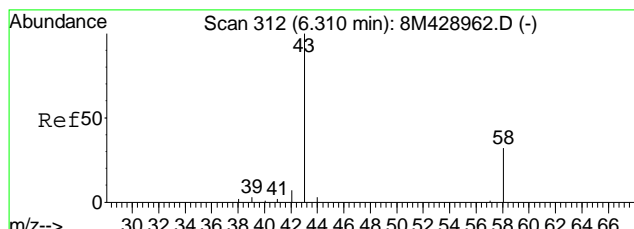
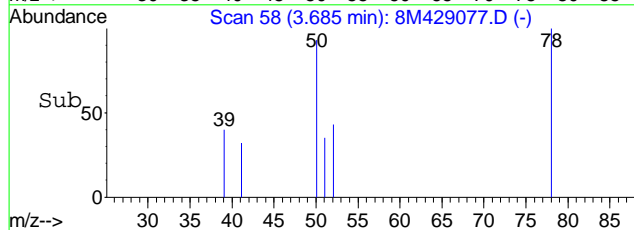
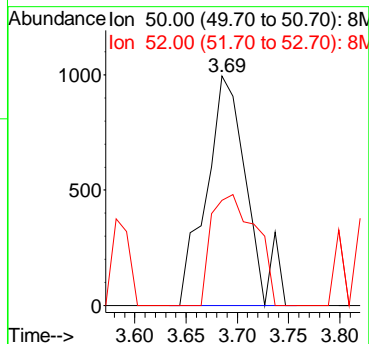
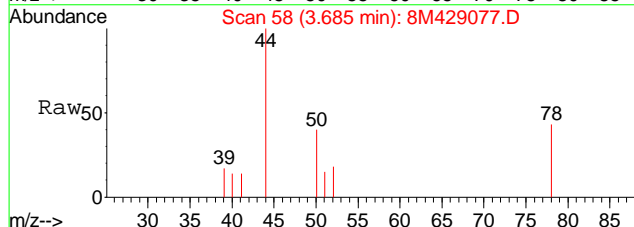
8M429077.D 8260WTR.M Tue Mar 12 09:00:38 2019

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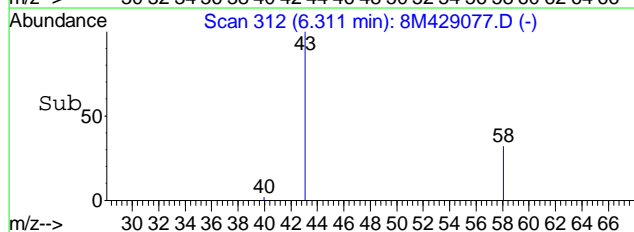
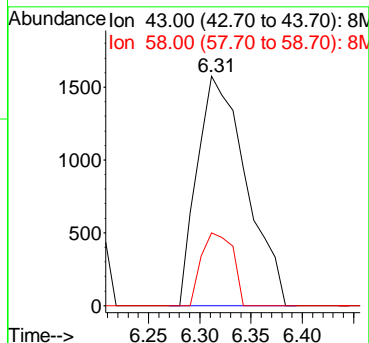
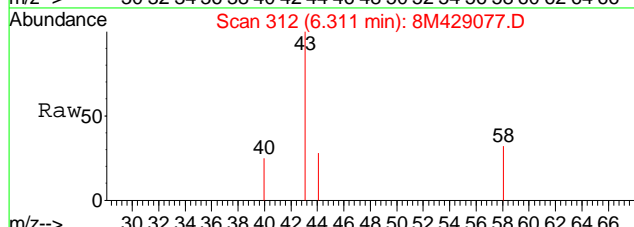
#3
 Chloromethane
 Concen: 0.24 ug/L
 RT: 3.69 min Scan# 58
 Delta R.T. 0.00 min
 Lab File: 8M429077.D
 Acq: 11 Mar 2019 17:30

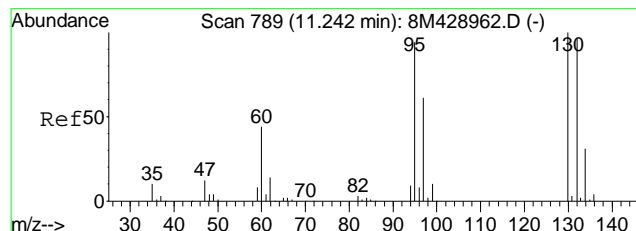
Tgt Ion: 50 Resp: 2748
 Ion Ratio Lower Upper
 50 100
 52 53.1 19.7 46.1#



#13
 Acetone
 Concen: 3.65 ug/L
 RT: 6.31 min Scan# 312
 Delta R.T. 0.00 min
 Lab File: 8M429077.D
 Acq: 11 Mar 2019 17:30

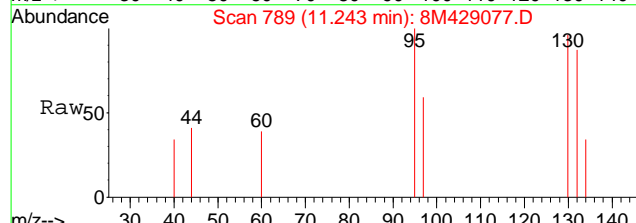
Tgt Ion: 43 Resp: 5246
 Ion Ratio Lower Upper
 43 100
 58 20.2 17.6 41.2



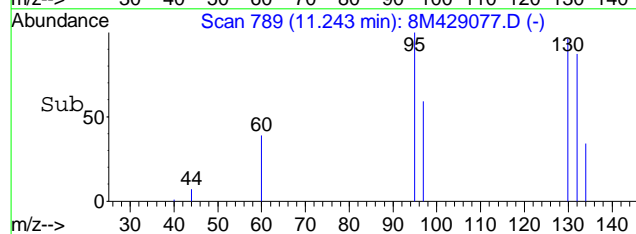
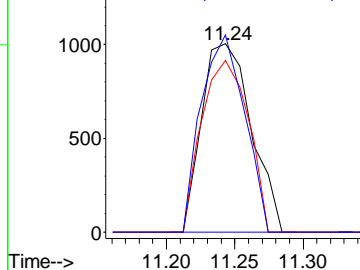


#47
 Trichloroethene
 Concen: 0.37 ug/L
 RT: 11.24 min Scan# 789
 Delta R.T. 0.00 min
 Lab File: 8M429077.D
 Acq: 11 Mar 2019 17:30

Tgt Ion	Ratio	Lower	Upper
130	100		
132	85.6	58.0	135.2
95	91.5	57.4	133.8



Abundance Ion 130.00 (129.70 to 130.70):
 Ion 132.00 (131.70 to 132.70):
 Ion 95.00 (94.70 to 95.70): 8N



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429078.D Vial: 18
 Acq On : 11 Mar 2019 17:59 Operator: EEA
 Sample : L19030638-06 A 826-SPE Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 09:00:39 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

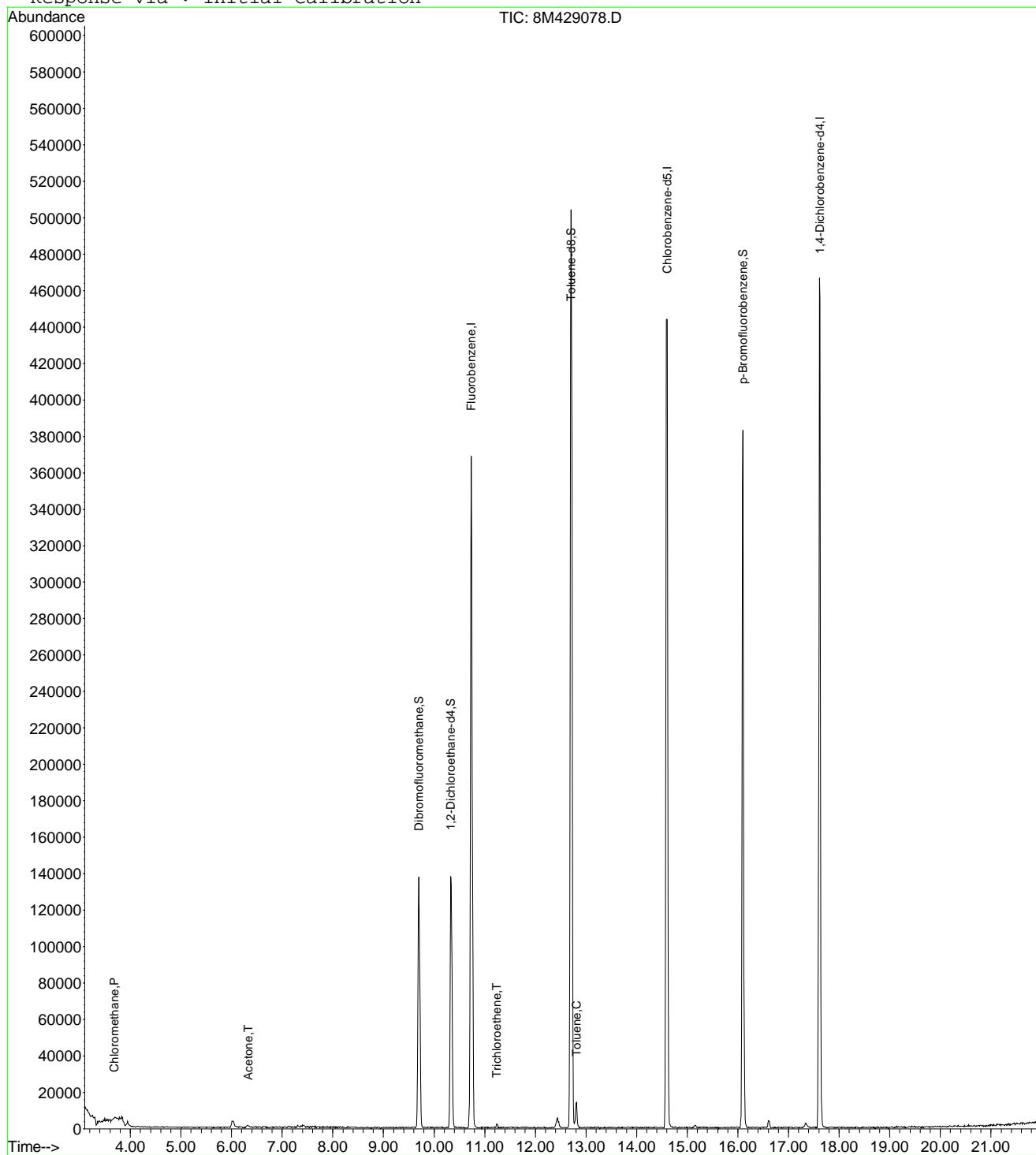
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.73	96	457216	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	341702	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	174153	25.00	ug/L	0.00
System Monitoring Compounds						
37) Dibromofluoromethane	9.70	111	118738	26.0532	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	104.20%	
43) 1,2-Dichloroethane-d4	10.33	65	122035	27.1234	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	108.48%	
58) Toluene-d8	12.71	98	450320	25.8720	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	103.48%	
80) p-Bromofluorobenzene	16.10	95	175118	27.8612	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	111.44%	
Target Compounds						
						Qvalue
3) Chloromethane	3.68	50	3338	0.2919	ug/L	87
13) Acetone	6.33	43	3143	2.2316	ug/L #	45
47) Trichloroethene	11.24	130	1116	0.1677	ug/L	79
59) Toluene	12.81	91	15796	0.6191	ug/L	98

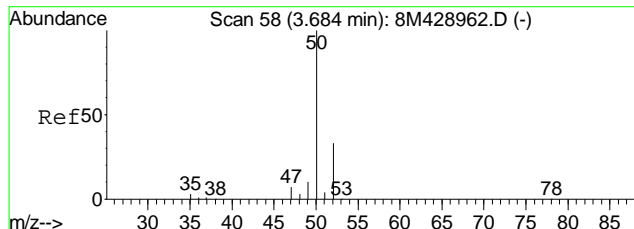
(#) = qualifier out of range (m) = manual integration
 8M429078.D 8260WTR.M Tue Mar 12 09:00:40 2019

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429078.D Vial: 18
 Acq On : 11 Mar 2019 17:59 Operator: EEA
 Sample : L19030638-06 A 826-SPE Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 9:00 2019 Quant Results File: 8260WTR.RES

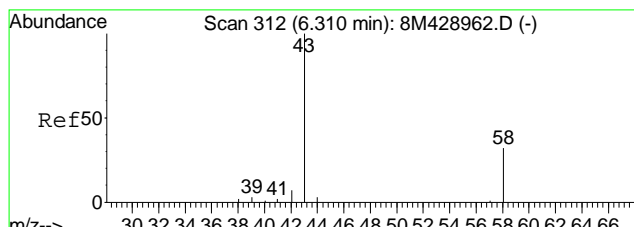
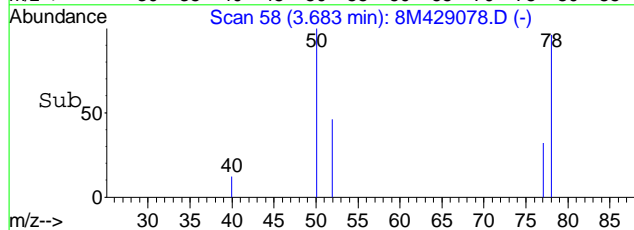
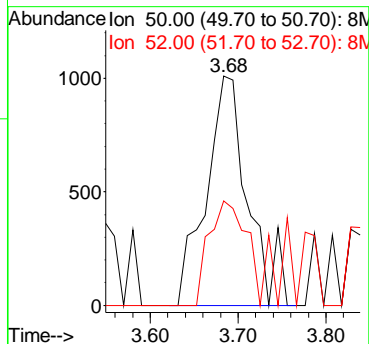
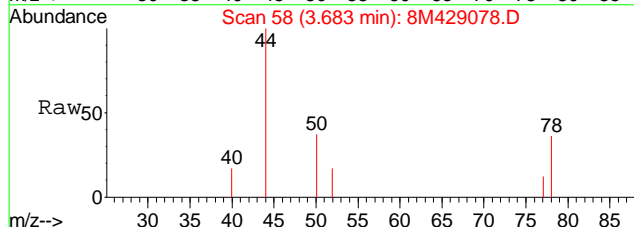
Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration





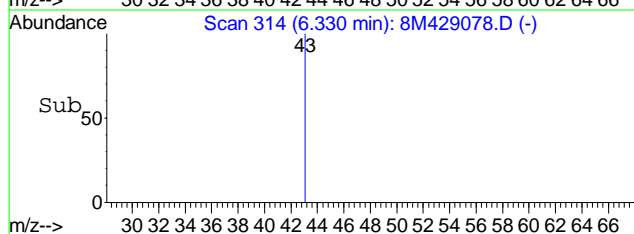
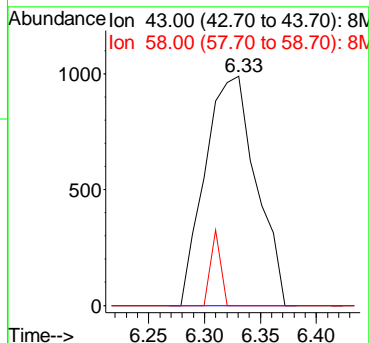
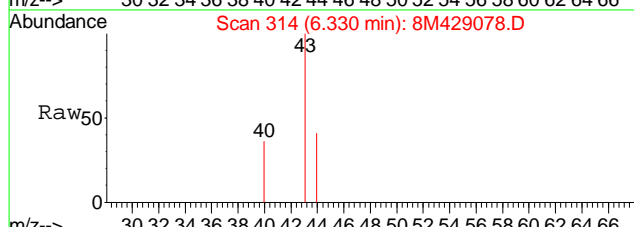
#3
 Chloromethane
 Concen: 0.29 ug/L
 RT: 3.68 min Scan# 58
 Delta R.T. -0.00 min
 Lab File: 8M429078.D
 Acq: 11 Mar 2019 17:59

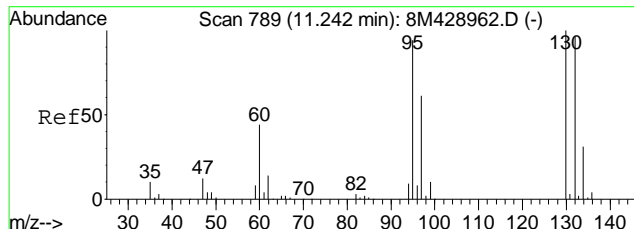
Tgt Ion	Resp	Ion	Ratio	Lower	Upper
50	3338	50	100		
52		52	40.4	19.7	46.1



#13
 Acetone
 Concen: 2.23 ug/L
 RT: 6.33 min Scan# 314
 Delta R.T. 0.02 min
 Lab File: 8M429078.D
 Acq: 11 Mar 2019 17:59

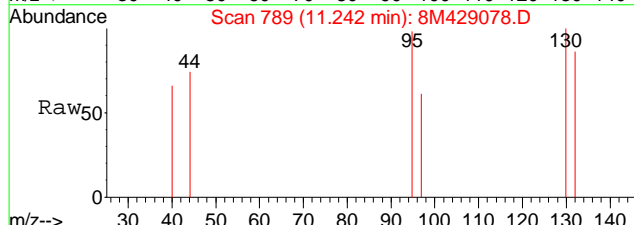
Tgt Ion	Resp	Ion	Ratio	Lower	Upper
43	3143	43	100		
58		58	0.0	17.6	41.2#



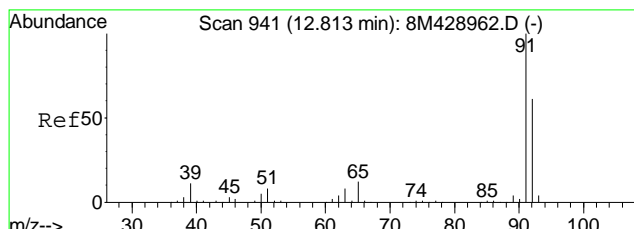
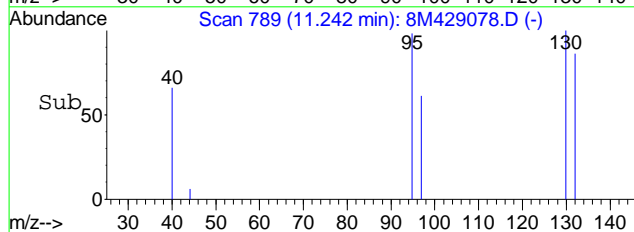
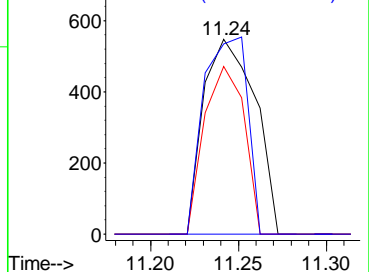


#47
 Trichloroethene
 Concen: 0.17 ug/L
 RT: 11.24 min Scan# 789
 Delta R.T. -0.00 min
 Lab File: 8M429078.D
 Acq: 11 Mar 2019 17:59

Tgt Ion	Resp	Ion	Ratio	Lower	Upper
130	1116	130	100		
132		132	66.5	58.0	135.2
95		95	85.7	57.4	133.8

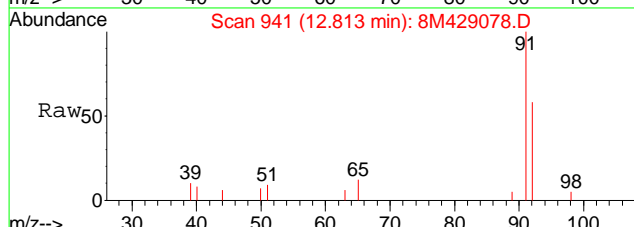


Abundance Ion 130.00 (129.70 to 130.70):
 Ion 132.00 (131.70 to 132.70): 8N
 Ion 95.00 (94.70 to 95.70): 8N

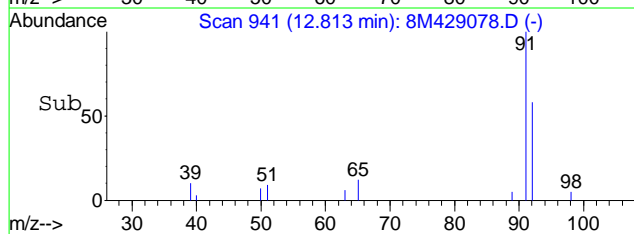
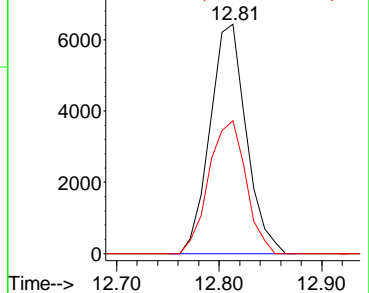


#59
 Toluene
 Concen: 0.62 ug/L
 RT: 12.81 min Scan# 941
 Delta R.T. -0.00 min
 Lab File: 8M429078.D
 Acq: 11 Mar 2019 17:59

Tgt Ion	Resp	Ion	Ratio	Lower	Upper
91	15796	91	100		
92		92	59.1	36.6	85.4



Abundance Ion 91.00 (90.70 to 91.70): 8N
 Ion 92.00 (91.70 to 92.70): 8N



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429079.D Vial: 19
 Acq On : 11 Mar 2019 18:28 Operator: EEA
 Sample : L19030638-07 A 826-SPE Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 09:00:41 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

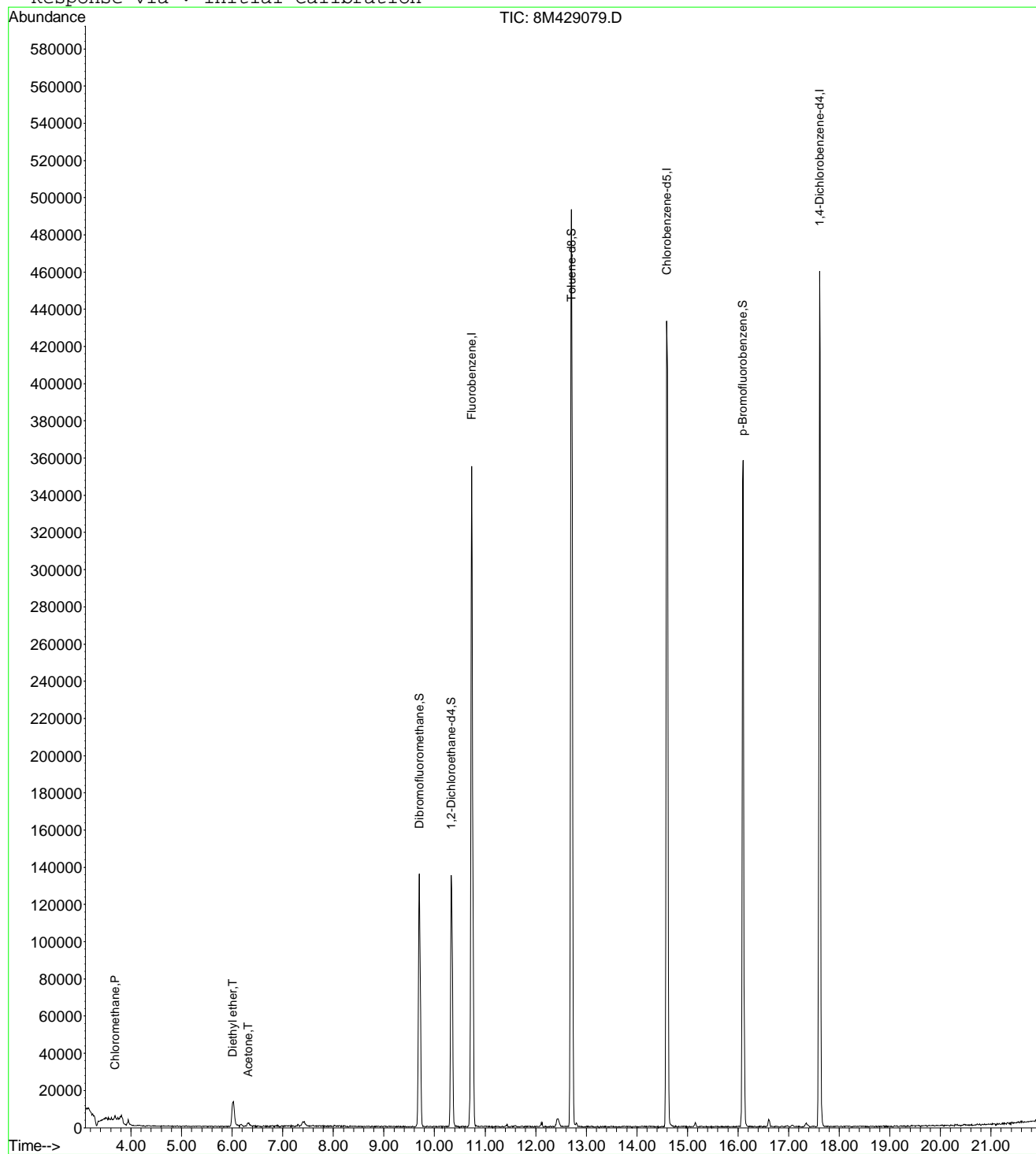
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	434603	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.59	117	327300	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.61	152	165281	25.00	ug/L	0.00
System Monitoring Compounds						
37) Dibromofluoromethane	9.70	111	114717	26.4806	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	105.92%	
43) 1,2-Dichloroethane-d4	10.33	65	118733	27.7625	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	111.04%	
58) Toluene-d8	12.71	98	442309	26.5300	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	106.12%	
80) p-Bromofluorobenzene	16.09	95	169582	28.4287	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	113.72%	
Target Compounds						
3) Chloromethane	3.68	50	2094	0.1926	ug/L	99
9) Diethyl ether	6.01	59	757	0.1578	ug/L #	1
13) Acetone	6.31	43	5028	3.7557	ug/L	78

(#) = qualifier out of range (m) = manual integration
 8M429079.D 8260WTR.M Tue Mar 12 09:00:42 2019

Page 1

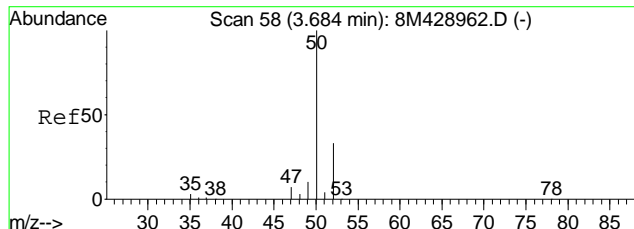
Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429079.D Vial: 19
Acq On : 11 Mar 2019 18:28 Operator: EEA
Sample : L19030638-07 A 826-SPE Inst : HPMS8
Misc : 1,1 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 12 9:00 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



8M429079.D 8260WTR.M Tue Mar 12 09:00:42 2019

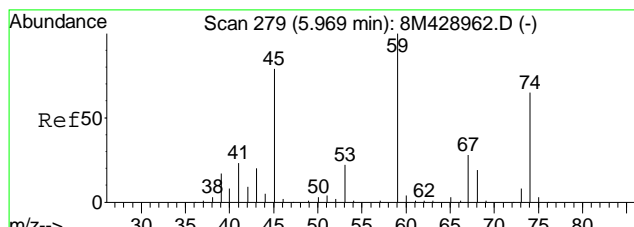
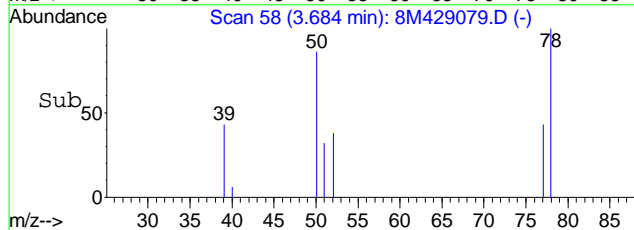
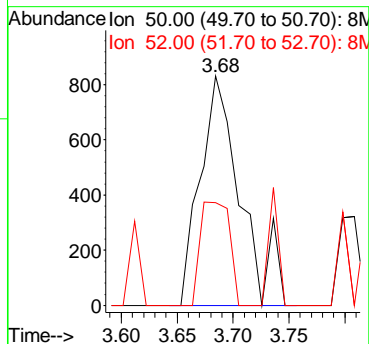
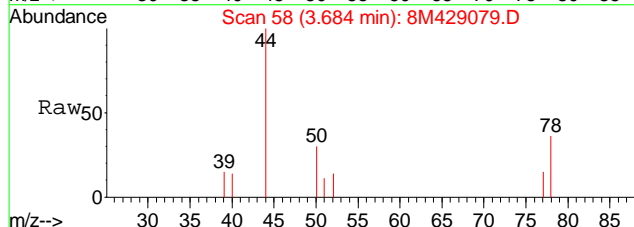
Page 2



#3
 Chloromethane
 Concen: 0.19 ug/L
 RT: 3.68 min Scan# 58
 Delta R.T. 0.00 min
 Lab File: 8M429079.D
 Acq: 11 Mar 2019 18:28

Tgt Ion: 50 Resp: 2094

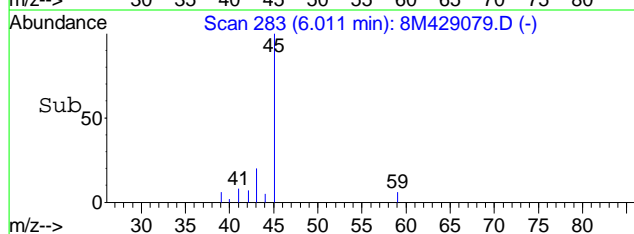
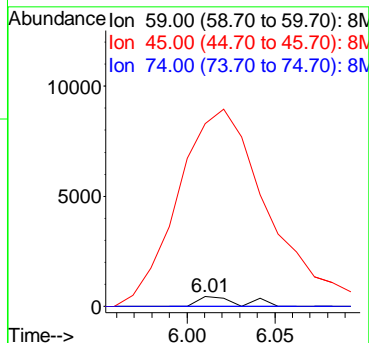
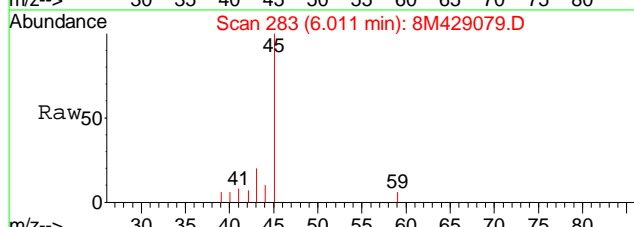
Ion	Ratio	Lower	Upper
50	100		
52	32.6	19.7	46.1

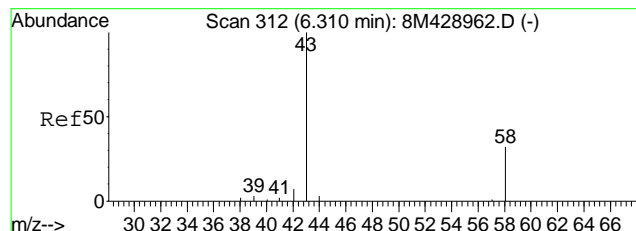


#9
 Diethyl ether
 Concen: 0.16 ug/L
 RT: 6.01 min Scan# 283
 Delta R.T. 0.04 min
 Lab File: 8M429079.D
 Acq: 11 Mar 2019 18:28

Tgt Ion: 59 Resp: 757

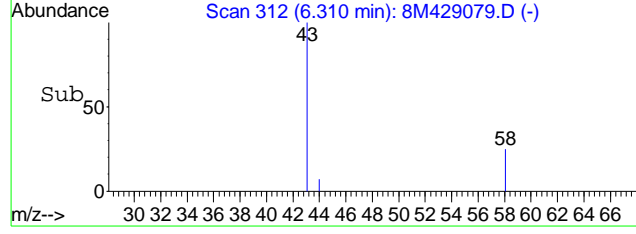
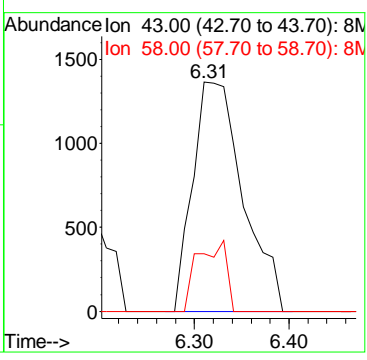
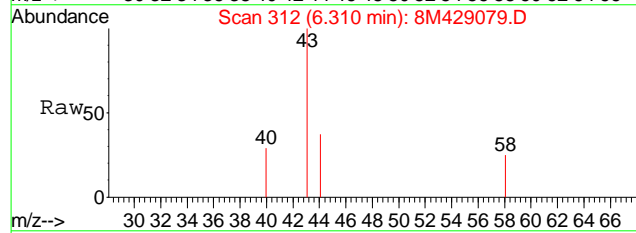
Ion	Ratio	Lower	Upper
59	100		
45	4317.3	47.6	111.2#
74	0.0	38.6	90.2#





#13
 Acetone
 Concen: 3.76 ug/L
 RT: 6.31 min Scan# 312
 Delta R.T. 0.00 min
 Lab File: 8M429079.D
 Acq: 11 Mar 2019 18:28

Tgt Ion	Ratio	Lower	Upper
43	100		
58	17.6	17.6	41.2



Data File : C:\MSDCHEM\1\DATA\031219\11M29836.D Vial: 8
 Acq On : 12 Mar 2019 14:59 Operator: KFR
 Sample : L19030638-08 A TBLK 826-SPE Inst : hpms11
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 16 10:12:51 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	375235	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0207	117	282943	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	140720	25.0000	ug/L	0.0000
System Monitoring Compounds						
37) Dibromofluoromethane	9.3989	111	107668	24.8572	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	99.4288%	
43) 1,2-Dichloroethane-d4	10.0089	65	128137	24.6558	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	98.6232%	
57) Toluene-d8	12.2423	98	350595	24.8984	ug/L	-0.0103
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	99.5936%	
78) p-Bromofluorobenzene	15.4165	95	132232	24.5552	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	98.2208%	
Target Compounds						
3) Chloromethane	3.5467	50	2094	0.1967	ug/L #	56
13) Acetone	6.1212	43	3660	2.6680	ug/L #	69
19) Methylene Chloride	7.0828	84	977	0.2442	ug/L	92
36) Tetrahydrofuran	9.3575	42	892	0.7879	ug/L #	37

(#) = qualifier out of range (m) = manual integration
 11M29836.D 8260WT.M Sat Mar 16 10:12:52 2019

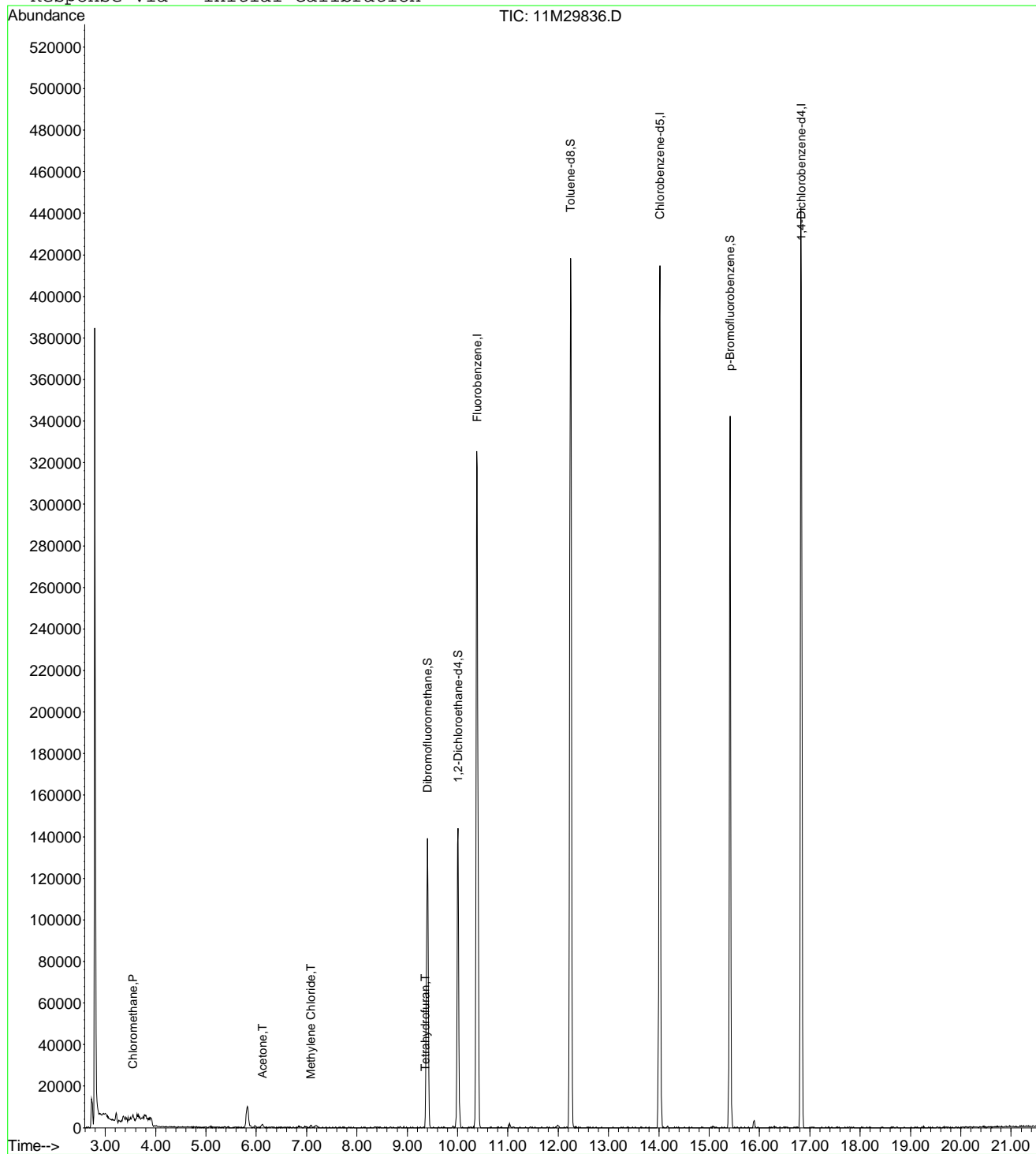
Page 1

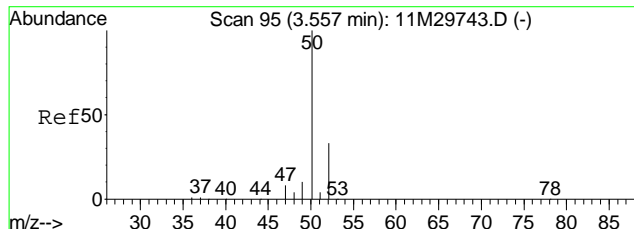
Data File : C:\MSDCHEM\1\DATA\031219\11M29836.D
 Acq On : 12 Mar 2019 14:59
 Sample : L19030638-08 A TBLK 826-SPE
 Misc : 1,1
 MS Integration Params: rteint.p
 Quant Time: Mar 16 10:12 2019

Vial: 8
 Operator: KFR
 Inst : hpms11
 Multiplr: 1.00

Quant Results File: 8260WT.RES

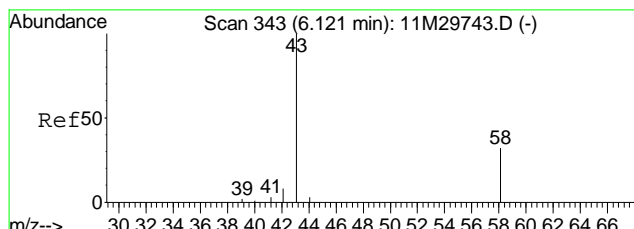
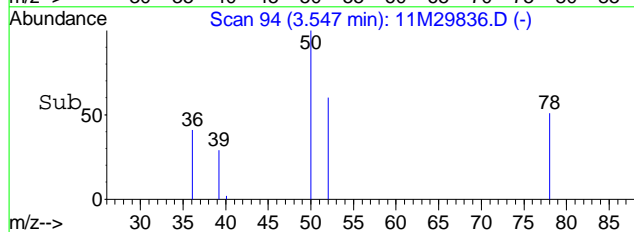
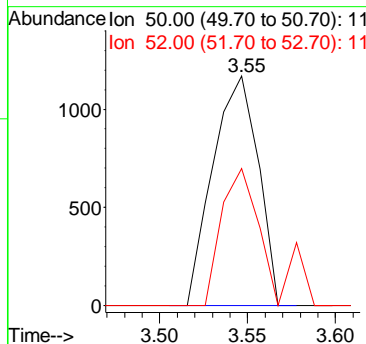
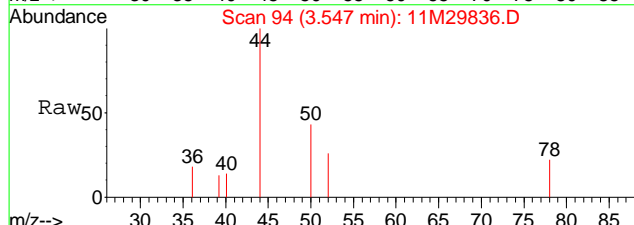
Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration





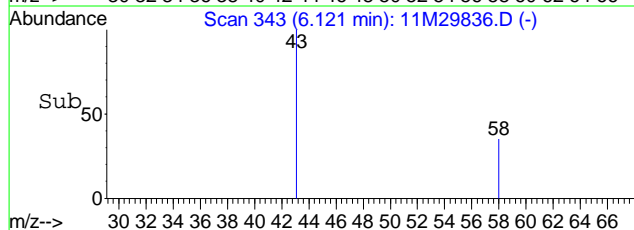
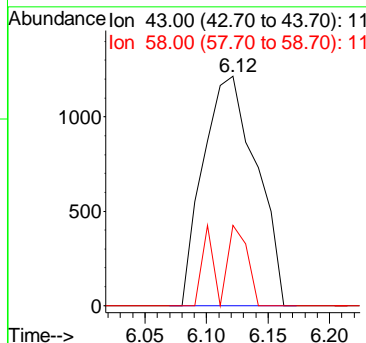
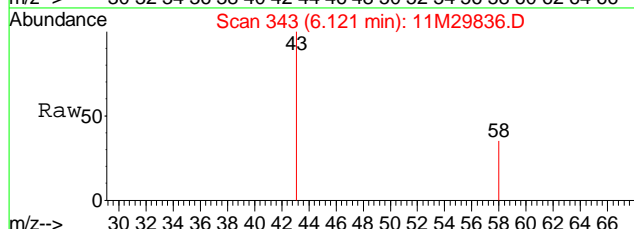
#3
 Chloromethane
 Concen: 0.1967 ug/L
 RT: 3.55 min Scan# 94
 Delta R.T. -0.01 min
 Lab File: 11M29836.D
 Acq: 12 Mar 2019 14:59

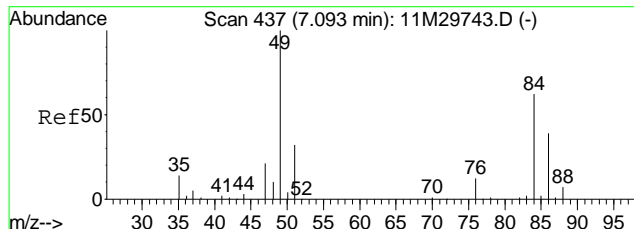
Tgt Ion	Resp	Lower	Upper
50	100		
52	57.4	19.6	45.6#



#13
 Acetone
 Concen: 2.6680 ug/L
 RT: 6.12 min Scan# 343
 Delta R.T. 0.00 min
 Lab File: 11M29836.D
 Acq: 12 Mar 2019 14:59

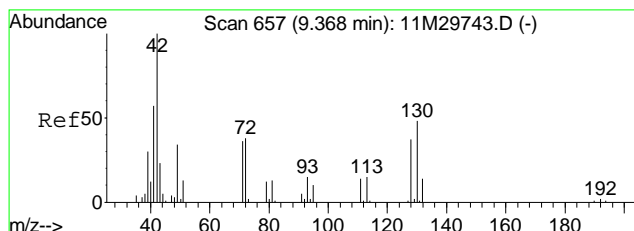
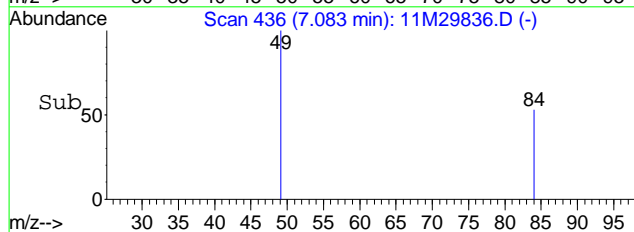
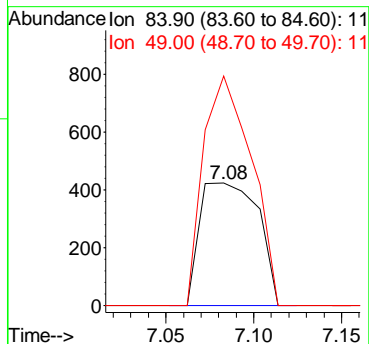
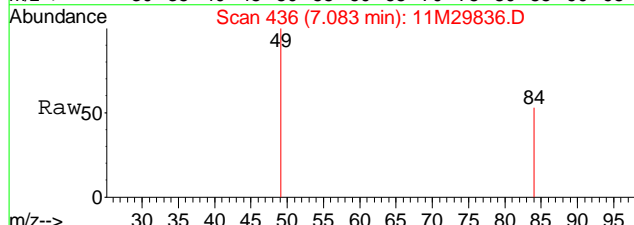
Tgt Ion	Resp	Lower	Upper
43	100		
58	12.7	17.8	41.4#





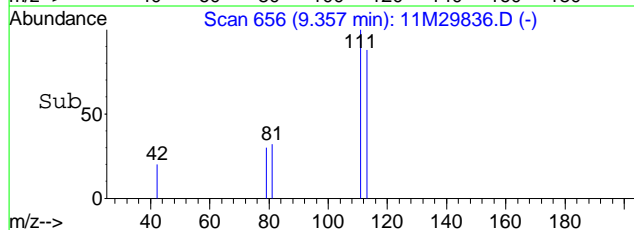
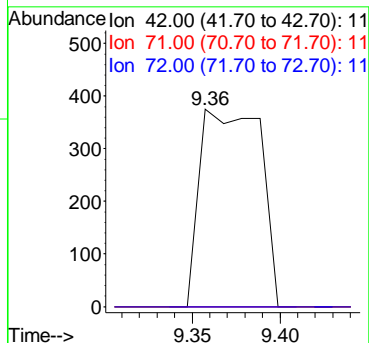
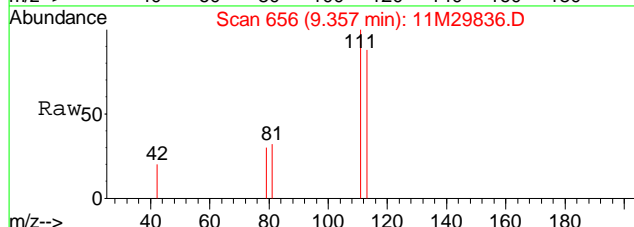
#19
Methylene Chloride
Concen: 0.2442 ug/L
RT: 7.08 min Scan# 436
Delta R.T. -0.01 min
Lab File: 11M29836.D
Acq: 12 Mar 2019 14:59

Tgt Ion	Resp	Lower	Upper
84	100		
49	154.5	99.4	231.8



#36
Tetrahydrofuran
Concen: 0.7879 ug/L
RT: 9.36 min Scan# 656
Delta R.T. -0.01 min
Lab File: 11M29836.D
Acq: 12 Mar 2019 14:59

Tgt Ion	Resp	Lower	Upper
42	100		
71	0.0	22.1	51.7#
72	0.0	22.9	53.3#



2.1.1.4 Standards Data

Data File : C:\MSDCHEM\1\DATA\021919\11M29628.D Vial: 4
 Acq On : 19 Feb 2019 13:17 Operator: KFR
 Sample : WG696624-03 5ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:20 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3812	96	493305	25.0000	ug/L	0.0000
12) Chlorobenzene-d5	14.0104	117	374544	25.0000	ug/L	0.0000
13) 1,4-Dichlorobenzene-d4	16.8227	152	189147	25.0000	ug/L	0.0000
						Qvalue
Target Compounds						
2) Acetonitrile	6.5452	41	3643	4.9714	ug/L	74
3) 3-Chloro-1-propene	6.9071	41	51893	5.3368	ug/L	99
4) 2-Chloro-1,3-butadiene	8.2513	53	58581	5.4502	ug/L	97
5) Methacrylonitrile	9.0060	41	24202	5.7049	ug/L	89
6) Isobutyl Alcohol	9.0164	43	4052	14.1171	ug/L	71
7) 1-Butanol	9.9056	56	628	3.7711	ug/L #	14
8) Cyclohexanone	15.1995	55	3200	7.7190	ug/L #	67
9) 2-Nitropropane	11.3738	43	9863	5.0857	ug/L	94
10) Ethyl Acetate	8.8509	43	29837	5.2588	ug/L	97
11) Methyl methacrylate	11.0636	41	30286	5.5638	ug/L	91

 (#) = qualifier out of range (m) = manual integration
 11M29628.D A9FOOWT.M Wed Feb 20 09:25:20 2019

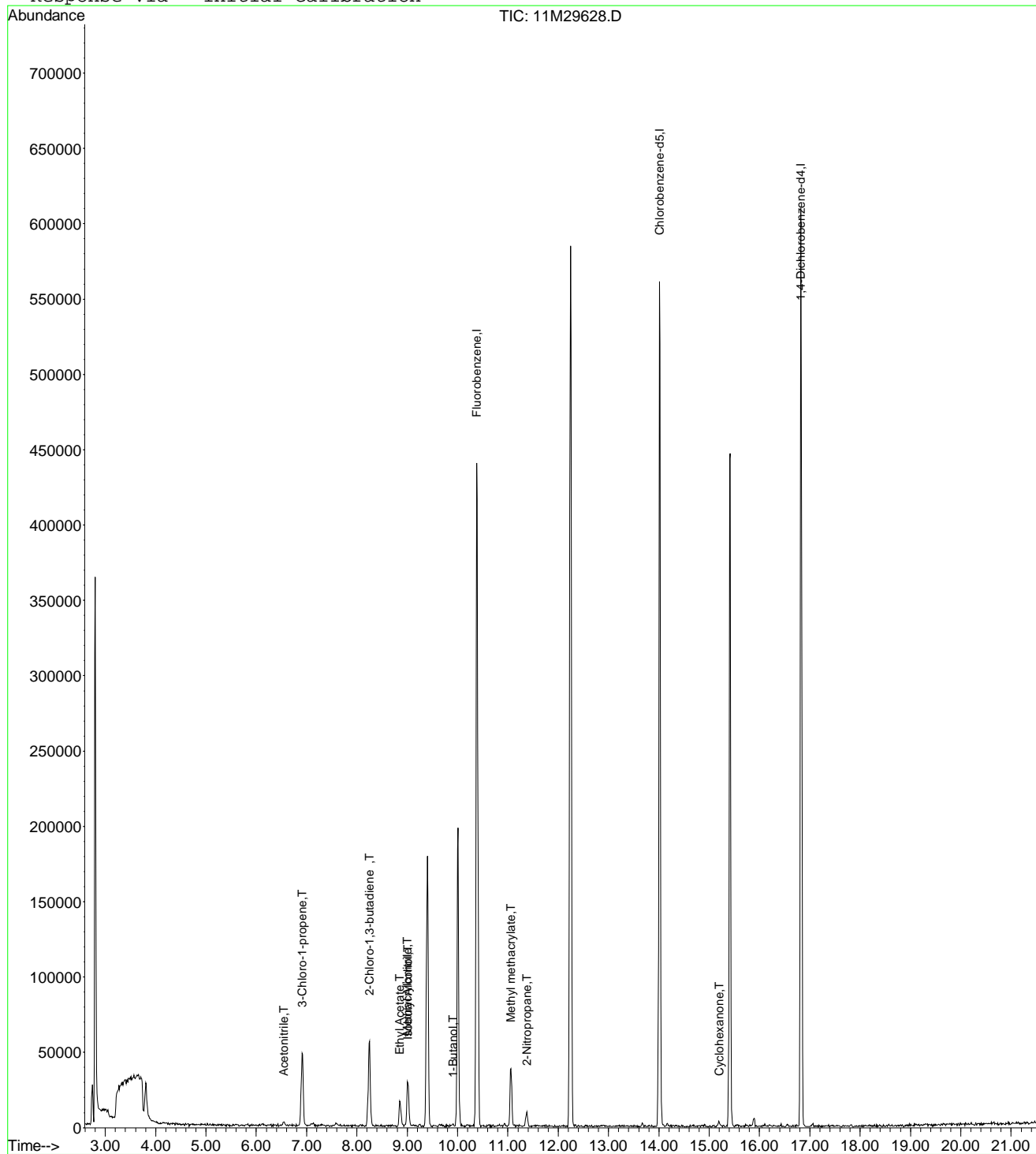
Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29628.D
 Acq On : 19 Feb 2019 13:17
 Sample : WG696624-03 5ug/L ICAL A9 8260
 Misc : 1,1 STD92151
 MS Integration Params: rteint.p
 Quant Time: Feb 20 9:25 2019

Vial: 4
 Operator: KFR
 Inst : hpms11
 Multiplr: 1.00

Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\021919\11M29628.D Vial: 4
 Acq On : 19 Feb 2019 13:17 Operator: KFR
 Sample : WG696624-03 5ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	5.000	4.971	0.6	100	0.00
3 T	3-Chloro-1-propene	5.000	5.337	-6.7	100	-0.01
4 T	2-Chloro-1,3-butadiene	5.000	5.450	-9.0	100	0.01
5 T	Methacrylonitrile	5.000	5.705	-14.1	100	0.00
6 T	Isobutyl Alcohol	-1.000	14.117	0.0	100	0.00
7 T	1-Butanol	5.000	3.771	24.6	100	0.01
8 T	Cyclohexanone	-1.000	7.719	0.0	0	0.01
9 T	2-Nitropropane	5.000	5.086	-1.7	100	0.00
10 T	Ethyl Acetate	5.000	5.259	-5.2	100	0.00
11 T	Methyl methacrylate	5.000	5.564	-11.3	100	0.01
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29628.D A9FOOWT.M Wed Feb 20 09:21:53 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29629.D Vial: 5
 Acq On : 19 Feb 2019 13:47 Operator: KFR
 Sample : WG696624-04 20ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:21 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QI	Ion	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3811	96		480707	25.0000	ug/L	0.0000
12) Chlorobenzene-d5	14.0206	117		363206	25.0000	ug/L	0.0102
13) 1,4-Dichlorobenzene-d4	16.8226	152		186142	25.0000	ug/L	0.0000
							Qvalue
Target Compounds							
2) Acetonitrile	6.5451	41		16699	23.3856	ug/L	99
3) 3-Chloro-1-propene	6.9173	41		191643	20.2255	ug/L	96
4) 2-Chloro-1,3-butadiene	8.2511	53		212583	20.2965	ug/L	99
5) Methacrylonitrile	9.0059	41		83846	20.2822	ug/L	96
6) Isobutyl Alcohol	9.0162	43		9203	32.9033	ug/L	85
7) 1-Butanol	9.9054	56		3151	19.4173	ug/L #	34
8) Cyclohexanone	15.1890	55		8986	22.2440	ug/L	96
9) 2-Nitropropane	11.3737	43		34565	18.2899	ug/L	97
10) Ethyl Acetate	8.8508	43		115887	20.9604	ug/L	97
11) Methyl methacrylate	11.0531	41		109713	20.6836	ug/L	98

 (#) = qualifier out of range (m) = manual integration
 11M29629.D A9FOOWT.M Wed Feb 20 09:25:21 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29629.D

Vial: 5

Acq On : 19 Feb 2019 13:47

Operator: KFR

Sample : WG696624-04 20ug/L ICAL A9 8260

Inst : hpms11

Misc : 1,1 STD92151

Multiplr: 1.00

MS Integration Params: rteint.p

Quant Time: Feb 20 9:25 2019

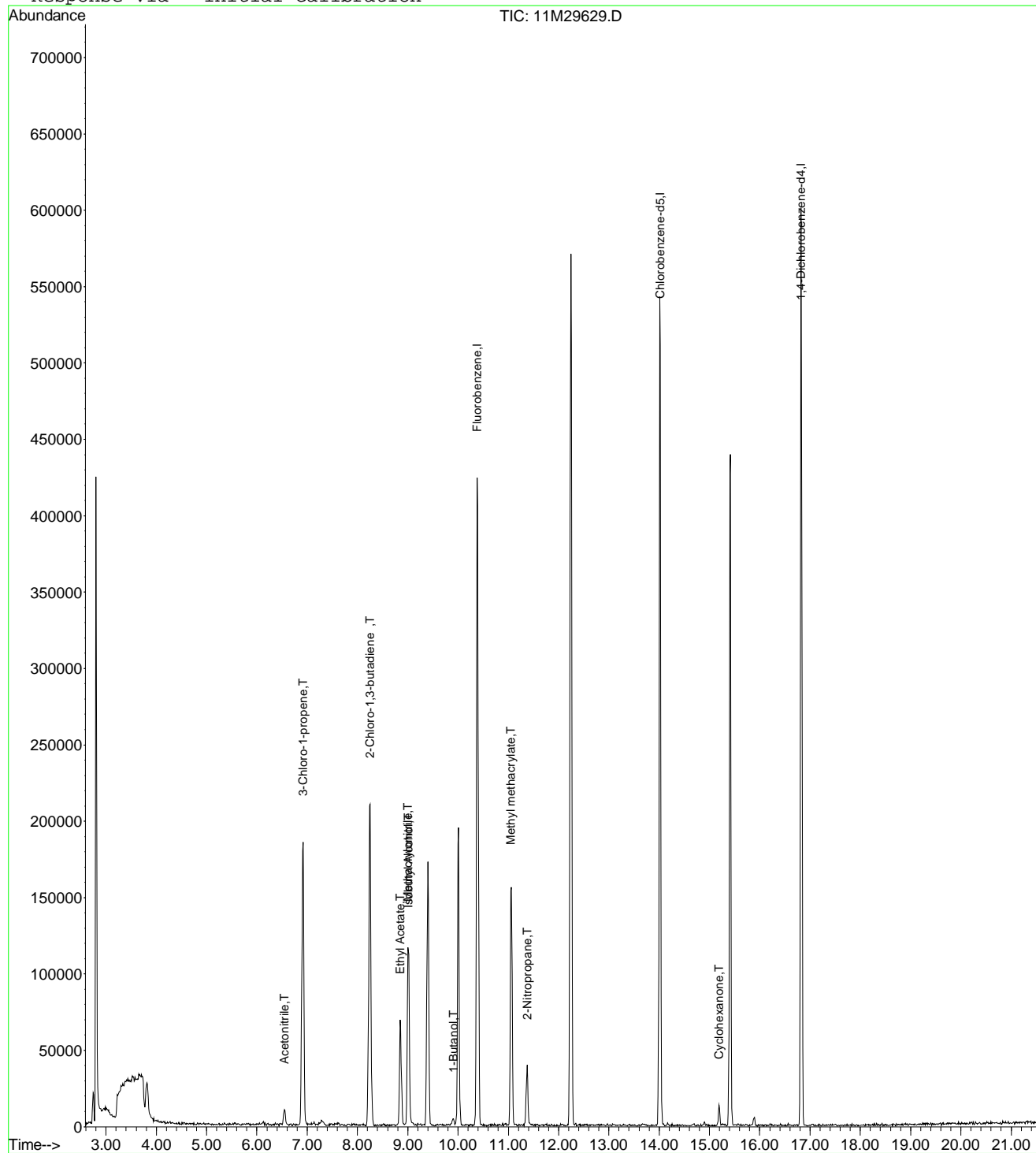
Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)

Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11

Last Update : Wed Feb 20 09:20:16 2019

Response via : Initial Calibration



11M29629.D A9FOOWT.M

Wed Feb 20 09:25:21 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\021919\11M29629.D Vial: 5
 Acq On : 19 Feb 2019 13:47 Operator: KFR
 Sample : WG696624-04 20ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	20.000	23.386	-16.9	100	0.00
3 T	3-Chloro-1-propene	20.000	20.226	-1.1	100	0.00
4 T	2-Chloro-1,3-butadiene	20.000	20.297	-1.5	100	0.01
5 T	Methacrylonitrile	20.000	20.282	-1.4	100	0.00
6 T	Isobutyl Alcohol	-1.000	32.903	0.0	100	0.00
7 T	1-Butanol	20.000	19.417	2.9	100	0.01
8 T	Cyclohexanone	20.000	22.244	-11.2	100	0.00
9 T	2-Nitropropane	20.000	18.290	8.6	100	0.00
10 T	Ethyl Acetate	20.000	20.960	-4.8	100	0.00
11 T	Methyl methacrylate	20.000	20.684	-3.4	100	0.00
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.01
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29629.D A9FOOWT.M Wed Feb 20 09:22:14 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29630.D Vial: 6
 Acq On : 19 Feb 2019 14:16 Operator: KFR
 Sample : WG696624-05 50ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:22 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	485902	25.0000	ug/L	0.0103
12) Chlorobenzene-d5	14.0207	117	364295	25.0000	ug/L	0.0103
13) 1,4-Dichlorobenzene-d4	16.8227	152	182799	25.0000	ug/L	0.0000
						Qvalue
Target Compounds						
2) Acetonitrile	6.5555	41	38349	53.1305	ug/L	98
3) 3-Chloro-1-propene	6.9174	41	493137	51.4880	ug/L	98
4) 2-Chloro-1,3-butadiene	8.2512	53	544956	51.4739	ug/L	98
5) Methacrylonitrile	9.0163	41	214843	51.4146	ug/L	100
6) Isobutyl Alcohol	9.0163	43	28972	102.4756	ug/L	91
7) 1-Butanol	9.8952	56	7907	48.2041	ug/L #	87
8) Cyclohexanone	15.1890	55	20882	51.1387	ug/L	95
9) 2-Nitropropane	11.3737	43	92750	48.5534	ug/L	99
10) Ethyl Acetate	8.8509	43	295121	52.8077	ug/L	99
11) Methyl methacrylate	11.0635	41	284315	53.0272	ug/L	98

 (#) = qualifier out of range (m) = manual integration
 11M29630.D A9FOOWT.M Wed Feb 20 09:25:22 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29630.D

Vial: 6

Acq On : 19 Feb 2019 14:16

Operator: KFR

Sample : WG696624-05 50ug/L ICAL A9 8260

Inst : hpms11

Misc : 1,1 STD92151

Multiplr: 1.00

MS Integration Params: rteint.p

Quant Time: Feb 20 9:25 2019

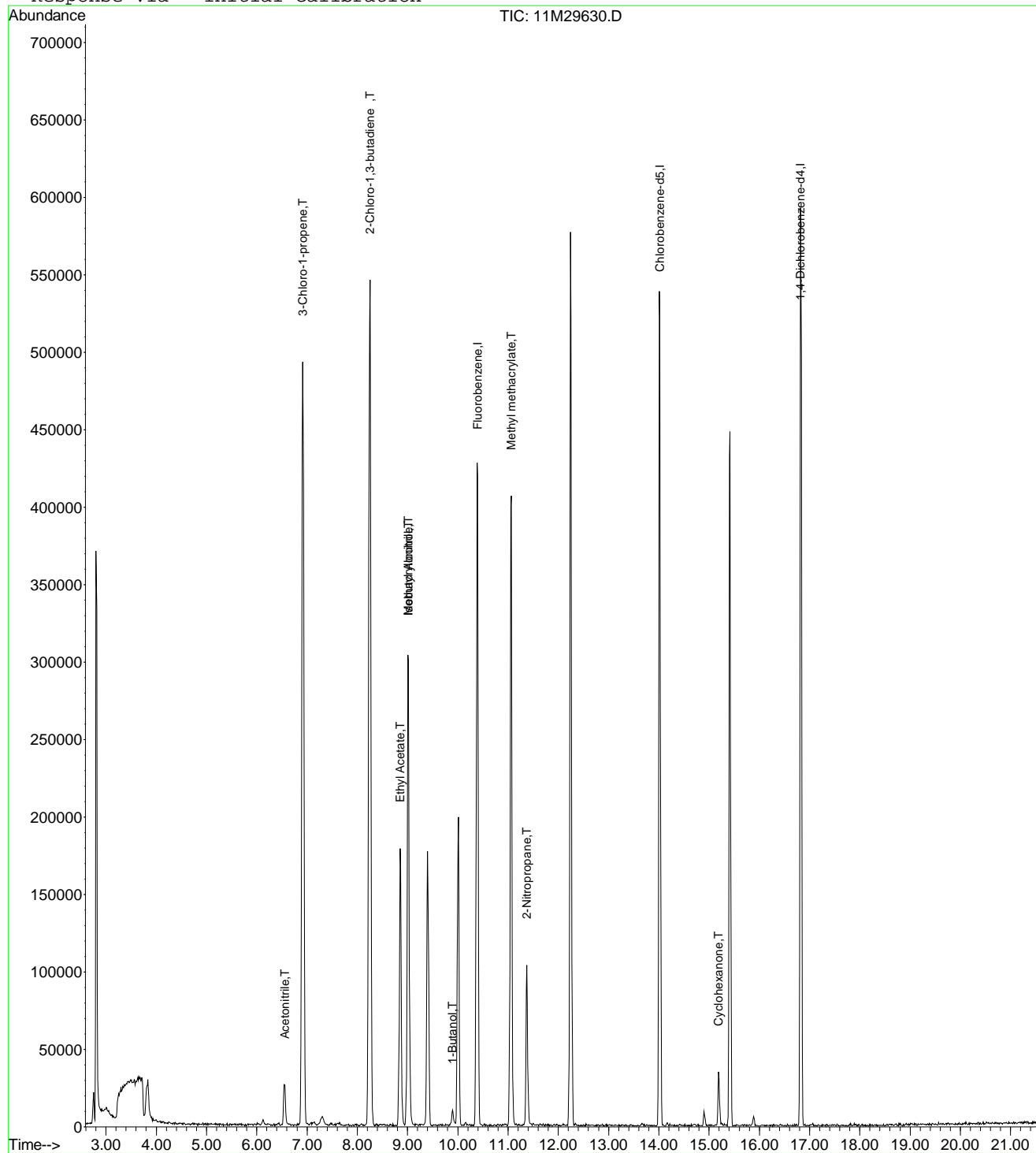
Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)

Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11

Last Update : Wed Feb 20 09:20:16 2019

Response via : Initial Calibration



11M29630.D A9FOOWT.M

Wed Feb 20 09:25:22 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29630.D Vial: 6
 Acq On : 19 Feb 2019 14:16 Operator: KFR
 Sample : WG696624-05 50ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.01
2 T	Acetonitrile	50.000	53.130	-6.3	100	0.01
3 T	3-Chloro-1-propene	50.000	51.488	-3.0	100	0.00
4 T	2-Chloro-1,3-butadiene	50.000	51.474	-2.9	100	0.01
5 T	Methacrylonitrile	50.000	51.415	-2.8	100	0.01
6 T	Isobutyl Alcohol	100.000	102.476	-2.5	100	0.00
7 T	1-Butanol	50.000	48.204	3.6	100	0.00
8 T	Cyclohexanone	50.000	51.139	-2.3	100	0.00
9 T	2-Nitropropane	50.000	48.553	2.9	100	0.00
10 T	Ethyl Acetate	50.000	52.808	-5.6	100	0.00
11 T	Methyl methacrylate	50.000	53.027	-6.1	100	0.01
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.01
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29630.D A9FOOWT.M Wed Feb 20 09:22:23 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29631.D Vial: 7
 Acq On : 19 Feb 2019 14:45 Operator: KFR
 Sample : WG696624-06 100ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:23 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3812	96	474275	25.0000	ug/L	0.0000
12) Chlorobenzene-d5	14.0104	117	359140	25.0000	ug/L	0.0000
13) 1,4-Dichlorobenzene-d4	16.8227	152	178860	25.0000	ug/L	0.0000
						Qvalue
Target Compounds	6.5452	41	68209	96.8166	ug/L	100
2) Acetonitrile	6.9174	41	974729	104.2656	ug/L	100
3) 3-Chloro-1-propene	8.2409	53	1129368	109.2897	ug/L	100
4) 2-Chloro-1,3-butadiene	9.0060	41	410236	100.5813	ug/L	100
5) Methacrylonitrile	9.0163	43	52091	188.7657	ug/L	100
6) Isobutyl Alcohol	9.8952	56	16192	101.1326	ug/L	100
7) 1-Butanol	15.1891	55	39101	98.1034	ug/L	100
8) Cyclohexanone	11.3738	43	182796	98.0372	ug/L	100
9) 2-Nitropropane	8.8509	43	565874	103.7374	ug/L	100
10) Ethyl Acetate	11.0532	41	544148	103.9763	ug/L	100
11) Methyl methacrylate						

 (#) = qualifier out of range (m) = manual integration
 11M29631.D A9FOOWT.M Wed Feb 20 09:25:23 2019

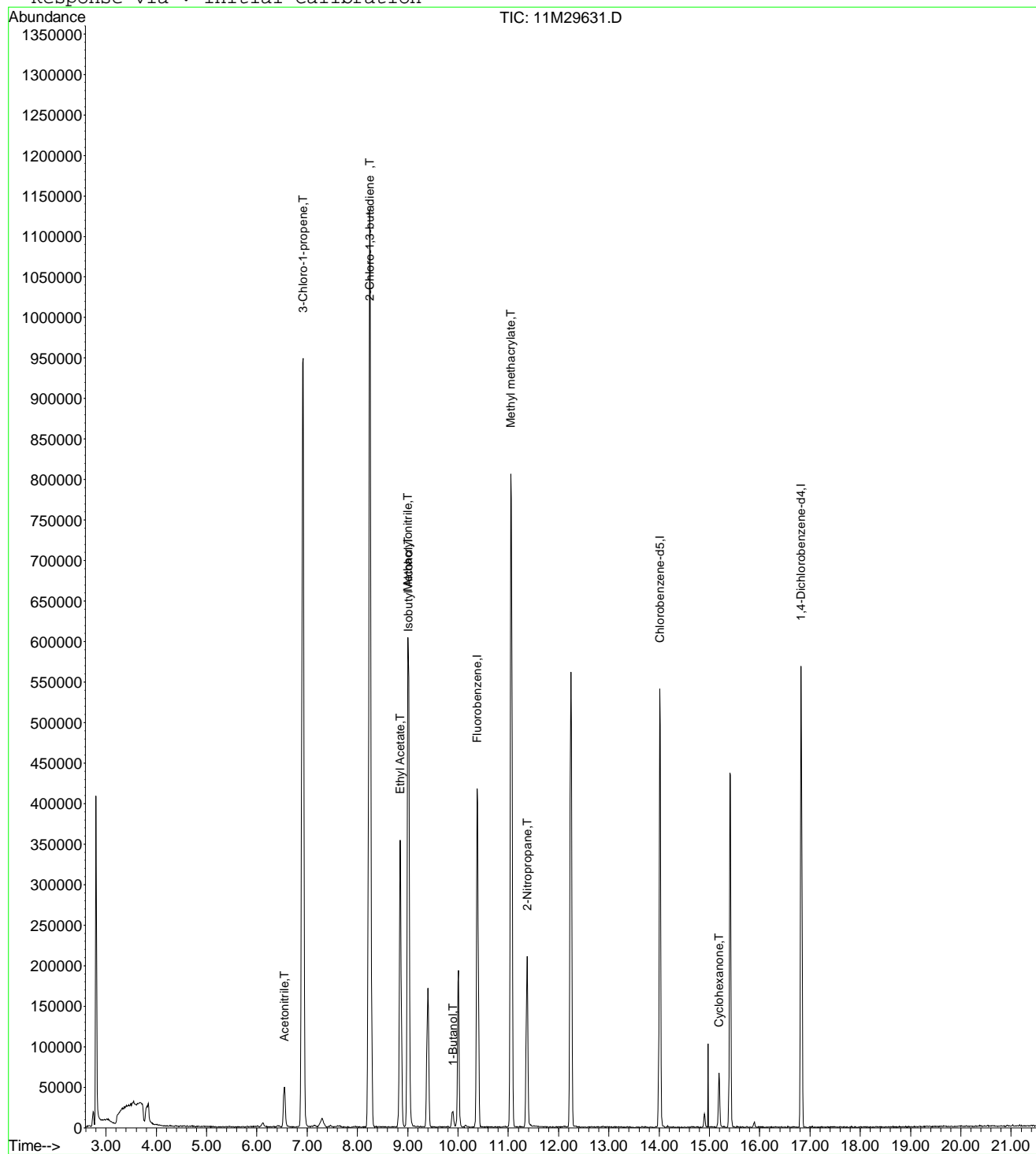
Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29631.D
Acq On : 19 Feb 2019 14:45
Sample : WG696624-06 100ug/L ICAL A9 8260
Misc : 1,1 STD92151
MS Integration Params: rteint.p
Quant Time: Feb 20 9:25 2019

Vial: 7
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
Last Update : Wed Feb 20 09:20:16 2019
Response via : Initial Calibration



11M29631.D A9FOOWT.M

Wed Feb 20 09:25:23 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29631.D Vial: 7
 Acq On : 19 Feb 2019 14:45 Operator: KFR
 Sample : WG696624-06 100ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	100.000	96.817	3.2	100	0.00
3 T	3-Chloro-1-propene	100.000	104.266	-4.3	100	0.00
4 T	2-Chloro-1,3-butadiene	100.000	109.290	-9.3	100	0.00
5 T	Methacrylonitrile	100.000	100.581	-0.6	100	0.00
6 T	Isobutyl Alcohol	200.000	188.766	5.6	100	0.00
7 T	1-Butanol	100.000	101.133	-1.1	100	0.00
8 T	Cyclohexanone	100.000	98.103	1.9	100	0.00
9 T	2-Nitropropane	100.000	98.037	2.0	100	0.00
10 T	Ethyl Acetate	100.000	103.737	-3.7	100	0.00
11 T	Methyl methacrylate	100.000	103.976	-4.0	100	0.00
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29631.D A9FOOWT.M Wed Feb 20 09:22:34 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29632.D Vial: 8
 Acq On : 19 Feb 2019 15:15 Operator: KFR
 Sample : WG696624-07 200ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:23 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3812	96	492245	25.0000	ug/L	0.0000
12) Chlorobenzene-d5	14.0104	117	371771	25.0000	ug/L	0.0000
13) 1,4-Dichlorobenzene-d4	16.8227	152	181875	25.0000	ug/L	0.0000

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.5452	41	133510	182.5875	ug/L	97
3) 3-Chloro-1-propene	6.9071	41	1836470	189.2735	ug/L	98
4) 2-Chloro-1,3-butadiene	8.2409	53	2100037	195.8032	ug/L	97
5) Methacrylonitrile	9.0060	41	781256	184.5549	ug/L	98
6) Isobutyl Alcohol	9.0060	43	106345	371.3012	ug/L	98
7) 1-Butanol	9.8952	56	32962	198.3596	ug/L	92
8) Cyclohexanone	15.1891	55	77205	186.6340	ug/L	98
9) 2-Nitropropane	11.3738	43	366954	189.6203	ug/L	98
10) Ethyl Acetate	8.8509	43	1072599	189.4531	ug/L	100
11) Methyl methacrylate	11.0533	41	1030275	189.6791	ug/L	99

 (#) = qualifier out of range (m) = manual integration
 11M29632.D A9FOOWT.M Wed Feb 20 09:25:24 2019

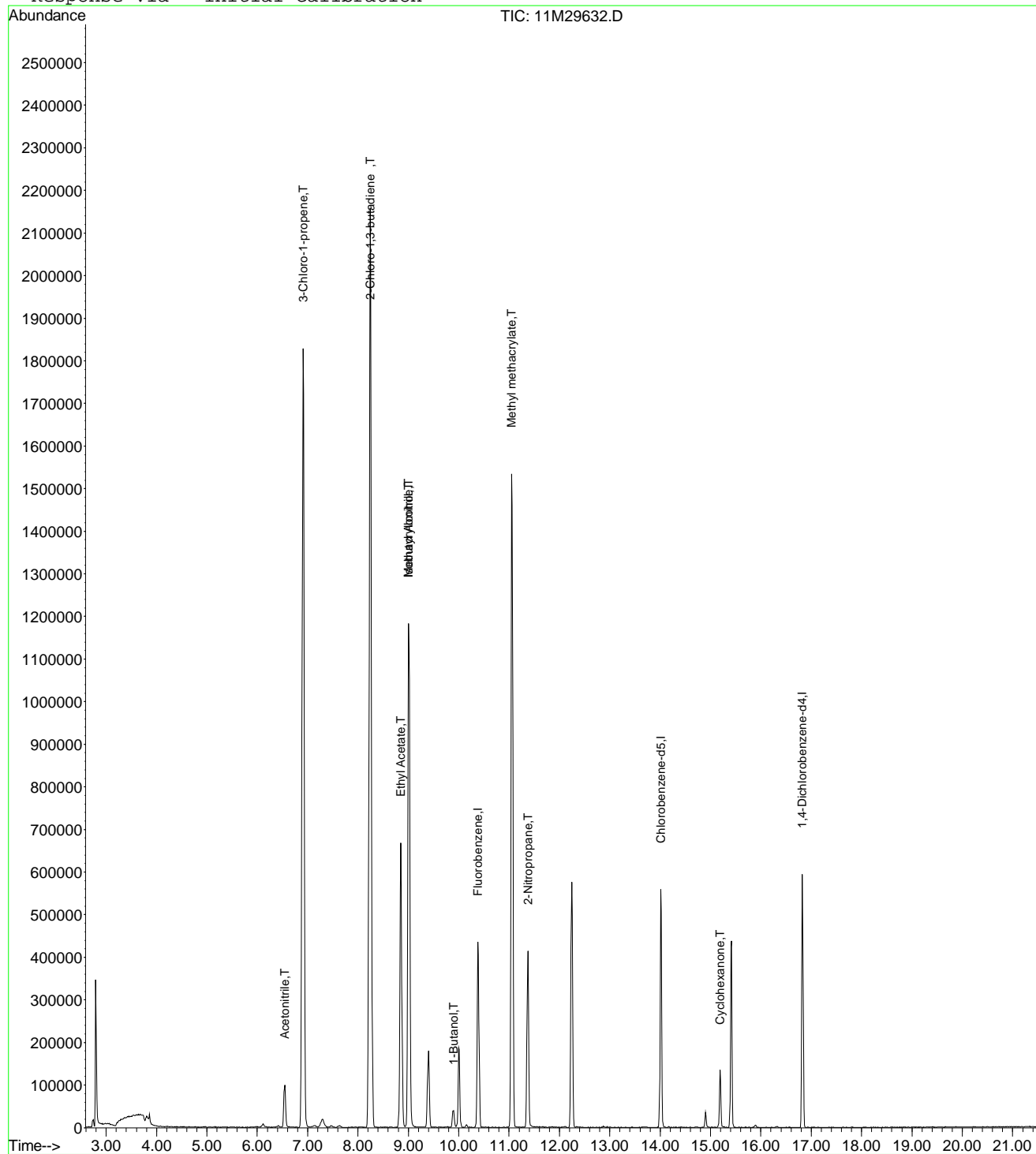
Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29632.D
Acq On : 19 Feb 2019 15:15
Sample : WG696624-07 200ug/L ICAL A9 8260
Misc : 1,1 STD92151
MS Integration Params: rteint.p
Quant Time: Feb 20 9:25 2019

Vial: 8
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
Last Update : Wed Feb 20 09:20:16 2019
Response via : Initial Calibration



11M29632.D A9FOOWT.M

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Data File : C:\MSDCHEM\1\DATA\021919\11M29632.D Vial: 8
 Acq On : 19 Feb 2019 15:15 Operator: KFR
 Sample : WG696624-07 200ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	200.000	182.587	8.7	100	0.00
3 T	3-Chloro-1-propene	200.000	189.274	5.4	100	-0.01
4 T	2-Chloro-1,3-butadiene	200.000	195.803	2.1	100	0.00
5 T	Methacrylonitrile	200.000	184.555	7.7	100	0.00
6 T	Isobutyl Alcohol	400.000	371.301	7.2	100	-0.01
7 T	1-Butanol	200.000	198.360	0.8	100	0.00
8 T	Cyclohexanone	200.000	186.634	6.7	100	0.00
9 T	2-Nitropropane	200.000	189.620	5.2	100	0.00
10 T	Ethyl Acetate	200.000	189.453	5.3	100	0.00
11 T	Methyl methacrylate	200.000	189.679	5.2	100	0.00
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29632.D A9FOOWT.M Wed Feb 20 09:22:43 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29633.D Vial: 9
 Acq On : 19 Feb 2019 15:44 Operator: KFR
 Sample : WG696624-08 300ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:24 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3812	96	474847	25.0000	ug/L	0.0000
12) Chlorobenzene-d5	14.0104	117	358911	25.0000	ug/L	0.0000
13) 1,4-Dichlorobenzene-d4	16.8228	152	182980	25.0000	ug/L	0.0000
						Qvalue
Target Compounds						
2) Acetonitrile	6.5452	41	207084	293.5833	ug/L	99
3) 3-Chloro-1-propene	6.9071	41	2534392	270.7743	ug/L	93
4) 2-Chloro-1,3-butadiene	8.2409	53	2894878	279.8019	ug/L	91
5) Methacrylonitrile	9.0061	41	1228345	300.8017	ug/L	97
6) Isobutyl Alcohol	9.0164	43	179274	648.8648	ug/L	96
7) 1-Butanol	9.8953	56	53953	336.5759	ug/L #	90
8) Cyclohexanone	15.1891	55	120994	303.2053	ug/L	94
9) 2-Nitropropane	11.3738	43	592773	317.5332	ug/L	97
10) Ethyl Acetate	8.8510	43	1650539	302.2162	ug/L	98
11) Methyl methacrylate	11.0533	41	1556331	297.0270	ug/L	96

 (#) = qualifier out of range (m) = manual integration
 11M29633.D A9FOOWT.M Wed Feb 20 09:25:24 2019

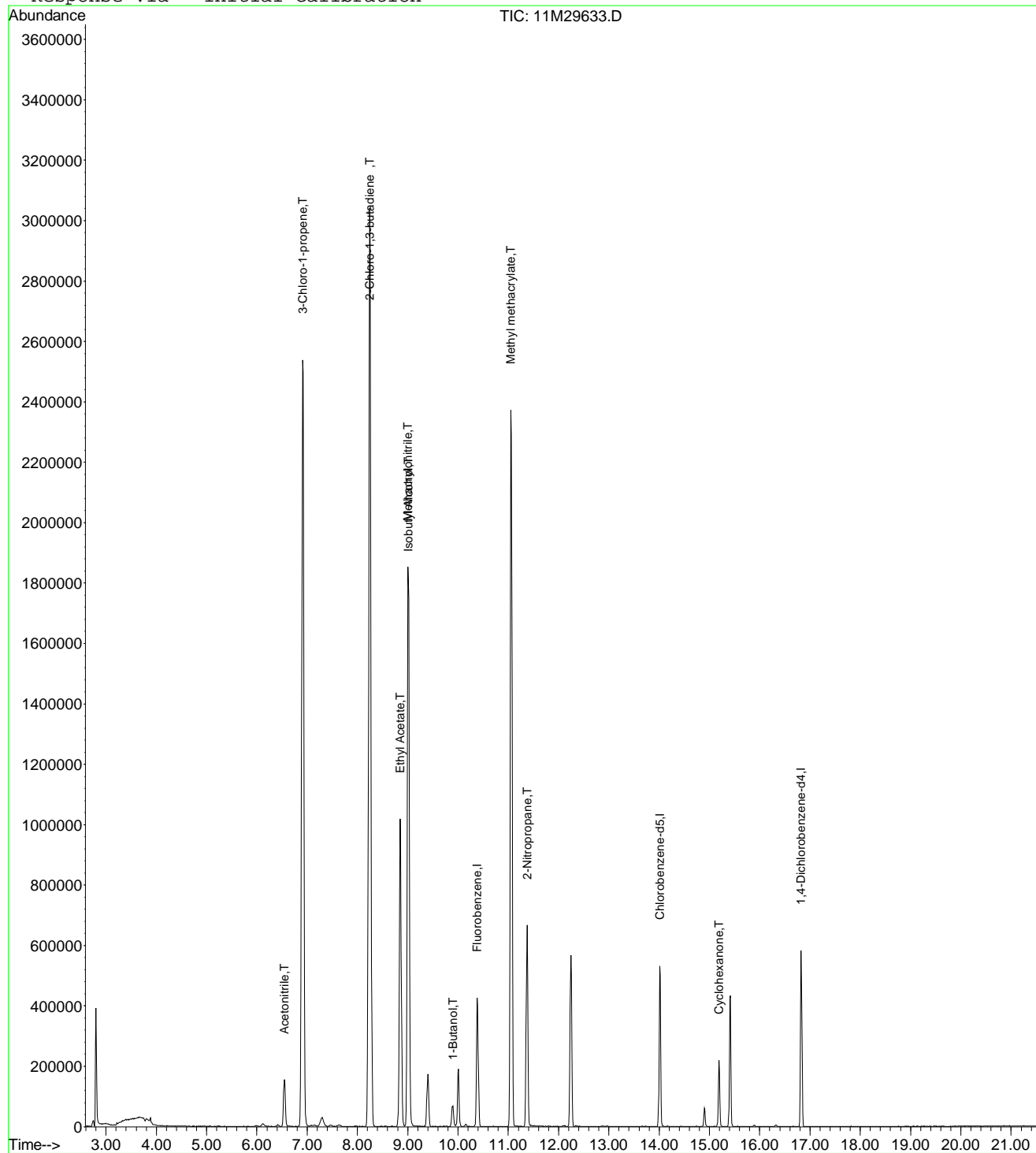
Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29633.D
Acq On : 19 Feb 2019 15:44
Sample : WG696624-08 300ug/L ICAL A9 8260
Misc : 1,1 STD92151
MS Integration Params: rteint.p
Quant Time: Feb 20 9:25 2019

Vial: 9
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
Last Update : Wed Feb 20 09:20:16 2019
Response via : Initial Calibration



11M29633.D A9FOOWT.M Wed Feb 20 09:25:25 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29633.D Vial: 9
 Acq On : 19 Feb 2019 15:44 Operator: KFR
 Sample : WG696624-08 300ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	300.000	293.583	2.1	100	0.00
3 T	3-Chloro-1-propene	300.000	270.774	9.7	100	-0.01
4 T	2-Chloro-1,3-butadiene	300.000	279.802	6.7	100	0.00
5 T	Methacrylonitrile	300.000	300.802	-0.3	100	0.00
6 T	Isobutyl Alcohol	600.000	648.865	-8.1	100	0.00
7 T	1-Butanol	300.000	336.576	-12.2	100	0.00
8 T	Cyclohexanone	300.000	303.205	-1.1	100	0.00
9 T	2-Nitropropane	300.000	317.533	-5.8	100	0.00
10 T	Ethyl Acetate	300.000	302.216	-0.7	100	0.00
11 T	Methyl methacrylate	300.000	297.027	1.0	100	0.00
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29633.D A9FOOWT.M Wed Feb 20 09:22:54 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29634.D Vial: 10
 Acq On : 19 Feb 2019 16:14 Operator: KFR
 Sample : WG696624-09 400ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:25 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3812	96	468154	25.0000	ug/L	0.0000
12) Chlorobenzene-d5	14.0104	117	356242	25.0000	ug/L	0.0000
13) 1,4-Dichlorobenzene-d4	16.8227	152	182659	25.0000	ug/L	0.0000
						Qvalue
Target Compounds						
2) Acetonitrile	6.5452	41	264059	379.7089	ug/L	100
3) 3-Chloro-1-propene	6.9071	41	3101308	336.0806	ug/L	88
4) 2-Chloro-1,3-butadiene	8.2409	53	3513249	344.4247	ug/L	85
5) Methacrylonitrile	9.0060	41	1525655	378.9494	ug/L	95
6) Isobutyl Alcohol	9.0164	43	222649	817.3773	ug/L	97
7) 1-Butanol	9.8952	56	68125	431.0611	ug/L #	88
8) Cyclohexanone	15.1891	55	149846	380.8755	ug/L	96
9) 2-Nitropropane	11.3738	43	779327	423.4338	ug/L	96
10) Ethyl Acetate	8.8509	43	2034170	377.7845	ug/L	97
11) Methyl methacrylate	11.0533	41	1917267	371.1432	ug/L	93

 (#) = qualifier out of range (m) = manual integration
 11M29634.D A9FOOWT.M Wed Feb 20 09:25:25 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29634.D

Vial: 10

Acq On : 19 Feb 2019 16:14

Operator: KFR

Sample : WG696624-09 400ug/L ICAL A9 8260

Inst : hpms11

Misc : 1,1 STD92151

Multiplr: 1.00

MS Integration Params: rteint.p

Quant Time: Feb 20 9:25 2019

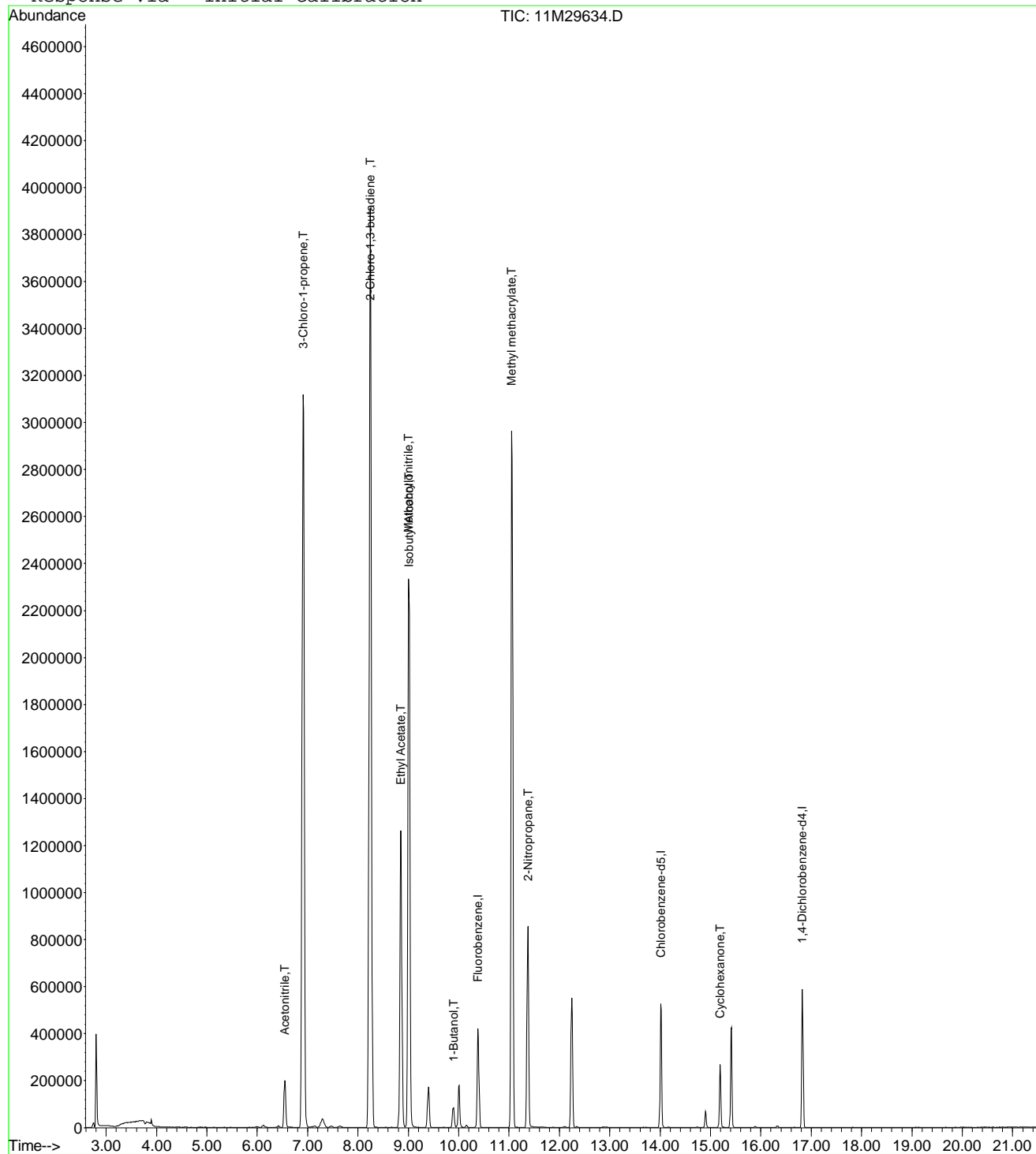
Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)

Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11

Last Update : Wed Feb 20 09:20:16 2019

Response via : Initial Calibration



11M29634.D A9FOOWT.M

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Data File : C:\MSDCHEM\1\DATA\021919\11M29634.D Vial: 10
 Acq On : 19 Feb 2019 16:14 Operator: KFR
 Sample : WG696624-09 400ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	400.000	379.709	5.1	100	0.00
3 T	3-Chloro-1-propene	-1.000	336.081	0.0	100	-0.01
4 T	2-Chloro-1,3-butadiene	400.000	344.425	13.9	100	0.00
5 T	Methacrylonitrile	400.000	378.949	5.3	100	0.00
6 T	Isobutyl Alcohol	800.000	817.377	-2.2	100	0.00
7 T	1-Butanol	400.000	431.061	-7.8	100	0.00
8 T	Cyclohexanone	400.000	380.876	4.8	100	0.00
9 T	2-Nitropropane	400.000	423.434	-5.9	100	0.00
10 T	Ethyl Acetate	400.000	377.785	5.6	100	0.00
11 T	Methyl methacrylate	400.000	371.143	7.2	100	0.00
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29634.D A9FOOWT.M Wed Feb 20 09:23:01 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29635.D Vial: 11
 Acq On : 19 Feb 2019 16:44 Operator: KFR
 Sample : WG696624-10 500ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:26 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	457625	25.0000	ug/L	0.0103
12) Chlorobenzene-d5	14.0206	117	342580	25.0000	ug/L	0.0103
13) 1,4-Dichlorobenzene-d4	16.8227	152	178100	25.0000	ug/L	0.0000
						Qvalue
Target Compounds						
2) Acetonitrile	6.5555	41	327940	482.4176	ug/L	100
3) 3-Chloro-1-propene	6.9173	41	3578473	396.7120	ug/L	83
4) 2-Chloro-1,3-butadiene	8.2512	53	4013475	402.5176	ug/L	79
5) Methacrylonitrile	9.0163	41	1845697	468.9907	ug/L	94
6) Isobutyl Alcohol	9.0163	43	278050	1044.2482	ug/L	98
7) 1-Butanol	9.8951	56	85596	554.0702	ug/L	91
8) Cyclohexanone	15.1890	55	189971	493.9741	ug/L	97
9) 2-Nitropropane	11.3737	43	946149	525.9013	ug/L	97
10) Ethyl Acetate	8.8508	43	2388468	453.7905	ug/L	95
11) Methyl methacrylate	11.0635	41	2238019	443.2020	ug/L	90

 (#) = qualifier out of range (m) = manual integration
 11M29635.D A9FOOWT.M Wed Feb 20 09:25:26 2019

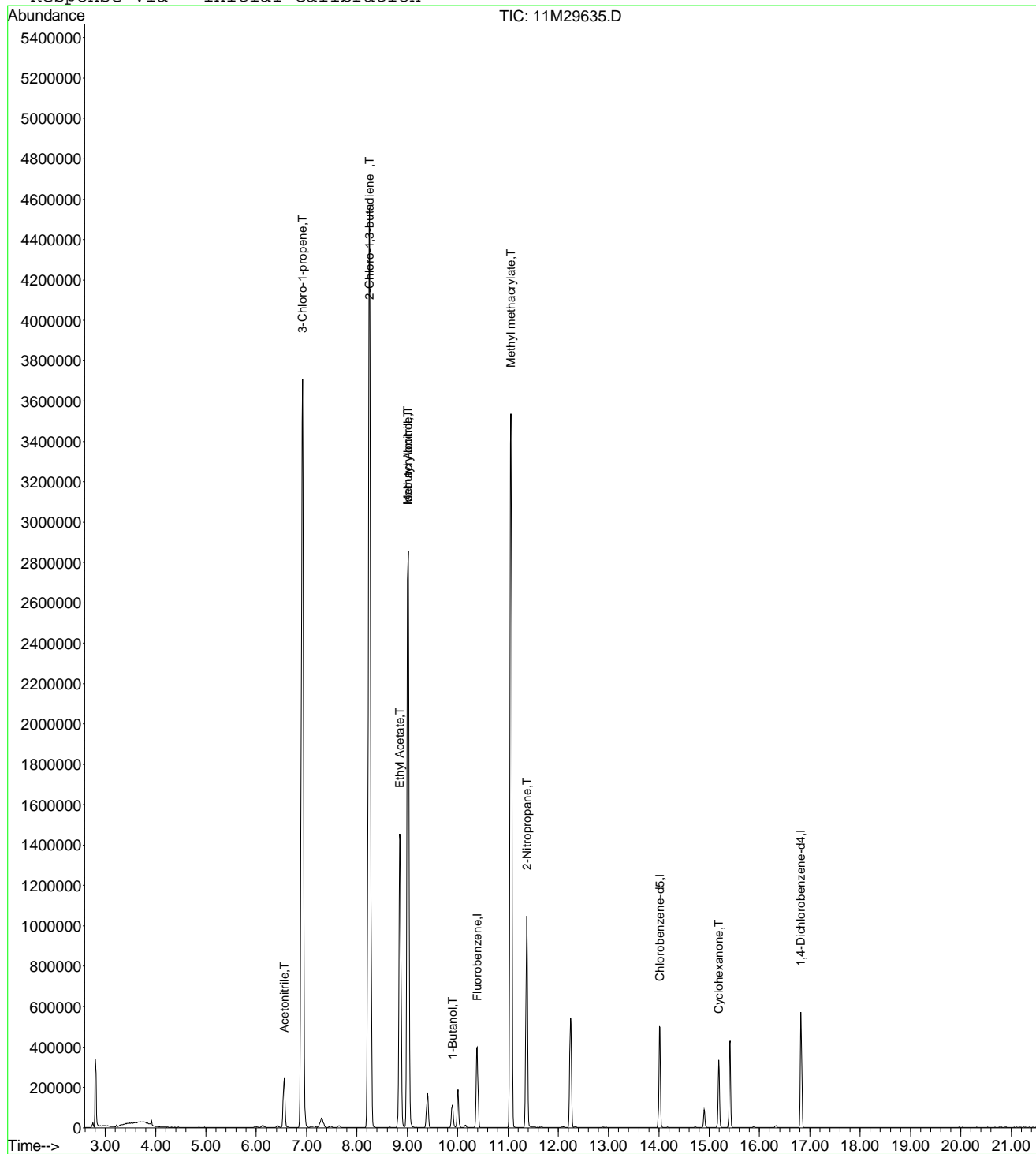
Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29635.D
 Acq On : 19 Feb 2019 16:44
 Sample : WG696624-10 500ug/L ICAL A9 8260
 Misc : 1,1 STD92151
 MS Integration Params: rteint.p
 Quant Time: Feb 20 9:25 2019

Vial: 11
 Operator: KFR
 Inst : hpms11
 Multiplr: 1.00

Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\021919\11M29635.D Vial: 11
 Acq On : 19 Feb 2019 16:44 Operator: KFR
 Sample : WG696624-10 500ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92151 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.01
2 T	Acetonitrile	500.000	482.418	3.5	100	0.01
3 T	3-Chloro-1-propene	-1.000	396.712	0.0	100	0.00
4 T	2-Chloro-1,3-butadiene	-1.000	402.518	0.0	100	0.01
5 T	Methacrylonitrile	500.000	468.991	6.2	100	0.01
6 T	Isobutyl Alcohol	-1.000	1044.248	0.0	100	0.00
7 T	1-Butanol	500.000	554.070	-10.8	100	0.00
8 T	Cyclohexanone	500.000	493.974	1.2	100	0.00
9 T	2-Nitropropane	500.000	525.901	-5.2	100	0.00
10 T	Ethyl Acetate	500.000	453.790	9.2	100	0.00
11 T	Methyl methacrylate	500.000	443.202	11.4	100	0.01
12 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.01
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29635.D A9FOOWT.M Wed Feb 20 09:23:09 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29637.D Vial: 13
 Acq On : 19 Feb 2019 17:43 Operator: KFR
 Sample : WG696624-11 ICV/ALT 100ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92076 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Feb 20 09:25:28 2019 Quant Results File: A9FOOWT.RES

Quant Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

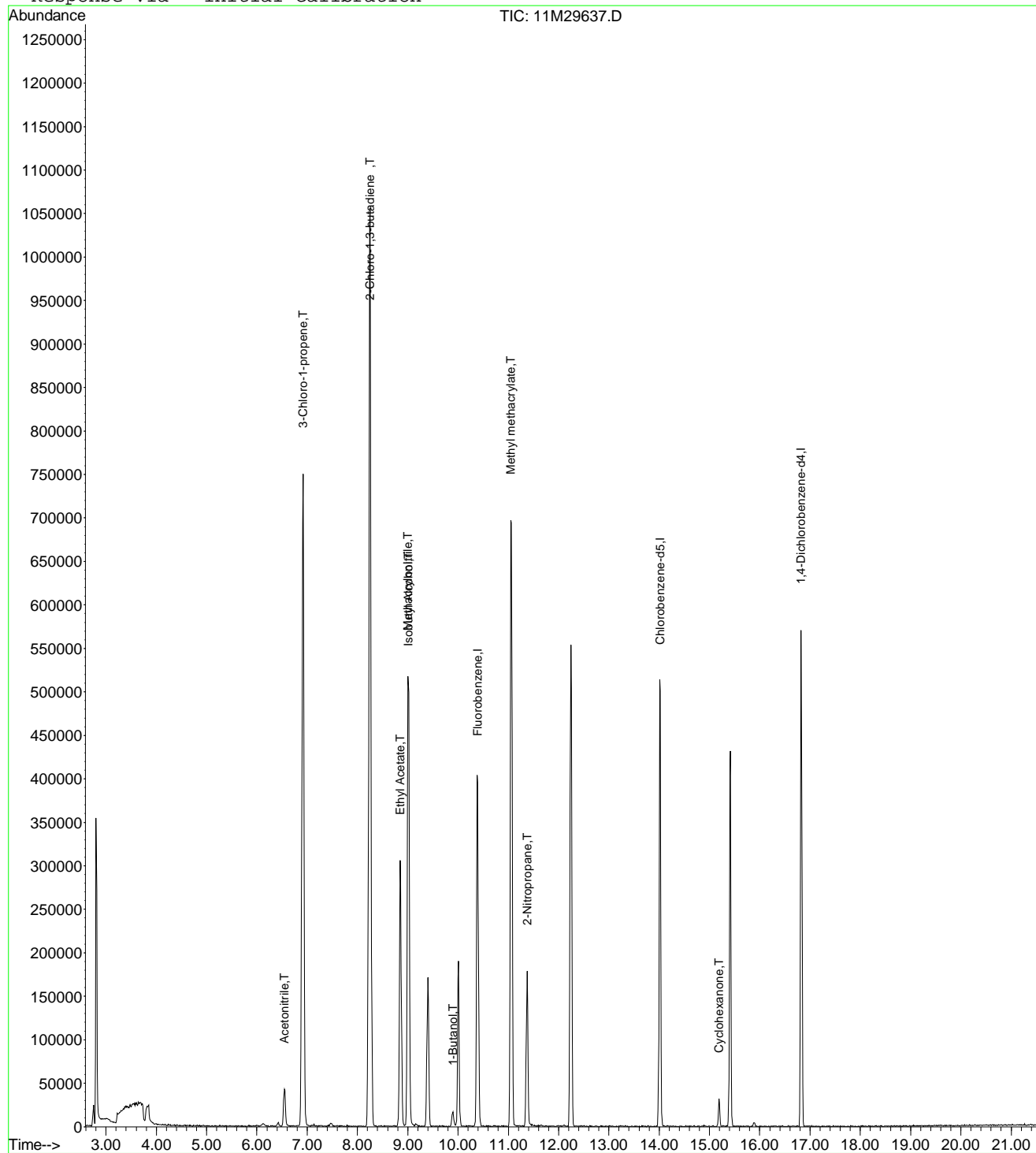
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3914	96	459316	25.0000	ug/L	0.0102
12) Chlorobenzene-d5	14.0206	117	349389	25.0000	ug/L	0.0102
13) 1,4-Dichlorobenzene-d4	16.8226	152	174186	25.0000	ug/L	0.0000
						Qvalue
Target Compounds						
2) Acetonitrile	6.5451	41	58636	85.9392	ug/L	98
3) 3-Chloro-1-propene	6.9173	41	745583	82.3515	ug/L	100
4) 2-Chloro-1,3-butadiene	8.2511	53	1057575	105.6753	ug/L	100
5) Methacrylonitrile	9.0059	41	356343	90.2132	ug/L	100
6) Isobutyl Alcohol	9.0162	43	46312	173.2896	ug/L	94
7) 1-Butanol	9.8951	56	12029	77.5781	ug/L	93
8) Cyclohexanone	15.1890	55	18983	49.1790	ug/L	98
9) 2-Nitropropane	11.3737	43	151551	83.9270	ug/L	100
10) Ethyl Acetate	8.8508	43	496274	93.9411	ug/L	100
11) Methyl methacrylate	11.0531	41	475432	93.8047	ug/L	100

 (#) = qualifier out of range (m) = manual integration
 11M29637.D A9FOOWT.M Wed Feb 20 09:25:28 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\021919\11M29637.D Vial: 13
Acq On : 19 Feb 2019 17:43 Operator: KFR
Sample : WG696624-11 ICV/ALT 100ug/L ICAL A9 8260 Inst : hpms11
Misc : 1,1 STD92076 Multiplr: 1.00
MS Integration Params: rteint.p
Quant Time: Feb 20 9:25 2019 Quant Results File: A9FOOWT.RES

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
Last Update : Wed Feb 20 09:20:16 2019
Response via : Initial Calibration



11M29637.D A9FOOWT.M Wed Feb 20 09:25:28 2019

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Data File : C:\MSDCHEM\1\DATA\021919\11M29637.D Vial: 13
 Acq On : 19 Feb 2019 17:43 Operator: KFR
 Sample : WG696624-11 ICV/ALT 100ug/L ICAL A9 8260 Inst : hpms11
 Misc : 1,1 STD92076 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\A9FOOWT.M (RTE Integrator)
 Title : Appendix IX (SOP:OVL MSV01) Water 021919 HPMS11
 Last Update : Wed Feb 20 09:20:16 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	97	0.01
2 T	Acetonitrile	100.000	85.939	14.1	86	0.00
3 T	3-Chloro-1-propene	100.000	82.352	17.6	76	0.00
4 T	2-Chloro-1,3-butadiene	100.000	105.675	-5.7	94	0.01
5 T	Methacrylonitrile	100.000	90.213	9.8	87	0.00
6 T	Isobutyl Alcohol	200.000	173.290	13.4	89	0.00
7 T	1-Butanol	100.000	77.578	22.4	74	0.00
8 T	Cyclohexanone	100.000	49.179	50.8	49	0.00
9 T	2-Nitropropane	100.000	83.927	16.1	83	0.00
10 T	Ethyl Acetate	100.000	93.941	6.1	88	0.00
11 T	Methyl methacrylate	100.000	93.805	6.2	87	0.00
12 I	Chlorobenzene-d5	25.000	25.000	0.0	97	0.01
13 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	97	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29637.D A9FOOWT.M Wed Feb 20 09:26:02 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29737.D Vial: 2
 Acq On : 5 Mar 2019 16:39 Operator: KFR
 Sample : WG698387-02 STD 0.3ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:19 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	386161	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0206	117	286256	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	138590	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	0.0000	111	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	0.0000%#	
43) 1,2-Dichloroethane-d4	0.0000	65	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	0.0000%#	
57) Toluene-d8	0.0000	98	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	0.0000%#	
78) p-Bromofluorobenzene	0.0000	95	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	0.0000%#	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1227	85	1412	0.2475	ug/L #	62
3) Chloromethane	3.5570	50	4628	0.4224	ug/L	94
4) Vinyl Chloride	3.7845	62	2722	0.3328	ug/L #	1
5) 1,3-Butadiene	3.8258	54	2961	Below Cal		83
6) Bromomethane	4.6633	94	749	0.2390	ug/L	93
8) Trichlorofluoromethane	5.2733	101	2030	0.2623	ug/L #	67
10) Isoprene	5.8213	67	1068	0.1752	ug/L	74
13) Acetone	6.1315	43	452	0.3202	ug/L #	45
14) 1,1-Dichloroethene	6.3487	61	1896	0.2566	ug/L	74
16) Dimethyl Sulfide	6.5968	62	1144	0.2342	ug/L	98
17) Iodomethane	6.8450	142	202	1.5324	ug/L #	29
19) Methylene Chloride	7.0931	84	1255	0.3048	ug/L	96
20) Carbon Disulfide	7.1345	76	4500	0.3965	ug/L #	74
22) Methyl Tert Butyl Ether	7.2999	73	3610	0.3091	ug/L #	48
23) trans-1,2-Dichloroethene	7.5377	96	1045	0.2611	ug/L	69
24) n-Hexane	7.6101	57	1883	0.3054	ug/L #	72
26) Vinyl Acetate	8.0961	43	1638	2.2761	ug/L #	75
27) 1,1-Dichloroethane	8.1374	63	2672	0.3059	ug/L #	52
31) 2,2-Dichloropropane	8.8612	77	1293	0.2141	ug/L #	43
32) cis-1,2-Dichloroethene	8.9336	96	1200	0.2641	ug/L	88
33) Chloroform	9.1300	83	2562	0.3231	ug/L	94
35) Bromochloromethane	9.3575	130	481	0.1805	ug/L #	21
36) Tetrahydrofuran	9.3782	42	1373	1.1785	ug/L #	37
38) 1,1,1-Trichloroethane	9.6160	97	2230	0.3106	ug/L #	35
39) Cyclohexane	9.6573	56	2370	0.2788	ug/L #	91
40) 1,1-Dichloropropene	9.8124	75	1521	0.2818	ug/L #	39
41) Carbon Tetrachloride	9.9572	117	1207	0.1977	ug/L #	77
44) 1,2-Dichloroethane	10.1226	62	1882	0.2731	ug/L #	53
45) Benzene	10.1536	78	5004	0.3119	ug/L	94
46) Trichloroethene	10.8567	130	1328	0.2919	ug/L	92
47) Methylcyclohexane	10.9498	83	1957	0.3053	ug/L #	77
48) 1,2-Dichloropropane	11.0739	63	1291	0.2709	ug/L #	64
50) Bromodichloromethane	11.3530	83	1329	0.2360	ug/L #	89
51) Dibromomethane	11.4358	93	463	0.1701	ug/L #	53
54) cis-1,3-Dichloropropene	11.9527	75	1834	0.2889	ug/L #	65
55) Dimethyl Disulfide	12.1905	79	220	2.4564	ug/L #	1
58) Toluene	12.3456	91	5233	0.3086	ug/L	94
59) Ethyl Methacrylate	12.4283	69	742	0.1549	ug/L	77
60) trans-1,3-Dichloropropene	12.5110	75	1066	0.1920	ug/L #	51
61) 1,1,2-Trichloroethane	12.7179	97	844	0.2338	ug/L	67

(#) = qualifier out of range (m) = manual integration
 11M29737.D 8260WT.M Wed Mar 06 14:03:22 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29737.D Vial: 2
 Acq On : 5 Mar 2019 16:39 Operator: KFR
 Sample : WG698387-02 STD 0.3ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:19 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
63) 1,3-Dichloropropane	12.9970	76	1537	0.2639	ug/L	99
64) Tetrachloroethene	13.1211	164	934	0.2597	ug/L	87
65) Dibromochloromethane	13.3589	129	832	0.2025	ug/L #	38
66) 1,2-Dibromoethane	13.5967	107	883	0.2552	ug/L	94
67) 1-Chlorohexane	13.6691	91	1266	0.2261	ug/L	92
68) Chlorobenzene	14.0723	112	3738	0.3166	ug/L	98
69) 1,1,1,2-Tetrachloroethane	14.0930	131	806	0.8276	ug/L	97
70) Ethylbenzene	14.0827	106	1567	0.2609	ug/L	61
71) m-,p-Xylene	14.1654	106	3991	0.5504	ug/L	83
72) o-Xylene	14.7030	106	1800	0.2559	ug/L	72
73) Styrene	14.7341	104	3323	0.2856	ug/L	77
74) Bromoform	15.1994	173	446	0.8783	ug/L #	29
75) Isopropylbenzene	15.0856	105	5224	0.2966	ug/L	89
77) 1,1,2,2-Tetrachloroethane	15.2924	83	903	0.2498	ug/L	87
81) n-Propylbenzene	15.5612	91	6486	0.3174	ug/L	91
82) Bromobenzene	15.6853	156	1377	0.2759	ug/L	87
83) 1,3,5-Trimethylbenzene	15.7370	105	4065	0.2802	ug/L	100
84) 2-Chlorotoluene	15.8197	91	4532	0.3116	ug/L	95
85) 4-Chlorotoluene	15.8611	91	3558	0.2977	ug/L	94
86) a-Methylstyrene	16.1092	118	2026	0.2452	ug/L	93
87) tert-Butylbenzene	16.1713	134	701	0.1962	ug/L #	17
88) 1,2,4-Trimethylbenzene	16.2126	105	4360	0.2912	ug/L	82
89) sec-Butylbenzene	16.4194	105	5766	0.3201	ug/L #	78
90) p-Isopropyltoluene	16.5642	119	4518	0.2885	ug/L	90
91) 1,3-Dichlorobenzene	16.7503	146	2748	0.2984	ug/L	90
92) 1,4-Dichlorobenzene	16.8744	146	2688	0.2923	ug/L #	34
93) n-Butylbenzene	17.0501	91	4116	0.2916	ug/L #	85
94) 1,2-Dichlorobenzene	17.3293	146	2622	0.2971	ug/L	95
96) 1,2,4-Trichlorobenzene	19.3145	180	1324	0.2319	ug/L #	69
97) Hexachlorobutadiene	19.4593	225	462	0.1606	ug/L #	23
98) Naphthalene	19.6660	128	2997	0.2452	ug/L #	86
99) 1,2,3-Trichlorobenzene	19.9452	180	1235	0.2303	ug/L #	68

(#) = qualifier out of range (m) = manual integration
 11M29737.D 8260WT.M Wed Mar 06 14:03:22 2019

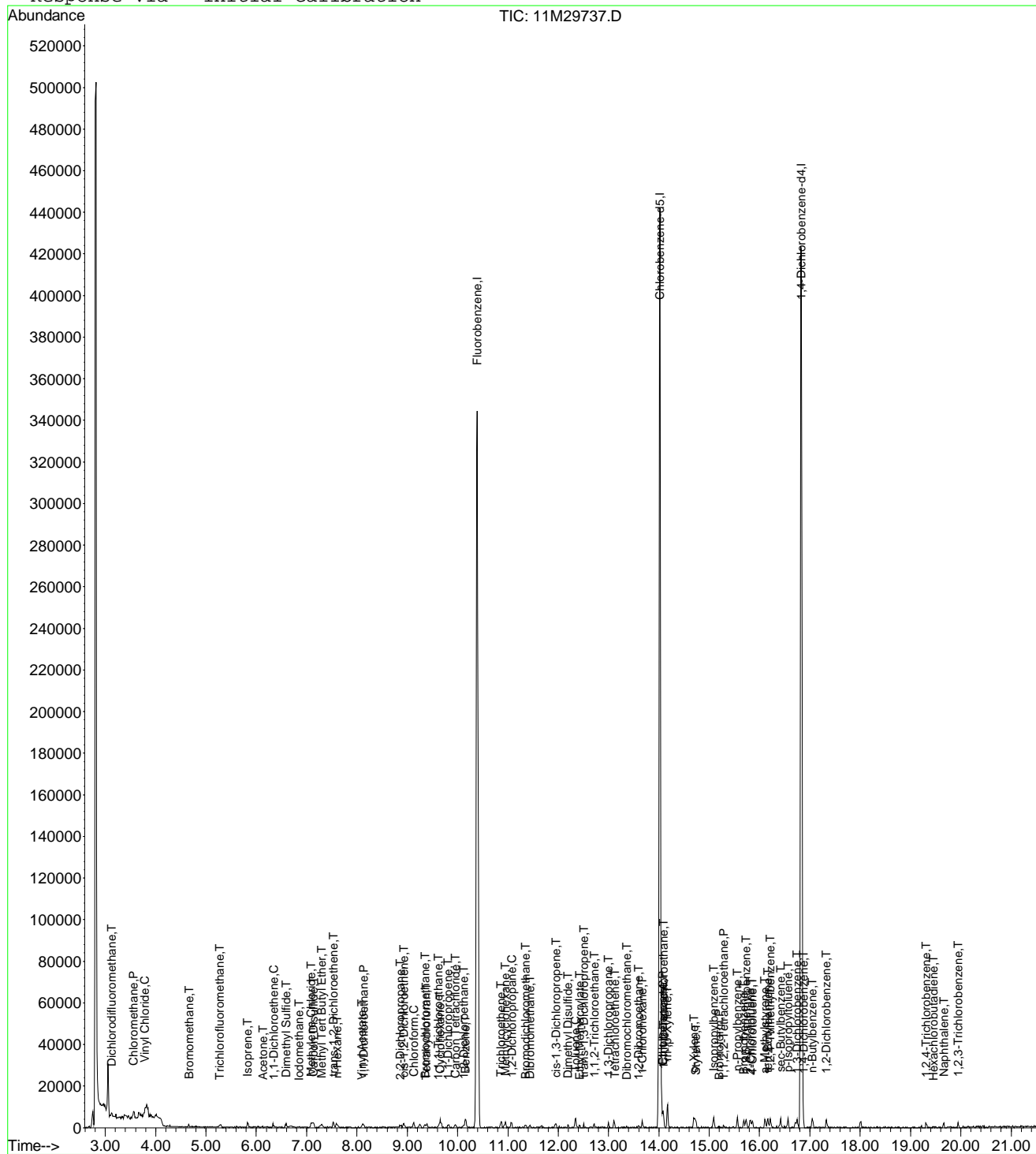
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29737.D
 Acq On : 5 Mar 2019 16:39
 Sample : WG698387-02 STD 0.3ug/L 8260
 Misc : 1,1 STD92320
 MS Integration Params: rteint.p
 Quant Time: Mar 6 14:03 2019

Vial: 2
 Operator: KFR
 Inst : hpms11
 Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29737.D Vial: 2
 Acq On : 5 Mar 2019 16:39 Operator: KFR
 Sample : WG698387-02 STD 0.3ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	-1.000	0.248	0.0	100	0.00
3 P	Chloromethane	-1.000	0.422	0.0	0	0.00
4 C	Vinyl Chloride	-1.000	0.333	0.0	100	0.01
5 T	1,3-Butadiene	-1.000	-4.635	0.0	100	0.01
6 T	Bromomethane	-1.000	0.239	0.0	100	0.02
7 T	Chloroethane	-1.000	0.067	0.0	100	-0.01
8 T	Trichlorofluoromethane	-1.000	0.262	0.0	100	0.00
9 T	Diethyl ether	-1.000	0.000	0.0	0	-5.79#
10 T	Isoprene	-1.000	0.175	0.0	100	0.00
11 T	Acrolein	-1.000	0.000	0.0	0	-6.02#
12 T	1,1,2-Trichloro-1,2,2-Trifl	-1.000	0.061	0.0	100	-0.01
13 T	Acetone	-1.000	0.320	0.0	100	0.01
14 C	1,1-Dichloroethene	-1.000	0.257	0.0	100	0.02
15 T	Tert-Butyl Alcohol	-1.000	0.000	0.0	0	-6.44#
16 T	Dimethyl Sulfide	-1.000	0.234	0.0	100	0.01
17 T	Iodomethane	-1.000	1.532	0.0	0	0.01
18 T	Methyl acetate	-1.000	0.062	0.0	100	0.01
19 T	Methylene Chloride	-1.000	0.305	0.0	100	0.00
20 T	Carbon Disulfide	-1.000	0.396	0.0	100	0.00
21 T	Acrylonitrile	-1.000	0.000	0.0	0	-7.27#
22 T	Methyl Tert Butyl Ether	-1.000	0.309	0.0	100	0.00
23 T	trans-1,2-Dichloroethene	-1.000	0.261	0.0	100	0.01
24 T	n-Hexane	-1.000	0.305	0.0	100	0.01
25 T	Diisopropyl ether	-1.000	0.000	0.0	0	-7.93#
26 T	Vinyl Acetate	-1.000	2.276	0.0	0	0.00
27 P	1,1-Dichloroethane	-1.000	0.306	0.0	100	0.01
28 T	Ethyl-Tert-Butyl ether	-1.000	0.000	0.0	0	-8.48#
29 T	2-Butanone	-1.000	0.000	0.0	0	-8.65#
30 T	Propionitrile	-1.000	0.000	0.0	0	-8.76#
31 T	2,2-Dichloropropane	-1.000	0.214	0.0	100	0.00
32 T	cis-1,2-Dichloroethene	-1.000	0.264	0.0	100	0.00
33 C	Chloroform	-1.000	0.323	0.0	100	0.00
34 T	1-Bromopropane	-1.000	0.000	0.0	0	-9.25#
35 T	Bromochloromethane	-1.000	0.181	0.0	100	0.01
36 T	Tetrahydrofuran	-1.000	1.178	0.0	100	0.01
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.40#
38 T	1,1,1-Trichloroethane	-1.000	0.311	0.0	100	-0.01
39 T	Cyclohexane	-1.000	0.279	0.0	100	0.00
40 T	1,1-Dichloropropene	-1.000	0.282	0.0	100	0.00
41 T	Carbon Tetrachloride	-1.000	0.198	0.0	0	0.00
42 T	Tert-Amyl-Methyl ether	-1.000	0.000	0.0	0	-9.92#
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.01#
44 T	1,2-Dichloroethane	0.300	0.273	9.0	100	0.00
45 T	Benzene	-1.000	0.312	0.0	100	0.00
46 T	Trichloroethene	0.300	0.292	2.7	100	-0.01
47 T	Methylcyclohexane	-1.000	0.305	0.0	100	0.01
48 C	1,2-Dichloropropane	-1.000	0.271	0.0	100	0.01
49 T	1,4-Dioxane	-1.000	0.000	0.0	0	-11.34#
50 T	Bromodichloromethane	-1.000	0.236	0.0	100	0.00
51 T	Dibromomethane	-1.000	0.170	0.0	100	0.00
52 T	2-Chloroethyl Vinyl Ether	-1.000	0.000	0.0	0	-11.62#
53 T	4-Methyl-2-Pentanone	-1.000	0.000	0.0	0	-11.65#
54 T	cis-1,3-Dichloropropene	-1.000	0.289	0.0	100	0.01

(#) = Out of Range

11M29737.D 8260WT.M Wed Mar 06 13:56:07 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29737.D Vial: 2
 Acq On : 5 Mar 2019 16:39 Operator: KFR
 Sample : WG698387-02 STD 0.3ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	-1.000	2.456	0.0	0	0.00
56 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
57 S	Toluene-d8	-1.000	0.000	0.0	0	-12.25#
58 C	Toluene	-1.000	0.309	0.0	0	0.00
59 T	Ethyl Methacrylate	-1.000	0.155	0.0	100	0.00
60 T	trans-1,3-Dichloropropene	-1.000	0.192	0.0	100	0.01
61 T	1,1,2-Trichloroethane	-1.000	0.234	0.0	100	0.01
62 T	2-Hexanone	-1.000	0.000	0.0	0	-12.65#
63 T	1,3-Dichloropropane	0.300	0.264	12.0	100	0.00
64 T	Tetrachloroethene	-1.000	0.272	0.0	100	0.01
65 T	Dibromochloromethane	-1.000	0.202	0.0	0	0.00
66 T	1,2-Dibromoethane	-1.000	0.255	0.0	100	0.00
67 T	1-Chlorohexane	-1.000	0.226	0.0	100	0.00
68 P	Chlorobenzene	0.300	0.317	-5.7	100	0.01
69 T	1,1,1,2-Tetrachloroethane	-1.000	0.828	0.0	0	0.00
70 C	Ethylbenzene	0.300	0.261	13.0	100	0.00
71 T	m-,p-Xylene	0.600	0.550	8.3	100	0.00
72 T	o-Xylene	0.300	0.256	14.7	100	0.01
73 T	Styrene	-1.000	0.286	0.0	100	0.00
74 P	Bromoform	-1.000	0.878	0.0	100	0.00
75 T	Isopropylbenzene	0.300	0.297	1.0	100	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
77 P	1,1,2,2-Tetrachloroethane	-1.000	0.250	0.0	100	0.00
78 S	p-Bromofluorobenzene	-1.000	0.000	0.0	0	-15.42#
79 T	1,2,3-Trichloropropane	-1.000	0.000	0.0	0	-15.47#
80 T	trans-1,4-Dichloro-2-Butene	-1.000	0.000	0.0	0	-15.51#
81 T	n-Propylbenzene	-1.000	0.317	0.0	0	0.00
82 T	Bromobenzene	-1.000	0.276	0.0	100	0.00
83 T	1,3,5-Trimethylbenzene	0.300	0.280	6.7	100	0.00
84 T	2-Chlorotoluene	-1.000	0.312	0.0	100	0.00
85 T	4-Chlorotoluene	-1.000	0.298	0.0	100	0.00
86 T	a-Methylstyrene	-1.000	0.245	0.0	100	0.00
87 T	tert-Butylbenzene	-1.000	0.196	0.0	100	0.00
88 T	1,2,4-Trimethylbenzene	0.300	0.291	3.0	100	0.00
89 T	sec-Butylbenzene	-1.000	0.320	0.0	0	0.00
90 T	p-Isopropyltoluene	0.300	0.288	4.0	100	0.00
91 T	1,3-Dichlorobenzene	0.300	0.298	0.7	100	0.00
92 T	1,4-Dichlorobenzene	0.300	0.292	2.7	100	0.01
93 T	n-Butylbenzene	-1.000	0.292	0.0	0	0.00
94 T	1,2-Dichlorobenzene	0.300	0.297	1.0	100	0.00
95 T	1,2-Dibromo-3-Chloropropane	-1.000	0.000	0.0	0	-18.25#
96 T	1,2,4-Trichlorobenzene	-1.000	0.232	0.0	100	0.00
97 T	Hexachlorobutadiene	-1.000	0.161	0.0	100	0.01
98 T	Naphthalene	-1.000	0.245	0.0	0	0.01
99 T	1,2,3-Trichlorobenzene	-1.000	0.230	0.0	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29737.D 8260WT.M Wed Mar 06 13:56:07 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29738.D Vial: 3
 Acq On : 5 Mar 2019 17:09 Operator: KFR
 Sample : WG698387-03 STD 0.4ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:22 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	377565	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0207	117	278973	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	132375	25.0000	ug/L	0.0000

System Monitoring Compounds

37) Dibromofluoromethane	0.0000	111	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	0.0000%#	
43) 1,2-Dichloroethane-d4	0.0000	65	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	0.0000%#	
57) Toluene-d8	0.0000	98	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	0.0000%#	
78) p-Bromofluorobenzene	0.0000	95	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	0.0000%#	

Target Compounds

					Qvalue	
2) Dichlorodifluoromethane	3.1124	85	2037	0.3652	ug/L #	62
3) Chloromethane	3.5570	50	5445	0.5083	ug/L	86
4) Vinyl Chloride	3.7741	62	3600	0.4502	ug/L #	1
5) 1,3-Butadiene	3.8362	54	3453	Below Cal	#	73
6) Bromomethane	4.6426	94	1237	0.4037	ug/L	87
7) Chloroethane	4.7977	64	1149	0.3716	ug/L #	45
8) Trichlorofluoromethane	5.2734	101	2805	0.3706	ug/L	91
10) Isoprene	5.8110	67	1801	0.3021	ug/L	82
13) Acetone	6.1212	43	615	0.4455	ug/L #	45
14) 1,1-Dichloroethene	6.3280	61	2520	0.3488	ug/L	77
16) Dimethyl Sulfide	6.5762	62	1596	0.3341	ug/L	87
17) Iodomethane	6.8346	142	413	1.5744	ug/L #	29
18) Methyl acetate	6.8450	43	691	0.2321	ug/L #	64
19) Methylene Chloride	7.0931	84	1475	0.3663	ug/L	81
20) Carbon Disulfide	7.1242	76	5151	0.4642	ug/L #	74
22) Methyl Tert Butyl Ether	7.2896	73	4630	0.4055	ug/L #	64
23) trans-1,2-Dichloroethene	7.5274	96	1309	0.3345	ug/L	81
24) n-Hexane	7.5998	57	2206	0.3659	ug/L #	72
26) Vinyl Acetate	8.0857	43	388	2.0546	ug/L #	75
27) 1,1-Dichloroethane	8.1167	63	3396	0.3976	ug/L #	88
29) 2-Butanone	8.6751	43	221	0.1257	ug/L #	54
31) 2,2-Dichloropropane	8.8612	77	2020	0.3421	ug/L #	62
32) cis-1,2-Dichloroethene	8.9232	96	1411	0.3176	ug/L	85
33) Chloroform	9.1197	83	3509	0.4527	ug/L	87
35) Bromochloromethane	9.3471	130	684	0.2625	ug/L #	46
36) Tetrahydrofuran	9.3782	42	1575	1.3826	ug/L #	37
38) 1,1,1-Trichloroethane	9.6263	97	2565	0.3654	ug/L #	82
39) Cyclohexane	9.6677	56	2783	0.3349	ug/L #	84
40) 1,1-Dichloropropene	9.8124	75	1675	0.3174	ug/L #	39
41) Carbon Tetrachloride	9.9572	117	1830	0.3066	ug/L #	84
44) 1,2-Dichloroethane	10.1226	62	2617	0.3884	ug/L #	60
45) Benzene	10.1537	78	6582	0.4196	ug/L	93
46) Trichloroethene	10.8671	130	1557	0.3500	ug/L	91
47) Methylcyclohexane	10.9395	83	2570	0.4101	ug/L #	84
48) 1,2-Dichloropropane	11.0739	63	1631	0.3500	ug/L #	60
50) Bromodichloromethane	11.3530	83	1927	0.3499	ug/L #	64
51) Dibromomethane	11.4254	93	730	0.2743	ug/L #	70
54) cis-1,3-Dichloropropene	11.9527	75	2382	0.3838	ug/L #	58
55) Dimethyl Disulfide	12.2009	79	547	2.5320	ug/L #	32
58) Toluene	12.3353	91	6944	0.4202	ug/L	92

(#) = qualifier out of range (m) = manual integration
 11M29738.D 8260WT.M Wed Mar 06 14:03:24 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29738.D Vial: 3
 Acq On : 5 Mar 2019 17:09 Operator: KFR
 Sample : WG698387-03 STD 0.4ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:22 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
59) Ethyl Methacrylate	12.4180	69	1232	0.2639	ug/L	90
60) trans-1,3-Dichloropropene	12.5111	75	1692	0.3127	ug/L #	51
61) 1,1,2-Trichloroethane	12.7075	97	1624	0.4616	ug/L	75
62) 2-Hexanone	12.6558	43	645	0.2467	ug/L #	13
63) 1,3-Dichloropropane	12.9970	76	2249	0.3962	ug/L	93
64) Tetrachloroethene	13.1108	164	895	0.2553	ug/L	91
65) Dibromochloromethane	13.3589	129	939	0.2345	ug/L	95
66) 1,2-Dibromoethane	13.6071	107	1006	0.2984	ug/L	93
67) 1-Chlorohexane	13.6691	91	1871	0.3429	ug/L	89
68) Chlorobenzene	14.0620	112	4896	0.4255	ug/L	89
69) 1,1,1,2-Tetrachloroethane	14.0930	131	1296	0.9306	ug/L	92
70) Ethylbenzene	14.0827	106	2058	0.3516	ug/L	66
71) m-,p-Xylene	14.1654	106	5214	0.7379	ug/L	80
72) o-Xylene	14.7031	106	2678	0.3906	ug/L	95
73) Styrene	14.7341	104	4225	0.3726	ug/L	95
74) Bromoform	15.1890	173	399	0.8661	ug/L #	29
75) Isopropylbenzene	15.0856	105	7237	0.4217	ug/L	87
77) 1,1,2,2-Tetrachloroethane	15.2924	83	1015	0.2939	ug/L	80
79) 1,2,3-Trichloropropane	15.4682	110	203	0.1761	ug/L #	1
80) trans-1,4-Dichloro-2-Buten	15.5199	53	212	0.1458	ug/L #	1
81) n-Propylbenzene	15.5612	91	8102	0.4150	ug/L	92
82) Bromobenzene	15.6853	156	1841	0.3862	ug/L	84
83) 1,3,5-Trimethylbenzene	15.7267	105	5846	0.4218	ug/L	98
84) 2-Chlorotoluene	15.8197	91	5911	0.4255	ug/L	95
85) 4-Chlorotoluene	15.8611	91	4679	0.4098	ug/L	89
86) a-Methylstyrene	16.1092	118	2822	0.3576	ug/L	77
87) tert-Butylbenzene	16.1713	134	1433	0.4198	ug/L	86
88) 1,2,4-Trimethylbenzene	16.2126	105	5997	0.4193	ug/L	78
89) sec-Butylbenzene	16.4194	105	6824	0.3966	ug/L	98
90) p-Isopropyltoluene	16.5642	119	5834	0.3900	ug/L	85
91) 1,3-Dichlorobenzene	16.7503	146	3553	0.4040	ug/L	100
92) 1,4-Dichlorobenzene	16.8640	146	3543	0.4034	ug/L #	6
93) n-Butylbenzene	17.0501	91	5310	0.3938	ug/L #	85
94) 1,2-Dichlorobenzene	17.3293	146	3212	0.3811	ug/L	97
96) 1,2,4-Trichlorobenzene	19.3145	180	1993	0.3655	ug/L	98
97) Hexachlorobutadiene	19.4386	225	775	0.2820	ug/L #	57
98) Naphthalene	19.6660	128	4804	0.4115	ug/L #	87
99) 1,2,3-Trichlorobenzene	19.9452	180	1868	0.3646	ug/L #	40

(#) = qualifier out of range (m) = manual integration
 11M29738.D 8260WT.M Wed Mar 06 14:03:24 2019

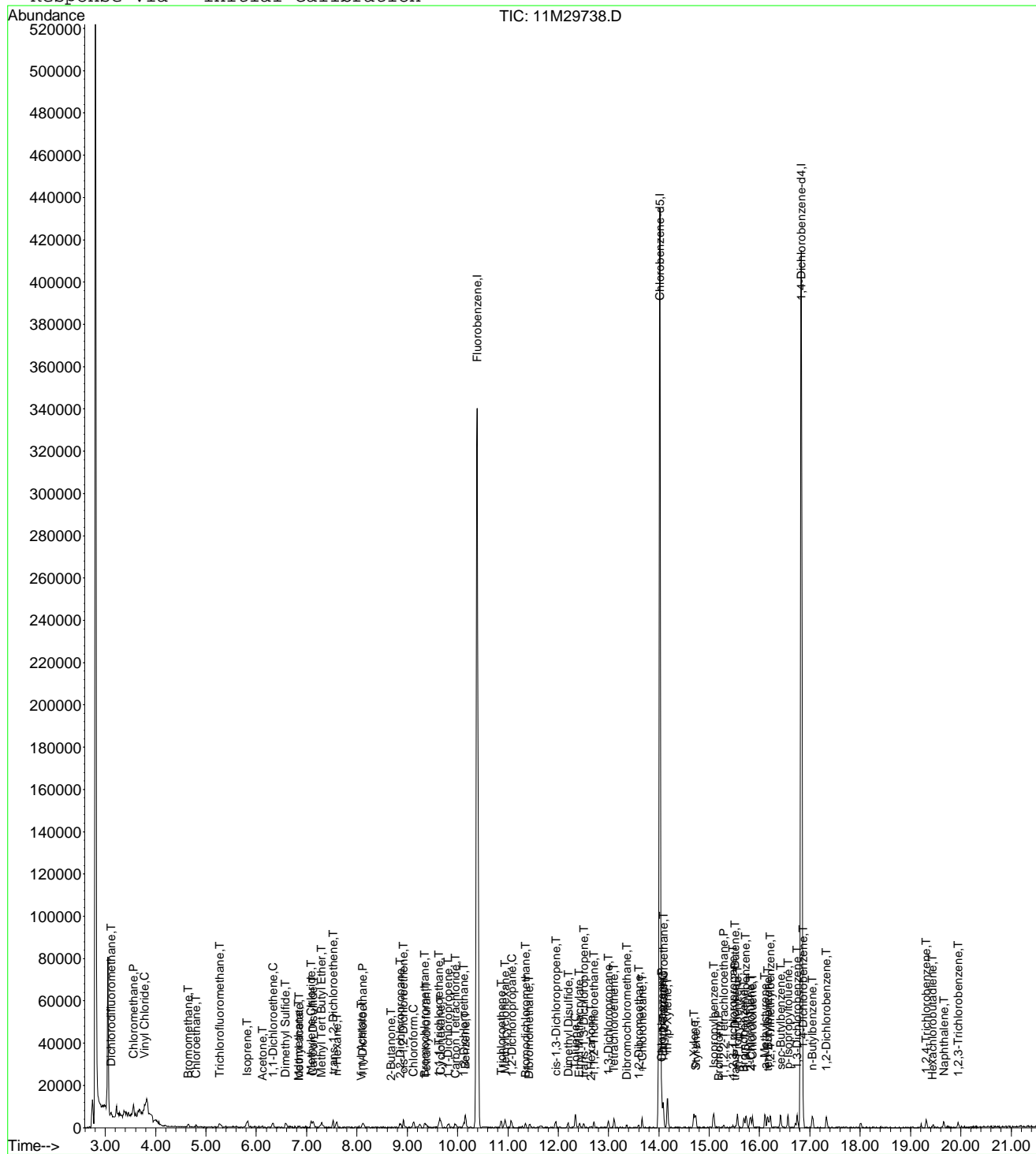
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29738.D
 Acq On : 5 Mar 2019 17:09
 Sample : WG698387-03 STD 0.4ug/L 8260
 Misc : 1,1 STD92320
 MS Integration Params: rteint.p
 Quant Time: Mar 6 14:03 2019

Vial: 3
 Operator: KFR
 Inst : hpms11
 Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29738.D Vial: 3
 Acq On : 5 Mar 2019 17:09 Operator: KFR
 Sample : WG698387-03 STD 0.4ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	-1.000	0.365	0.0	100	-0.01
3 P	Chloromethane	0.400	0.508	-27.0#	100	0.00
4 C	Vinyl Chloride	0.400	0.450	-12.5	100	0.00
5 T	1,3-Butadiene	-1.000	-4.414	0.0	100	0.02
6 T	Bromomethane	-1.000	0.404	0.0	100	0.00
7 T	Chloroethane	-1.000	0.372	0.0	100	0.00
8 T	Trichlorofluoromethane	0.400	0.371	7.3	100	0.00
9 T	Diethyl ether	-1.000	0.000	0.0	0	-5.79#
10 T	Isoprene	-1.000	0.302	0.0	100	-0.01
11 T	Acrolein	-1.000	0.000	0.0	0	-6.02#
12 T	1,1,2-Trichloro-1,2,2-Trifl	-1.000	0.060	0.0	100	0.01
13 T	Acetone	-1.000	0.446	0.0	100	0.00
14 C	1,1-Dichloroethene	0.400	0.349	12.8	100	0.00
15 T	Tert-Butyl Alcohol	-1.000	0.000	0.0	0	-6.44#
16 T	Dimethyl Sulfide	-1.000	0.334	0.0	100	-0.01
17 T	Iodomethane	-1.000	1.574	0.0	0	0.00
18 T	Methyl acetate	-1.000	0.232	0.0	100	0.00
19 T	Methylene Chloride	-1.000	0.366	0.0	100	0.00
20 T	Carbon Disulfide	-1.000	0.464	0.0	100	-0.01
21 T	Acrylonitrile	-1.000	0.000	0.0	0	-7.27#
22 T	Methyl Tert Butyl Ether	0.400	0.406	-1.5	100	-0.01
23 T	trans-1,2-Dichloroethene	0.400	0.334	16.5	100	0.00
24 T	n-Hexane	-1.000	0.366	0.0	100	0.00
25 T	Diisopropyl ether	-1.000	0.000	0.0	0	-7.93#
26 T	Vinyl Acetate	-1.000	2.055	0.0	0	-0.01
27 P	1,1-Dichloroethane	0.400	0.398	0.5	100	-0.01
28 T	Ethyl-Tert-Butyl ether	-1.000	0.000	0.0	0	-8.48#
29 T	2-Butanone	-1.000	0.126	0.0	100	0.02
30 T	Propionitrile	-1.000	0.000	0.0	0	-8.76#
31 T	2,2-Dichloropropane	-1.000	0.342	0.0	100	0.00
32 T	cis-1,2-Dichloroethene	0.400	0.318	20.5	100	-0.01
33 C	Chloroform	0.400	0.453	-13.2	100	-0.01
34 T	1-Bromopropane	-1.000	0.000	0.0	0	-9.25#
35 T	Bromochloromethane	-1.000	0.263	0.0	100	0.00
36 T	Tetrahydrofuran	-1.000	1.383	0.0	100	0.01
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.40#
38 T	1,1,1-Trichloroethane	0.400	0.365	8.8	100	0.00
39 T	Cyclohexane	-1.000	0.335	0.0	100	0.01
40 T	1,1-Dichloropropene	-1.000	0.317	0.0	100	0.00
41 T	Carbon Tetrachloride	-1.000	0.307	0.0	0	0.00
42 T	Tert-Amyl-Methyl ether	-1.000	0.000	0.0	0	-9.92#
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.01#
44 T	1,2-Dichloroethane	0.400	0.388	3.0	100	0.00
45 T	Benzene	0.400	0.420	-5.0	100	0.00
46 T	Trichloroethene	0.400	0.350	12.5	100	0.00
47 T	Methylcyclohexane	-1.000	0.410	0.0	100	0.00
48 C	1,2-Dichloropropane	0.400	0.350	12.5	100	0.01
49 T	1,4-Dioxane	-1.000	0.000	0.0	0	-11.34#
50 T	Bromodichloromethane	0.400	0.350	12.5	100	0.00
51 T	Dibromomethane	-1.000	0.274	0.0	100	-0.01
52 T	2-Chloroethyl Vinyl Ether	-1.000	0.000	0.0	0	-11.62#
53 T	4-Methyl-2-Pentanone	-1.000	0.118	0.0	100	0.01
54 T	cis-1,3-Dichloropropene	0.400	0.384	4.0	100	0.01

(#) = Out of Range

11M29738.D 8260WT.M

Wed Mar 06 14:02:13 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29738.D Vial: 3
 Acq On : 5 Mar 2019 17:09 Operator: KFR
 Sample : WG698387-03 STD 0.4ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	-1.000	2.532	0.0	0	0.01
56 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
57 S	Toluene-d8	-1.000	0.000	0.0	0	-12.25#
58 C	Toluene	0.400	0.420	-5.0	100	0.00
59 T	Ethyl Methacrylate	-1.000	0.264	0.0	100	-0.01
60 T	trans-1,3-Dichloropropene	-1.000	0.313	0.0	100	0.01
61 T	1,1,2-Trichloroethane	0.400	0.462	-15.5	100	0.00
62 T	2-Hexanone	-1.000	0.247	0.0	100	0.01
63 T	1,3-Dichloropropane	0.400	0.396	1.0	100	0.00
64 T	Tetrachloroethene	-1.000	0.267	0.0	0	0.00
65 T	Dibromochloromethane	-1.000	0.234	0.0	0	0.00
66 T	1,2-Dibromoethane	-1.000	0.298	0.0	100	0.01
67 T	1-Chlorohexane	-1.000	0.343	0.0	100	0.00
68 P	Chlorobenzene	0.400	0.426	-6.5	100	0.00
69 T	1,1,1,2-Tetrachloroethane	-1.000	0.931	0.0	0	0.00
70 C	Ethylbenzene	0.400	0.352	12.0	100	0.00
71 T	m-,p-Xylene	0.800	0.738	7.8	100	0.00
72 T	o-Xylene	0.400	0.391	2.3	100	0.01
73 T	Styrene	0.400	0.373	6.8	100	0.00
74 P	Bromoform	-1.000	0.866	0.0	100	0.00
75 T	Isopropylbenzene	0.400	0.422	-5.5	100	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
77 P	1,1,2,2-Tetrachloroethane	0.400	0.294	26.5#	100	0.00
78 S	p-Bromofluorobenzene	-1.000	0.000	0.0	0	-15.42#
79 T	1,2,3-Trichloropropane	-1.000	0.176	0.0	100	0.00
80 T	trans-1,4-Dichloro-2-Butene	-1.000	0.146	0.0	100	0.01
81 T	n-Propylbenzene	0.400	0.415	-3.7	100	0.00
82 T	Bromobenzene	0.400	0.386	3.5	100	0.00
83 T	1,3,5-Trimethylbenzene	0.400	0.422	-5.5	100	-0.01
84 T	2-Chlorotoluene	0.400	0.426	-6.5	100	0.00
85 T	4-Chlorotoluene	0.400	0.410	-2.5	100	0.00
86 T	a-Methylstyrene	-1.000	0.358	0.0	100	0.00
87 T	tert-Butylbenzene	-1.000	0.420	0.0	100	0.00
88 T	1,2,4-Trimethylbenzene	0.400	0.419	-4.7	100	0.00
89 T	sec-Butylbenzene	0.400	0.397	0.8	100	0.00
90 T	p-Isopropyltoluene	0.400	0.390	2.5	100	0.00
91 T	1,3-Dichlorobenzene	0.400	0.404	-1.0	100	0.00
92 T	1,4-Dichlorobenzene	0.400	0.403	-0.8	100	0.00
93 T	n-Butylbenzene	0.400	0.394	1.5	100	0.00
94 T	1,2-Dichlorobenzene	0.400	0.381	4.8	100	0.00
95 T	1,2-Dibromo-3-Chloropropane	-1.000	0.000	0.0	0	-18.25#
96 T	1,2,4-Trichlorobenzene	-1.000	0.366	0.0	100	0.00
97 T	Hexachlorobutadiene	-1.000	0.282	0.0	100	-0.01
98 T	Naphthalene	0.400	0.411	-2.7	100	0.01
99 T	1,2,3-Trichlorobenzene	0.400	0.365	8.8	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29738.D 8260WT.M Wed Mar 06 14:02:13 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29739.D Vial: 4
 Acq On : 5 Mar 2019 17:38 Operator: KFR
 Sample : WG698387-04 STD lug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:24 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3914	96	384700	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0206	117	284086	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	139198	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	0.0000	111	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	0.0000%#	
43) 1,2-Dichloroethane-d4	0.0000	65	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	0.0000%#	
57) Toluene-d8	0.0000	98	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	0.0000%#	
78) p-Bromofluorobenzene	0.0000	95	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	0.0000%#	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1123	85	5353	0.9419	ug/L	98
3) Chloromethane	3.5569	50	11863	1.0868	ug/L	99
4) Vinyl Chloride	3.7741	62	8135	0.9983	ug/L #	37
5) 1,3-Butadiene	3.8258	54	8383	Below Cal		88
6) Bromomethane	4.6426	94	3541	1.1343	ug/L	94
7) Chloroethane	4.7874	64	3156	1.0019	ug/L #	68
8) Trichlorofluoromethane	5.2836	101	7392	0.9586	ug/L	98
9) Diethyl ether	5.7800	59	20415	5.0640	ug/L	99
10) Isoprene	5.8213	67	5610	0.9237	ug/L	87
11) Acrolein	6.0178	56	1845	2.6355	ug/L	83
12) 1,1,2-Trichloro-1,2,2-Trif	6.0281	101	3450	0.9121	ug/L	100
13) Acetone	6.1108	43	1412	1.0040	ug/L #	45
14) 1,1-Dichloroethene	6.3280	61	7652	1.0394	ug/L	90
15) Tert-Butyl Alcohol	6.4314	59	2669	6.8036	ug/L #	72
16) Dimethyl Sulfide	6.5864	62	4585	0.9421	ug/L	96
17) Iodomethane	6.8346	142	1869	1.8517	ug/L	96
18) Methyl acetate	6.8553	43	2719	0.8963	ug/L #	64
19) Methylene Chloride	7.0931	84	3907	0.9524	ug/L	82
20) Carbon Disulfide	7.1344	76	10930	0.9666	ug/L	96
21) Acrylonitrile	7.2688	53	3210	2.0704	ug/L	78
22) Methyl Tert Butyl Ether	7.3102	73	11394	0.9794	ug/L	89
23) trans-1,2-Dichloroethene	7.5273	96	3681	0.9231	ug/L	96
24) n-Hexane	7.5997	57	5259	0.8562	ug/L #	72
25) Diisopropyl ether	7.9306	45	78591	5.2425	ug/L	98
26) Vinyl Acetate	8.1064	43	1790	2.3044	ug/L #	75
27) 1,1-Dichloroethane	8.1270	63	8573	0.9852	ug/L	96
28) Ethyl-Tert-Butyl ether	8.4786	59	78527	5.1825	ug/L	99
29) 2-Butanone	8.6647	43	1260	0.7033	ug/L #	54
30) Propionitrile	8.7577	54	2233	4.2662	ug/L #	61
31) 2,2-Dichloropropane	8.8612	77	5262	0.8746	ug/L	95
32) cis-1,2-Dichloroethene	8.9335	96	4374	0.9661	ug/L	89
33) Chloroform	9.1300	83	7424	0.9399	ug/L	97
34) 1-Bromopropane	9.2644	122	437	0.4456	ug/L	45
35) Bromochloromethane	9.3471	130	1899	0.7154	ug/L	68
36) Tetrahydrofuran	9.3781	42	7056	6.0794	ug/L	98
38) 1,1,1-Trichloroethane	9.6263	97	6335	0.8858	ug/L #	93
39) Cyclohexane	9.6573	56	8321	0.9827	ug/L	98
40) 1,1-Dichloropropene	9.8124	75	4731	0.8798	ug/L	95
41) Carbon Tetrachloride	9.9571	117	4680	0.7695	ug/L	89
42) Tert-Amyl-Methyl ether	9.9158	73	56129	4.9665	ug/L	99

(#) = qualifier out of range (m) = manual integration
 11M29739.D 8260WT.M Wed Mar 06 14:03:25 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29739.D Vial: 4
 Acq On : 5 Mar 2019 17:38 Operator: KFR
 Sample : WG698387-04 STD lug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:24 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1226	62	6571	0.9572	ug/L	94
45) Benzene	10.1639	78	15399	0.9635	ug/L	94
46) Trichloroethene	10.8670	130	4299	0.9484	ug/L	95
47) Methylcyclohexane	10.9497	83	5922	0.9274	ug/L	96
48) 1,2-Dichloropropane	11.0738	63	4564	0.9613	ug/L	96
50) Bromodichloromethane	11.3530	83	4854	0.8651	ug/L #	96
51) Dibromomethane	11.4357	93	2301	0.8485	ug/L	92
52) 2-Chloroethyl Vinyl Ether	11.6218	63	1169	0.5088	ug/L #	52
53) 4-Methyl-2-Pentanone	11.6735	58	1053	0.6433	ug/L #	60
54) cis-1,3-Dichloropropene	11.9423	75	5466	0.8644	ug/L	99
55) Dimethyl Disulfide	12.1905	79	1853	2.8216	ug/L	82
58) Toluene	12.3456	91	16901	1.0043	ug/L	91
59) Ethyl Methacrylate	12.4283	69	3811	0.8018	ug/L	96
60) trans-1,3-Dichloropropene	12.5007	75	4648	0.8436	ug/L	100
61) 1,1,2-Trichloroethane	12.7178	97	3306	0.9228	ug/L	90
62) 2-Hexanone	12.6454	43	2125	0.7980	ug/L #	62
63) 1,3-Dichloropropane	12.9970	76	6082	1.0522	ug/L	96
64) Tetrachloroethene	13.1004	164	3452	0.9671	ug/L	99
65) Dibromochloromethane	13.3589	129	2960	0.7258	ug/L	96
66) 1,2-Dibromoethane	13.5967	107	3256	0.9483	ug/L	90
67) 1-Chlorohexane	13.6691	91	5020	0.9035	ug/L	95
68) Chlorobenzene	14.0620	112	11177	0.9540	ug/L	96
69) 1,1,1,2-Tetrachloroethane	14.0930	131	3601	1.3826	ug/L	99
70) Ethylbenzene	14.0930	106	5651	0.9481	ug/L	85
71) m-,p-Xylene	14.1654	106	13797	1.9174	ug/L	91
72) o-Xylene	14.6927	106	6415	0.9188	ug/L	80
73) Styrene	14.7340	104	10433	0.9034	ug/L	99
74) Bromoform	15.1993	173	1683	1.2940	ug/L	87
75) Isopropylbenzene	15.0856	105	17441	0.9979	ug/L	94
77) 1,1,2,2-Tetrachloroethane	15.2924	83	3484	0.9594	ug/L	100
79) 1,2,3-Trichloropropane	15.4681	110	947	0.7810	ug/L	58
80) trans-1,4-Dichloro-2-Buten	15.5198	53	1289	0.8432	ug/L #	91
81) n-Propylbenzene	15.5612	91	20991	1.0226	ug/L	94
82) Bromobenzene	15.6956	156	4704	0.9383	ug/L	97
83) 1,3,5-Trimethylbenzene	15.7370	105	14295	0.9809	ug/L	100
84) 2-Chlorotoluene	15.8197	91	14437	0.9884	ug/L	95
85) 4-Chlorotoluene	15.8610	91	12567	1.0468	ug/L	91
86) a-Methylstyrene	16.1092	118	7163	0.8632	ug/L	85
87) tert-Butylbenzene	16.1712	134	3458	0.9634	ug/L	89
88) 1,2,4-Trimethylbenzene	16.2126	105	15075	1.0023	ug/L	98
89) sec-Butylbenzene	16.4194	105	17862	0.9873	ug/L	99
90) p-Isopropyltoluene	16.5641	119	16110	1.0241	ug/L	93
91) 1,3-Dichlorobenzene	16.7502	146	8551	0.9246	ug/L	99
92) 1,4-Dichlorobenzene	16.8640	146	8959	0.9701	ug/L #	66
93) n-Butylbenzene	17.0501	91	13479	0.9507	ug/L	97
94) 1,2-Dichlorobenzene	17.3292	146	8675	0.9788	ug/L	97
95) 1,2-Dibromo-3-Chloropropan	18.2495	75	206	0.2746	ug/L	92
96) 1,2,4-Trichlorobenzene	19.3145	180	4843	0.8447	ug/L	94
97) Hexachlorobutadiene	19.4489	225	2172	0.7516	ug/L	94
98) Naphthalene	19.6660	128	11494	0.9362	ug/L #	93
99) 1,2,3-Trichlorobenzene	19.9555	180	5087	0.9443	ug/L	90

(#) = qualifier out of range (m) = manual integration
 11M29739.D 8260WT.M Wed Mar 06 14:03:26 2019

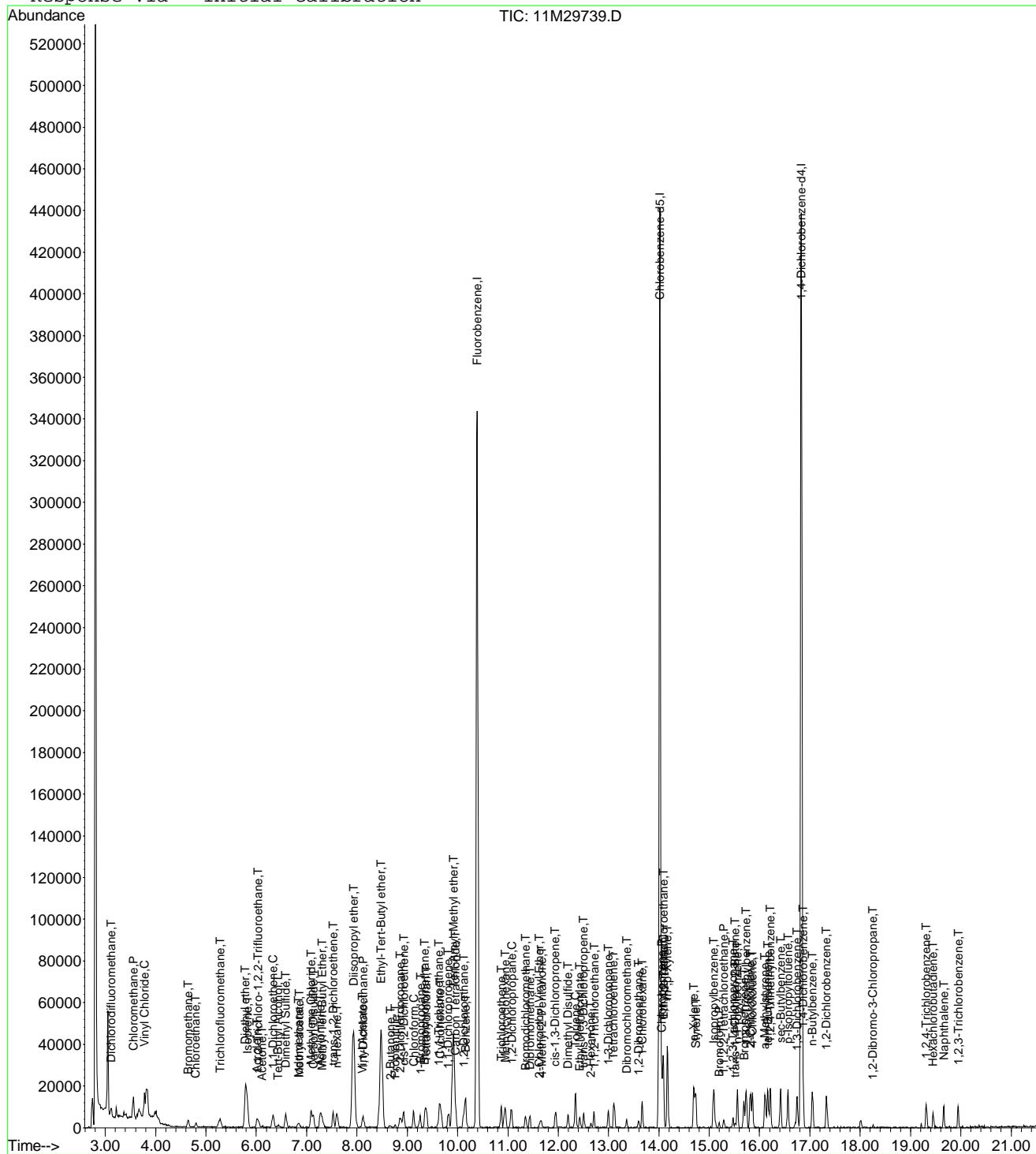
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29739.D
Acq On : 5 Mar 2019 17:38
Sample : WG698387-04 STD Iug/L 8260
Misc : 1,1 STD92320
MS Integration Params: rteint.p
Quant Time: Mar 6 14:03 2019

Vial: 4
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:00:17 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29739.D Vial: 4
 Acq On : 5 Mar 2019 17:38 Operator: KFR
 Sample : WG698387-04 STD lug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	1.000	0.942	5.8	100	-0.01
3 P	Chloromethane	1.000	1.087	-8.7	100	0.00
4 C	Vinyl Chloride	1.000	0.998	0.2	100	0.00
5 T	1,3-Butadiene	-1.000	-2.522	0.0	100	0.01
6 T	Bromomethane	1.000	1.134	-13.4	100	0.00
7 T	Chloroethane	1.000	1.002	-0.2	100	-0.01
8 T	Trichlorofluoromethane	1.000	0.959	4.1	100	0.01
9 T	Diethyl ether	5.000	5.064	-1.3	100	-0.01
10 T	Isoprene	1.000	0.924	7.6	100	0.00
11 T	Acrolein	-1.000	2.635	0.0	100	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	1.000	0.912	8.8	100	-0.01
13 T	Acetone	-1.000	1.004	0.0	100	-0.01
14 C	1,1-Dichloroethene	1.000	1.039	-3.9	100	0.00
15 T	Tert-Butyl Alcohol	-1.000	6.804	0.0	100	-0.01
16 T	Dimethyl Sulfide	1.000	0.942	5.8	100	0.00
17 T	Iodomethane	-1.000	1.852	0.0	0	0.00
18 T	Methyl acetate	-1.000	0.896	0.0	100	0.01
19 T	Methylene Chloride	1.000	0.952	4.8	100	0.00
20 T	Carbon Disulfide	1.000	0.967	3.3	100	0.00
21 T	Acrylonitrile	2.500	2.070	17.2	100	0.00
22 T	Methyl Tert Butyl Ether	1.000	0.979	2.1	100	0.01
23 T	trans-1,2-Dichloroethene	1.000	0.923	7.7	100	0.00
24 T	n-Hexane	-1.000	0.856	0.0	100	0.00
25 T	Diisopropyl ether	5.000	5.242	-4.8	100	0.00
26 T	Vinyl Acetate	-1.000	2.304	0.0	0	0.01
27 P	1,1-Dichloroethane	1.000	0.985	1.5	100	0.00
28 T	Ethyl-Tert-Butyl ether	5.000	5.182	-3.6	100	0.00
29 T	2-Butanone	-1.000	0.703	0.0	100	0.01
30 T	Propionitrile	5.000	4.266	14.7	100	0.00
31 T	2,2-Dichloropropane	1.000	0.875	12.5	100	0.00
32 T	cis-1,2-Dichloroethene	1.000	0.966	3.4	100	0.00
33 C	Chloroform	1.000	0.940	6.0	100	0.00
34 T	1-Bromopropane	-1.000	0.446	0.0	100	0.01
35 T	Bromochloromethane	1.000	0.715	28.5#	100	0.00
36 T	Tetrahydrofuran	5.000	6.079	-21.6	100	0.01
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.40#
38 T	1,1,1-Trichloroethane	1.000	0.886	11.4	100	0.00
39 T	Cyclohexane	1.000	0.983	1.7	100	0.00
40 T	1,1-Dichloropropene	1.000	0.880	12.0	100	0.00
41 T	Carbon Tetrachloride	1.000	0.770	23.0	100	0.00
42 T	Tert-Amyl-Methyl ether	5.000	4.967	0.7	100	0.00
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.01#
44 T	1,2-Dichloroethane	1.000	0.957	4.3	100	0.00
45 T	Benzene	1.000	0.963	3.7	100	0.01
46 T	Trichloroethene	1.000	0.948	5.2	100	0.00
47 T	Methylcyclohexane	1.000	0.927	7.3	100	0.01
48 C	1,2-Dichloropropane	1.000	0.961	3.9	100	0.01
49 T	1,4-Dioxane	-1.000	0.000	0.0	0	-11.34#
50 T	Bromodichloromethane	1.000	0.865	13.5	100	0.00
51 T	Dibromomethane	1.000	0.848	15.2	100	0.00
52 T	2-Chloroethyl Vinyl Ether	-1.000	0.509	0.0	0	0.00
53 T	4-Methyl-2-Pentanone	-1.000	0.643	0.0	100	0.02
54 T	cis-1,3-Dichloropropene	1.000	0.864	13.6	100	0.00

(#) = Out of Range

11M29739.D 8260WT.M Wed Mar 06 13:57:31 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29739.D Vial: 4
 Acq On : 5 Mar 2019 17:38 Operator: KFR
 Sample : WG698387-04 STD lug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	-1.000	2.822	0.0	0	0.00
56 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
57 S	Toluene-d8	-1.000	0.000	0.0	0	-12.25#
58 C	Toluene	1.000	1.004	-0.4	100	0.00
59 T	Ethyl Methacrylate	1.000	0.802	19.8	100	0.00
60 T	trans-1,3-Dichloropropene	1.000	0.844	15.6	100	0.00
61 T	1,1,2-Trichloroethane	1.000	0.923	7.7	100	0.01
62 T	2-Hexanone	-1.000	0.798	0.0	100	0.00
63 T	1,3-Dichloropropane	1.000	1.052	-5.2	100	0.00
64 T	Tetrachloroethene	1.000	1.013	-1.3	100	-0.01
65 T	Dibromochloromethane	-1.000	0.726	0.0	0	0.00
66 T	1,2-Dibromoethane	1.000	0.948	5.2	100	0.00
67 T	1-Chlorohexane	1.000	0.904	9.6	100	0.00
68 P	Chlorobenzene	1.000	0.954	4.6	100	0.00
69 T	1,1,1,2-Tetrachloroethane	-1.000	1.383	0.0	0	0.00
70 C	Ethylbenzene	1.000	0.948	5.2	100	0.01
71 T	m-,p-Xylene	2.000	1.917	4.1	100	0.00
72 T	o-Xylene	1.000	0.919	8.1	100	0.00
73 T	Styrene	1.000	0.903	9.7	100	0.00
74 P	Bromoform	1.000	1.294	-29.4#	100	0.00
75 T	Isopropylbenzene	1.000	0.998	0.2	100	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
77 P	1,1,2,2-Tetrachloroethane	1.000	0.959	4.1	100	0.00
78 S	p-Bromofluorobenzene	-1.000	0.000	0.0	0	-15.42#
79 T	1,2,3-Trichloropropane	1.000	0.781	21.9	100	0.00
80 T	trans-1,4-Dichloro-2-Butene	1.000	0.843	15.7	100	0.01
81 T	n-Propylbenzene	1.000	1.023	-2.3	100	0.00
82 T	Bromobenzene	1.000	0.938	6.2	100	0.01
83 T	1,3,5-Trimethylbenzene	1.000	0.981	1.9	100	0.00
84 T	2-Chlorotoluene	1.000	0.988	1.2	100	0.00
85 T	4-Chlorotoluene	1.000	1.047	-4.7	100	0.00
86 T	a-Methylstyrene	1.000	0.863	13.7	100	0.00
87 T	tert-Butylbenzene	1.000	0.963	3.7	100	0.00
88 T	1,2,4-Trimethylbenzene	1.000	1.002	-0.2	100	0.00
89 T	sec-Butylbenzene	1.000	0.987	1.3	100	0.00
90 T	p-Isopropyltoluene	1.000	1.024	-2.4	100	0.00
91 T	1,3-Dichlorobenzene	1.000	0.925	7.5	100	0.00
92 T	1,4-Dichlorobenzene	1.000	0.970	3.0	100	0.00
93 T	n-Butylbenzene	1.000	0.951	4.9	100	0.00
94 T	1,2-Dichlorobenzene	1.000	0.979	2.1	100	0.00
95 T	1,2-Dibromo-3-Chloropropane	-1.000	0.275	0.0	0	0.00
96 T	1,2,4-Trichlorobenzene	1.000	0.845	15.5	100	0.00
97 T	Hexachlorobutadiene	1.000	0.752	24.8	100	0.00
98 T	Naphthalene	1.000	0.936	6.4	100	0.01
99 T	1,2,3-Trichlorobenzene	1.000	0.944	5.6	100	0.01

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29739.D 8260WT.M Wed Mar 06 13:57:32 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29740.D Vial: 5
 Acq On : 5 Mar 2019 18:08 Operator: KFR
 Sample : WG698387-05 STD 2ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:28 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3811	96	378991	25.0000	ug/L	-0.0104
56) Chlorobenzene-d5	14.0206	117	282579	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	138944	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	0.0000	111	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	0.0000%#	
43) 1,2-Dichloroethane-d4	0.0000	65	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	0.0000%#	
57) Toluene-d8	0.0000	98	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	0.0000%#	
78) p-Bromofluorobenzene	0.0000	95	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	0.0000%#	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1124	85	10625	1.8977	ug/L	97
3) Chloromethane	3.5466	50	22686	2.1097	ug/L	100
4) Vinyl Chloride	3.7741	62	16908	2.1063	ug/L #	66
5) 1,3-Butadiene	3.8258	54	15068	0.1664	ug/L	93
6) Bromomethane	4.6426	94	6619	2.1521	ug/L	96
7) Chloroethane	4.7977	64	5844	1.8831	ug/L	87
8) Trichlorofluoromethane	5.2733	101	15552	2.0472	ug/L	97
9) Diethyl ether	5.7800	59	81473	20.5141	ug/L	99
10) Isoprene	5.8213	67	11094	1.8542	ug/L	99
11) Acrolein	6.0074	56	6790	9.8451	ug/L	99
12) 1,1,2-Trichloro-1,2,2-Trif	6.0281	101	7157	1.9206	ug/L	93
13) Acetone	6.1212	43	3297	2.3795	ug/L #	67
14) 1,1-Dichloroethene	6.3383	61	13742	1.8948	ug/L	99
15) Tert-Butyl Alcohol	6.4314	59	13890	35.9405	ug/L #	95
16) Dimethyl Sulfide	6.5864	62	8647	1.8036	ug/L	99
17) Iodomethane	6.8346	142	5095	2.4842	ug/L	87
18) Methyl acetate	6.8449	43	4915	1.6445	ug/L #	70
19) Methylene Chloride	7.0827	84	7726	1.9116	ug/L	89
20) Carbon Disulfide	7.1241	76	21916	1.9674	ug/L	94
21) Acrylonitrile	7.2585	53	15196	9.9487	ug/L	92
22) Methyl Tert Butyl Ether	7.2999	73	22380	1.9528	ug/L	98
23) trans-1,2-Dichloroethene	7.5273	96	7684	1.9559	ug/L	92
24) n-Hexane	7.5997	57	11469	1.8953	ug/L #	85
25) Diisopropyl ether	7.9306	45	315707	21.3768	ug/L	99
26) Vinyl Acetate	8.0960	43	5449	2.9745	ug/L #	81
27) 1,1-Dichloroethane	8.1270	63	16376	1.9102	ug/L	97
28) Ethyl-Tert-Butyl ether	8.4786	59	318557	21.3401	ug/L	98
29) 2-Butanone	8.6647	43	3219	1.8239	ug/L #	80
30) Propionitrile	8.7577	54	11015	21.3614	ug/L	96
31) 2,2-Dichloropropane	8.8612	77	11781	1.9875	ug/L	99
32) cis-1,2-Dichloroethene	8.9232	96	8733	1.9580	ug/L	99
33) Chloroform	9.1196	83	14912	1.9164	ug/L	98
34) 1-Bromopropane	9.2540	122	1286	1.3312	ug/L	82
35) Bromochloromethane	9.3471	130	4887	1.8687	ug/L	95
36) Tetrahydrofuran	9.3678	42	22315	19.5159	ug/L	98
38) 1,1,1-Trichloroethane	9.6263	97	13456	1.9099	ug/L #	94
39) Cyclohexane	9.6470	56	16282	1.9519	ug/L	97
40) 1,1-Dichloropropene	9.8124	75	10481	1.9785	ug/L	97
41) Carbon Tetrachloride	9.9572	117	10773	1.7981	ug/L	97
42) Tert-Amyl-Methyl ether	9.9158	73	235381	21.1411	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29740.D 8260WT.M Wed Mar 06 14:03:30 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29740.D Vial: 5
 Acq On : 5 Mar 2019 18:08 Operator: KFR
 Sample : WG698387-05 STD 2ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:28 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1226	62	13667	2.0209	ug/L	97
45) Benzene	10.1639	78	33141	2.1048	ug/L	93
46) Trichloroethene	10.8670	130	9146	2.0481	ug/L	93
47) Methylcyclohexane	10.9394	83	11788	1.8738	ug/L	96
48) 1,2-Dichloropropane	11.0738	63	9712	2.0765	ug/L	94
49) 1,4-Dioxane	11.3323	88	735	22.9025	ug/L	74
50) Bromodichloromethane	11.3530	83	9790	1.7711	ug/L	97
51) Dibromomethane	11.4254	93	5012	1.8760	ug/L	93
52) 2-Chloroethyl Vinyl Ether	11.6218	63	2669	1.1792	ug/L #	74
53) 4-Methyl-2-Pentanone	11.6528	58	2783	1.7258	ug/L #	64
54) cis-1,3-Dichloropropene	11.9423	75	11273	1.8096	ug/L	94
55) Dimethyl Disulfide	12.2008	79	4445	3.4159	ug/L	94
58) Toluene	12.3456	91	32770	1.9576	ug/L	96
59) Ethyl Methacrylate	12.4283	69	8224	1.7394	ug/L	100
60) trans-1,3-Dichloropropene	12.5110	75	9761	1.7810	ug/L	93
61) 1,1,2-Trichloroethane	12.7075	97	6630	1.8605	ug/L	93
62) 2-Hexanone	12.6558	43	4277	1.6147	ug/L #	77
63) 1,3-Dichloropropane	12.9970	76	11407	1.9840	ug/L	100
64) Tetrachloroethene	13.1107	164	6478	1.8244	ug/L	100
65) Dibromochloromethane	13.3589	129	6749	1.6637	ug/L	93
66) 1,2-Dibromoethane	13.5967	107	6352	1.8599	ug/L	100
67) 1-Chlorohexane	13.6691	91	10068	1.8218	ug/L	99
68) Chlorobenzene	14.0620	112	21643	1.8571	ug/L	98
69) 1,1,1,2-Tetrachloroethane	14.0930	131	7202	2.1037	ug/L	99
70) Ethylbenzene	14.0826	106	10939	1.8451	ug/L	82
71) m-,p-Xylene	14.1654	106	27179	3.7973	ug/L	85
72) o-Xylene	14.6927	106	13955	2.0095	ug/L	99
73) Styrene	14.7340	104	22207	1.9332	ug/L	99
74) Bromoform	15.1993	173	3881	2.0377	ug/L	93
75) Isopropylbenzene	15.0856	105	34361	1.9765	ug/L	97
77) 1,1,2,2-Tetrachloroethane	15.2924	83	7039	1.9420	ug/L	98
79) 1,2,3-Trichloropropane	15.4681	110	2527	2.0880	ug/L	78
80) trans-1,4-Dichloro-2-Butene	15.5095	53	2315	1.5171	ug/L	66
81) n-Propylbenzene	15.5612	91	41272	2.0142	ug/L	98
82) Bromobenzene	15.6853	156	9342	1.8669	ug/L	89
83) 1,3,5-Trimethylbenzene	15.7370	105	28921	1.9882	ug/L	99
84) 2-Chlorotoluene	15.8197	91	29474	2.0215	ug/L	99
85) 4-Chlorotoluene	15.8610	91	24240	2.0228	ug/L	97
86) a-Methylstyrene	16.1092	118	15618	1.8856	ug/L	94
87) tert-Butylbenzene	16.1712	134	6692	1.8679	ug/L	85
88) 1,2,4-Trimethylbenzene	16.2126	105	29847	1.9881	ug/L	98
89) sec-Butylbenzene	16.4194	105	38024	2.1056	ug/L	96
90) p-Isopropyltoluene	16.5641	119	31128	1.9823	ug/L	100
91) 1,3-Dichlorobenzene	16.7503	146	18388	1.9920	ug/L	97
92) 1,4-Dichlorobenzene	16.8743	146	17794	1.9303	ug/L	81
93) n-Butylbenzene	17.0501	91	27074	1.9131	ug/L	99
94) 1,2-Dichlorobenzene	17.3293	146	17273	1.9524	ug/L	99
95) 1,2-Dibromo-3-Chloropropane	18.2598	75	1087	1.4517	ug/L	79
96) 1,2,4-Trichlorobenzene	19.3145	180	10335	1.8058	ug/L	99
97) Hexachlorobutadiene	19.4592	225	5249	1.8197	ug/L	98
98) Naphthalene	19.6557	128	23379	1.9077	ug/L #	95
99) 1,2,3-Trichlorobenzene	19.9452	180	9758	1.8148	ug/L	95

(#) = qualifier out of range (m) = manual integration
 11M29740.D 8260WT.M Wed Mar 06 14:03:31 2019

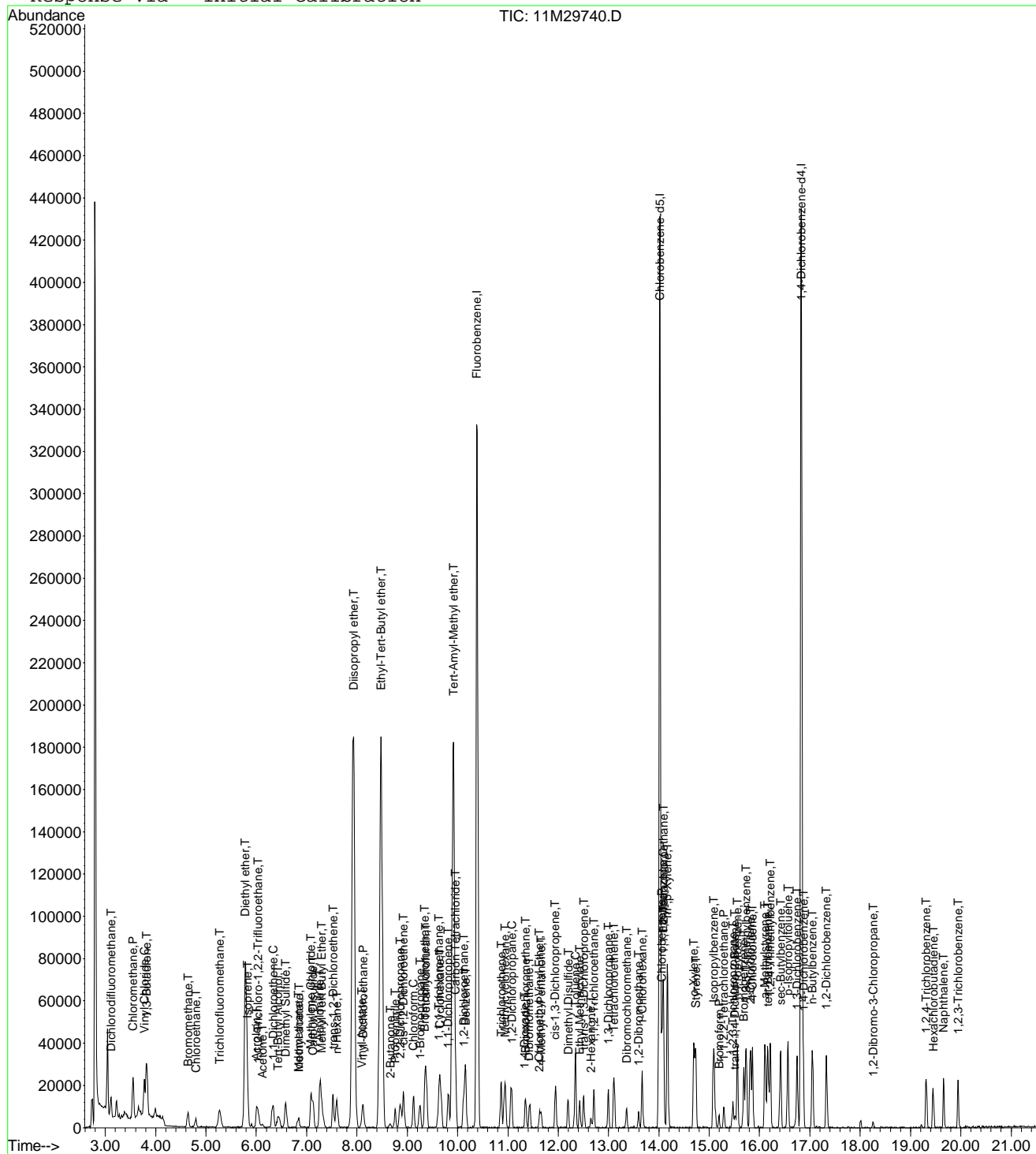
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29740.D
Acq On : 5 Mar 2019 18:08
Sample : WG698387-05 STD 2ug/L 8260
Misc : 1,1 STD92320
MS Integration Params: rteint.p
Quant Time: Mar 6 14:03 2019

Vial: 5
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:00:17 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29740.D Vial: 5
 Acq On : 5 Mar 2019 18:08 Operator: KFR
 Sample : WG698387-05 STD 2ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	-0.01
2 T	Dichlorodifluoromethane	2.000	1.898	5.1	100	-0.01
3 P	Chloromethane	2.000	2.110	-5.5	100	-0.01
4 C	Vinyl Chloride	2.000	2.106	-5.3	100	0.00
5 T	1,3-Butadiene	-1.000	0.166	0.0	100	0.01
6 T	Bromomethane	2.000	2.152	-7.6	100	0.00
7 T	Chloroethane	2.000	1.883	5.8	100	0.00
8 T	Trichlorofluoromethane	2.000	2.047	-2.4	100	0.00
9 T	Diethyl ether	20.000	20.514	-2.6	100	-0.01
10 T	Isoprene	2.000	1.854	7.3	100	0.00
11 T	Acrolein	10.000	9.845	1.5	100	-0.01
12 T	1,1,2-Trichloro-1,2,2-Trifl	2.000	1.921	3.9	100	-0.01
13 T	Acetone	-1.000	2.380	0.0	100	0.00
14 C	1,1-Dichloroethene	2.000	1.895	5.2	100	0.01
15 T	Tert-Butyl Alcohol	40.000	35.940	10.2	100	-0.01
16 T	Dimethyl Sulfide	2.000	1.804	9.8	100	0.00
17 T	Iodomethane	2.000	2.484	-24.2	100	0.00
18 T	Methyl acetate	2.000	1.645	17.8	100	0.00
19 T	Methylene Chloride	2.000	1.912	4.4	100	-0.01
20 T	Carbon Disulfide	2.000	1.967	1.6	100	-0.01
21 T	Acrylonitrile	10.000	9.949	0.5	100	-0.01
22 T	Methyl Tert Butyl Ether	2.000	1.953	2.3	100	0.00
23 T	trans-1,2-Dichloroethene	2.000	1.956	2.2	100	0.00
24 T	n-Hexane	2.000	1.895	5.2	100	0.00
25 T	Diisopropyl ether	20.000	21.377	-6.9	100	0.00
26 T	Vinyl Acetate	-1.000	2.974	0.0	0	0.00
27 P	1,1-Dichloroethane	2.000	1.910	4.5	100	0.00
28 T	Ethyl-Tert-Butyl ether	20.000	21.340	-6.7	100	0.00
29 T	2-Butanone	2.000	1.824	8.8	100	0.01
30 T	Propionitrile	20.000	21.361	-6.8	100	0.00
31 T	2,2-Dichloropropane	2.000	1.988	0.6	100	0.00
32 T	cis-1,2-Dichloroethene	2.000	1.958	2.1	100	-0.01
33 C	Chloroform	2.000	1.916	4.2	100	-0.01
34 T	1-Bromopropane	-1.000	1.331	0.0	100	0.00
35 T	Bromochloromethane	2.000	1.869	6.6	100	0.00
36 T	Tetrahydrofuran	20.000	19.516	2.4	100	0.00
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.40#
38 T	1,1,1-Trichloroethane	2.000	1.910	4.5	100	0.00
39 T	Cyclohexane	2.000	1.952	2.4	100	-0.01
40 T	1,1-Dichloropropene	2.000	1.979	1.0	100	0.00
41 T	Carbon Tetrachloride	2.000	1.798	10.1	100	0.00
42 T	Tert-Amyl-Methyl ether	20.000	21.141	-5.7	100	0.00
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.01#
44 T	1,2-Dichloroethane	2.000	2.021	-1.0	100	0.00
45 T	Benzene	2.000	2.105	-5.2	100	0.01
46 T	Trichloroethene	2.000	2.048	-2.4	100	0.00
47 T	Methylcyclohexane	2.000	1.874	6.3	100	0.00
48 C	1,2-Dichloropropane	2.000	2.076	-3.8	100	0.01
49 T	1,4-Dioxane	-1.000	22.902	0.0	100	-0.01
50 T	Bromodichloromethane	2.000	1.771	11.5	100	0.00
51 T	Dibromomethane	2.000	1.876	6.2	100	-0.01
52 T	2-Chloroethyl Vinyl Ether	-1.000	1.179	0.0	0	0.00
53 T	4-Methyl-2-Pentanone	-1.000	1.726	0.0	100	0.00
54 T	cis-1,3-Dichloropropene	2.000	1.810	9.5	100	0.00

(#) = Out of Range

11M29740.D 8260WT.M Wed Mar 06 13:57:55 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29740.D Vial: 5
 Acq On : 5 Mar 2019 18:08 Operator: KFR
 Sample : WG698387-05 STD 2ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	-1.000	3.416	0.0	0	0.01
56 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
57 S	Toluene-d8	-1.000	0.000	0.0	0	-12.25#
58 C	Toluene	2.000	1.958	2.1	100	0.00
59 T	Ethyl Methacrylate	2.000	1.739	13.0	100	0.00
60 T	trans-1,3-Dichloropropene	2.000	1.781	11.0	100	0.01
61 T	1,1,2-Trichloroethane	2.000	1.860	7.0	100	0.00
62 T	2-Hexanone	2.000	1.615	19.3	100	0.01
63 T	1,3-Dichloropropane	2.000	1.984	0.8	100	0.00
64 T	Tetrachloroethene	2.000	1.911	4.4	100	0.00
65 T	Dibromochloromethane	2.000	1.664	16.8	100	0.00
66 T	1,2-Dibromoethane	2.000	1.860	7.0	100	0.00
67 T	1-Chlorohexane	2.000	1.822	8.9	100	0.00
68 P	Chlorobenzene	2.000	1.857	7.2	100	0.00
69 T	1,1,1,2-Tetrachloroethane	2.000	2.104	-5.2	100	0.00
70 C	Ethylbenzene	2.000	1.845	7.8	100	0.00
71 T	m-,p-Xylene	4.000	3.797	5.1	100	0.00
72 T	o-Xylene	2.000	2.009	-0.4	100	0.00
73 T	Styrene	2.000	1.933	3.3	100	0.00
74 P	Bromoform	2.000	2.038	-1.9	100	0.00
75 T	Isopropylbenzene	2.000	1.976	1.2	100	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
77 P	1,1,2,2-Tetrachloroethane	2.000	1.942	2.9	100	0.00
78 S	p-Bromofluorobenzene	1.000	0.000	100.0#	0	-15.42#
79 T	1,2,3-Trichloropropane	2.000	2.088	-4.4	100	0.00
80 T	trans-1,4-Dichloro-2-Butene	2.000	1.517	24.2	100	0.00
81 T	n-Propylbenzene	2.000	2.014	-0.7	100	0.00
82 T	Bromobenzene	2.000	1.867	6.7	100	0.00
83 T	1,3,5-Trimethylbenzene	2.000	1.988	0.6	100	0.00
84 T	2-Chlorotoluene	2.000	2.022	-1.1	100	0.00
85 T	4-Chlorotoluene	2.000	2.023	-1.2	100	0.00
86 T	a-Methylstyrene	2.000	1.886	5.7	100	0.00
87 T	tert-Butylbenzene	2.000	1.868	6.6	100	0.00
88 T	1,2,4-Trimethylbenzene	2.000	1.988	0.6	100	0.00
89 T	sec-Butylbenzene	2.000	2.106	-5.3	100	0.00
90 T	p-Isopropyltoluene	2.000	1.982	0.9	100	0.00
91 T	1,3-Dichlorobenzene	2.000	1.992	0.4	100	0.00
92 T	1,4-Dichlorobenzene	2.000	1.930	3.5	100	0.01
93 T	n-Butylbenzene	2.000	1.913	4.3	100	0.00
94 T	1,2-Dichlorobenzene	2.000	1.952	2.4	100	0.00
95 T	1,2-Dibromo-3-Chloropropane	-1.000	1.452	0.0	0	0.01
96 T	1,2,4-Trichlorobenzene	2.000	1.806	9.7	100	0.00
97 T	Hexachlorobutadiene	2.000	1.820	9.0	100	0.01
98 T	Naphthalene	2.000	1.908	4.6	100	0.00
99 T	1,2,3-Trichlorobenzene	2.000	1.815	9.3	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29740.D 8260WT.M Wed Mar 06 13:57:55 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29741.D Vial: 6
 Acq On : 5 Mar 2019 18:38 Operator: KFR
 Sample : WG698387-06 STD 5ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:34 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3914	96	392987	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0206	117	289755	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	139899	25.0000	ug/L	0.0000
System Monitoring Compounds						
37) Dibromofluoromethane	9.3988	111	22696	5.0031	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	20.0124%#	
43) 1,2-Dichloroethane-d4	10.0089	65	28500	5.2362	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	20.9448%#	
57) Toluene-d8	12.2525	98	76134	5.2797	ug/L	0.0000
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	21.1188%#	
78) p-Bromofluorobenzene	15.4164	95	27613	5.1578	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	20.6312%#	
Target Compounds						
					Qvalue	
2) Dichlorodifluoromethane	3.1124	85	25923	4.4652	ug/L	98
3) Chloromethane	3.5466	50	53957	4.8390	ug/L	100
4) Vinyl Chloride	3.7741	62	41361	4.9689	ug/L	88
5) 1,3-Butadiene	3.8258	54	24006	3.3577	ug/L	92
6) Bromomethane	4.6426	94	15504	4.8615	ug/L	100
7) Chloroethane	4.7977	64	14961	4.6491	ug/L	95
8) Trichlorofluoromethane	5.2733	101	38366	4.8706	ug/L	99
9) Diethyl ether	5.7800	59	209116	50.7782	ug/L	99
10) Isoprene	5.8213	67	28561	4.6034	ug/L	98
11) Acrolein	6.0074	56	17920	25.0577	ug/L	100
12) 1,1,2-Trichloro-1,2,2-Trif	6.0281	101	17984	4.6542	ug/L	98
13) Acetone	6.1212	43	7821	5.4436	ug/L	97
14) 1,1-Dichloroethene	6.3280	61	37262	4.9548	ug/L	97
15) Tert-Butyl Alcohol	6.4314	59	37162	92.7324	ug/L	98
16) Dimethyl Sulfide	6.5865	62	24073	4.8422	ug/L	98
17) Iodomethane	6.8346	142	14449	4.2023	ug/L	98
18) Methyl acetate	6.8346	43	14821	4.7825	ug/L	98
19) Methylene Chloride	7.0931	84	20465	4.8833	ug/L	99
20) Carbon Disulfide	7.1241	76	56967	4.9319	ug/L	98
21) Acrylonitrile	7.2689	53	39559	24.9766	ug/L	94
22) Methyl Tert Butyl Ether	7.2999	73	57453	4.8346	ug/L	99
23) trans-1,2-Dichloroethene	7.5274	96	19820	4.8653	ug/L	100
24) n-Hexane	7.5997	57	29004	4.6224	ug/L #	96
25) Diisopropyl ether	7.9306	45	812924	53.0833	ug/L	100
26) Vinyl Acetate	8.0960	43	17244	5.0073	ug/L	98
27) 1,1-Dichloroethane	8.1167	63	42561	4.7879	ug/L	98
28) Ethyl-Tert-Butyl ether	8.4786	59	812824	52.5119	ug/L	99
29) 2-Butanone	8.6544	43	8490	4.6391	ug/L	92
30) Propionitrile	8.7578	54	27847	52.0803	ug/L	99
31) 2,2-Dichloropropane	8.8612	77	28125	4.5759	ug/L	100
32) cis-1,2-Dichloroethene	8.9232	96	22324	4.8269	ug/L	96
33) Chloroform	9.1197	83	38862	4.8165	ug/L	96
34) 1-Bromopropane	9.2541	122	4320	4.3125	ug/L	93
35) Bromochloromethane	9.3471	130	13508	4.9812	ug/L	99
36) Tetrahydrofuran	9.3678	42	56348	47.5249	ug/L	98
38) 1,1,1-Trichloroethane	9.6263	97	34541	4.7279	ug/L	99
39) Cyclohexane	9.6573	56	40501	4.6825	ug/L	99
40) 1,1-Dichloropropene	9.8124	75	25937	4.7218	ug/L	97
41) Carbon Tetrachloride	9.9572	117	27882	4.4880	ug/L	99
42) Tert-Amyl-Methyl ether	9.9158	73	598486	51.8396	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29741.D 8260WT.M Wed Mar 06 14:03:35 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29741.D Vial: 6
 Acq On : 5 Mar 2019 18:38 Operator: KFR
 Sample : WG698387-06 STD 5ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:34 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1226	62	35675	5.0872	ug/L	98
45) Benzene	10.1536	78	79739	4.8839	ug/L	100
46) Trichloroethene	10.8670	130	22287	4.8130	ug/L	100
47) Methylcyclohexane	10.9498	83	30480	4.6725	ug/L	98
48) 1,2-Dichloropropane	11.0635	63	22894	4.7205	ug/L	97
49) 1,4-Dioxane	11.3323	88	2997	90.0601	ug/L	90
50) Bromodichloromethane	11.3530	83	27396	4.7798	ug/L	99
51) Dibromomethane	11.4357	93	13935	5.0300	ug/L	95
52) 2-Chloroethyl Vinyl Ether	11.6218	63	8940	3.8091	ug/L	97
53) 4-Methyl-2-Pentanone	11.6529	58	7495	4.4824	ug/L	99
54) cis-1,3-Dichloropropene	11.9423	75	30690	4.7510	ug/L	99
55) Dimethyl Disulfide	12.1905	79	13399	5.3390	ug/L	86
58) Toluene	12.3456	91	89627	5.2216	ug/L	95
59) Ethyl Methacrylate	12.4283	69	22563	4.6540	ug/L	98
60) trans-1,3-Dichloropropene	12.5007	75	26133	4.6501	ug/L	97
61) 1,1,2-Trichloroethane	12.7075	97	17300	4.7344	ug/L	96
62) 2-Hexanone	12.6455	43	13329	4.9076	ug/L	92
63) 1,3-Dichloropropane	12.9970	76	29055	4.9282	ug/L	97
64) Tetrachloroethene	13.1107	164	17118	4.7017	ug/L	99
65) Dibromochloromethane	13.3589	129	17620	4.2359	ug/L	99
66) 1,2-Dibromoethane	13.5967	107	16570	4.7316	ug/L	96
67) 1-Chlorohexane	13.6691	91	26371	4.6535	ug/L	99
68) Chlorobenzene	14.0620	112	57129	4.7806	ug/L	100
69) 1,1,1,2-Tetrachloroethane	14.0930	131	19294	4.4172	ug/L	99
70) Ethylbenzene	14.0826	106	29387	4.8340	ug/L	94
71) m-,p-Xylene	14.1654	106	69659	9.4913	ug/L	86
72) o-Xylene	14.6927	106	34481	4.8422	ug/L	97
73) Styrene	14.7340	104	58036	4.9272	ug/L	100
74) Bromoform	15.1993	173	9642	3.8985	ug/L	93
75) Isopropylbenzene	15.0856	105	90391	5.0706	ug/L	97
77) 1,1,2,2-Tetrachloroethane	15.2924	83	18603	5.0974	ug/L	100
79) 1,2,3-Trichloropropane	15.4682	110	6006	4.9287	ug/L	74
80) trans-1,4-Dichloro-2-Butene	15.5095	53	7392	4.8111	ug/L	96
81) n-Propylbenzene	15.5612	91	111335	5.3965	ug/L	97
82) Bromobenzene	15.6853	156	25563	5.0736	ug/L	98
83) 1,3,5-Trimethylbenzene	15.7370	105	76366	5.2141	ug/L	98
84) 2-Chlorotoluene	15.8197	91	77542	5.2820	ug/L	94
85) 4-Chlorotoluene	15.8611	91	62041	5.1419	ug/L	99
86) a-Methylstyrene	16.1092	118	40145	4.8137	ug/L	97
87) tert-Butylbenzene	16.1712	134	17911	4.9652	ug/L	97
88) 1,2,4-Trimethylbenzene	16.2126	105	76458	5.0581	ug/L	100
89) sec-Butylbenzene	16.4194	105	93501	5.1423	ug/L	97
90) p-Isopropyltoluene	16.5642	119	82364	5.2094	ug/L	97
91) 1,3-Dichlorobenzene	16.7503	146	46292	4.9805	ug/L	97
92) 1,4-Dichlorobenzene	16.8743	146	45984	4.9544	ug/L	94
93) n-Butylbenzene	17.0501	91	71407	5.0114	ug/L	97
94) 1,2-Dichlorobenzene	17.3293	146	44856	5.0355	ug/L	100
95) 1,2-Dibromo-3-Chloropropane	18.2495	75	3084	4.0905	ug/L	97
96) 1,2,4-Trichlorobenzene	19.3145	180	26680	4.6300	ug/L	99
97) Hexachlorobutadiene	19.4489	225	14319	4.9301	ug/L	97
98) Naphthalene	19.6557	128	59627	4.8323	ug/L	99
99) 1,2,3-Trichlorobenzene	19.9452	180	24480	4.5217	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29741.D 8260WT.M Wed Mar 06 14:03:35 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29741.D Vial: 6
 Acq On : 5 Mar 2019 18:38 Operator: KFR
 Sample : WG698387-06 STD 5ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	5.000	4.465	10.7	100	-0.01
3 P	Chloromethane	5.000	4.839	3.2	100	-0.01
4 C	Vinyl Chloride	5.000	4.969	0.6	100	0.00
5 T	1,3-Butadiene	5.000	3.358	32.8#	100	0.01
6 T	Bromomethane	5.000	4.862	2.8	100	0.00
7 T	Chloroethane	5.000	4.649	7.0	100	0.00
8 T	Trichlorofluoromethane	5.000	4.871	2.6	100	0.00
9 T	Diethyl ether	50.000	50.778	-1.6	100	-0.01
10 T	Isoprene	5.000	4.603	7.9	100	0.00
11 T	Acrolein	25.000	25.058	-0.2	100	-0.01
12 T	1,1,2-Trichloro-1,2,2-Trifl	5.000	4.654	6.9	100	-0.01
13 T	Acetone	5.000	5.444	-8.9	100	0.00
14 C	1,1-Dichloroethene	5.000	4.955	0.9	100	0.00
15 T	Tert-Butyl Alcohol	100.000	92.732	7.3	100	-0.01
16 T	Dimethyl Sulfide	5.000	4.842	3.2	100	0.00
17 T	Iodomethane	5.000	4.202	16.0	100	0.00
18 T	Methyl acetate	5.000	4.782	4.4	100	-0.01
19 T	Methylene Chloride	5.000	4.883	2.3	100	0.00
20 T	Carbon Disulfide	5.000	4.932	1.4	100	-0.01
21 T	Acrylonitrile	25.000	24.977	0.1	100	0.00
22 T	Methyl Tert Butyl Ether	5.000	4.835	3.3	100	0.00
23 T	trans-1,2-Dichloroethene	5.000	4.865	2.7	100	0.00
24 T	n-Hexane	5.000	4.622	7.6	100	0.00
25 T	Diisopropyl ether	50.000	53.083	-6.2	100	0.00
26 T	Vinyl Acetate	5.000	5.007	-0.1	100	0.00
27 P	1,1-Dichloroethane	5.000	4.788	4.2	100	-0.01
28 T	Ethyl-Tert-Butyl ether	50.000	52.512	-5.0	100	0.00
29 T	2-Butanone	5.000	4.639	7.2	100	0.00
30 T	Propionitrile	50.000	52.080	-4.2	100	0.00
31 T	2,2-Dichloropropane	5.000	4.576	8.5	100	0.00
32 T	cis-1,2-Dichloroethene	5.000	4.827	3.5	100	-0.01
33 C	Chloroform	5.000	4.817	3.7	100	-0.01
34 T	1-Bromopropane	5.000	4.312	13.8	100	0.00
35 T	Bromochloromethane	5.000	4.981	0.4	100	0.00
36 T	Tetrahydrofuran	50.000	47.525	5.0	100	0.00
37 S	Dibromofluoromethane	5.000	5.003	-0.1	100	0.00
38 T	1,1,1-Trichloroethane	5.000	4.728	5.4	100	0.00
39 T	Cyclohexane	5.000	4.682	6.4	100	0.00
40 T	1,1-Dichloropropene	5.000	4.722	5.6	100	0.00
41 T	Carbon Tetrachloride	5.000	4.488	10.2	100	0.00
42 T	Tert-Amyl-Methyl ether	50.000	51.840	-3.7	100	0.00
43 S	1,2-Dichloroethane-d4	5.000	5.236	-4.7	100	0.00
44 T	1,2-Dichloroethane	5.000	5.087	-1.7	100	0.00
45 T	Benzene	5.000	4.884	2.3	100	0.00
46 T	Trichloroethene	5.000	4.813	3.7	100	0.00
47 T	Methylcyclohexane	5.000	4.672	6.6	100	0.01
48 C	1,2-Dichloropropane	5.000	4.721	5.6	100	0.00
49 T	1,4-Dioxane	100.000	90.060	9.9	100	-0.01
50 T	Bromodichloromethane	5.000	4.780	4.4	100	0.00
51 T	Dibromomethane	5.000	5.030	-0.6	100	0.00
52 T	2-Chloroethyl Vinyl Ether	5.000	3.809	23.8	100	0.00
53 T	4-Methyl-2-Pentanone	5.000	4.482	10.4	100	0.00
54 T	cis-1,3-Dichloropropene	5.000	4.751	5.0	100	0.00

(#) = Out of Range

11M29741.D 8260WT.M Wed Mar 06 13:58:12 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29741.D Vial: 6
 Acq On : 5 Mar 2019 18:38 Operator: KFR
 Sample : WG698387-06 STD 5ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 13:50:26 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	5.000	5.339	-6.8	100	0.00
56 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
57 S	Toluene-d8	5.000	5.280	-5.6	100	0.00
58 C	Toluene	5.000	5.222	-4.4	100	0.00
59 T	Ethyl Methacrylate	5.000	4.654	6.9	100	0.00
60 T	trans-1,3-Dichloropropene	5.000	4.650	7.0	100	0.00
61 T	1,1,2-Trichloroethane	5.000	4.734	5.3	100	0.00
62 T	2-Hexanone	5.000	4.908	1.8	100	0.00
63 T	1,3-Dichloropropane	5.000	4.928	1.4	100	0.00
64 T	Tetrachloroethene	5.000	4.924	1.5	100	0.00
65 T	Dibromochloromethane	5.000	4.236	15.3	100	0.00
66 T	1,2-Dibromoethane	5.000	4.732	5.4	100	0.00
67 T	1-Chlorohexane	5.000	4.654	6.9	100	0.00
68 P	Chlorobenzene	5.000	4.781	4.4	100	0.00
69 T	1,1,1,2-Tetrachloroethane	5.000	4.417	11.7	100	0.00
70 C	Ethylbenzene	5.000	4.834	3.3	100	0.00
71 T	m-,p-Xylene	10.000	9.491	5.1	100	0.00
72 T	o-Xylene	5.000	4.842	3.2	100	0.00
73 T	Styrene	5.000	4.927	1.5	100	0.00
74 P	Bromoform	5.000	3.898	22.0	100	0.00
75 T	Isopropylbenzene	5.000	5.071	-1.4	100	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
77 P	1,1,2,2-Tetrachloroethane	5.000	5.097	-1.9	100	0.00
78 S	p-Bromofluorobenzene	5.000	5.158	-3.2	100	0.00
79 T	1,2,3-Trichloropropane	5.000	4.929	1.4	100	0.00
80 T	trans-1,4-Dichloro-2-Butene	5.000	4.811	3.8	100	0.00
81 T	n-Propylbenzene	5.000	5.397	-7.9	100	0.00
82 T	Bromobenzene	5.000	5.074	-1.5	100	0.00
83 T	1,3,5-Trimethylbenzene	5.000	5.214	-4.3	100	0.00
84 T	2-Chlorotoluene	5.000	5.282	-5.6	100	0.00
85 T	4-Chlorotoluene	5.000	5.142	-2.8	100	0.00
86 T	a-Methylstyrene	5.000	4.814	3.7	100	0.00
87 T	tert-Butylbenzene	5.000	4.965	0.7	100	0.00
88 T	1,2,4-Trimethylbenzene	5.000	5.058	-1.2	100	0.00
89 T	sec-Butylbenzene	5.000	5.142	-2.8	100	0.00
90 T	p-Isopropyltoluene	5.000	5.209	-4.2	100	0.00
91 T	1,3-Dichlorobenzene	5.000	4.981	0.4	100	0.00
92 T	1,4-Dichlorobenzene	5.000	4.954	0.9	100	0.01
93 T	n-Butylbenzene	5.000	5.011	-0.2	100	0.00
94 T	1,2-Dichlorobenzene	5.000	5.036	-0.7	100	0.00
95 T	1,2-Dibromo-3-Chloropropane	5.000	4.091	18.2	100	0.00
96 T	1,2,4-Trichlorobenzene	5.000	4.630	7.4	100	0.00
97 T	Hexachlorobutadiene	5.000	4.930	1.4	100	0.00
98 T	Naphthalene	5.000	4.832	3.4	100	0.00
99 T	1,2,3-Trichlorobenzene	5.000	4.522	9.6	100	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29741.D 8260WT.M Wed Mar 06 13:58:12 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29742.D Vial: 7
 Acq On : 5 Mar 2019 19:08 Operator: KFR
 Sample : WG698387-07 STD 20ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:36 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	390701	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0207	117	294074	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	148164	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3989	111	42025	9.3182	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	37.2728%#	
43) 1,2-Dichloroethane-d4	10.0089	65	50464	9.3258	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	37.3032%#	
57) Toluene-d8	12.2526	98	137326	9.3834	ug/L	0.0000
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	37.5336%#	
78) p-Bromofluorobenzene	15.4165	95	51838	9.1426	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	36.5704%#	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1124	85	125308	21.7106	ug/L	99
3) Chloromethane	3.5467	50	224404	20.2429	ug/L	100
4) Vinyl Chloride	3.7638	62	176474	21.3247	ug/L	99
5) 1,3-Butadiene	3.8051	54	79748	24.7594	ug/L	96
6) Bromomethane	4.6427	94	61649	19.4441	ug/L	98
7) Chloroethane	4.7874	64	66614	20.8215	ug/L	99
8) Trichlorofluoromethane	5.2630	101	161131	20.5754	ug/L	100
9) Diethyl ether	5.7800	59	308356	75.3141	ug/L	99
10) Isoprene	5.8214	67	125862	20.4051	ug/L	99
11) Acrolein	6.0075	56	27463	38.6264	ug/L	98
12) 1,1,2-Trichloro-1,2,2-Trif	6.0282	101	79889	20.7960	ug/L	99
13) Acetone	6.1109	43	31044	21.7338	ug/L	96
14) 1,1-Dichloroethene	6.3280	61	157212	21.0272	ug/L	98
15) Tert-Butyl Alcohol	6.4314	59	64696	162.3841	ug/L	99
16) Dimethyl Sulfide	6.5865	62	102463	20.7307	ug/L	98
17) Iodomethane	6.8243	142	88521	18.1838	ug/L	100
18) Methyl acetate	6.8450	43	62615	20.3229	ug/L	99
19) Methylene Chloride	7.0828	84	85684	20.5652	ug/L	99
20) Carbon Disulfide	7.1242	76	250026	21.7726	ug/L	99
21) Acrylonitrile	7.2586	53	63669	40.4343	ug/L	93
22) Methyl Tert Butyl Ether	7.2896	73	243518	20.6117	ug/L	100
23) trans-1,2-Dichloroethene	7.5274	96	85524	21.1168	ug/L	99
24) n-Hexane	7.5894	57	125092	20.0529	ug/L	99
25) Diisopropyl ether	7.9307	45	1201067	78.8877	ug/L	100
26) Vinyl Acetate	8.0961	43	101860	19.9478	ug/L	99
27) 1,1-Dichloroethane	8.1167	63	183782	20.7953	ug/L	99
28) Ethyl-Tert-Butyl ether	8.4787	59	1222739	79.4563	ug/L	100
29) 2-Butanone	8.6544	43	39021	21.4468	ug/L	96
30) Propionitrile	8.7578	54	46276	87.0532	ug/L	98
31) 2,2-Dichloropropane	8.8612	77	127123	20.8037	ug/L	99
32) cis-1,2-Dichloroethene	8.9232	96	98859	21.5006	ug/L	99
33) Chloroform	9.1197	83	163362	20.3655	ug/L	99
34) 1-Bromopropane	9.2541	122	20388	20.4717	ug/L	92
35) Bromochloromethane	9.3472	130	57852	21.4584	ug/L	98
36) Tetrahydrofuran	9.3679	42	90012	76.3620	ug/L	99
38) 1,1,1-Trichloroethane	9.6263	97	150552	20.7280	ug/L	99
39) Cyclohexane	9.6573	56	174247	20.2632	ug/L	98
40) 1,1-Dichloropropene	9.8125	75	111283	20.3777	ug/L	99
41) Carbon Tetrachloride	9.9572	117	126215	20.4351	ug/L	99
42) Tert-Amyl-Methyl ether	9.9159	73	906856	79.0095	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29742.D 8260WT.M Wed Mar 06 14:03:37 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29742.D Vial: 7
 Acq On : 5 Mar 2019 19:08 Operator: KFR
 Sample : WG698387-07 STD 20ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:36 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1123	62	145731	20.9027	ug/L	98
45) Benzene	10.1537	78	339062	20.8887	ug/L	99
46) Trichloroethene	10.8671	130	95118	20.6615	ug/L	98
47) Methylcyclohexane	10.9395	83	133133	20.5282	ug/L	99
48) 1,2-Dichloropropane	11.0635	63	100929	20.9323	ug/L	99
49) 1,4-Dioxane	11.3427	88	5301	160.2274	ug/L	99
50) Bromodichloromethane	11.3531	83	122680	21.5292	ug/L	100
51) Dibromomethane	11.4358	93	57070	20.7207	ug/L	99
52) 2-Chloroethyl Vinyl Ether	11.6219	63	44520	19.0800	ug/L	100
53) 4-Methyl-2-Pentanone	11.6529	58	34506	20.7572	ug/L	98
54) cis-1,3-Dichloropropene	11.9424	75	136570	21.2655	ug/L	99
55) Dimethyl Disulfide	12.1906	79	76353	19.2103	ug/L	96
58) Toluene	12.3456	91	373915	21.4641	ug/L	97
59) Ethyl Methacrylate	12.4284	69	106101	21.5636	ug/L	97
60) trans-1,3-Dichloropropene	12.5007	75	119252	20.9079	ug/L	100
61) 1,1,2-Trichloroethane	12.7075	97	76173	20.5399	ug/L	98
62) 2-Hexanone	12.6455	43	59069	21.4290	ug/L	99
63) 1,3-Dichloropropane	12.9970	76	126137	21.0809	ug/L	98
64) Tetrachloroethene	13.1108	164	75608	20.4616	ug/L	99
65) Dibromochloromethane	13.3589	129	85987	20.3681	ug/L	98
66) 1,2-Dibromoethane	13.5967	107	73437	20.6619	ug/L	99
67) 1-Chlorohexane	13.6691	91	118090	20.5326	ug/L	97
68) Chlorobenzene	14.0620	112	249566	20.5772	ug/L	99
69) 1,1,1,2-Tetrachloroethane	14.0930	131	93650	18.5949	ug/L	98
70) Ethylbenzene	14.0827	106	126947	20.5753	ug/L	92
71) m-,p-Xylene	14.1654	106	311240	41.7848	ug/L	91
72) o-Xylene	14.6927	106	152623	21.1181	ug/L	96
73) Styrene	14.7341	104	257910	21.5748	ug/L	99
74) Bromoform	15.1994	173	52978	17.8837	ug/L	98
75) Isopropylbenzene	15.0856	105	394369	21.7978	ug/L	98
77) 1,1,2,2-Tetrachloroethane	15.2924	83	84470	21.8543	ug/L	100
79) 1,2,3-Trichloropropane	15.4682	110	27663	21.4346	ug/L	94
80) trans-1,4-Dichloro-2-Butene	15.5096	53	33901	20.8338	ug/L	98
81) n-Propylbenzene	15.5612	91	486553	22.2681	ug/L	98
82) Bromobenzene	15.6853	156	108647	20.3609	ug/L	99
83) 1,3,5-Trimethylbenzene	15.7370	105	333787	21.5189	ug/L	99
84) 2-Chlorotoluene	15.8198	91	325566	20.9398	ug/L	99
85) 4-Chlorotoluene	15.8611	91	267721	20.9509	ug/L	98
86) a-Methylstyrene	16.1092	118	193534	21.9120	ug/L	98
87) tert-Butylbenzene	16.1713	134	77137	20.1906	ug/L	98
88) 1,2,4-Trimethylbenzene	16.2127	105	347387	21.6993	ug/L	97
89) sec-Butylbenzene	16.4194	105	424363	22.0367	ug/L	98
90) p-Isopropyltoluene	16.5642	119	368039	21.9793	ug/L	98
91) 1,3-Dichlorobenzene	16.7503	146	203238	20.6466	ug/L	99
92) 1,4-Dichlorobenzene	16.8640	146	204384	20.7923	ug/L	100
93) n-Butylbenzene	17.0501	91	329180	21.8134	ug/L	99
94) 1,2-Dichlorobenzene	17.3293	146	196258	20.8028	ug/L	99
95) 1,2-Dibromo-3-Chloropropane	18.2495	75	15193	19.0275	ug/L	95
96) 1,2,4-Trichlorobenzene	19.3145	180	124786	20.4470	ug/L	100
97) Hexachlorobutadiene	19.4489	225	63234	20.5573	ug/L	99
98) Naphthalene	19.6557	128	275722	21.0986	ug/L	99
99) 1,2,3-Trichlorobenzene	19.9452	180	114903	20.0397	ug/L	99

(#) = qualifier out of range (m) = manual integration
 11M29742.D 8260WT.M Wed Mar 06 14:03:37 2019

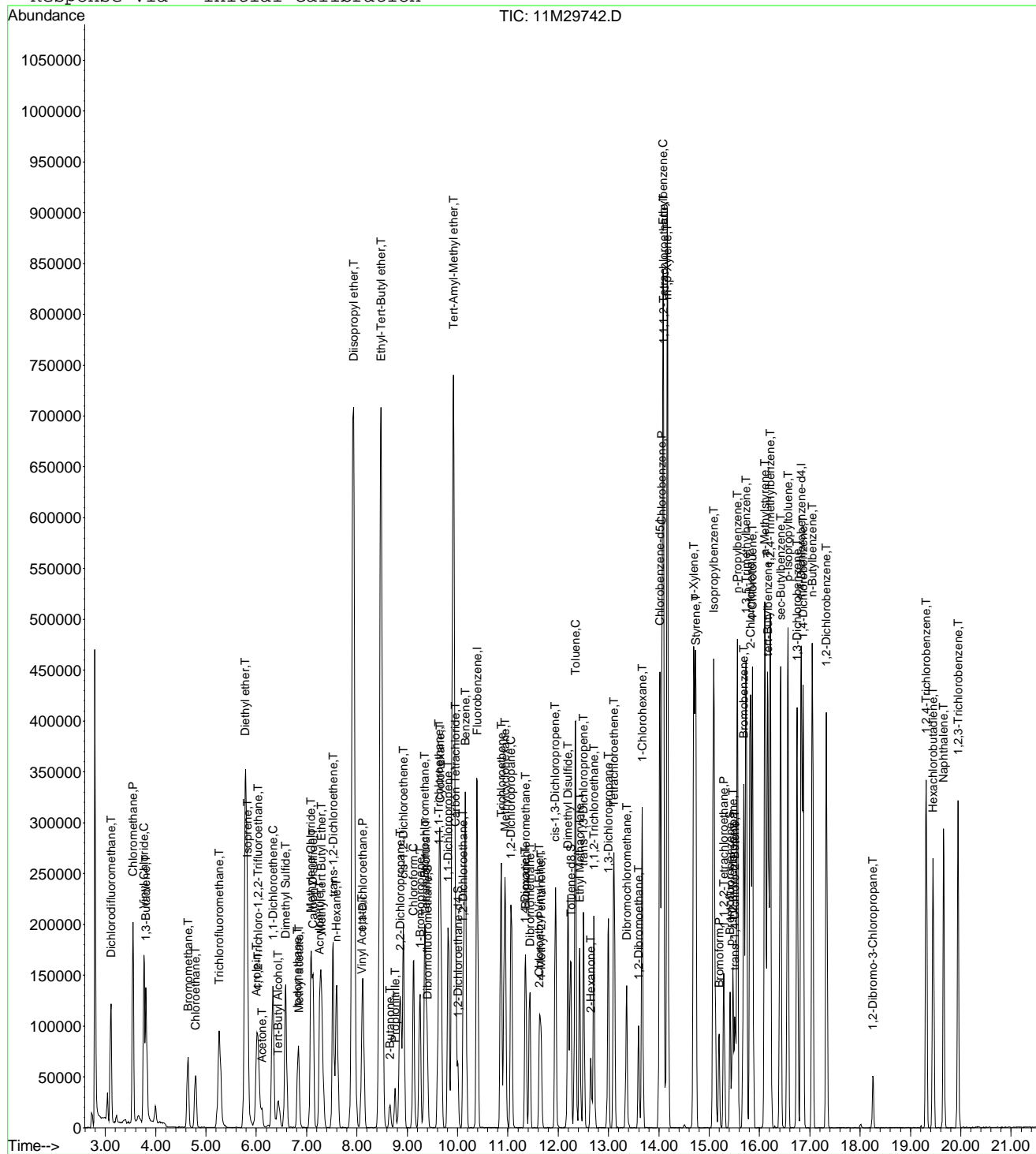
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29742.D
Acq On : 5 Mar 2019 19:08
Sample : WG698387-07 STD 20ug/L 8260
Misc : 1,1 STD92320
MS Integration Params: rteint.p
Quant Time: Mar 6 14:03 2019

Vial: 7
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:00:17 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29743.D Vial: 8
 Acq On : 5 Mar 2019 19:38 Operator: KFR
 Sample : WG698387-08 STD 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:37 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	395124	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0207	117	299517	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	148978	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3989	111	119072	26.1063	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	= 104.4252%		
43) 1,2-Dichloroethane-d4	10.0089	65	142217	25.9876	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	= 103.9504%		
57) Toluene-d8	12.2526	98	388036	26.0325	ug/L	0.0000
Spiked Amount	25.0000	Range 88 - 110	Recovery	= 104.1300%		
78) p-Bromofluorobenzene	15.4165	95	154561	27.1107	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	= 108.4428%		

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1227	85	319634	54.7593	ug/L	100
3) Chloromethane	3.5570	50	536525	47.8568	ug/L	100
4) Vinyl Chloride	3.7741	62	429474	51.3158	ug/L	100
5) 1,3-Butadiene	3.8155	54	159508	54.6222	ug/L	100
6) Bromomethane	4.6426	94	146832	45.7924	ug/L	100
7) Chloroethane	4.7977	64	168484	52.0735	ug/L	100
8) Trichlorofluoromethane	5.2734	101	416066	52.5342	ug/L	100
9) Diethyl ether	5.7904	59	378702	91.4602	ug/L	100
10) Isoprene	5.8214	67	323736	51.8974	ug/L	100
11) Acrolein	6.0178	56	35356	49.1712	ug/L	100
12) 1,1,2-Trichloro-1,2,2-Trif	6.0385	101	207472	53.4028	ug/L	100
13) Acetone	6.1212	43	73792	51.0832	ug/L	100
14) 1,1-Dichloroethene	6.3280	61	397617	52.5861	ug/L	100
15) Tert-Butyl Alcohol	6.4417	59	80396	199.5315	ug/L	100
16) Dimethyl Sulfide	6.5865	62	259928	52.0010	ug/L	100
17) Iodomethane	6.8346	142	264453	50.7961	ug/L	100
18) Methyl acetate	6.8450	43	161156	51.7207	ug/L	100
19) Methylene Chloride	7.0931	84	216377	51.3517	ug/L	100
20) Carbon Disulfide	7.1345	76	627208	54.0068	ug/L	100
21) Acrylonitrile	7.2689	53	86947	54.5993	ug/L	100
22) Methyl Tert Butyl Ether	7.2999	73	619562	51.8536	ug/L	100
23) trans-1,2-Dichloroethene	7.5274	96	220129	53.7438	ug/L	100
24) n-Hexane	7.5998	57	326174	51.7020	ug/L	100
25) Diisopropyl ether	7.9306	45	1437902	93.3861	ug/L	100
26) Vinyl Acetate	8.0961	43	275033	49.9454	ug/L	100
27) 1,1-Dichloroethane	8.1271	63	465254	52.0552	ug/L	100
28) Ethyl-Tert-Butyl ether	8.4786	59	1483214	95.3037	ug/L	100
29) 2-Butanone	8.6544	43	96483	52.4356	ug/L	100
30) Propionitrile	8.7578	54	58685	109.1609	ug/L	100
31) 2,2-Dichloropropane	8.8612	77	325588	52.6862	ug/L	100
32) cis-1,2-Dichloroethene	8.9336	96	249007	53.5497	ug/L	100
33) Chloroform	9.1300	83	411337	50.7052	ug/L	100
34) 1-Bromopropane	9.2541	122	51248	50.8823	ug/L	100
35) Bromochloromethane	9.3472	130	149707	54.9074	ug/L	100
36) Tetrahydrofuran	9.3678	42	115030	96.4937	ug/L	100
38) 1,1,1-Trichloroethane	9.6263	97	393600	53.5842	ug/L	100
39) Cyclohexane	9.6573	56	454208	52.2286	ug/L	100
40) 1,1-Dichloropropene	9.8125	75	292591	52.9783	ug/L	100
41) Carbon Tetrachloride	9.9572	117	347273	55.5964	ug/L	100
42) Tert-Amyl-Methyl ether	9.9158	73	1118994	96.4006	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29743.D 8260WT.M Wed Mar 06 14:03:38 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29743.D Vial: 8
 Acq On : 5 Mar 2019 19:38 Operator: KFR
 Sample : WG698387-08 STD 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:37 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1226	62	372401	52.8169	ug/L	100
45) Benzene	10.1537	78	859665	52.3689	ug/L	100
46) Trichloroethene	10.8671	130	246923	53.0361	ug/L	100
47) Methylcyclohexane	10.9395	83	342261	52.1834	ug/L	100
48) 1,2-Dichloropropane	11.0635	63	253911	52.0708	ug/L	100
49) 1,4-Dioxane	11.3427	88	6871	205.3573	ug/L	100
50) Bromodichloromethane	11.3530	83	322541	55.9694	ug/L	100
51) Dibromomethane	11.4358	93	148264	53.2284	ug/L	100
52) 2-Chloroethyl Vinyl Ether	11.6219	63	121055	51.2999	ug/L	100
53) 4-Methyl-2-Pentanone	11.6529	58	87502	52.0479	ug/L	100
54) cis-1,3-Dichloropropene	11.9424	75	356078	54.8246	ug/L	100
55) Dimethyl Disulfide	12.1906	79	211371	48.4027	ug/L	100
58) Toluene	12.3456	91	942130	53.0989	ug/L	100
59) Ethyl Methacrylate	12.4284	69	273140	54.5033	ug/L	100
60) trans-1,3-Dichloropropene	12.5007	75	321046	55.2645	ug/L	100
61) 1,1,2-Trichloroethane	12.7075	97	194552	51.5072	ug/L	100
62) 2-Hexanone	12.6455	43	151006	53.7863	ug/L	100
63) 1,3-Dichloropropane	12.9970	76	319726	52.4637	ug/L	100
64) Tetrachloroethene	13.1108	164	196155	52.1203	ug/L	100
65) Dibromochloromethane	13.3589	129	236412	54.9823	ug/L	100
66) 1,2-Dibromoethane	13.5967	107	193590	53.4779	ug/L	100
67) 1-Chlorohexane	13.6691	91	311199	53.1257	ug/L	100
68) Chlorobenzene	14.0620	112	654155	52.9562	ug/L	100
69) 1,1,1,2-Tetrachloroethane	14.0930	131	262859	50.0692	ug/L	100
70) Ethylbenzene	14.0827	106	345055	54.9095	ug/L	100
71) m-,p-Xylene	14.1654	106	820245	108.1189	ug/L	100
72) o-Xylene	14.6927	106	402899	54.7353	ug/L	100
73) Styrene	14.7341	104	685057	56.2652	ug/L	100
74) Bromoform	15.1994	173	156045	50.3375	ug/L	100
75) Isopropylbenzene	15.0856	105	1015763	55.1237	ug/L	100
77) 1,1,2,2-Tetrachloroethane	15.2924	83	225117	57.9246	ug/L	100
79) 1,2,3-Trichloropropane	15.4682	110	71864	55.3794	ug/L	100
80) trans-1,4-Dichloro-2-Butene	15.5096	53	91132	55.6989	ug/L	100
81) n-Propylbenzene	15.5612	91	1229035	55.9420	ug/L	100
82) Bromobenzene	15.6853	156	293574	54.7163	ug/L	100
83) 1,3,5-Trimethylbenzene	15.7370	105	875594	56.1401	ug/L	100
84) 2-Chlorotoluene	15.8197	91	849903	54.3656	ug/L	100
85) 4-Chlorotoluene	15.8611	91	707826	55.0893	ug/L	100
86) a-Methylstyrene	16.1092	118	496169	55.8694	ug/L	100
87) tert-Butylbenzene	16.1713	134	202064	52.6013	ug/L	100
88) 1,2,4-Trimethylbenzene	16.2126	105	901561	56.0078	ug/L	100
89) sec-Butylbenzene	16.4194	105	1092033	56.3983	ug/L	100
90) p-Isopropyltoluene	16.5642	119	957350	56.8606	ug/L	100
91) 1,3-Dichlorobenzene	16.7503	146	546574	55.2220	ug/L	100
92) 1,4-Dichlorobenzene	16.8640	146	549748	55.6213	ug/L	100
93) n-Butylbenzene	17.0501	91	884045	58.2620	ug/L	100
94) 1,2-Dichlorobenzene	17.3293	146	527196	55.5760	ug/L	100
95) 1,2-Dibromo-3-Chloropropane	18.2495	75	44949	55.9860	ug/L	100
96) 1,2,4-Trichlorobenzene	19.3145	180	343464	55.9713	ug/L	100
97) Hexachlorobutadiene	19.4489	225	169902	54.9333	ug/L	100
98) Naphthalene	19.6557	128	759495	57.7999	ug/L	100
99) 1,2,3-Trichlorobenzene	19.9452	180	324460	56.2784	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29743.D 8260WT.M Wed Mar 06 14:03:40 2019

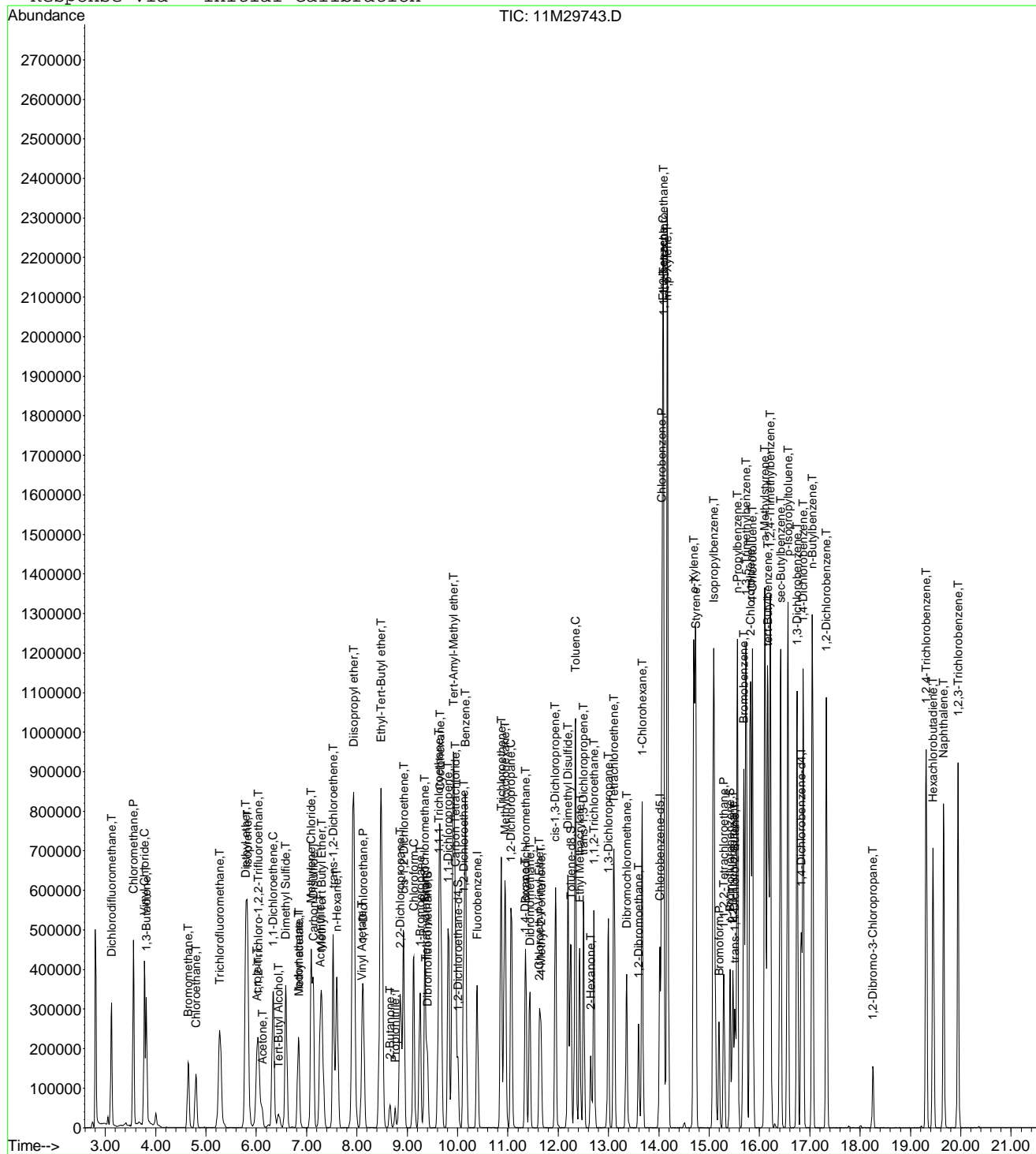
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29743.D
Acq On : 5 Mar 2019 19:38
Sample : WG698387-08 STD 50ug/L 8260
Misc : 1,1 STD92320
MS Integration Params: rteint.p
Quant Time: Mar 6 14:03 2019

Vial: 8
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:00:17 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29744.D Vial: 9
 Acq On : 5 Mar 2019 20:08 Operator: KFR
 Sample : WG698387-09 STD 100ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:41 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3914	96	415385	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0206	117	311453	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	157604	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3988	111	247388	51.5937	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	= 206.3748%#		
43) 1,2-Dichloroethane-d4	10.0089	65	291480	50.6647	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	= 202.6588%#		
57) Toluene-d8	12.2422	98	789829	50.9572	ug/L	-0.0104
Spiked Amount	25.0000	Range 88 - 110	Recovery	= 203.8288%#		
78) p-Bromofluorobenzene	15.4165	95	304258	50.4473	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	= 201.7892%#		

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1124	85	632796	103.1219	ug/L	100
3) Chloromethane	3.5570	50	1030983	87.4757	ug/L	97
4) Vinyl Chloride	3.7741	62	814275	92.5481	ug/L	99
5) 1,3-Butadiene	3.8051	54	301777	102.9245	ug/L	98
6) Bromomethane	4.6426	94	314715	93.3626	ug/L	99
7) Chloroethane	4.7977	64	343303	100.9294	ug/L	100
8) Trichlorofluoromethane	5.2630	101	851180	102.2313	ug/L	100
9) Diethyl ether	5.7800	59	893245	205.2050	ug/L	99
10) Isoprene	5.8213	67	691488	105.4440	ug/L	99
11) Acrolein	6.0178	56	77209	102.1405	ug/L	99
12) 1,1,2-Trichloro-1,2,2-Trif	6.0281	101	421860	103.2893	ug/L	99
13) Acetone	6.1108	43	141631	93.2631	ug/L	99
14) 1,1-Dichloroethene	6.3280	61	821836	103.3889	ug/L	99
15) Tert-Butyl Alcohol	6.4417	59	176386	416.4125	ug/L	100
16) Dimethyl Sulfide	6.5865	62	541839	103.1125	ug/L	100
17) Iodomethane	6.8243	142	539332	97.1378	ug/L	100
18) Methyl acetate	6.8449	43	337408	103.0045	ug/L	99
19) Methylene Chloride	7.0828	84	452640	102.1832	ug/L	96
20) Carbon Disulfide	7.1241	76	1289051	105.5820	ug/L	100
21) Acrylonitrile	7.2689	53	182195	108.8308	ug/L	97
22) Methyl Tert Butyl Ether	7.2896	73	1272204	101.2824	ug/L	100
23) trans-1,2-Dichloroethene	7.5274	96	459624	106.7422	ug/L	100
24) n-Hexane	7.5997	57	687134	103.6054	ug/L	100
25) Diisopropyl ether	7.9306	45	3237070	199.9805	ug/L	96
26) Vinyl Acetate	8.0960	43	563662	95.4835	ug/L	99
27) 1,1-Dichloroethane	8.1167	63	968353	103.0600	ug/L	99
28) Ethyl-Tert-Butyl ether	8.4786	59	3270320	199.8843	ug/L	95
29) 2-Butanone	8.6544	43	194976	100.7950	ug/L	99
30) Propionitrile	8.7578	54	117851	208.5238	ug/L	97
31) 2,2-Dichloropropane	8.8612	77	694068	106.8349	ug/L	99
32) cis-1,2-Dichloroethene	8.9232	96	524813	107.3575	ug/L	98
33) Chloroform	9.1197	83	855826	100.3513	ug/L	100
34) 1-Bromopropane	9.2541	122	106621	100.6966	ug/L	98
35) Bromochloromethane	9.3471	130	308016	107.4596	ug/L	99
36) Tetrahydrofuran	9.3678	42	243433	194.2451	ug/L	99
38) 1,1,1-Trichloroethane	9.6263	97	845315	109.4669	ug/L	99
39) Cyclohexane	9.6573	56	950015	103.9122	ug/L	99
40) 1,1-Dichloropropene	9.8124	75	614532	105.8434	ug/L	100
41) Carbon Tetrachloride	9.9572	117	765034	116.5036	ug/L	100
42) Tert-Amyl-Methyl ether	9.9158	73	2495067	204.4640	ug/L	98

(#) = qualifier out of range (m) = manual integration
 11M29744.D 8260WT.M Wed Mar 06 14:03:42 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29744.D Vial: 9
 Acq On : 5 Mar 2019 20:08 Operator: KFR
 Sample : WG698387-09 STD 100ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:41 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1226	62	766097	103.3544	ug/L	99
45) Benzene	10.1536	78	1727579	100.1070	ug/L	98
46) Trichloroethene	10.8671	130	521978	106.6462	ug/L	99
47) Methylcyclohexane	10.9394	83	723217	104.8882	ug/L	99
48) 1,2-Dichloropropane	11.0635	63	534603	104.2861	ug/L	100
49) 1,4-Dioxane	11.3427	88	14544	413.4821	ug/L	92
50) Bromodichloromethane	11.3530	83	675450	111.4914	ug/L	99
51) Dibromomethane	11.4357	93	305931	104.4754	ug/L	99
52) 2-Chloroethyl Vinyl Ether	11.6218	63	256622	103.4451	ug/L	99
53) 4-Methyl-2-Pentanone	11.6529	58	173401	98.1114	ug/L	98
54) cis-1,3-Dichloropropene	11.9424	75	744841	109.0880	ug/L	99
55) Dimethyl Disulfide	12.1905	79	465102	98.6792	ug/L	96
58) Toluene	12.3456	91	1836635	99.5467	ug/L	96
59) Ethyl Methacrylate	12.4283	69	561442	107.7387	ug/L	100
60) trans-1,3-Dichloropropene	12.5007	75	669902	110.8968	ug/L	100
61) 1,1,2-Trichloroethane	12.7075	97	391481	99.6716	ug/L	99
62) 2-Hexanone	12.6455	43	296376	101.5195	ug/L	99
63) 1,3-Dichloropropane	12.9970	76	645987	101.9374	ug/L	100
64) Tetrachloroethene	13.1107	164	408078	104.2748	ug/L	99
65) Dibromochloromethane	13.3589	129	489986	109.5887	ug/L	100
66) 1,2-Dibromoethane	13.5967	107	391461	103.9941	ug/L	100
67) 1-Chlorohexane	13.6691	91	645592	105.9872	ug/L	100
68) Chlorobenzene	14.0620	112	1326225	103.2482	ug/L	97
69) 1,1,1,2-Tetrachloroethane	14.0930	131	572327	104.1068	ug/L	100
70) Ethylbenzene	14.0826	106	737666	112.8879	ug/L	82
71) m-,p-Xylene	14.1654	106	1624155	205.8801	ug/L	80
72) o-Xylene	14.6927	106	821160	107.2822	ug/L	91
73) Styrene	14.7341	104	1354826	107.0102	ug/L	97
74) Bromoform	15.1993	173	320623	98.7515	ug/L	99
75) Isopropylbenzene	15.0856	105	1885676	98.4106	ug/L	96
77) 1,1,2,2-Tetrachloroethane	15.2924	83	420749	102.3370	ug/L	100
79) 1,2,3-Trichloropropane	15.4682	110	139039	101.2812	ug/L	75
80) trans-1,4-Dichloro-2-Butene	15.5095	53	186906	107.9826	ug/L	93
81) n-Propylbenzene	15.5612	91	2168390	93.2966	ug/L	94
82) Bromobenzene	15.6853	156	583544	102.8081	ug/L	99
83) 1,3,5-Trimethylbenzene	15.7370	105	1636117	99.1607	ug/L	96
84) 2-Chlorotoluene	15.8197	91	1595448	96.4699	ug/L	97
85) 4-Chlorotoluene	15.8611	91	1310872	96.4398	ug/L	96
86) a-Methylstyrene	16.1092	118	1012308	107.7487	ug/L	100
87) tert-Butylbenzene	16.1713	134	408998	100.6430	ug/L	93
88) 1,2,4-Trimethylbenzene	16.2126	105	1669192	98.0200	ug/L	94
89) sec-Butylbenzene	16.4194	105	1973074	96.3226	ug/L	95
90) p-Isopropyltoluene	16.5642	119	1752347	98.3820	ug/L	94
91) 1,3-Dichlorobenzene	16.7503	146	1063781	101.5945	ug/L	98
92) 1,4-Dichlorobenzene	16.8640	146	1063224	101.6850	ug/L	98
93) n-Butylbenzene	17.0501	91	1641647	102.2695	ug/L	96
94) 1,2-Dichlorobenzene	17.3293	146	1013857	101.0291	ug/L	98
95) 1,2-Dibromo-3-Chloropropane	18.2495	75	87196	102.6623	ug/L	98
96) 1,2,4-Trichlorobenzene	19.3145	180	701877	108.1184	ug/L	98
97) Hexachlorobutadiene	19.4489	225	349257	106.7424	ug/L	100
98) Naphthalene	19.6557	128	1438608	103.4903	ug/L	98
99) 1,2,3-Trichlorobenzene	19.9452	180	665078	109.0455	ug/L	98

(#) = qualifier out of range (m) = manual integration
 11M29744.D 8260WT.M Wed Mar 06 14:03:42 2019

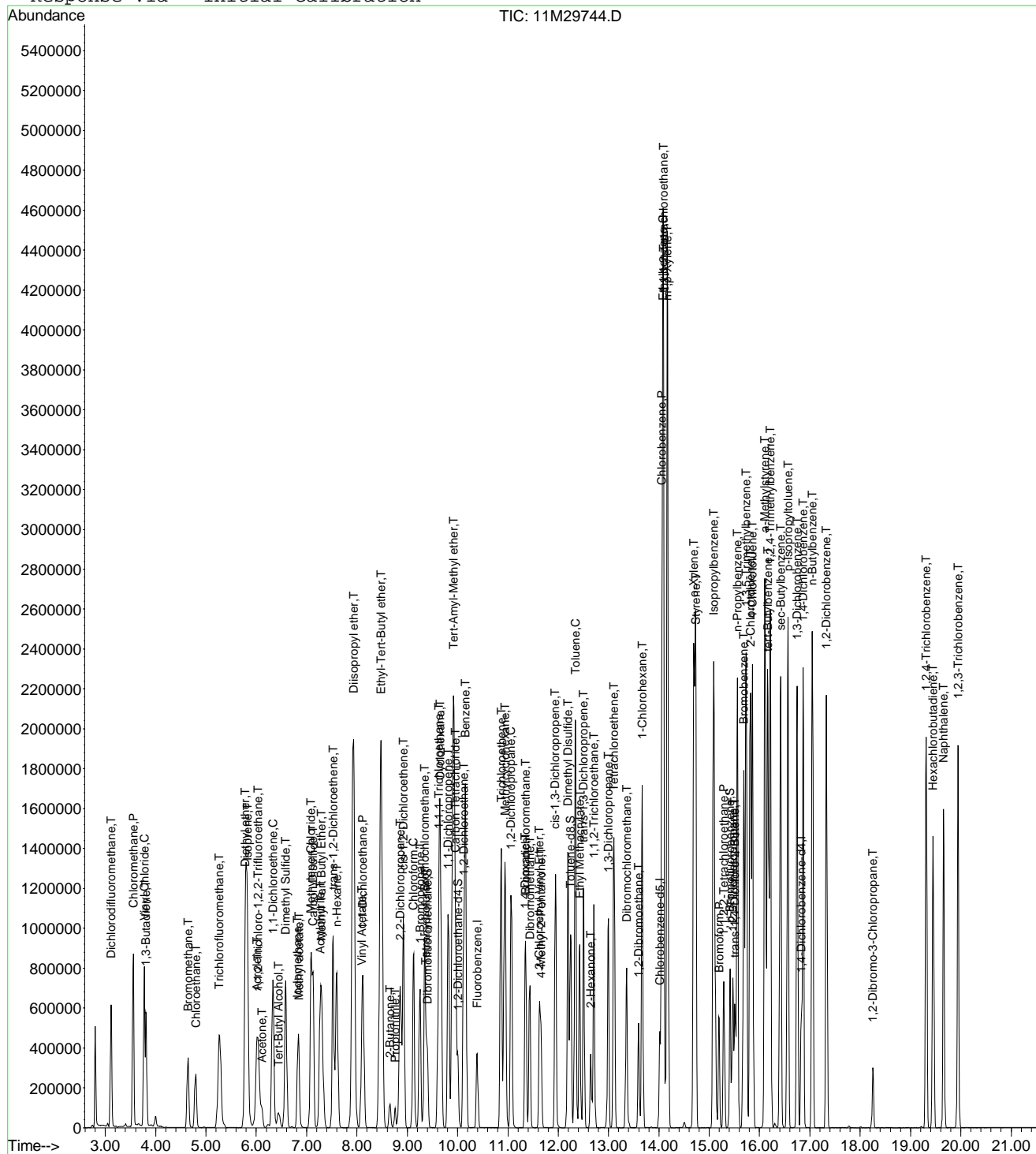
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29744.D
Acq On : 5 Mar 2019 20:08
Sample : WG698387-09 STD 100ug/L 8260
Misc : 1,1 STD92320
MS Integration Params: rteint.p
Quant Time: Mar 6 14:03 2019

Vial: 9
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:00:17 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29745.D Vial: 10
 Acq On : 5 Mar 2019 20:37 Operator: KFR
 Sample : WG698387-10 STD 200ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:42 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3913	96	428361	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0205	117	327954	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8329	152	168167	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3987	111	490236	99.1434	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	= 396.5736%#		
43) 1,2-Dichloroethane-d4	10.0087	65	573937	96.7391	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	= 386.9564%#		
57) Toluene-d8	12.2421	98	1542774	94.5267	ug/L	-0.0105
Spiked Amount	25.0000	Range 88 - 110	Recovery	= 378.1068%#		
78) p-Bromofluorobenzene	15.4164	95	618324	96.0812	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	= 384.3248%#		

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1019	85	1270961	200.8447	ug/L	99
3) Chloromethane	3.5465	50	1885900	155.1656	ug/L	90
4) Vinyl Chloride	3.7637	62	1472282	162.2664	ug/L	95
5) 1,3-Butadiene	3.8050	54	596563	202.6036	ug/L	98
6) Bromomethane	4.6322	94	692246	199.1392	ug/L	99
7) Chloroethane	4.7873	64	726036	206.9852	ug/L	99
8) Trichlorofluoromethane	5.2629	101	1743409	203.0499	ug/L	98
10) Isoprene	5.8109	67	1413331	208.9882	ug/L	100
11) Acrolein	6.0177	56	485	0.6222	ug/L	85
12) 1,1,2-Trichloro-1,2,2-Trif	6.0280	101	889556	211.2035	ug/L	99
13) Acetone	6.1107	43	303071	193.5249	ug/L	99
14) 1,1-Dichloroethene	6.3279	61	1661278	202.6618	ug/L	97
16) Dimethyl Sulfide	6.5864	62	1131090	208.7272	ug/L	99
17) Iodomethane	6.8242	142	1178735	204.1959	ug/L	99
18) Methyl acetate	6.8449	43	715518	211.8177	ug/L	98
19) Methylene Chloride	7.0827	84	948379	207.6105	ug/L	93
20) Carbon Disulfide	7.1240	76	2465515	195.8251	ug/L	96
21) Acrylonitrile	7.2895	53	33298	19.2874	ug/L #	34
22) Methyl Tert Butyl Ether	7.2895	73	2546443	196.5857	ug/L	97
23) trans-1,2-Dichloroethene	7.5169	96	969505	218.3356	ug/L	95
24) n-Hexane	7.5893	57	1438700	210.3545	ug/L	99
26) Vinyl Acetate	8.0959	43	1350499	219.2172	ug/L	99
27) 1,1-Dichloroethane	8.1166	63	1931262	199.3144	ug/L	96
29) 2-Butanone	8.6543	43	399832	200.4363	ug/L	99
31) 2,2-Dichloropropane	8.8611	77	1412579	210.8457	ug/L	98
32) cis-1,2-Dichloroethene	8.9231	96	1084862	215.2005	ug/L	95
33) Chloroform	9.1196	83	1707865	194.1923	ug/L	98
34) 1-Bromopropane	9.2540	122	225233	206.2743	ug/L	98
35) Bromochloromethane	9.3470	130	655255	221.6782	ug/L	96
36) Tetrahydrofuran	9.3677	42	1370	1.0601	ug/L #	37
38) 1,1,1-Trichloroethane	9.6262	97	1747688	219.4668	ug/L	100
39) Cyclohexane	9.6572	56	1958321	207.7117	ug/L	97
40) 1,1-Dichloropropene	9.8123	75	1256934	209.9291	ug/L	97
41) Carbon Tetrachloride	9.9571	117	1537293	227.0158	ug/L	99
44) 1,2-Dichloroethane	10.1225	62	1526389	199.6876	ug/L	96
45) Benzene	10.1535	78	3080240	173.0820	ug/L	89
46) Trichloroethene	10.8669	130	1066911	211.3791	ug/L	99
47) Methylcyclohexane	10.9393	83	1499216	210.8449	ug/L	99
48) 1,2-Dichloropropane	11.0634	63	1110542	210.0733	ug/L	100
49) 1,4-Dioxane	11.3529	88	1143	31.5108	ug/L #	11

(#) = qualifier out of range (m) = manual integration
 11M29745.D 8260WT.M Wed Mar 06 14:03:43 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29745.D Vial: 10
 Acq On : 5 Mar 2019 20:37 Operator: KFR
 Sample : WG698387-10 STD 200ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:42 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
50) Bromodichloromethane	11.3529	83	1384146	221.5497	ug/L	98
51) Dibromomethane	11.4356	93	641505	212.4376	ug/L	98
52) 2-Chloroethyl Vinyl Ether	11.6217	63	559641	218.7593	ug/L	99
53) 4-Methyl-2-Pentanone	11.6527	58	363640	199.5172	ug/L	99
54) cis-1,3-Dichloropropene	11.9423	75	1507465	214.0925	ug/L	96
55) Dimethyl Disulfide	12.1904	79	1001200	203.3688	ug/L	90
58) Toluene	12.3455	91	3075154	158.2889	ug/L	84
59) Ethyl Methacrylate	12.4282	69	1165365	212.3774	ug/L	99
60) trans-1,3-Dichloropropene	12.5006	75	1369222	215.2590	ug/L	97
61) 1,1,2-Trichloroethane	12.7074	97	820697	198.4373	ug/L	99
62) 2-Hexanone	12.6454	43	621972	202.3284	ug/L	100
63) 1,3-Dichloropropane	12.9969	76	1304284	195.4617	ug/L	99
64) Tetrachloroethene	13.1106	164	883638	214.4323	ug/L	99
65) Dibromochloromethane	13.3588	129	1042382	221.4055	ug/L	99
66) 1,2-Dibromoethane	13.5966	107	818783	206.5706	ug/L	99
67) 1-Chlorohexane	13.6690	91	1364610	212.7569	ug/L	99
68) Chlorobenzene	14.0619	112	2493144	184.3284	ug/L	86
69) 1,1,1,2-Tetrachloroethane	14.0929	131	1269144	218.5028	ug/L	99
70) Ethylbenzene	14.0825	106	1593549	231.5969	ug/L	48
71) m-,p-Xylene	14.1653	106	2860586	344.3669	ug/L	61
72) o-Xylene	14.6926	106	1700846	211.0302	ug/L	71
73) Styrene	14.7340	104	2503558	187.7929	ug/L	86
74) Bromoform	15.1992	173	699412	203.7971	ug/L	98
75) Isopropylbenzene	15.0855	105	3148871	156.0664	ug/L	79
77) 1,1,2,2-Tetrachloroethane	15.2923	83	913096	208.1385	ug/L	99
79) 1,2,3-Trichloropropane	15.4681	110	292113	199.4203	ug/L	76
80) trans-1,4-Dichloro-2-Butene	15.5094	53	412571	223.3859	ug/L	89
81) n-Propylbenzene	15.5611	91	3413475	137.6422	ug/L #	75
82) Bromobenzene	15.6852	156	1220551	201.5284	ug/L	70
83) 1,3,5-Trimethylbenzene	15.7369	105	2828431	160.6562	ug/L	81
84) 2-Chlorotoluene	15.8196	91	2758682	156.3283	ug/L	83
85) 4-Chlorotoluene	15.8610	91	2248327	155.0179	ug/L	78
86) a-Methylstyrene	16.1091	118	1990995	198.6077	ug/L	92
87) tert-Butylbenzene	16.1712	134	903332	208.3225	ug/L	73
88) 1,2,4-Trimethylbenzene	16.2125	105	2863604	157.5970	ug/L	76
89) sec-Butylbenzene	16.4193	105	3264819	149.3724	ug/L #	78
90) p-Isopropyltoluene	16.5641	119	2989514	157.2977	ug/L #	77
91) 1,3-Dichlorobenzene	16.7502	146	2068616	185.1503	ug/L	90
92) 1,4-Dichlorobenzene	16.8742	146	2057069	184.3773	ug/L	89
93) n-Butylbenzene	17.0500	91	2833678	165.4410	ug/L #	82
94) 1,2-Dichlorobenzene	17.3292	146	1997899	186.5820	ug/L	90
95) 1,2-Dibromo-3-Chloropropane	18.2494	75	196511	216.8342	ug/L	99
96) 1,2,4-Trichlorobenzene	19.3144	180	1528702	220.6928	ug/L	93
97) Hexachlorobutadiene	19.4488	225	808980	231.7160	ug/L	99
98) Naphthalene	19.6556	128	2578020	173.8080	ug/L #	89
99) 1,2,3-Trichlorobenzene	19.9451	180	1450350	222.8612	ug/L	94

(#) = qualifier out of range (m) = manual integration
 11M29745.D 8260WT.M Wed Mar 06 14:03:43 2019

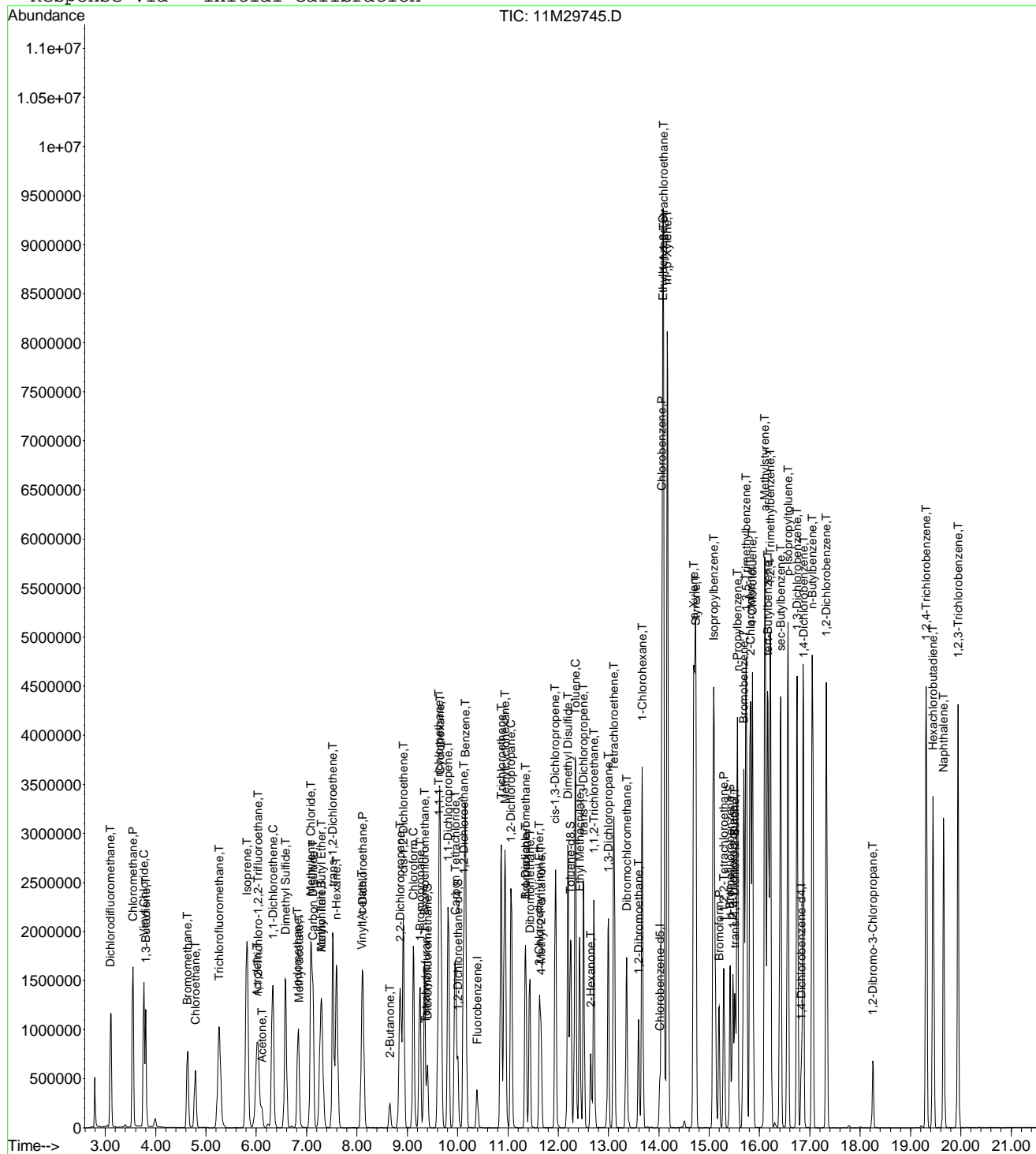
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29745.D
Acq On : 5 Mar 2019 20:37
Sample : WG698387-10 STD 200ug/L 8260
Misc : 1,1 STD92320
MS Integration Params: rteint.p
Quant Time: Mar 6 14:03 2019

Vial: 10
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:00:17 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29746.D Vial: 11
 Acq On : 5 Mar 2019 21:06 Operator: KFR
 Sample : WG698387-11 STD 300ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:44 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3813	96	427056	25.0000	ug/L	-0.0102
56) Chlorobenzene-d5	14.0208	117	316685	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8332	152	153710	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	0.0000	111	0	0.0000	ug/L	
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	0.0000%#	
43) 1,2-Dichloroethane-d4	9.9987	65	490	0.0828	ug/L	-0.0102
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	0.3312%#	
57) Toluene-d8	12.1907	98	8364	0.5307	ug/L	-0.0619
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	2.1228%#	
78) p-Bromofluorobenzene	15.4166	95	2545	0.4327	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	1.7308%#	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1125	85	5406	0.8569	ug/L	96
3) Chloromethane	3.5468	50	58494	4.8274	ug/L	98
4) Vinyl Chloride	3.7639	62	2436	0.2693	ug/L #	16
5) 1,3-Butadiene	3.8053	54	834854	286.7325	ug/L	98
6) Bromomethane	4.6325	94	106548	30.7445	ug/L	100
8) Trichlorofluoromethane	5.2632	101	2791	0.3261	ug/L	85
9) Diethyl ether	5.7801	59	1428327	319.1619	ug/L	98
10) Isoprene	5.8112	67	2166154	321.2864	ug/L	99
11) Acrolein	6.0076	56	121549	156.4038	ug/L	95
12) 1,1,2-Trichloro-1,2,2-Trif	6.0490	101	1522	0.3625	ug/L	77
13) Acetone	6.1110	43	422798	270.8011	ug/L	98
14) 1,1-Dichloroethene	6.3178	61	1985	0.2429	ug/L #	27
15) Tert-Butyl Alcohol	6.4419	59	292797	672.3451	ug/L	99
16) Dimethyl Sulfide	6.5763	62	1679640	310.9018	ug/L	99
17) Iodomethane	6.8244	142	1404276	243.7192	ug/L	99
18) Methyl acetate	6.8451	43	1092692	324.4626	ug/L	98
19) Methylene Chloride	7.0829	84	1415	0.3107	ug/L	92
20) Carbon Disulfide	7.1243	76	3236660	257.8593	ug/L	91
21) Acrylonitrile	7.2587	53	254755	148.0145	ug/L	90
22) Methyl Tert Butyl Ether	7.2897	73	2841	0.2200	ug/L #	1
23) trans-1,2-Dichloroethene	7.5275	96	2279	0.5148	ug/L	84
24) n-Hexane	7.5896	57	2052381	300.9988	ug/L	97
25) Diisopropyl ether	7.9308	45	4498959	270.3420	ug/L	90
26) Vinyl Acetate	8.0859	43	1756570	285.3988	ug/L	99
28) Ethyl-Tert-Butyl ether	8.4788	59	4544638	270.1803	ug/L	88
29) 2-Butanone	8.6546	43	613943	308.7109	ug/L	99
30) Propionitrile	8.7579	54	189351	325.8787	ug/L	99
32) cis-1,2-Dichloroethene	8.9234	96	1148	0.2284	ug/L	93
34) 1-Bromopropane	9.2542	122	345495	317.3803	ug/L	97
36) Tetrahydrofuran	9.3680	42	373832	290.1436	ug/L	99
39) Cyclohexane	9.6575	56	2733471	290.8149	ug/L	95
40) 1,1-Dichloropropene	9.8126	75	2367	0.3965	ug/L #	53
41) Carbon Tetrachloride	9.9470	117	960	0.1422	ug/L #	69
42) Tert-Amyl-Methyl ether	9.9160	73	3533814	281.6724	ug/L	93
44) 1,2-Dichloroethane	10.1124	62	1085	0.1424	ug/L #	40
45) Benzene	10.1538	78	3944	0.2223	ug/L	88
46) Trichloroethene	10.8672	130	1750	0.3478	ug/L	91
47) Methylcyclohexane	10.9396	83	2186326	308.4174	ug/L	97
49) 1,4-Dioxane	11.3428	88	22511	622.4917	ug/L	93
51) Dibromomethane	11.4255	93	486	0.1614	ug/L #	57

(#) = qualifier out of range (m) = manual integration
 11M29746.D 8260WT.M Wed Mar 06 14:03:45 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29746.D Vial: 11
 Acq On : 5 Mar 2019 21:06 Operator: KFR
 Sample : WG698387-11 STD 300ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:03:44 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
52) 2-Chloroethyl Vinyl Ether	11.6220	63	864552	338.9793	ug/L	98
53) 4-Methyl-2-Pentanone	11.6530	58	570189	313.7998	ug/L	99
54) cis-1,3-Dichloropropene	11.9425	75	1075	0.1531	ug/L #	43
55) Dimethyl Disulfide	12.1907	79	1475968	299.5697	ug/L	79
58) Toluene	12.3354	91	4554	0.2428	ug/L	97
59) Ethyl Methacrylate	12.4285	69	1732998	327.0618	ug/L	99
60) trans-1,3-Dichloropropene	12.5009	75	2259	0.3678	ug/L #	51
62) 2-Hexanone	12.6456	43	923582	311.1336	ug/L	97
64) Tetrachloroethene	13.1212	164	2035	0.5114	ug/L	95
66) 1,2-Dibromoethane	13.5969	107	566	0.1479	ug/L #	41
67) 1-Chlorohexane	13.6692	91	1936253	312.6242	ug/L	99
68) Chlorobenzene	14.0622	112	4481	0.3431	ug/L	86
70) Ethylbenzene	14.0828	106	2492	0.3751	ug/L	95
71) m-,p-Xylene	14.1759	106	5977	0.7451	ug/L	90
72) o-Xylene	14.6929	106	2419	0.3108	ug/L	99
73) Styrene	14.7342	104	4055	0.3150	ug/L	80
75) Isopropylbenzene	15.0858	105	14600	0.7494	ug/L	97
80) trans-1,4-Dichloro-2-Buten	15.5097	53	548795	325.0918	ug/L #	12
81) n-Propylbenzene	15.5097	91	118053	5.2080	ug/L #	59
82) Bromobenzene	15.6854	156	1717	0.3102	ug/L	80
83) 1,3,5-Trimethylbenzene	15.7372	105	6615	0.4111	ug/L	95
84) 2-Chlorotoluene	15.8612	91	13520	0.8382	ug/L	98
85) 4-Chlorotoluene	15.8612	91	13517	1.0196	ug/L	96
86) a-Methylstyrene	16.1094	118	2605115	284.3095	ug/L	88
87) tert-Butylbenzene	16.1714	134	1554	0.3921	ug/L #	1
88) 1,2,4-Trimethylbenzene	16.2128	105	7192	0.4330	ug/L	97
89) sec-Butylbenzene	16.4196	105	16306	0.8162	ug/L	95
90) p-Isopropyltoluene	16.5643	119	9508	0.5473	ug/L	95
92) 1,4-Dichlorobenzene	16.8642	146	7658	0.7510	ug/L #	47
93) n-Butylbenzene	17.0503	91	12522	0.7998	ug/L	94
94) 1,2-Dichlorobenzene	17.3295	146	4341	0.4435	ug/L	91
96) 1,2,4-Trichlorobenzene	19.3147	180	8523	1.3462	ug/L	94
97) Hexachlorobutadiene	19.4491	225	2841	0.8903	ug/L	84
98) Naphthalene	19.6559	128	12052	0.8890	ug/L #	94
99) 1,2,3-Trichlorobenzene	19.9454	180	6215	1.0448	ug/L	94

(#) = qualifier out of range (m) = manual integration
 11M29746.D 8260WT.M Wed Mar 06 14:03:45 2019

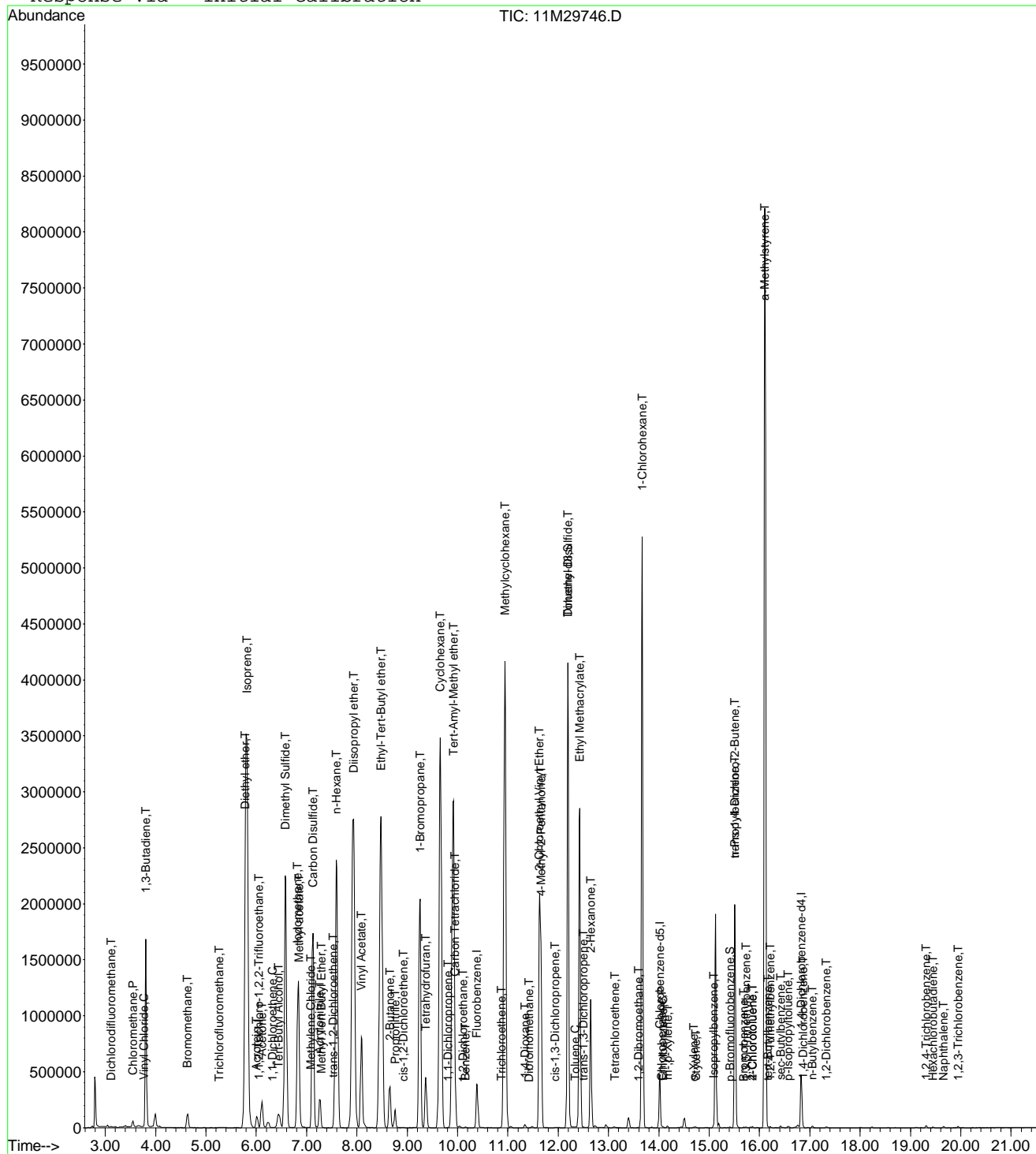
Page 2

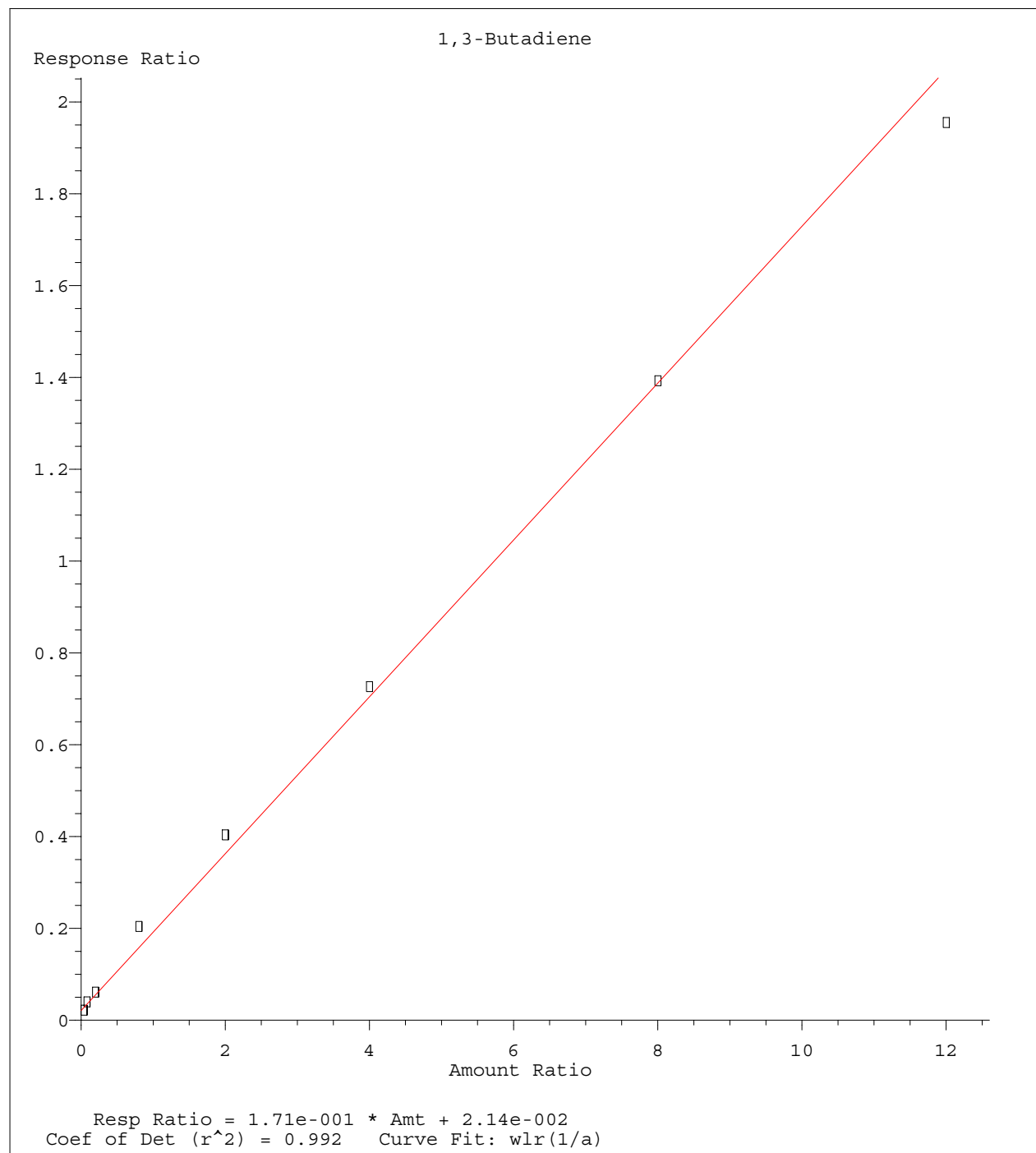
Data File : C:\MSDCHEM\1\DATA\030519\11M29746.D
 Acq On : 5 Mar 2019 21:06
 Sample : WG698387-11 STD 300ug/L 8260
 Misc : 1,1 STD92320
 MS Integration Params: rteint.p
 Quant Time: Mar 6 14:03 2019

Vial: 11
 Operator: KFR
 Inst : hpms11
 Multiplr: 1.00

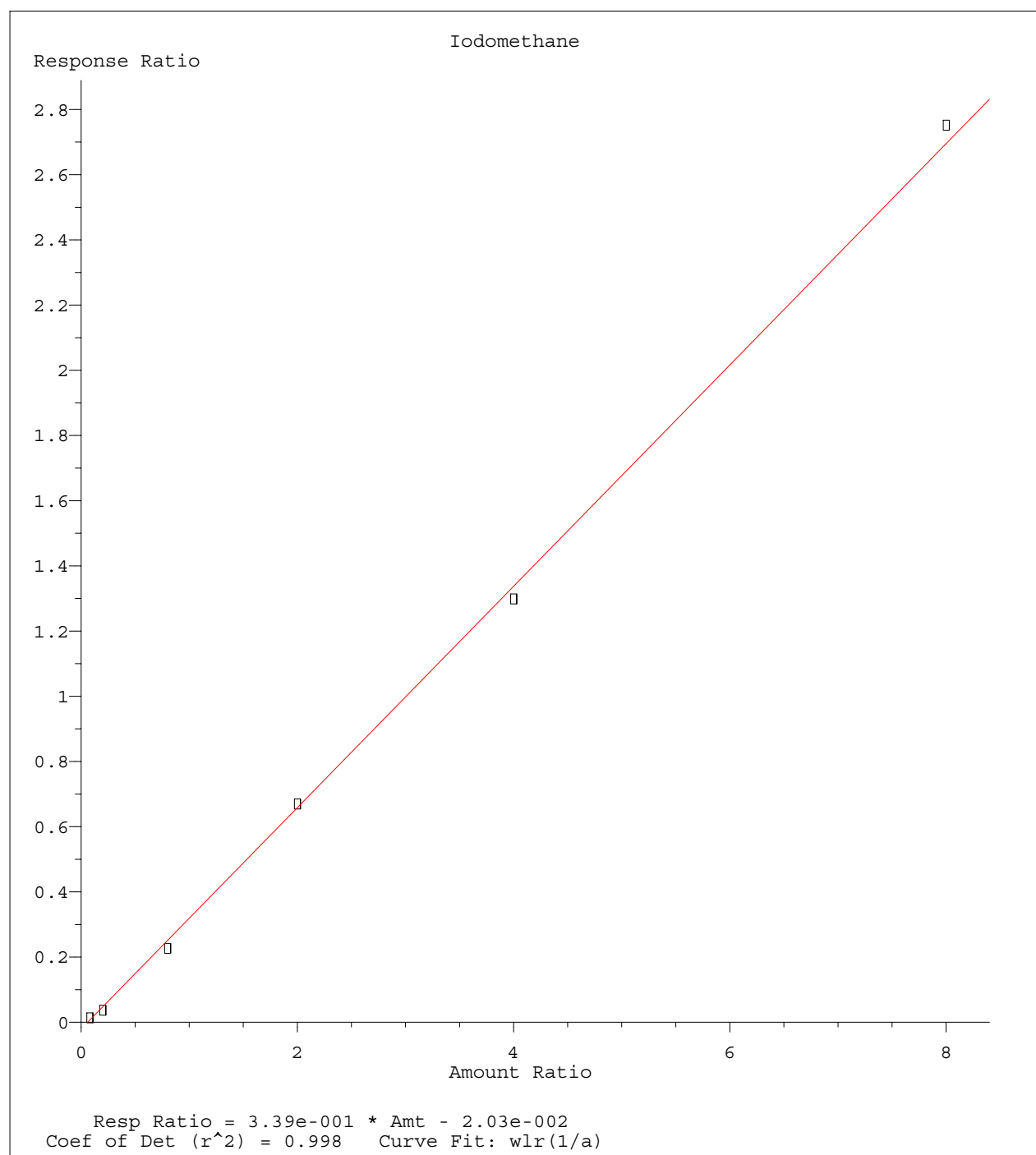
Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration

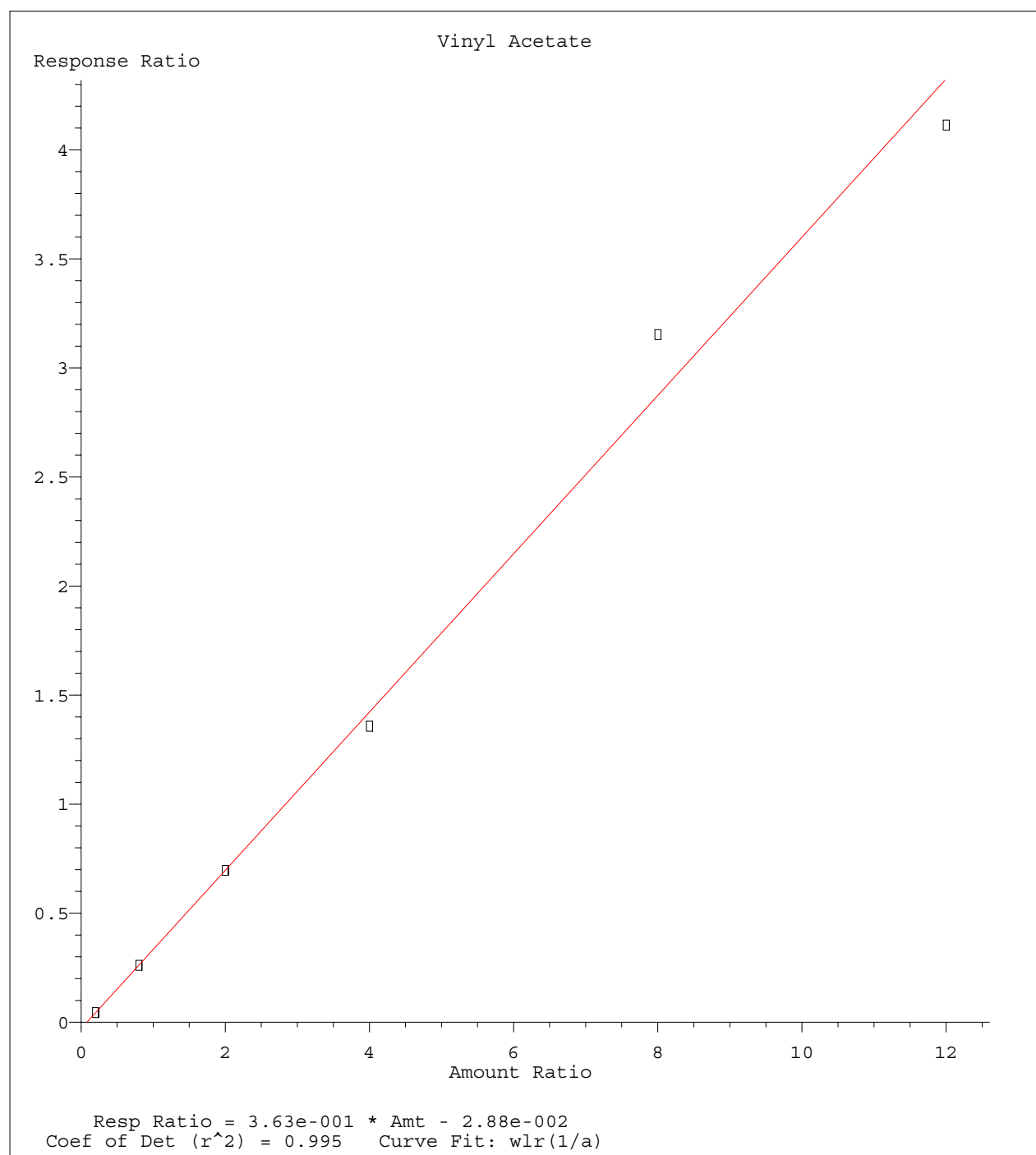




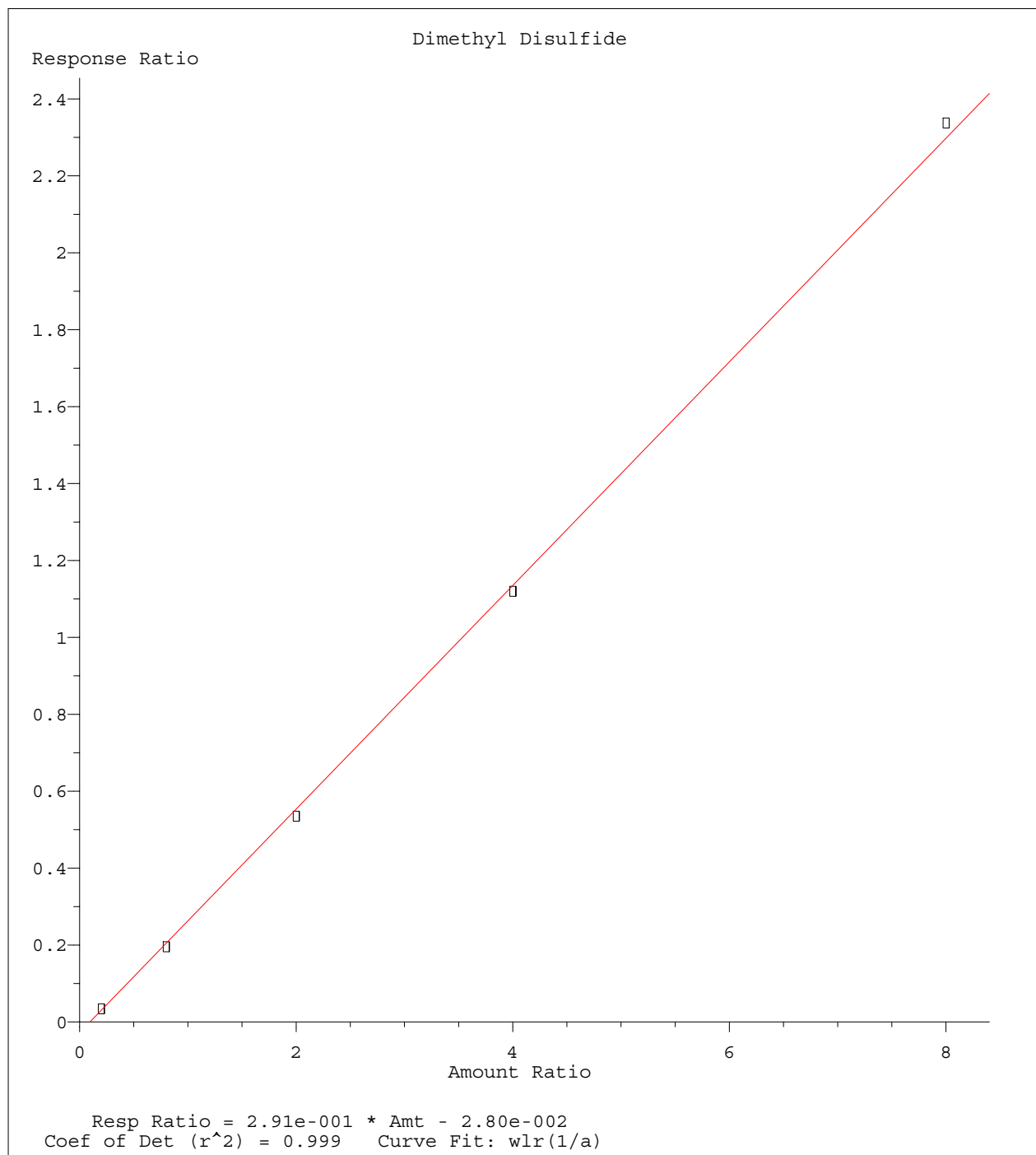
Method Name: C:\MSDCHEM\1\METHODS\8260WT.M
Calibration Table Last Updated: Wed Mar 06 14:13:24 2019



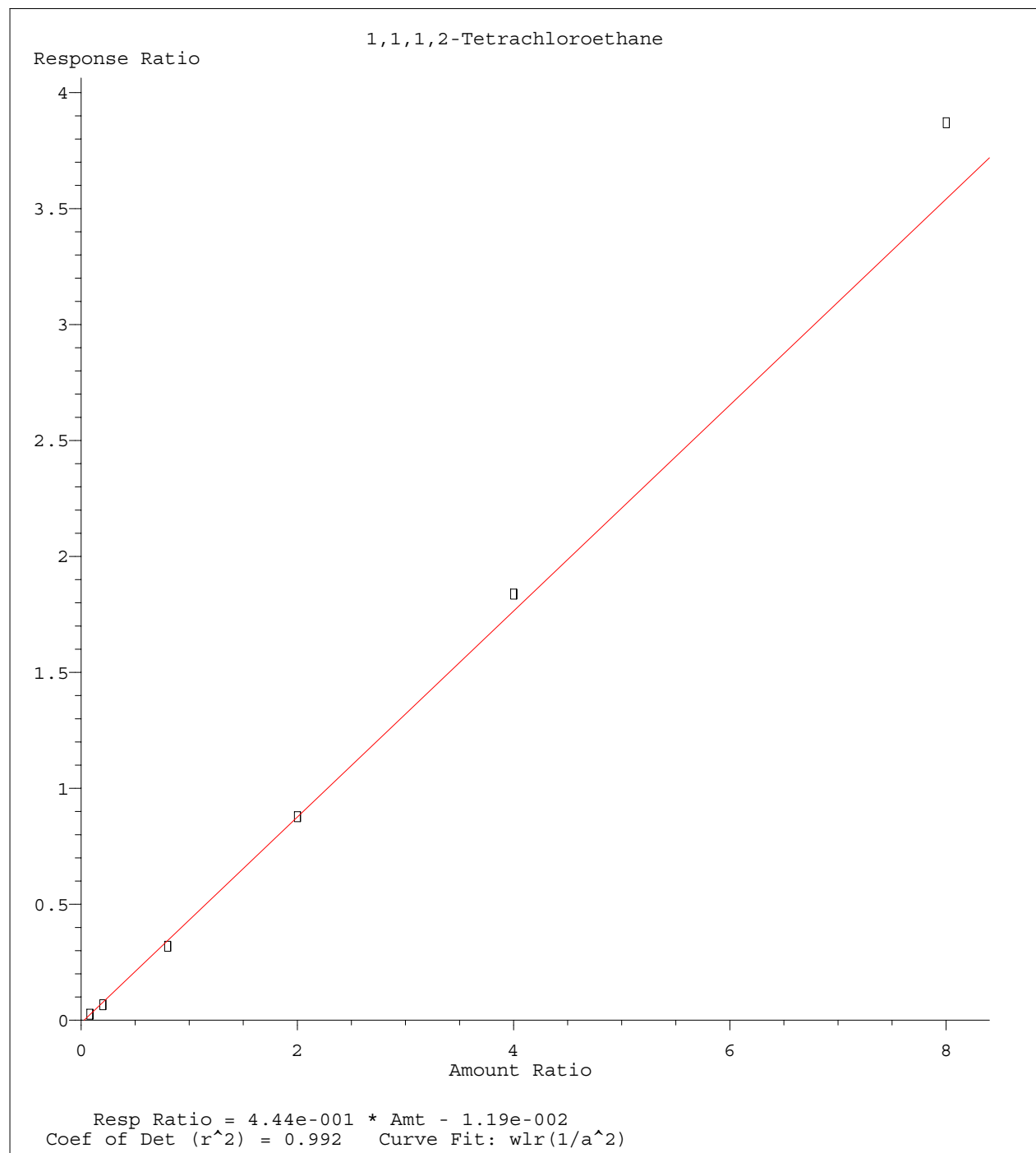
Method Name: C:\MSDCHEM\1\METHODS\8260WT.M
Calibration Table Last Updated: Wed Mar 06 14:13:24 2019



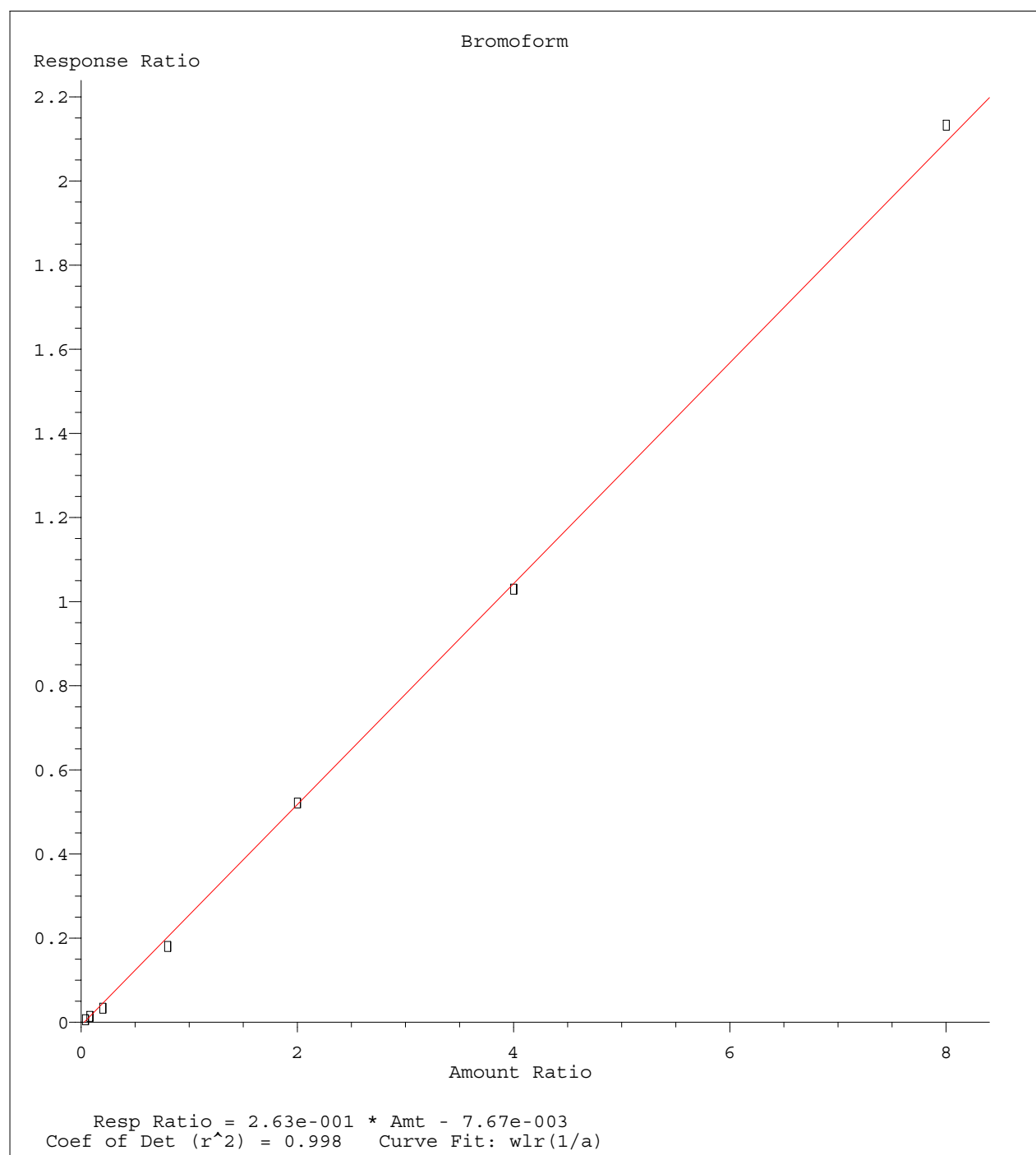
Method Name: C:\MSDCHEM\1\METHODS\8260WT.M
Calibration Table Last Updated: Wed Mar 06 14:13:24 2019



Method Name: C:\MSDCHEM\1\METHODS\8260WT.M
Calibration Table Last Updated: Wed Mar 06 14:13:24 2019



Method Name: C:\MSDCHEM\1\METHODS\8260WT.M
Calibration Table Last Updated: Wed Mar 06 14:13:24 2019



Method Name: C:\MSDCHEM\1\METHODS\8260WT.M
Calibration Table Last Updated: Wed Mar 06 14:13:24 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29748.D Vial: 13
 Acq On : 5 Mar 2019 22:04 Operator: KFR
 Sample : WG698387-12 ICV 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:04:07 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3811	96	386715	25.0000	ug/L	-0.0104
56) Chlorobenzene-d5	14.0206	117	292297	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	155467	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3988	111	111894	25.0660	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	100.2640%	
43) 1,2-Dichloroethane-d4	10.0089	65	135183	25.2394	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	100.9576%	
57) Toluene-d8	12.2422	98	362421	24.9146	ug/L	-0.0104
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	99.6584%	
78) p-Bromofluorobenzene	15.4165	95	145455	24.4486	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	97.7944%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1227	85	289735	50.7164	ug/L	100
3) Chloromethane	3.5570	50	475907	43.3728	ug/L	100
4) Vinyl Chloride	3.7741	62	369939	45.1634	ug/L	99
5) 1,3-Butadiene	3.8154	54	138904	47.9634	ug/L	99
6) Bromomethane	4.6323	94	132979	42.3739	ug/L	99
7) Chloroethane	4.7874	64	150096	47.3990	ug/L	98
8) Trichlorofluoromethane	5.2630	101	362897	46.8172	ug/L	100
9) Diethyl ether	5.7903	59	403945	99.6780	ug/L	99
10) Isoprene	5.8213	67	305673	50.0673	ug/L	99
11) Acrolein	6.0178	56	62254	88.4621	ug/L	98
12) 1,1,2-Trichloro-1,2,2-Trif	6.0281	101	193938	51.0046	ug/L	100
13) Acetone	6.1212	43	62519	44.2205	ug/L	98
14) 1,1-Dichloroethene	6.3280	61	378767	51.1824	ug/L	100
15) Tert-Butyl Alcohol	6.4624	59	93422	236.9019	ug/L	98
16) Dimethyl Sulfide	6.5865	62	271232	55.4424	ug/L	100
17) Iodomethane	6.8346	142	337280	65.7407	ug/L	97
18) Methyl acetate	6.8449	43	163842	53.7262	ug/L	98
19) Methylene Chloride	7.0931	84	209684	50.8454	ug/L	98
20) Carbon Disulfide	7.1241	76	536742	47.2221	ug/L	100
21) Acrylonitrile	7.2689	53	81286	52.1544	ug/L	96
22) Methyl Tert Butyl Ether	7.2999	73	565588	48.3657	ug/L	98
23) trans-1,2-Dichloroethene	7.5274	96	210986	52.6317	ug/L	99
24) n-Hexane	7.5997	57	271496	43.9708	ug/L	99
25) Diisopropyl ether	7.9306	45	1580423	104.8742	ug/L	100
26) Vinyl Acetate	8.0960	43	171935	32.6187	ug/L	99
27) 1,1-Dichloroethane	8.1167	63	449642	51.4024	ug/L	99
28) Ethyl-Tert-Butyl ether	8.4786	59	1625562	106.7214	ug/L	99
29) 2-Butanone	8.6544	43	87976	48.8520	ug/L	97
30) Propionitrile	8.7578	54	59303	112.7091	ug/L	100
31) 2,2-Dichloropropane	8.8612	77	301940	49.9220	ug/L	98
32) cis-1,2-Dichloroethene	8.9232	96	241911	53.1549	ug/L	99
33) Chloroform	9.1300	83	405475	51.0695	ug/L	99
34) 1-Bromopropane	9.2541	122	49023	49.7315	ug/L	95
35) Bromochloromethane	9.3471	130	143553	53.7952	ug/L	99
36) Tetrahydrofuran	9.3678	42	116935	100.2247	ug/L	98
38) 1,1,1-Trichloroethane	9.6263	97	396924	55.2117	ug/L	98
39) Cyclohexane	9.6573	56	352416	41.4049	ug/L	98
40) 1,1-Dichloropropene	9.8124	75	293301	54.2616	ug/L	99
41) Carbon Tetrachloride	9.9572	117	336039	54.9678	ug/L	99
42) Tert-Amyl-Methyl ether	9.9158	73	1230422	108.3050	ug/L	98

(#) = qualifier out of range (m) = manual integration
 11M29748.D 8260WT.M Wed Mar 06 14:04:08 2019

Data File : C:\MSDCHEM\1\DATA\030519\11M29748.D Vial: 13
 Acq On : 5 Mar 2019 22:04 Operator: KFR
 Sample : WG698387-12 ICV 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 06 14:04:07 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1226	62	359321	52.0700	ug/L	99
45) Benzene	10.1536	78	837480	52.1268	ug/L	100
46) Trichloroethene	10.8671	130	247418	54.2980	ug/L	100
47) Methylcyclohexane	10.9394	83	297902	46.4078	ug/L	98
48) 1,2-Dichloropropane	11.0635	63	247364	51.8312	ug/L	99
49) 1,4-Dioxane	11.3427	88	8323	254.1631	ug/L	98
50) Bromodichloromethane	11.3530	83	301185	53.4000	ug/L	99
51) Dibromomethane	11.4357	93	140548	51.5555	ug/L	98
52) 2-Chloroethyl Vinyl Ether	11.6218	63	117689	50.9579	ug/L	99
53) 4-Methyl-2-Pentanone	11.6529	58	82421	50.0917	ug/L	98
54) cis-1,3-Dichloropropene	11.9424	75	351329	55.2697	ug/L	99
55) Dimethyl Disulfide	12.1905	79	183574	43.2226	ug/L	99
58) Toluene	12.3353	91	918334	53.0363	ug/L	99
59) Ethyl Methacrylate	12.4283	69	261494	53.4683	ug/L	99
60) trans-1,3-Dichloropropene	12.5007	75	304243	53.6657	ug/L	100
61) 1,1,2-Trichloroethane	12.7075	97	186431	50.5763	ug/L	100
62) 2-Hexanone	12.6455	43	141874	51.7818	ug/L	99
63) 1,3-Dichloropropane	12.9970	76	319634	53.7441	ug/L	100
64) Tetrachloroethene	13.1107	164	189951	51.7185	ug/L	98
65) Dibromochloromethane	13.3589	129	224538	53.5106	ug/L	99
66) 1,2-Dibromoethane	13.5967	107	186111	52.6818	ug/L	99
67) 1-Chlorohexane	13.6691	91	284180	49.7115	ug/L	99
68) Chlorobenzene	14.0620	112	633960	52.5890	ug/L	100
69) 1,1,1,2-Tetrachloroethane	14.0930	131	249378	48.6934	ug/L	99
70) Ethylbenzene	14.0826	106	330037	53.8169	ug/L	100
71) m-,p-Xylene	14.1654	106	794611	107.3271	ug/L	100
72) o-Xylene	14.6927	106	393911	54.8361	ug/L	99
73) Styrene	14.7341	104	667526	56.1795	ug/L	100
74) Bromoform	15.1993	173	141028	46.6710	ug/L	99
75) Isopropylbenzene	15.0856	105	953203	53.0064	ug/L	99
77) 1,1,2,2-Tetrachloroethane	15.2924	83	202309	49.8831	ug/L	100
79) 1,2,3-Trichloropropane	15.4682	110	72854	53.7990	ug/L	98
80) trans-1,4-Dichloro-2-Butene	15.5095	53	82302	48.2026	ug/L	92
81) n-Propylbenzene	15.5612	91	1203296	52.4844	ug/L	100
82) Bromobenzene	15.6853	156	278745	49.7840	ug/L	100
83) 1,3,5-Trimethylbenzene	15.7370	105	842593	51.7693	ug/L	100
84) 2-Chlorotoluene	15.8197	91	825177	50.5808	ug/L	100
85) 4-Chlorotoluene	15.8611	91	675304	50.3645	ug/L	99
86) a-Methylstyrene	16.1092	118	489194	52.7849	ug/L	100
87) tert-Butylbenzene	16.1609	134	200021	49.8961	ug/L	98
88) 1,2,4-Trimethylbenzene	16.2126	105	870375	51.8136	ug/L	100
89) sec-Butylbenzene	16.4194	105	1060708	52.4940	ug/L	100
90) p-Isopropyltoluene	16.5642	119	930561	52.9626	ug/L	100
91) 1,3-Dichlorobenzene	16.7503	146	528121	51.1306	ug/L	99
92) 1,4-Dichlorobenzene	16.8640	146	541106	52.4618	ug/L	99
93) n-Butylbenzene	17.0501	91	853955	53.9300	ug/L	99
94) 1,2-Dichlorobenzene	17.3293	146	531745	53.7158	ug/L	100
95) 1,2-Dibromo-3-Chloropropane	18.2495	75	45751	54.6065	ug/L	99
96) 1,2,4-Trichlorobenzene	19.3145	180	352342	55.0215	ug/L	99
97) Hexachlorobutadiene	19.4489	225	179393	55.5810	ug/L	99
98) Naphthalene	19.6557	128	796590	58.0926	ug/L	100
99) 1,2,3-Trichlorobenzene	19.9452	180	338519	56.2661	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29748.D 8260WT.M Wed Mar 06 14:04:08 2019

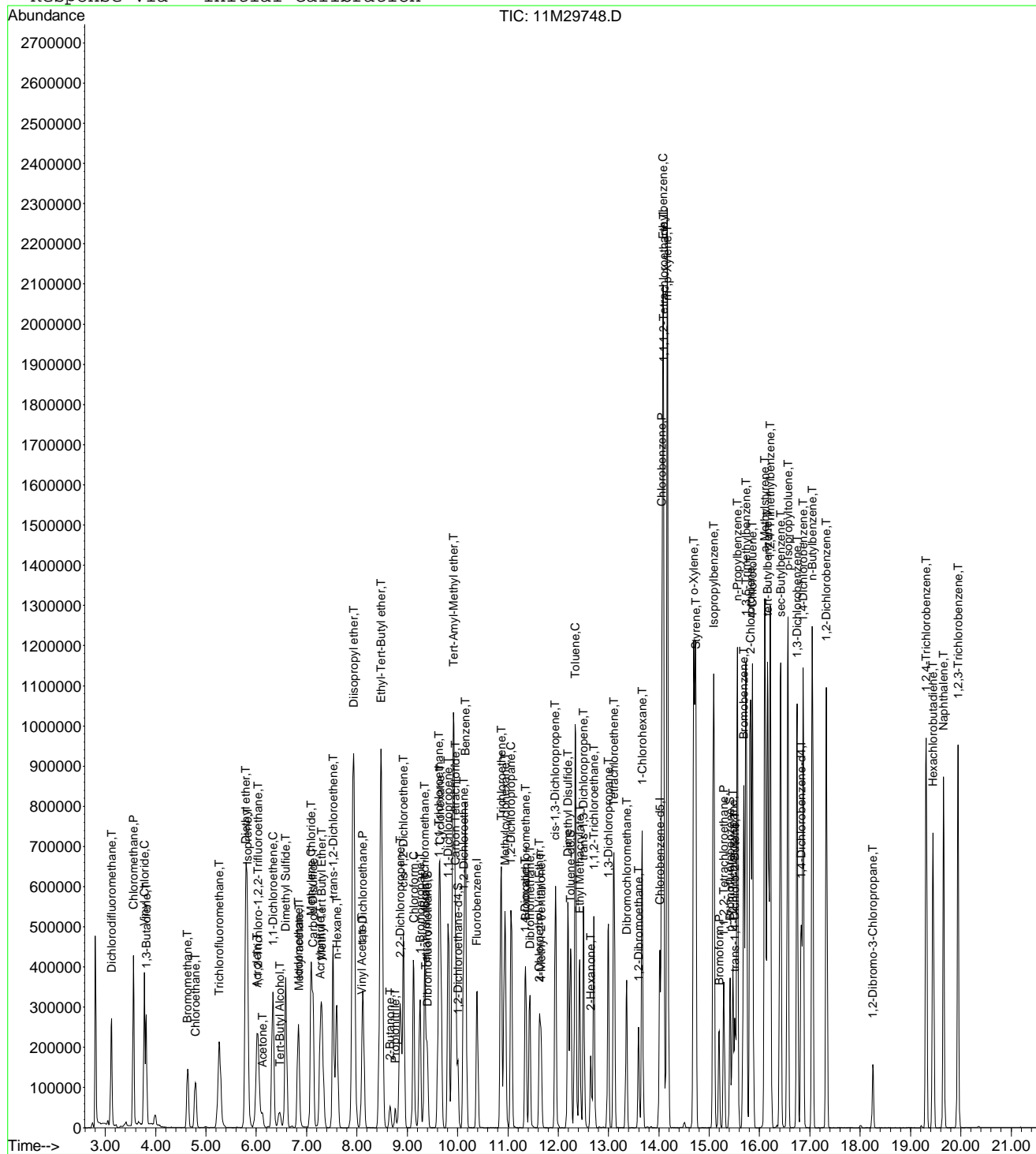
Page 2

Data File : C:\MSDCHEM\1\DATA\030519\11M29748.D
Acq On : 5 Mar 2019 22:04
Sample : WG698387-12 ICV 50ug/L 8260
Misc : 1,1 STD92316
MS Integration Params: rteint.p
Quant Time: Mar 6 14:04 2019

Vial: 13
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:00:17 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\030519\11M29748.D Vial: 13
 Acq On : 5 Mar 2019 22:04 Operator: KFR
 Sample : WG698387-12 ICV 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	98	-0.01
2 T	Dichlorodifluoromethane	50.000	50.716	-1.4	91	0.00
3 P	Chloromethane	50.000	43.373	13.3	89	0.00
4 C	Vinyl Chloride	50.000	45.163	9.7	86	0.00
5 T	1,3-Butadiene	50.000	47.963	4.1	87	0.00
6 T	Bromomethane	50.000	42.374	15.3	91	-0.01
7 T	Chloroethane	50.000	47.399	5.2	89	-0.01
8 T	Trichlorofluoromethane	50.000	46.817	6.4	87	-0.01
9 T	Diethyl ether	100.000	99.678	0.3	107	0.00
10 T	Isoprene	50.000	50.067	-0.1	94	0.00
11 T	Acrolein	50.000	88.462	-76.9#	176	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	50.000	51.005	-2.0	93	-0.01
13 T	Acetone	50.000	44.220	11.6	85	0.00
14 C	1,1-Dichloroethene	50.000	51.182	-2.4	95	0.00
15 T	Tert-Butyl Alcohol	200.000	236.902	-18.5	116	0.02
16 T	Dimethyl Sulfide	50.000	55.442	-10.9	104	0.00
17 T	Iodomethane	50.000	65.741	-31.5#	128	0.00
18 T	Methyl acetate	50.000	53.726	-7.5	102	0.00
19 T	Methylene Chloride	50.000	50.845	-1.7	97	0.00
20 T	Carbon Disulfide	50.000	47.222	5.6	86	-0.01
21 T	Acrylonitrile	50.000	52.154	-4.3	93	0.00
22 T	Methyl Tert Butyl Ether	50.000	48.366	3.3	91	0.00
23 T	trans-1,2-Dichloroethene	50.000	52.632	-5.3	96	0.00
24 T	n-Hexane	50.000	43.971	12.1	83	0.00
25 T	Diisopropyl ether	100.000	104.874	-4.9	110	0.00
26 T	Vinyl Acetate	50.000	32.619	34.8#	63	0.00
27 P	1,1-Dichloroethane	50.000	51.402	-2.8	97	-0.01
28 T	Ethyl-Tert-Butyl ether	100.000	106.721	-6.7	110	0.00
29 T	2-Butanone	50.000	48.852	2.3	91	0.00
30 T	Propionitrile	100.000	112.709	-12.7	101	0.00
31 T	2,2-Dichloropropane	50.000	49.922	0.2	93	0.00
32 T	cis-1,2-Dichloroethene	50.000	53.155	-6.3	97	-0.01
33 C	Chloroform	50.000	51.069	-2.1	99	0.00
34 T	1-Bromopropane	50.000	49.732	0.5	96	0.00
35 T	Bromochloromethane	50.000	53.795	-7.6	96	0.00
36 T	Tetrahydrofuran	100.000	100.225	-0.2	102	0.00
37 S	Dibromofluoromethane	25.000	25.066	-0.3	94	0.00
38 T	1,1,1-Trichloroethane	50.000	55.212	-10.4	101	0.00
39 T	Cyclohexane	50.000	41.405	17.2	78	0.00
40 T	1,1-Dichloropropene	50.000	54.262	-8.5	100	0.00
41 T	Carbon Tetrachloride	50.000	54.968	-9.9	97	0.00
42 T	Tert-Amyl-Methyl ether	100.000	108.305	-8.3	110	0.00
43 S	1,2-Dichloroethane-d4	25.000	25.239	-1.0	95	0.00
44 T	1,2-Dichloroethane	50.000	52.070	-4.1	96	0.00
45 T	Benzene	50.000	52.127	-4.3	97	0.00
46 T	Trichloroethene	50.000	54.298	-8.6	100	0.00
47 T	Methylcyclohexane	50.000	46.408	7.2	87	0.00
48 C	1,2-Dichloropropane	50.000	51.831	-3.7	97	0.00
49 T	1,4-Dioxane	200.000	254.163	-27.1#	121	0.00
50 T	Bromodichloromethane	50.000	53.400	-6.8	93	0.00
51 T	Dibromomethane	50.000	51.556	-3.1	95	0.00
52 T	2-Chloroethyl Vinyl Ether	50.000	50.958	-1.9	97	0.00
53 T	4-Methyl-2-Pentanone	50.000	50.092	-0.2	94	0.00
54 T	cis-1,3-Dichloropropene	50.000	55.270	-10.5	99	0.00

(#) = Out of Range

11M29748.D 8260WT.M Wed Mar 06 14:04:11 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\030519\11M29748.D Vial: 13
 Acq On : 5 Mar 2019 22:04 Operator: KFR
 Sample : WG698387-12 ICV 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:00:17 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	50.000	43.223	13.6	87	0.00
56 I	Chlorobenzene-d5	25.000	25.000	0.0	98	0.00
57 S	Toluene-d8	25.000	24.915	0.3	93	-0.01
58 C	Toluene	50.000	53.036	-6.1	97	0.00
59 T	Ethyl Methacrylate	50.000	53.468	-6.9	96	0.00
60 T	trans-1,3-Dichloropropene	50.000	53.666	-7.3	95	0.00
61 T	1,1,2-Trichloroethane	50.000	50.576	-1.2	96	0.00
62 T	2-Hexanone	50.000	51.782	-3.6	94	0.00
63 T	1,3-Dichloropropane	50.000	53.744	-7.5	100	0.00
64 T	Tetrachloroethene	50.000	51.719	-3.4	97	0.00
65 T	Dibromochloromethane	50.000	53.511	-7.0	95	0.00
66 T	1,2-Dibromoethane	50.000	52.682	-5.4	96	0.00
67 T	1-Chlorohexane	50.000	49.712	0.6	91	0.00
68 P	Chlorobenzene	50.000	52.589	-5.2	97	0.00
69 T	1,1,1,2-Tetrachloroethane	50.000	48.693	2.6	95	0.00
70 C	Ethylbenzene	50.000	53.817	-7.6	96	0.00
71 T	m-,p-Xylene	100.000	107.327	-7.3	97	0.00
72 T	o-Xylene	50.000	54.836	-9.7	98	0.00
73 T	Styrene	50.000	56.180	-12.4	97	0.00
74 P	Bromoform	50.000	46.671	6.7	90	0.00
75 T	Isopropylbenzene	50.000	53.006	-6.0	94	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	104	0.00
77 P	1,1,2,2-Tetrachloroethane	50.000	49.883	0.2	90	0.00
78 S	p-Bromofluorobenzene	25.000	24.449	2.2	94	0.00
79 T	1,2,3-Trichloropropane	50.000	53.799	-7.6	101	0.00
80 T	trans-1,4-Dichloro-2-Butene	50.000	48.203	3.6	90	0.00
81 T	n-Propylbenzene	50.000	52.484	-5.0	98	0.00
82 T	Bromobenzene	50.000	49.784	0.4	95	0.00
83 T	1,3,5-Trimethylbenzene	50.000	51.769	-3.5	96	0.00
84 T	2-Chlorotoluene	50.000	50.581	-1.2	97	0.00
85 T	4-Chlorotoluene	50.000	50.364	-0.7	95	0.00
86 T	a-Methylstyrene	50.000	52.785	-5.6	99	0.00
87 T	tert-Butylbenzene	50.000	49.896	0.2	99	-0.01
88 T	1,2,4-Trimethylbenzene	50.000	51.814	-3.6	97	0.00
89 T	sec-Butylbenzene	50.000	52.494	-5.0	97	0.00
90 T	p-Isopropyltoluene	50.000	52.963	-5.9	97	0.00
91 T	1,3-Dichlorobenzene	50.000	51.131	-2.3	97	0.00
92 T	1,4-Dichlorobenzene	50.000	52.462	-4.9	98	0.00
93 T	n-Butylbenzene	50.000	53.930	-7.9	97	0.00
94 T	1,2-Dichlorobenzene	50.000	53.716	-7.4	101	0.00
95 T	1,2-Dibromo-3-Chloropropane	50.000	54.606	-9.2	102	0.00
96 T	1,2,4-Trichlorobenzene	50.000	55.021	-10.0	103	0.00
97 T	Hexachlorobutadiene	50.000	55.581	-11.2	106	0.00
98 T	Naphthalene	50.000	58.093	-16.2	105	0.00
99 T	1,2,3-Trichlorobenzene	50.000	56.266	-12.5	104	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29748.D 8260WT.M Wed Mar 06 14:04:12 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427930.D Vial: 5
 Acq On : 12 Nov 2018 13:37 Operator: EEA
 Sample : WG684281-02 5 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:37 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.77	96	905676	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.64	117	676804	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.65	152	343581	25.00	ug/L	0.00

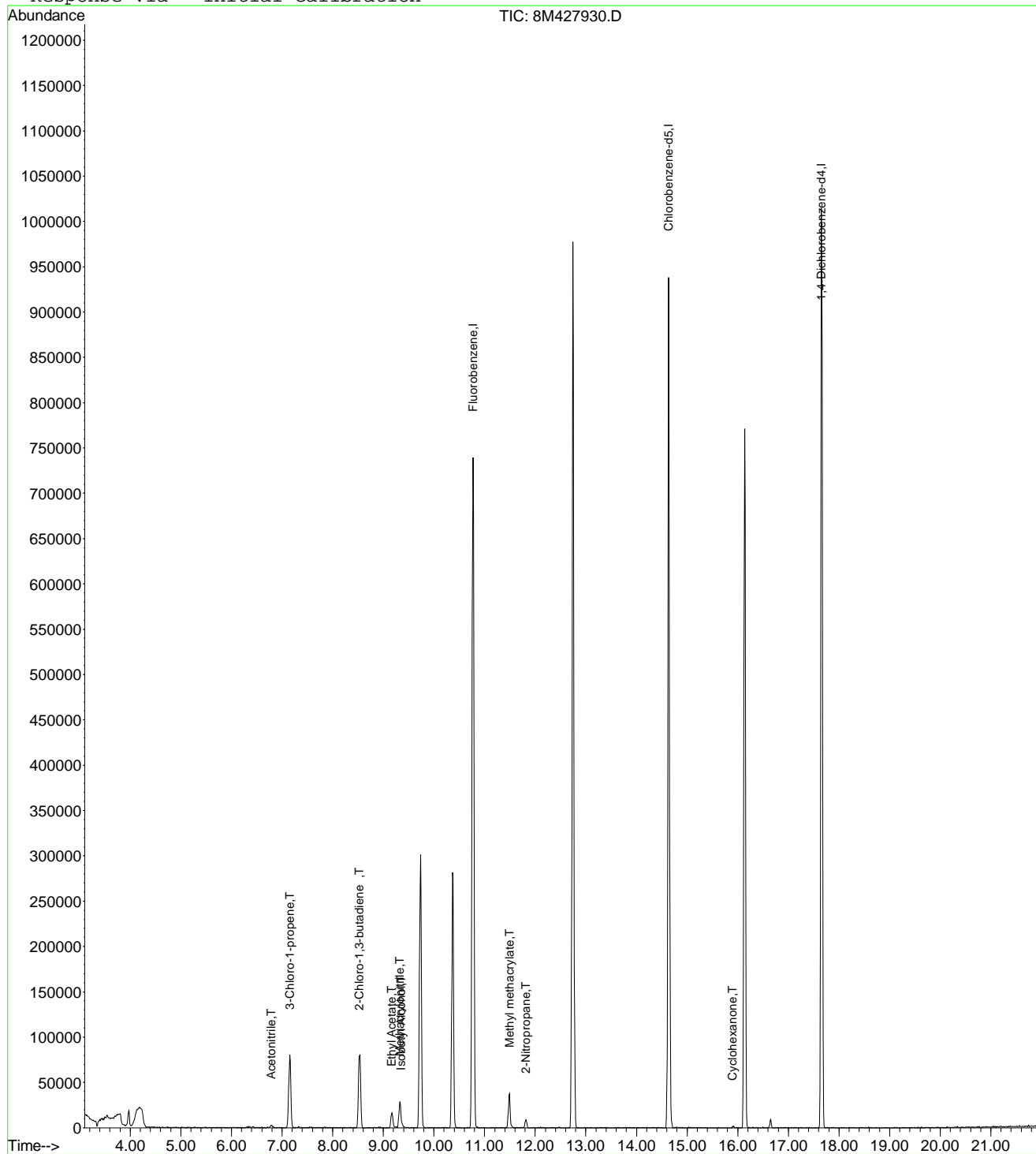
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.79	41	3867	4.5939	ug/L	65
3) 3-Chloro-1-propene	7.15	41	79744	4.8119	ug/L	96
4) 2-Chloro-1,3-butadiene	8.52	53	75225	4.4974	ug/L	99
5) Ethyl Acetate	9.17	43	29157	3.8316	ug/L	97
6) Methacrylonitrile	9.32	67	13803	3.9932	ug/L	95
7) Isobutyl Alcohol	9.35	43	1018	3.9730	ug/L #	41
9) Methyl methacrylate	11.49	41	27219	3.6869	ug/L	98
10) 2-Nitropropane	11.81	43	7409	2.9142	ug/L	96
13) Cyclohexanone	15.90	55	1169	3.2976	ug/L	94

 (#) = qualifier out of range (m) = manual integration
 8M427930.D A9FOOWT.M Tue Nov 13 12:19:38 2018

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427930.D Vial: 5
 Acq On : 12 Nov 2018 13:37 Operator: EEA
 Sample : WG684281-02 5 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427930.D Vial: 5
 Acq On : 12 Nov 2018 13:37 Operator: EEA
 Sample : WG684281-02 5 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	-1.000	4.594	0.0	100	0.00
3 T	3-Chloro-1-propene	5.000	4.812	3.8	100	0.00
4 T	2-Chloro-1,3-butadiene	5.000	4.497	10.1	100	0.00
5 T	Ethyl Acetate	-1.000	3.832	0.0	100	0.00
6 T	Methacrylonitrile	5.000	3.993	20.1	100	0.00
7 T	Isobutyl Alcohol	-1.000	3.973	0.0	100	0.01
8 T	1-Butanol	-1.000	0.000	0.0	0	-10.29#
9 T	Methyl methacrylate	5.000	3.687	26.3	100	0.00
10 T	2-Nitropropane	-1.000	2.914	0.0	100	0.00
11 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
12 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
13 T	Cyclohexanone	-1.000	3.298	0.0	100	0.00

(#) = Out of Range
 8M427930.D A9FOOWT.M

SPCC's out = 0 CCC's out = 0
 Tue Nov 13 12:17:03 2018

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427931.D Vial: 3
 Acq On : 12 Nov 2018 14:06 Operator: EEA
 Sample : WG684281-03 20 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:39 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.78	96	874045	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.63	117	672711	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.66	152	343559	25.00	ug/L	0.00

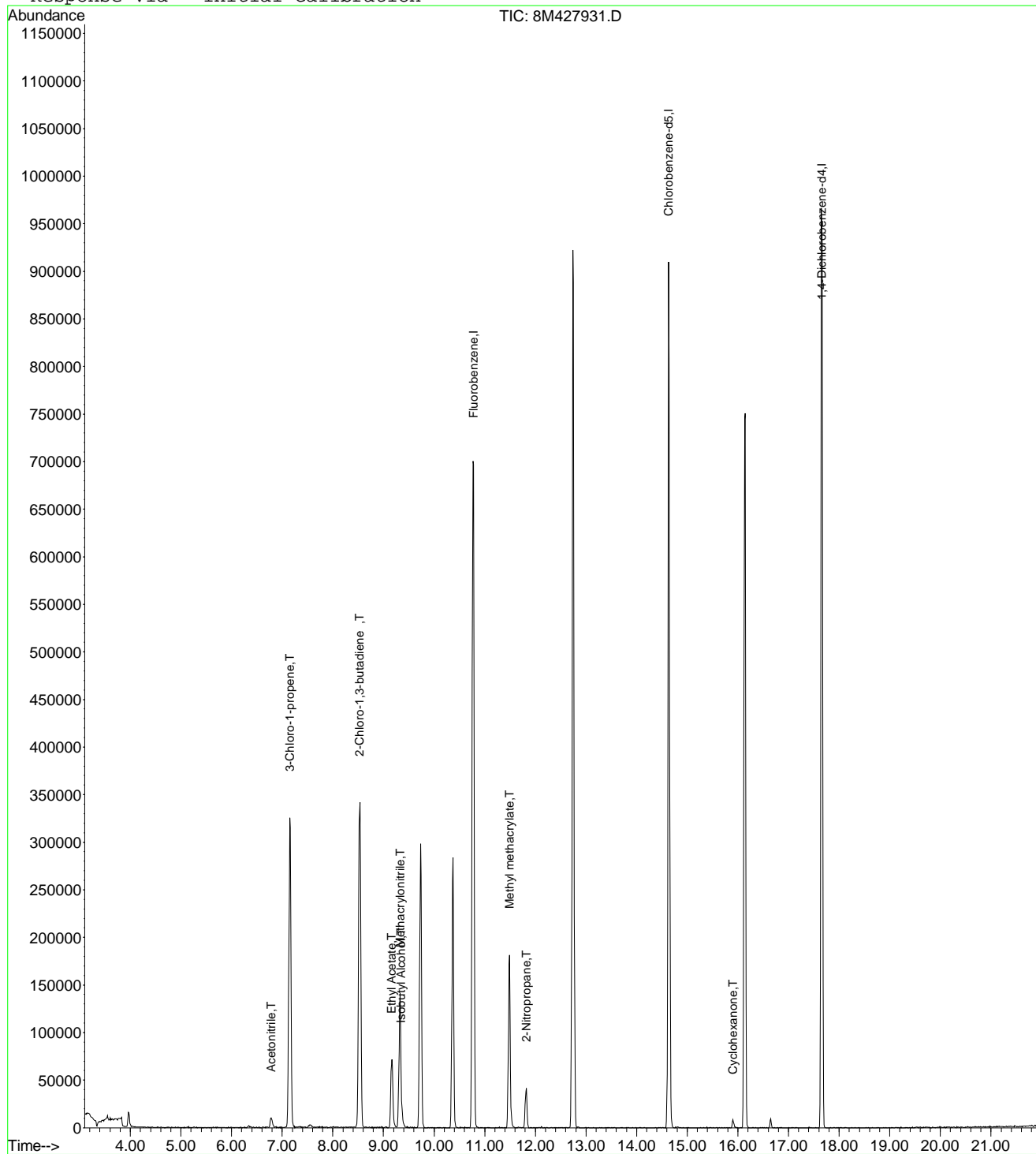
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.79	41	14838	18.2650	ug/L	94
3) 3-Chloro-1-propene	7.15	41	320423	20.0345	ug/L	100
4) 2-Chloro-1,3-butadiene	8.53	53	321084	19.8908	ug/L	99
5) Ethyl Acetate	9.16	43	127962	17.4245	ug/L	98
6) Methacrylonitrile	9.33	67	59985	17.9816	ug/L	96
7) Isobutyl Alcohol	9.35	43	9196	37.1883	ug/L #	79
9) Methyl methacrylate	11.49	41	124457	17.4683	ug/L	100
10) 2-Nitropropane	11.82	43	36091	14.7094	ug/L	94
13) Cyclohexanone	15.90	55	5307	14.9715	ug/L	96

 (#) = qualifier out of range (m) = manual integration
 8M427931.D A9FOOWT.M Tue Nov 13 12:19:39 2018

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427931.D Vial: 3
 Acq On : 12 Nov 2018 14:06 Operator: EEA
 Sample : WG684281-03 20 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427931.D Vial: 3
 Acq On : 12 Nov 2018 14:06 Operator: EEA
 Sample : WG684281-03 20 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	20.000	18.265	8.7	100	0.00
3 T	3-Chloro-1-propene	20.000	20.034	-0.2	100	0.00
4 T	2-Chloro-1,3-butadiene	20.000	19.891	0.5	100	0.00
5 T	Ethyl Acetate	20.000	17.424	12.9	100	0.00
6 T	Methacrylonitrile	20.000	17.982	10.1	100	0.00
7 T	Isobutyl Alcohol	-1.000	37.188	0.0	100	0.01
8 T	1-Butanol	-1.000	0.000	0.0	0	-10.29#
9 T	Methyl methacrylate	20.000	17.468	12.7	100	0.00
10 T	2-Nitropropane	20.000	14.709	26.5	100	0.00
11 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
12 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
13 T	Cyclohexanone	-1.000	14.971	0.0	100	0.00

(#) = Out of Range
 8M427931.D A9FOOWT.M

SPCC's out = 0 CCC's out = 0
 Tue Nov 13 12:17:43 2018

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427932.D Vial: 4
 Acq On : 12 Nov 2018 14:34 Operator: EEA
 Sample : WG684281-04 50 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:40 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.78	96	885100	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.63	117	662296	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.66	152	334189	25.00	ug/L	0.00

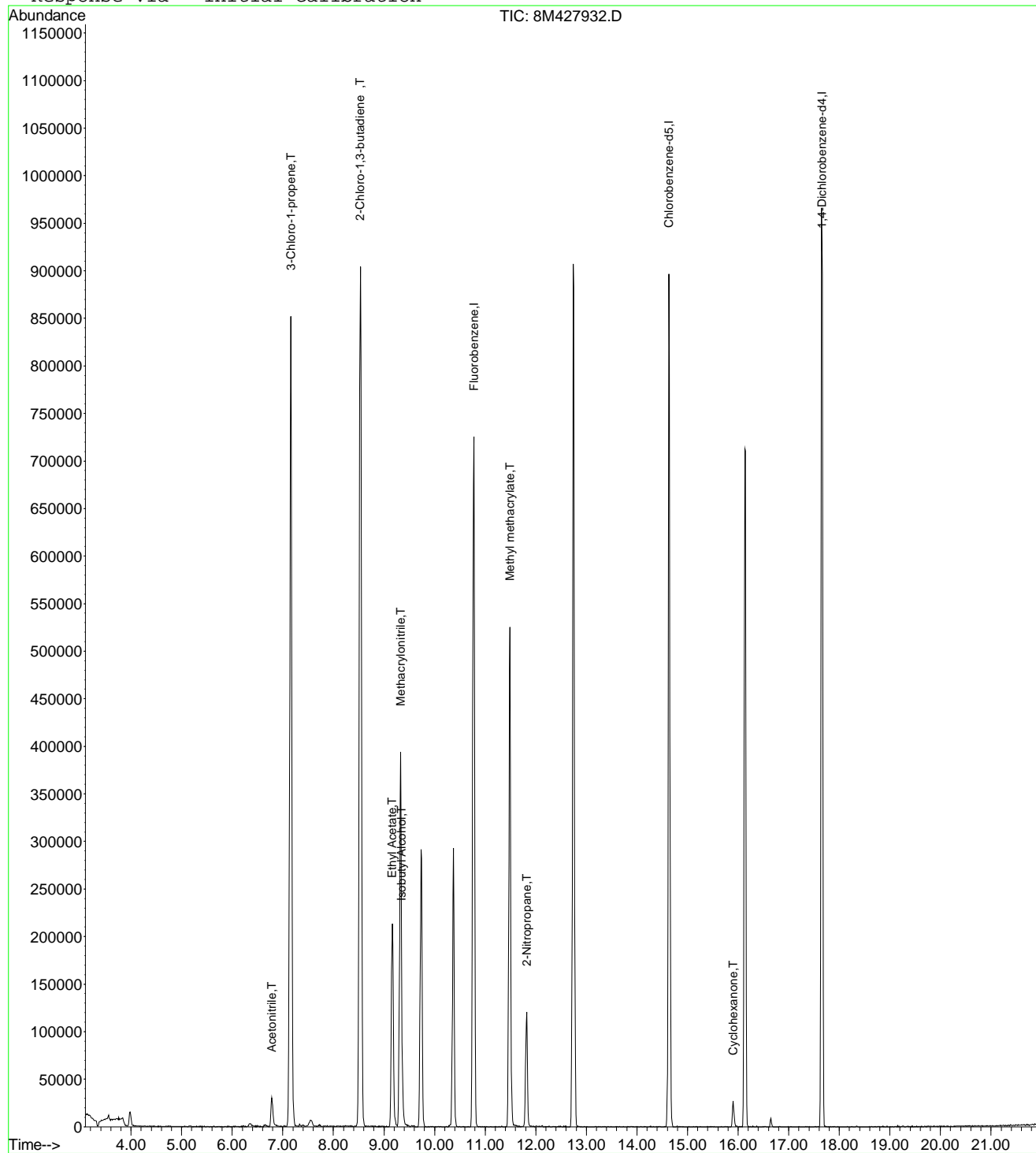
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.78	41	41148	50.0190	ug/L	96
3) 3-Chloro-1-propene	7.16	41	814844	50.3118	ug/L	99
4) 2-Chloro-1,3-butadiene	8.53	53	824807	50.4577	ug/L	100
5) Ethyl Acetate	9.16	43	357052	48.0122	ug/L	99
6) Methacrylonitrile	9.33	67	165999	49.1397	ug/L	97
7) Isobutyl Alcohol	9.35	43	25778	102.9433	ug/L	88
9) Methyl methacrylate	11.49	41	346219	47.9870	ug/L	100
10) 2-Nitropropane	11.82	43	109997	44.2709	ug/L	99
13) Cyclohexanone	15.90	55	15640	45.3588	ug/L	99

 (#) = qualifier out of range (m) = manual integration
 8M427932.D A9FOOWT.M Tue Nov 13 12:19:41 2018

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427932.D Vial: 4
Acq On : 12 Nov 2018 14:34 Operator: EEA
Sample : WG684281-04 50 ug/L ICAL A9 8260 Inst : HPMS8
Misc : 1,1 STD90626 Multiplr: 1.00
MS Integration Params: rteint.p
Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
Title : A9-FOO Water SOP:MSV01 111218 HPMS8
Last Update : Tue Nov 13 11:28:33 2018
Response via : Initial Calibration



8M427932.D A9FOOWT.M

Tue Nov 13 12:19:41 2018

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427932.D Vial: 4
 Acq On : 12 Nov 2018 14:34 Operator: EEA
 Sample : WG684281-04 50 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	50.000	50.019	-0.0	100	0.00
3 T	3-Chloro-1-propene	50.000	50.312	-0.6	100	0.00
4 T	2-Chloro-1,3-butadiene	50.000	50.458	-0.9	100	0.00
5 T	Ethyl Acetate	50.000	48.012	4.0	100	0.00
6 T	Methacrylonitrile	50.000	49.140	1.7	100	0.00
7 T	Isobutyl Alcohol	100.000	102.943	-2.9	100	0.01
8 T	1-Butanol	-1.000	0.000	0.0	100	0.02
9 T	Methyl methacrylate	50.000	47.987	4.0	100	0.00
10 T	2-Nitropropane	50.000	44.271	11.5	100	0.00
11 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
12 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
13 T	Cyclohexanone	50.000	45.359	9.3	100	0.00

(#) = Out of Range
 8M427932.D A9FOOWT.M

SPCC's out = 0 CCC's out = 0
 Tue Nov 13 12:18:19 2018

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427933.D Vial: 5
 Acq On : 12 Nov 2018 15:03 Operator: EEA
 Sample : WG684281-05 100 ug/L ICAL A9 826 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:41 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.78	96	893935	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.63	117	667066	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.66	152	341780	25.00	ug/L	0.00

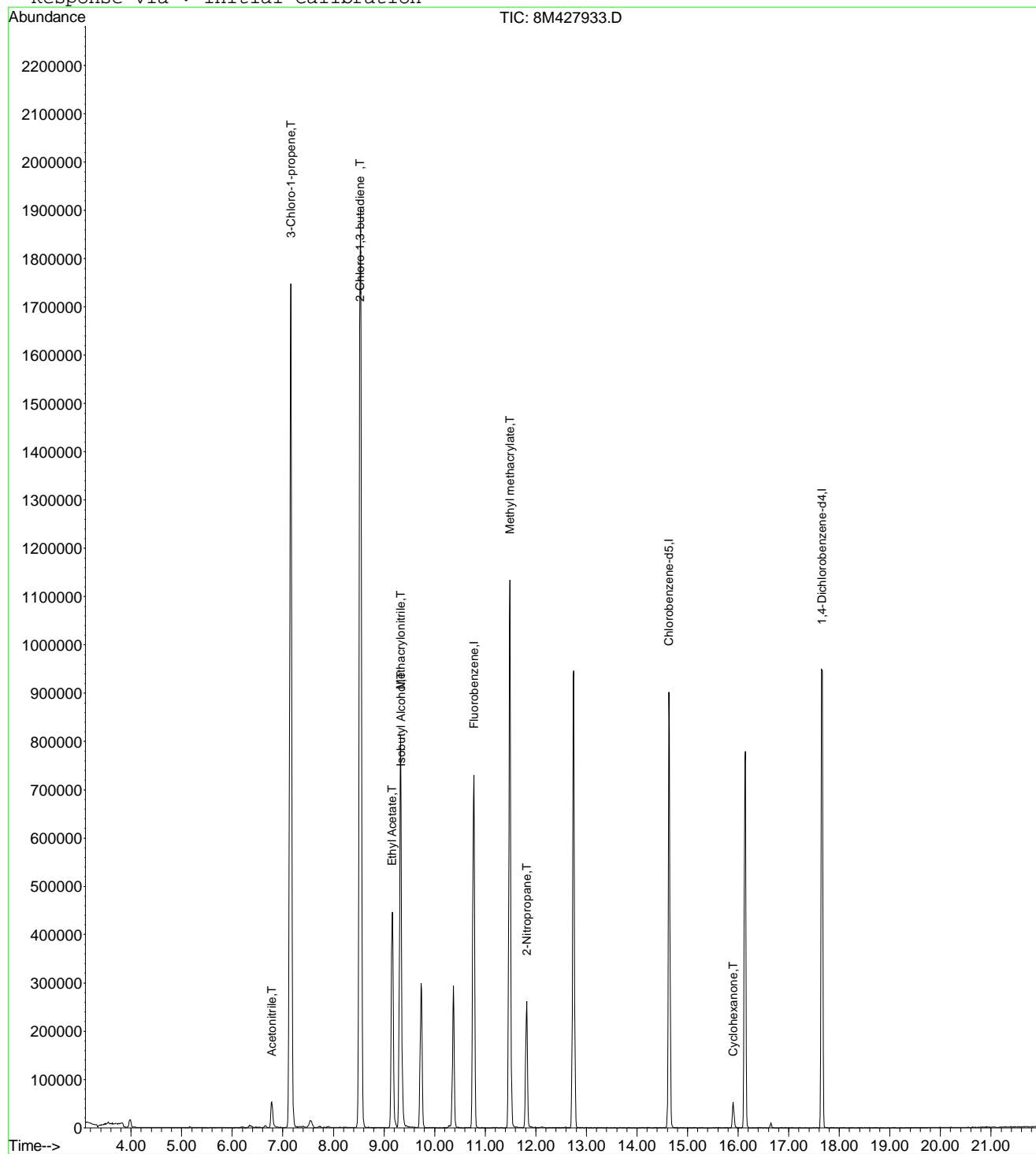
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.78	41	75612	91.0045	ug/L	100
3) 3-Chloro-1-propene	7.16	41	1695520	103.6537	ug/L	100
4) 2-Chloro-1,3-butadiene	8.53	53	1714203	103.8302	ug/L	100
5) Ethyl Acetate	9.16	43	745480	99.2528	ug/L	100
6) Methacrylonitrile	9.33	67	347443	101.8351	ug/L	100
7) Isobutyl Alcohol	9.34	43	41468	163.9639	ug/L	100
9) Methyl methacrylate	11.49	41	748715	102.7486	ug/L	100
10) 2-Nitropropane	11.82	43	241099	96.0769	ug/L	100
13) Cyclohexanone	15.90	55	30575	86.7036	ug/L	100

 (#) = qualifier out of range (m) = manual integration
 8M427933.D A9FOOWT.M Tue Nov 13 12:19:42 2018

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427933.D Vial: 5
Acq On : 12 Nov 2018 15:03 Operator: EEA
Sample : WG684281-05 100 ug/L ICAL A9 826 Inst : HPMS8
Misc : 1,1 STD90626 Multiplr: 1.00
MS Integration Params: rteint.p
Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
Title : A9-FOO Water SOP:MSV01 111218 HPMS8
Last Update : Tue Nov 13 11:28:33 2018
Response via : Initial Calibration



8M427933.D A9FOOWT.M

Tue Nov 13 12:19:42 2018

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427933.D Vial: 5
 Acq On : 12 Nov 2018 15:03 Operator: EEA
 Sample : WG684281-05 100 ug/L ICAL A9 826 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Acetonitrile	100.000	91.005	9.0	100	0.00
3 T	3-Chloro-1-propene	100.000	103.654	-3.7	100	0.00
4 T	2-Chloro-1,3-butadiene	100.000	103.830	-3.8	100	0.00
5 T	Ethyl Acetate	100.000	99.253	0.7	100	0.00
6 T	Methacrylonitrile	100.000	101.835	-1.8	100	0.00
7 T	Isobutyl Alcohol	200.000	163.964	18.0	100	0.00
8 T	1-Butanol	-1.000	0.000	0.0	100	0.00
9 T	Methyl methacrylate	100.000	102.749	-2.7	100	0.00
10 T	2-Nitropropane	100.000	96.077	3.9	100	0.00
11 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00
12 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
13 T	Cyclohexanone	100.000	86.704	13.3	100	0.00

(#) = Out of Range
 8M427933.D A9FOOWT.M

SPCC's out = 0 CCC's out = 0
 Tue Nov 13 12:18:56 2018

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427934.D Vial: 6
 Acq On : 12 Nov 2018 15:31 Operator: EEA
 Sample : WG684281-06 200 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:43 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.77	96	888856	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.64	117	662279	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.65	152	337695	25.00	ug/L	0.00

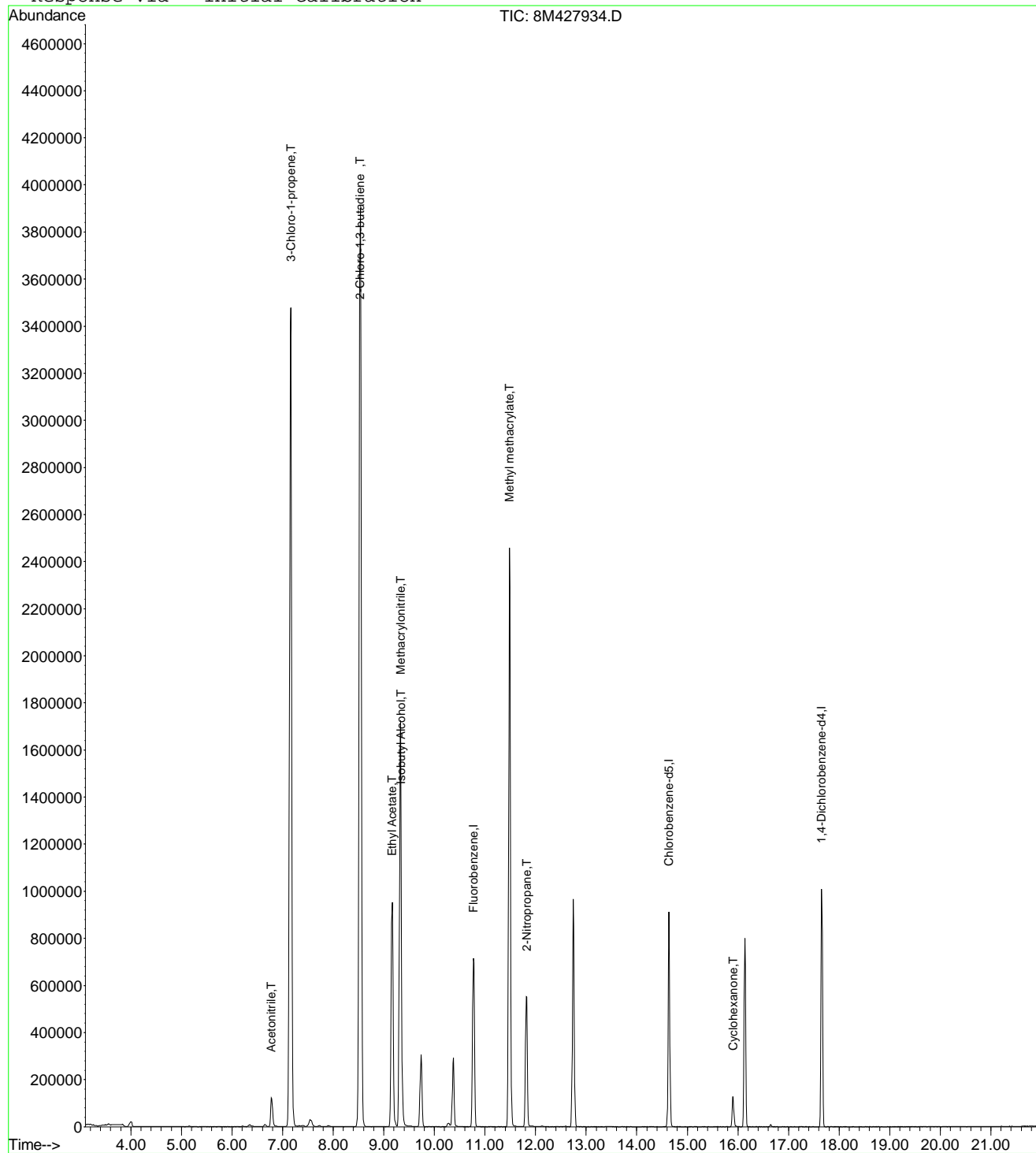
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.78	41	176351	213.4640	ug/L	95
3) 3-Chloro-1-propene	7.16	41	3397396	208.8827	ug/L	98
4) 2-Chloro-1,3-butadiene	8.54	53	3479174	211.9397	ug/L	99
5) Ethyl Acetate	9.17	43	1581923	211.8198	ug/L	100
6) Methacrylonitrile	9.33	67	728151	214.6396	ug/L	99
7) Isobutyl Alcohol	9.34	43	104801	416.7495	ug/L	98
9) Methyl methacrylate	11.48	41	1598234	220.5841	ug/L	99
10) 2-Nitropropane	11.82	43	539348	216.1560	ug/L	97
13) Cyclohexanone	15.91	55	73604	211.2488	ug/L	93

 (#) = qualifier out of range (m) = manual integration
 8M427934.D A9FOOWT.M Tue Nov 13 12:19:43 2018

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427934.D Vial: 6
Acq On : 12 Nov 2018 15:31 Operator: EEA
Sample : WG684281-06 200 ug/L ICAL A9 8260 Inst : HPMS8
Misc : 1,1 STD90626 Multiplr: 1.00
MS Integration Params: rteint.p
Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
Title : A9-FOO Water SOP:MSV01 111218 HPMS8
Last Update : Tue Nov 13 11:28:33 2018
Response via : Initial Calibration



8M427934.D A9FOOWT.M

Tue Nov 13 12:19:43 2018

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427935.D Vial: 7
 Acq On : 12 Nov 2018 15:59 Operator: EEA
 Sample : WG684281-07 300 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:44 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.77	96	906816	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.63	117	674675	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.65	152	342902	25.00	ug/L	0.00

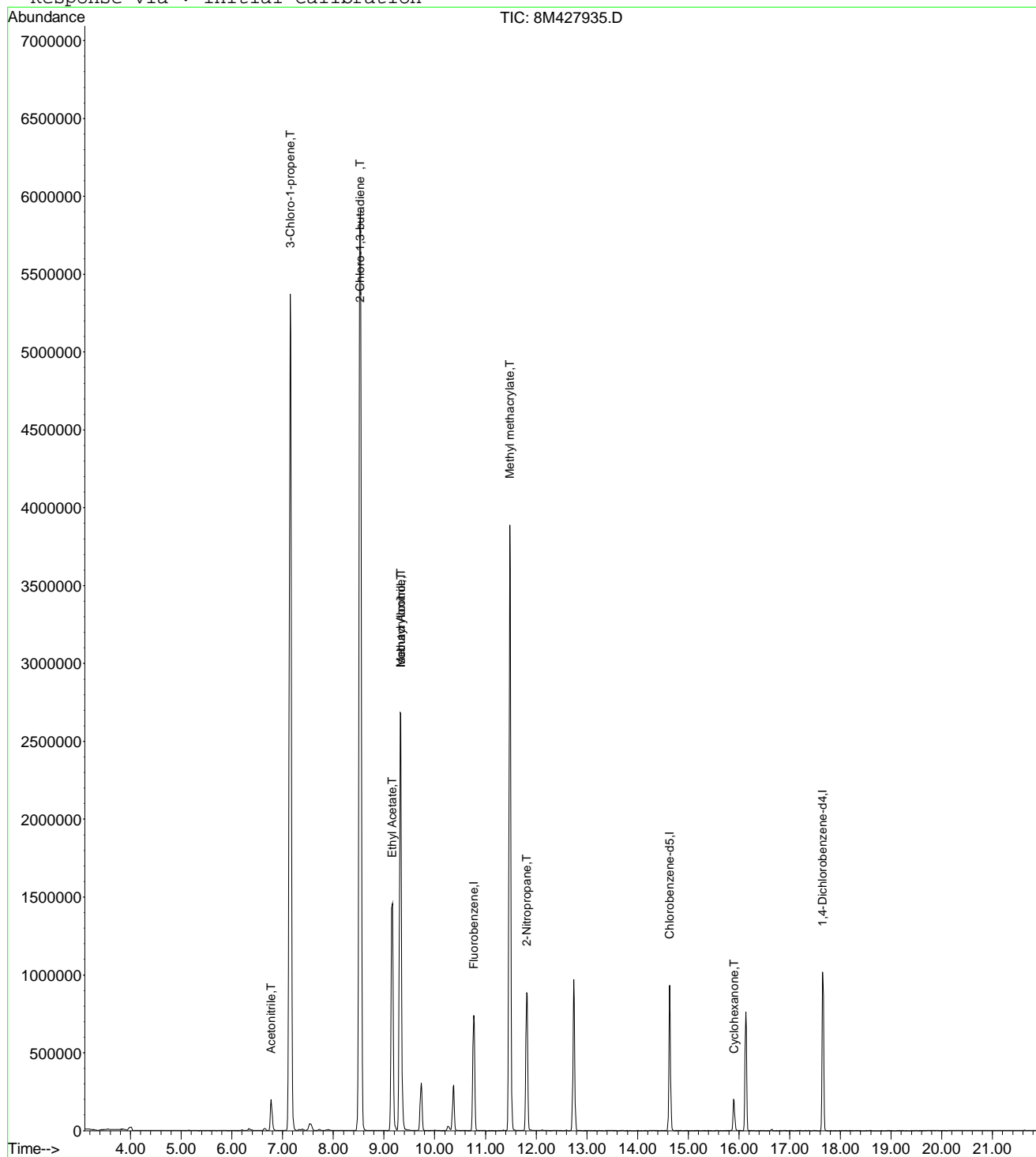
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.78	41	270445	320.8766	ug/L	95
3) 3-Chloro-1-propene	7.15	41	5177043	311.9970	ug/L	97
4) 2-Chloro-1,3-butadiene	8.53	53	5299143	316.4128	ug/L	97
5) Ethyl Acetate	9.16	43	2468218	323.9492	ug/L	99
6) Methacrylonitrile	9.33	67	1142367	330.0701	ug/L	99
7) Isobutyl Alcohol	9.33	43	162083	631.7706	ug/L	100
9) Methyl methacrylate	11.48	41	2521386	341.1028	ug/L	99
10) 2-Nitropropane	11.81	43	861383	338.3815	ug/L	97
13) Cyclohexanone	15.91	55	116314	328.7603	ug/L	96

 (#) = qualifier out of range (m) = manual integration
 8M427935.D A9FOOWT.M Tue Nov 13 12:19:45 2018

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427935.D Vial: 7
Acq On : 12 Nov 2018 15:59 Operator: EEA
Sample : WG684281-07 300 ug/L ICAL A9 8260 Inst : HPMS8
Misc : 1,1 STD90626 Multiplr: 1.00
MS Integration Params: rteint.p
Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
Title : A9-FOO Water SOP:MSV01 111218 HPMS8
Last Update : Tue Nov 13 11:28:33 2018
Response via : Initial Calibration



8M427935.D A9FOOWT.M

Tue Nov 13 12:19:45 2018

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427936.D Vial: 8
 Acq On : 12 Nov 2018 16:28 Operator: EEA
 Sample : WG684281-08 400 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:45 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.78	96	920293	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.63	117	690560	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.66	152	356915	25.00	ug/L	0.00

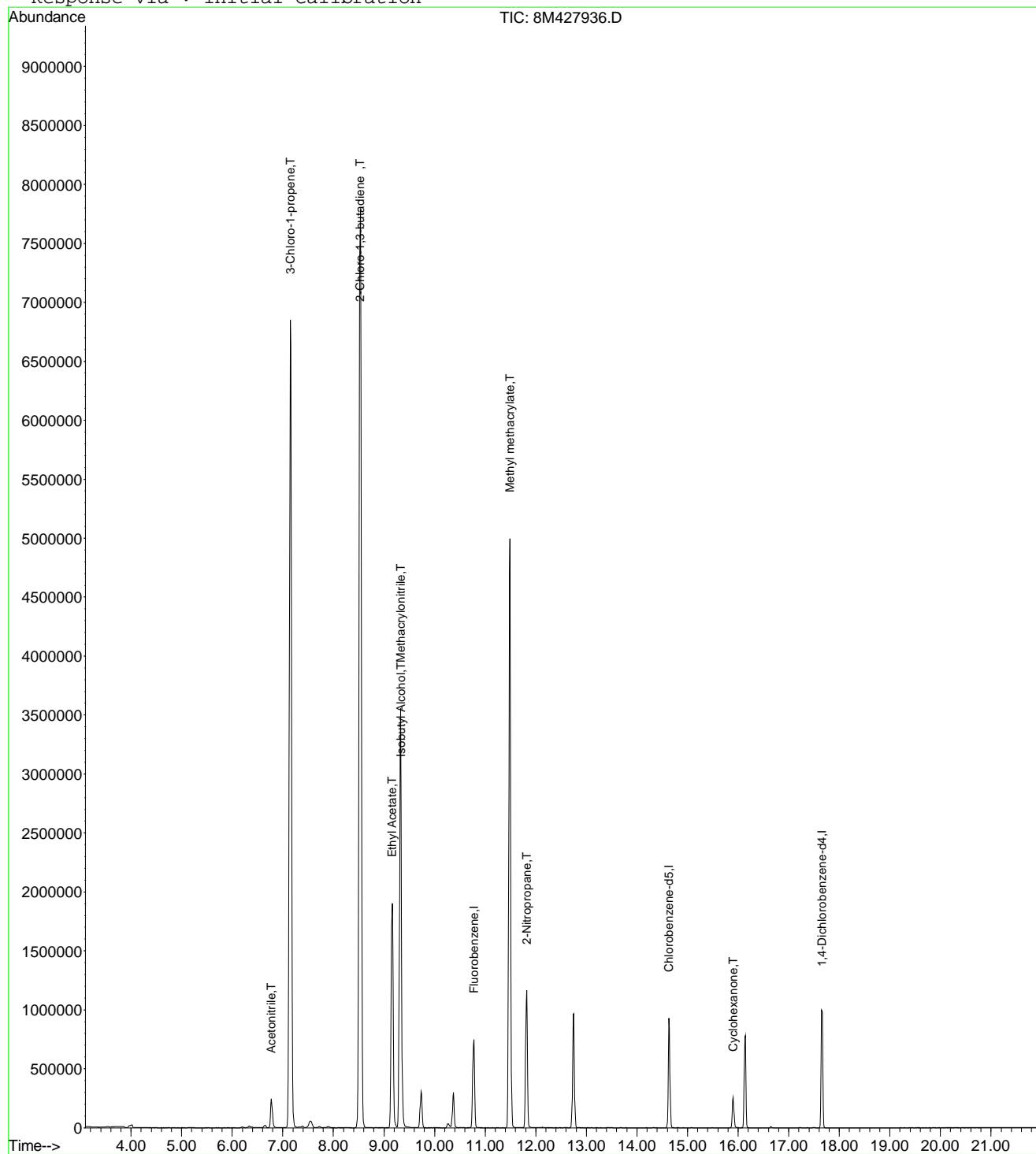
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.77	41	334101	390.5979	ug/L	94
3) 3-Chloro-1-propene	7.15	41	6583975	390.9758	ug/L	96
4) 2-Chloro-1,3-butadiene	8.53	53	6768931	398.2552	ug/L	94
5) Ethyl Acetate	9.16	43	3123513	403.9521	ug/L	99
6) Methacrylonitrile	9.33	67	1474353	419.7542	ug/L	99
7) Isobutyl Alcohol	9.34	43	198441	762.1604	ug/L	97
9) Methyl methacrylate	11.49	41	3222257	429.5357	ug/L	98
10) 2-Nitropropane	11.82	43	1113942	431.1873	ug/L	98
13) Cyclohexanone	15.90	55	144776	393.1417	ug/L	95

 (#) = qualifier out of range (m) = manual integration
 8M427936.D A9FOOWT.M Tue Nov 13 12:19:46 2018

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427936.D Vial: 8
Acq On : 12 Nov 2018 16:28 Operator: EEA
Sample : WG684281-08 400 ug/L ICAL A9 8260 Inst : HPMS8
Misc : 1,1 STD90626 Multiplr: 1.00
MS Integration Params: rteint.p
Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
Title : A9-FOO Water SOP:MSV01 111218 HPMS8
Last Update : Tue Nov 13 11:28:33 2018
Response via : Initial Calibration



8M427936.D A9FOOWT.M

Tue Nov 13 12:19:46 2018

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427937.D Vial: 9
 Acq On : 12 Nov 2018 16:57 Operator: EEA
 Sample : WG684281-09 500 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19:47 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.77	96	947276	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.63	117	711576	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.65	152	362181	25.00	ug/L	0.00

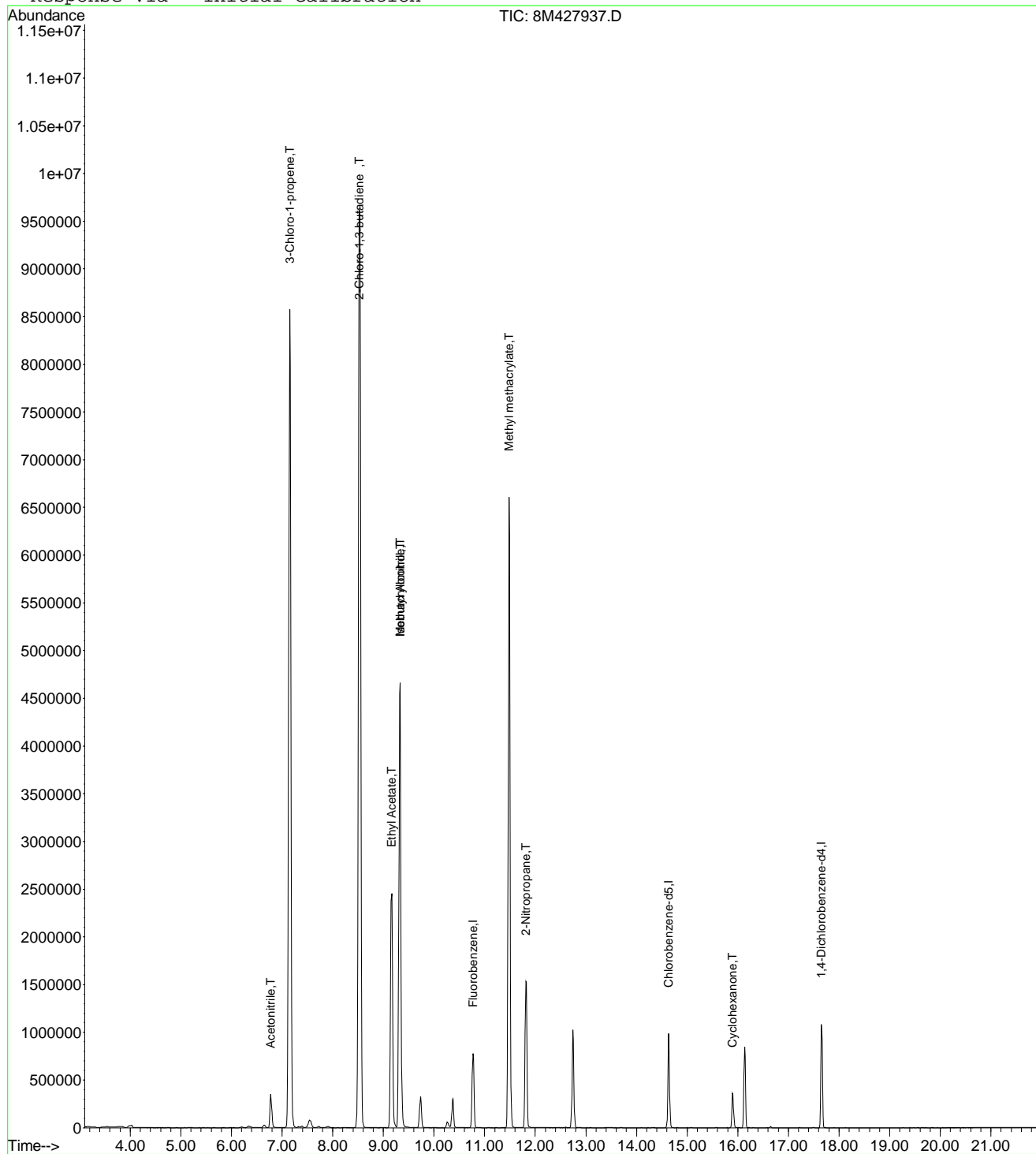
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.78	41	467918	531.4610	ug/L	96
3) 3-Chloro-1-propene	7.15	41	8071291	465.6443	ug/L	93
4) 2-Chloro-1,3-butadiene	8.52	53	8296799	474.2437	ug/L	90
5) Ethyl Acetate	9.17	43	4087769	513.5969	ug/L	98
6) Methacrylonitrile	9.33	67	1949278	539.1591	ug/L	100
7) Isobutyl Alcohol	9.33	43	295664	1103.2223	ug/L	97
9) Methyl methacrylate	11.48	41	4201381	544.1025	ug/L	96
10) 2-Nitropropane	11.81	43	1504638	565.8288	ug/L	96
13) Cyclohexanone	15.90	55	203813	545.4106	ug/L	94

 (#) = qualifier out of range (m) = manual integration
 8M427937.D A9FOOWT.M Tue Nov 13 12:19:47 2018

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427937.D Vial: 9
 Acq On : 12 Nov 2018 16:57 Operator: EEA
 Sample : WG684281-09 500 ug/L ICAL A9 8260 Inst : HPMS8
 Misc : 1,1 STD90626 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:19 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427939.D Vial: 11
 Acq On : 12 Nov 2018 17:54 Operator: EEA
 Sample : WG684 Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Nov 13 12:21:16 2018 Quant Results File: A9FOOWT.RES

Quant Method : K:\ORGANICS\V...\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.78	96	867230	25.00	ug/L	0.00
11) Chlorobenzene-d5	14.63	117	659296	25.00	ug/L	0.00
12) 1,4-Dichlorobenzene-d4	17.65	152	337570	25.00	ug/L	0.00

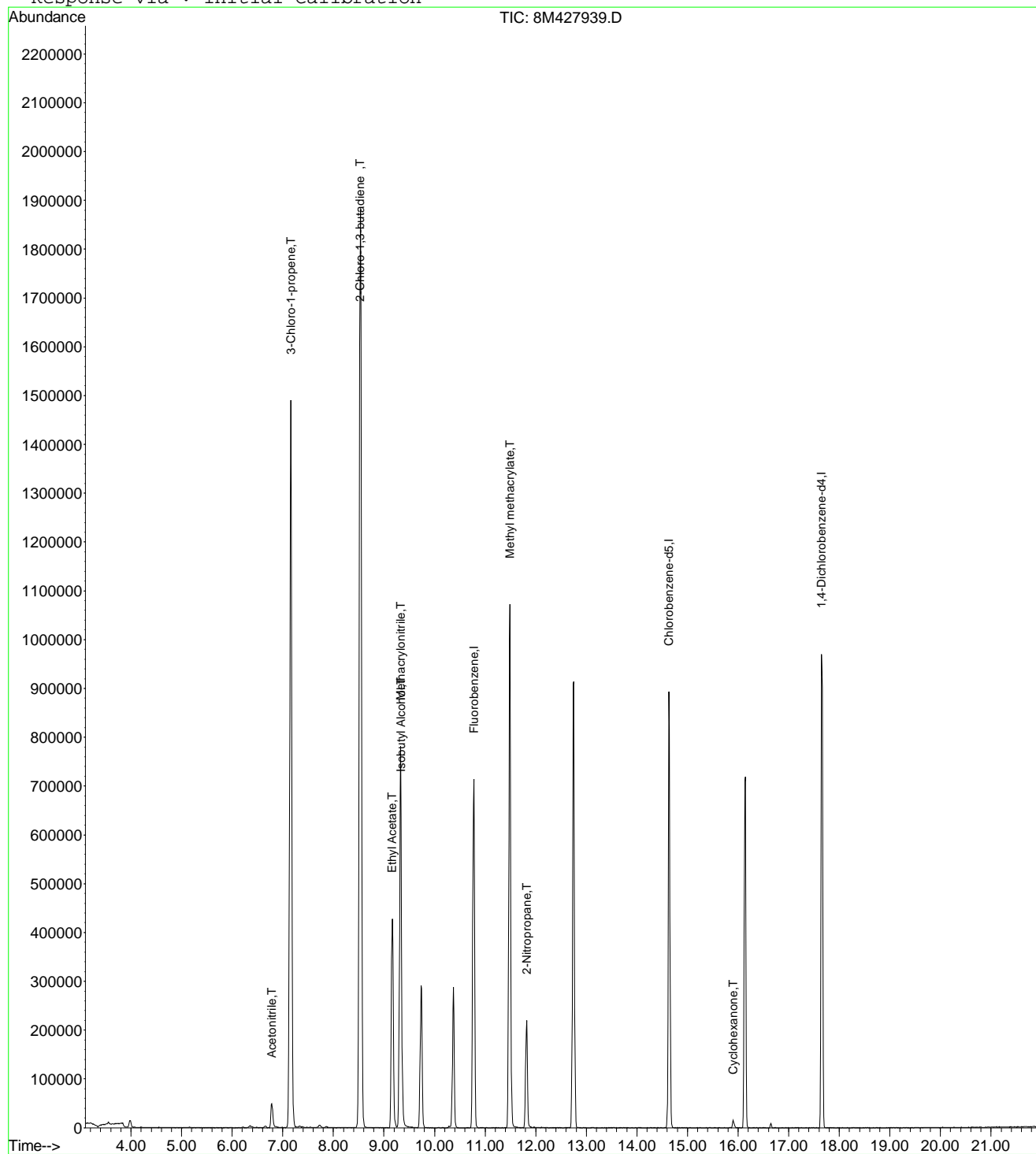
Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Acetonitrile	6.79	41	69875	86.6893	ug/L	96
3) 3-Chloro-1-propene	7.16	41	1445633	91.0986	ug/L	100
4) 2-Chloro-1,3-butadiene	8.53	53	1706775	106.5637	ug/L	99
5) Ethyl Acetate	9.16	43	722113	99.1022	ug/L	99
6) Methacrylonitrile	9.33	67	329117	99.4342	ug/L	98
7) Isobutyl Alcohol	9.34	43	40832	166.4207	ug/L #	8
9) Methyl methacrylate	11.49	41	709327	100.3408	ug/L	100
10) 2-Nitropropane	11.82	43	204866	84.1521	ug/L	100
13) Cyclohexanone	15.90	55	9261	26.5896	ug/L	99

 (#) = qualifier out of range (m) = manual integration
 8M427939.D A9FOOWT.M Tue Nov 13 12:21:17 2018

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427939.D Vial: 11
Acq On : 12 Nov 2018 17:54 Operator: EEA
Sample : WG684 Inst : HPMS8
Misc : 1,1 Multiplr: 1.00
MS Integration Params: rteint.p
Quant Time: Nov 13 12:20 2018 Quant Results File: A9FOOWT.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
Title : A9-FOO Water SOP:MSV01 111218 HPMS8
Last Update : Tue Nov 13 11:28:33 2018
Response via : Initial Calibration



8M427939.D A9FOOWT.M

Tue Nov 13 12:21:17 2018

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427939.D Vial: 11
 Acq On : 12 Nov 2018 17:54 Operator: EEA
 Sample : WG684 Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8
 Last Update : Tue Nov 13 11:28:33 2018
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 1% Max. R.T. Dev 0.50min
 Max. RRF Dev : 75% Max. Rel. Area : 200%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	97	0.00
2 T	Acetonitrile	100.000	86.689	13.3	92	0.00
3 T	3-Chloro-1-propene	100.000	91.099	8.9	85	0.00
4 T	2-Chloro-1,3-butadiene	100.000	106.564	-6.6	100	0.00
5 T	Ethyl Acetate	100.000	99.102	0.9	97	0.00
6 T	Methacrylonitrile	100.000	99.434	0.6	95	0.00
7 T	Isobutyl Alcohol	200.000	166.421	16.8	98	0.00
8 T	1-Butanol	-1.000	0.000	0.0	64	0.00
9 T	Methyl methacrylate	100.000	100.341	-0.3	95	0.00
10 T	2-Nitropropane	100.000	84.152	15.8	85	0.00
11 I	Chlorobenzene-d5	25.000	25.000	0.0	99	0.00
12 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	99	0.00
13 T	Cyclohexanone	100.000	26.590	73.4	30	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 8M427939.D A9FOOWT.M Tue Nov 13 12:22:00 2018

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428956.D Vial: 2
 Acq On : 4 Mar 2019 12:45 Operator: EEA
 Sample : WG698192-02 0.3 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:33 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	556454	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.59	117	412849	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.61	152	214737	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	0.00	111	0	0.0000	ug/L	
Spiked Amount	25.000	Range 86 - 118	Recovery	=	0.00%#	
43) 1,2-Dichloroethane-d4	0.00	65	0	0.0000	ug/L	
Spiked Amount	25.000	Range 80 - 120	Recovery	=	0.00%#	
58) Toluene-d8	12.71	98	5830	0.2772	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	1.12%#	
80) p-Bromofluorobenzene	16.09	95	3136	0.4046	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	1.60%#	

Target Compounds

						Qvalue
2) Dichlorodifluoromethane	3.23	85	2603	0.2689	ug/L	94
3) Chloromethane	3.69	50	6260	0.4497	ug/L #	59
4) Vinyl Chloride	3.91	62	3220	0.2839	ug/L	95
5) 1,3-Butadiene	3.97	54	3386	Below Cal	#	72
6) Bromomethane	4.80	94	2456	0.3600	ug/L	87
7) Chloroethane	4.97	64	1400	0.2306	ug/L #	45
8) Trichlorofluoromethane	5.45	101	3061	0.2658	ug/L #	81
10) Isoprene	6.00	67	2845	0.2472	ug/L	99
12) 1,1,2-Trichloro-1,2,2-Trif	6.22	101	1756	0.2647	ug/L	79
14) 1,1-Dichloroethene	6.52	61	3658	0.3082	ug/L	91
16) Dimethyl Sulfide	6.79	62	2020	0.2363	ug/L	84
19) Methylene Chloride	7.31	84	2445	0.3297	ug/L	94
20) Carbon Disulfide	7.34	76	7502	0.3381	ug/L	97
22) Methyl Tert Butyl Ether	7.51	73	4997	0.2831	ug/L	82
23) trans-1,2-Dichloroethene	7.75	61	3661	0.3229	ug/L	85
24) n-Hexane	7.82	57	3338	0.3000	ug/L	94
27) 1,1-Dichloroethane	8.37	63	4361	0.2975	ug/L #	73
31) 2,2-Dichloropropane	9.14	77	2983	0.2805	ug/L	82
32) cis-1,2-Dichloroethene	9.20	96	1953	0.2413	ug/L	79
33) Chloroform	9.40	83	4369	0.3358	ug/L	96
35) Bromochloromethane	9.64	130	932	0.2019	ug/L	70
36) Tetrahydrofuran	9.69	42	1559	0.9176	ug/L #	40
38) 1,1,1-Trichloroethane	9.94	97	3064	0.2711	ug/L	95
39) Cyclohexane	9.96	56	4646	0.3087	ug/L #	74
40) 1,1-Dichloropropene	10.13	75	2837	0.2990	ug/L #	37
42) Carbon Tetrachloride	10.27	117	2488	0.2423	ug/L #	86
45) 1,2-Dichloroethane	10.46	62	2598	0.2920	ug/L #	75
46) Benzene	10.49	78	9038	0.3041	ug/L	93
47) Trichloroethene	11.24	130	2501	0.3088	ug/L	92
48) Methylcyclohexane	11.33	83	3683	0.2937	ug/L	93
49) 1,2-Dichloropropane	11.45	63	2210	0.2739	ug/L	88
50) Bromodichloromethane	11.75	83	2302	0.2464	ug/L #	90
55) cis-1,3-Dichloropropene	12.39	75	3125	0.2812	ug/L	85
56) Dimethyl Disulfide	12.64	79	1319	0.1933	ug/L	88
59) Toluene	12.81	91	9322	0.3024	ug/L	95
60) Ethyl Methacrylate	12.92	69	704	0.5646	ug/L #	9
62) trans-1,3-Dichloropropene	12.99	75	2419	0.2610	ug/L #	43
63) 1,1,2-Trichloroethane	13.21	97	1108	0.2056	ug/L	90
65) 1,3-Dichloropropane	13.50	76	2630	0.2834	ug/L	98
66) Tetrachloroethene	13.63	164	1640	0.2525	ug/L	98

(#) = qualifier out of range (m) = manual integration
 8M428956.D 8260WTR.M Tue Mar 05 10:54:36 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428956.D Vial: 2
 Acq On : 4 Mar 2019 12:45 Operator: EEA
 Sample : WG698192-02 0.3 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:33 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

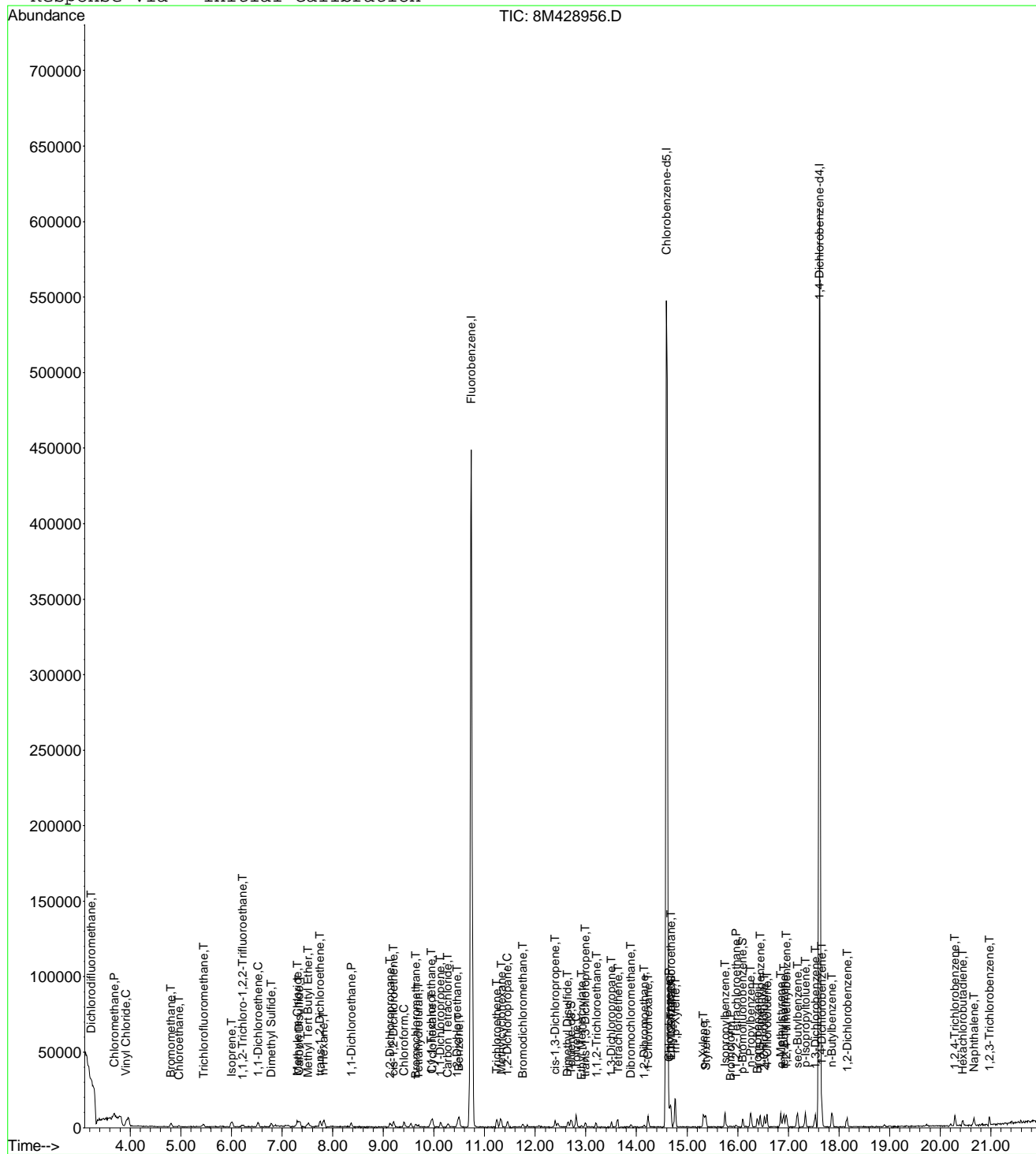
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
67) Dibromochloromethane	13.89	129	1608	0.2351	ug/L	75
68) 1,2-Dibromoethane	14.15	107	1081	0.2066	ug/L	94
69) 1-Chlorohexane	14.23	91	3109	0.2911	ug/L	97
70) Chlorobenzene	14.65	112	7026	0.3362	ug/L	84
71) 1,1,1,2-Tetrachloroethane	14.68	131	2050	0.2668	ug/L	83
72) Ethylbenzene	14.68	106	3105	0.2746	ug/L	91
73) m-,p-Xylene	14.77	106	8538	0.6198	ug/L	99
74) o-Xylene	15.32	106	3828	0.2864	ug/L	88
75) Styrene	15.37	104	5590	0.2526	ug/L	87
76) Bromoform	15.86	173	774	0.1825	ug/L #	27
77) Isopropylbenzene	15.75	105	10184	0.3016	ug/L	97
79) 1,1,2,2-Tetrachloroethane	15.97	83	1768	0.2918	ug/L #	77
83) n-Propylbenzene	16.26	91	12434	0.3200	ug/L	93
84) Bromobenzene	16.39	156	2466	0.2773	ug/L #	67
85) 1,3,5-Trimethylbenzene	16.45	105	8697	0.3105	ug/L	91
86) 2-Chlorotoluene	16.53	91	7138	0.2778	ug/L	95
87) 4-Chlorotoluene	16.58	91	8090	0.3603	ug/L	95
88) a-Methylstyrene	16.85	118	4037	0.2511	ug/L	98
89) tert-Butylbenzene	16.91	134	1609	0.2490	ug/L	60
90) 1,2,4-Trimethylbenzene	16.96	105	8318	0.2933	ug/L	84
91) sec-Butylbenzene	17.18	105	11206	0.3126	ug/L	92
92) p-Isopropyltoluene	17.33	119	9472	0.3082	ug/L	94
93) 1,3-Dichlorobenzene	17.53	146	5075	0.2928	ug/L	98
94) 1,4-Dichlorobenzene	17.67	146	5788	0.3297	ug/L #	26
95) n-Butylbenzene	17.86	91	9088	0.3245	ug/L #	88
96) 1,2-Dichlorobenzene	18.16	146	4801	0.2985	ug/L	89
98) 1,2,4-Trichlorobenzene	20.29	180	3746	0.3209	ug/L	96
99) Hexachlorobutadiene	20.45	225	1598	0.2906	ug/L #	67
100) Naphthalene	20.66	128	6216	0.2922	ug/L #	68
101) 1,2,3-Trichlorobenzene	20.97	180	3248	0.3029	ug/L #	90

(#) = qualifier out of range (m) = manual integration
 8M428956.D 8260WTR.M Tue Mar 05 10:54:36 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428956.D Vial: 2
 Acq On : 4 Mar 2019 12:45 Operator: EEA
 Sample : WG698192-02 0.3 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428956.D Vial: 2
 Acq On : 4 Mar 2019 12:45 Operator: EEA
 Sample : WG698192-02 0.3 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	-1.000	0.269	0.0	100	0.00
3 P	Chloromethane	-1.000	0.450	0.0	100	0.00
4 C	Vinyl Chloride	-1.000	0.284	0.0	100	0.00
5 T	1,3-Butadiene	-1.000	-4.402	0.0	0	0.01
6 T	Bromomethane	-1.000	0.360	0.0	100	0.00
7 T	Chloroethane	-1.000	0.231	0.0	100	0.00
8 T	Trichlorofluoromethane	-1.000	0.266	0.0	100	0.01
9 T	Diethyl ether	-1.000	0.000	0.0	0	-5.97#
10 T	Isoprene	-1.000	0.247	0.0	100	0.00
11 T	Acrolein	-1.000	0.000	0.0	0	-6.21#
12 T	1,1,2-Trichloro-1,2,2-Trifl	-1.000	0.265	0.0	100	0.00
13 T	Acetone	-1.000	0.000	0.0	0	-6.31#
14 C	1,1-Dichloroethene	-1.000	0.308	0.0	100	0.00
15 T	Tert-Butyl Alcohol	-1.000	0.000	0.0	0	-6.63#
16 T	Dimethyl Sulfide	-1.000	0.236	0.0	100	0.00
17 T	Iodomethane	-1.000	0.000	0.0	0	-7.03#
18 T	Methyl acetate	-1.000	0.000	0.0	0	-7.05#
19 T	Methylene Chloride	-1.000	0.330	0.0	100	0.00
20 T	Carbon Disulfide	-1.000	0.338	0.0	100	0.00
21 T	Acrylonitrile	-1.000	0.000	0.0	0	-7.49#
22 T	Methyl Tert Butyl Ether	-1.000	0.283	0.0	100	0.00
23 T	trans-1,2-Dichloroethene	-1.000	0.323	0.0	100	0.00
24 T	n-Hexane	-1.000	0.300	0.0	100	0.00
25 T	Diisopropyl ether	-1.000	0.000	0.0	0	-8.17#
26 T	Vinyl Acetate	-1.000	0.081	0.0	100	0.02
27 P	1,1-Dichloroethane	-1.000	0.298	0.0	100	0.00
28 T	Ethyl-Tert-Butyl ether	-1.000	0.000	0.0	0	-8.74#
29 T	2-Butanone	-1.000	0.000	0.0	0	-8.93#
30 T	Propionitrile	-1.000	0.000	0.0	0	-9.03#
31 T	2,2-Dichloropropane	-1.000	0.280	0.0	100	0.00
32 T	cis-1,2-Dichloroethene	-1.000	0.241	0.0	100	0.00
33 C	Chloroform	0.300	0.336	-12.0	100	0.00
34	1-Bromopropane	-1.000	0.000	0.0	0	-9.55#
35 T	Bromochloromethane	-1.000	0.202	0.0	100	0.00
36 T	Tetrahydrofuran	-1.000	0.918	0.0	100	0.02
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.70#
38 T	1,1,1-Trichloroethane	-1.000	0.271	0.0	100	0.00
39 T	Cyclohexane	-1.000	0.309	0.0	100	0.00
40 T	1,1-Dichloropropene	-1.000	0.299	0.0	100	0.00
41 T	Tert-Amyl-Methyl ether	-1.000	0.000	0.0	0	-10.24#
42 T	Carbon Tetrachloride	-1.000	0.242	0.0	100	0.00
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.33#
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	-1.000	0.292	0.0	100	0.00
46 T	Benzene	-1.000	0.304	0.0	100	0.00
47 T	Trichloroethene	0.300	0.309	-3.0	100	0.00
48 T	Methylcyclohexane	-1.000	0.294	0.0	100	0.00
49 C	1,2-Dichloropropane	-1.000	0.274	0.0	100	0.00
50 T	Bromodichloromethane	-1.000	0.246	0.0	100	0.00
51 T	1,4-Dioxane	-1.000	0.000	0.0	0	-11.75#
52 T	Dibromomethane	-1.000	0.000	0.0	0	-11.84#
53 T	2-Chloroethyl Vinyl Ether	-1.000	0.000	0.0	0	-12.05#
54 T	4-Methyl-2-Pentanone	-1.000	0.000	0.0	0	-12.08#
55 T	cis-1,3-Dichloropropene	-1.000	0.281	0.0	100	0.00
56 T	Dimethyl Disulfide	-1.000	0.193	0.0	100	0.00
57 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00

58	S	Toluene-d8	-1.000	0.277	0.0	100	0.00
59	C	Toluene	-1.000	0.302	0.0	100	0.00
60	T	Ethyl Methacrylate	-1.000	0.565	0.0	100	0.01
61		Paraldehyde	-1.000	0.000	0.0	0	-13.45#
62	T	trans-1,3-Dichloropropene	-1.000	0.261	0.0	100	0.00
63	T	1,1,2-Trichloroethane	-1.000	0.206	0.0	100	0.00
64	T	2-Hexanone	-1.000	0.000	0.0	0	-13.14#
65	T	1,3-Dichloropropane	-1.000	0.283	0.0	100	0.00
66	T	Tetrachloroethene	-1.000	0.252	0.0	100	0.00
67	T	Dibromochloromethane	-1.000	0.235	0.0	100	0.00
68	T	1,2-Dibromoethane	-1.000	0.207	0.0	100	0.00
69	T	1-Chlorohexane	-1.000	0.291	0.0	100	0.00
70	P	Chlorobenzene	0.300	0.336	-12.0	100	0.00
71	T	1,1,1,2-Tetrachloroethane	-1.000	0.267	0.0	100	0.00
72	C	Ethylbenzene	0.300	0.275	8.3	100	0.00
73	T	m-,p-Xylene	0.600	0.620	-3.3	100	0.00
74	T	o-Xylene	0.300	0.286	4.7	100	0.00
75	T	Styrene	-1.000	0.253	0.0	100	0.00
76	P	Bromoform	-1.000	0.183	0.0	100	0.00
77	T	Isopropylbenzene	-1.000	0.302	0.0	100	0.00
78	I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
79	P	1,1,2,2-Tetrachloroethane	-1.000	0.292	0.0	100	0.00
80	S	p-Bromofluorobenzene	-1.000	0.405	0.0	100	0.00
81	T	1,2,3-Trichloropropane	-1.000	0.000	0.0	0	-16.16#
82	T	trans-1,4-Dichloro-2-Butene	-1.000	0.000	0.0	0	-16.20#
83	T	n-Propylbenzene	-1.000	0.320	0.0	100	0.00
84	T	Bromobenzene	-1.000	0.277	0.0	100	0.00
85	T	1,3,5-Trimethylbenzene	-1.000	0.310	0.0	100	0.00
86	T	2-Chlorotoluene	-1.000	0.278	0.0	100	0.00
87	T	4-Chlorotoluene	-1.000	0.360	0.0	100	0.00
88	T	a-Methylstyrene	-1.000	0.251	0.0	100	0.00
89	T	tert-Butylbenzene	-1.000	0.249	0.0	100	0.00
90	T	1,2,4-Trimethylbenzene	-1.000	0.293	0.0	100	0.00
91	T	sec-Butylbenzene	-1.000	0.313	0.0	100	0.00
92	T	p-Isopropyltoluene	-1.000	0.308	0.0	100	0.00
93	T	1,3-Dichlorobenzene	-1.000	0.293	0.0	100	0.00
94	T	1,4-Dichlorobenzene	-1.000	0.330	0.0	100	0.00
95	T	n-Butylbenzene	-1.000	0.325	0.0	100	0.00
96	T	1,2-Dichlorobenzene	0.300	0.298	0.7	100	0.00
97	T	1,2-Dibromo-3-Chloropropane	-1.000	0.000	0.0	0	-19.15#
98	T	1,2,4-Trichlorobenzene	-1.000	0.321	0.0	100	0.00
99	T	Hexachlorobutadiene	-1.000	0.291	0.0	100	0.00
100	T	Naphthalene	-1.000	0.292	0.0	100	0.00
101	T	1,2,3-Trichlorobenzene	-1.000	0.303	0.0	100	0.00

(#) = Out of Range
8M428956.D 8260WTR.M

SPCC's out = 0 CCC's out = 0
Tue Mar 05 10:50:08 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428957.D Vial: 3
 Acq On : 4 Mar 2019 13:15 Operator: EEA
 Sample : WG698192-03 0.4 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:37 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	562289	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	415167	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	213977	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	0.00	111	0	0.0000	ug/L	
Spiked Amount	25.000	Range 86 - 118	Recovery	=	0.00%#	
43) 1,2-Dichloroethane-d4	0.00	65	0	0.0000	ug/L	
Spiked Amount	25.000	Range 80 - 120	Recovery	=	0.00%#	
58) Toluene-d8	12.71	98	3766	0.1781	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	0.72%#	
80) p-Bromofluorobenzene	16.10	95	2055	0.2661	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	1.08%#	

Target Compounds

	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.24	85	2978	0.3045	ug/L	96
3) Chloromethane	3.69	50	7913	0.5626	ug/L	# 60
4) Vinyl Chloride	3.92	62	4171	0.3639	ug/L	94
5) 1,3-Butadiene	3.98	54	3680	Below Cal	#	63
6) Bromomethane	4.81	94	2607	0.3781	ug/L	90
7) Chloroethane	4.97	64	1991	0.3246	ug/L	# 45
8) Trichlorofluoromethane	5.44	101	4235	0.3639	ug/L	# 89
10) Isoprene	6.01	67	3944	0.3391	ug/L	90
12) 1,1,2-Trichloro-1,2,2-Trif	6.21	101	1876	0.2798	ug/L	82
14) 1,1-Dichloroethene	6.54	61	4599	0.3835	ug/L	100
16) Dimethyl Sulfide	6.79	62	2470	0.2859	ug/L	95
19) Methylene Chloride	7.31	84	2981	0.3978	ug/L	90
20) Carbon Disulfide	7.35	76	9485	0.4230	ug/L	# 74
22) Methyl Tert Butyl Ether	7.51	73	6596	0.3698	ug/L	# 76
23) trans-1,2-Dichloroethene	7.75	61	4305	0.3757	ug/L	97
24) n-Hexane	7.83	57	4217	0.3750	ug/L	88
27) 1,1-Dichloroethane	8.37	63	5617	0.3792	ug/L	# 84
31) 2,2-Dichloropropane	9.13	77	4556	0.4239	ug/L	# 69
32) cis-1,2-Dichloroethene	9.21	96	2824	0.3453	ug/L	93
33) Chloroform	9.41	83	5232	0.3980	ug/L	96
35) Bromochloromethane	9.65	130	1640	0.3516	ug/L	87
36) Tetrahydrofuran	9.69	42	1180	0.6874	ug/L	# 40
38) 1,1,1-Trichloroethane	9.94	97	4196	0.3675	ug/L	99
39) Cyclohexane	9.97	56	5825	0.3830	ug/L	# 86
40) 1,1-Dichloropropene	10.14	75	3187	0.3324	ug/L	84
42) Carbon Tetrachloride	10.28	117	3577	0.3447	ug/L	# 85
45) 1,2-Dichloroethane	10.46	62	3047	0.3389	ug/L	# 81
46) Benzene	10.50	78	11649	0.3878	ug/L	92
47) Trichloroethene	11.23	130	2846	0.3478	ug/L	99
48) Methylcyclohexane	11.33	83	4638	0.3660	ug/L	98
49) 1,2-Dichloropropane	11.46	63	2858	0.3505	ug/L	85
50) Bromodichloromethane	11.76	83	3394	0.3595	ug/L	# 80
52) Dibromomethane	11.84	93	846	0.2254	ug/L	73
55) cis-1,3-Dichloropropene	12.39	75	3885	0.3459	ug/L	94
56) Dimethyl Disulfide	12.66	79	1797	0.2606	ug/L	92
59) Toluene	12.82	91	12108	0.3906	ug/L	97
60) Ethyl Methacrylate	12.91	69	1310	0.6387	ug/L	95
62) trans-1,3-Dichloropropene	12.99	75	2606	0.2796	ug/L	94
63) 1,1,2-Trichloroethane	13.20	97	1836	0.3388	ug/L	91
65) 1,3-Dichloropropane	13.51	76	3153	0.3379	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M428957.D 8260WTR.M Tue Mar 05 10:54:40 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428957.D Vial: 3
 Acq On : 4 Mar 2019 13:15 Operator: EEA
 Sample : WG698192-03 0.4 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:37 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

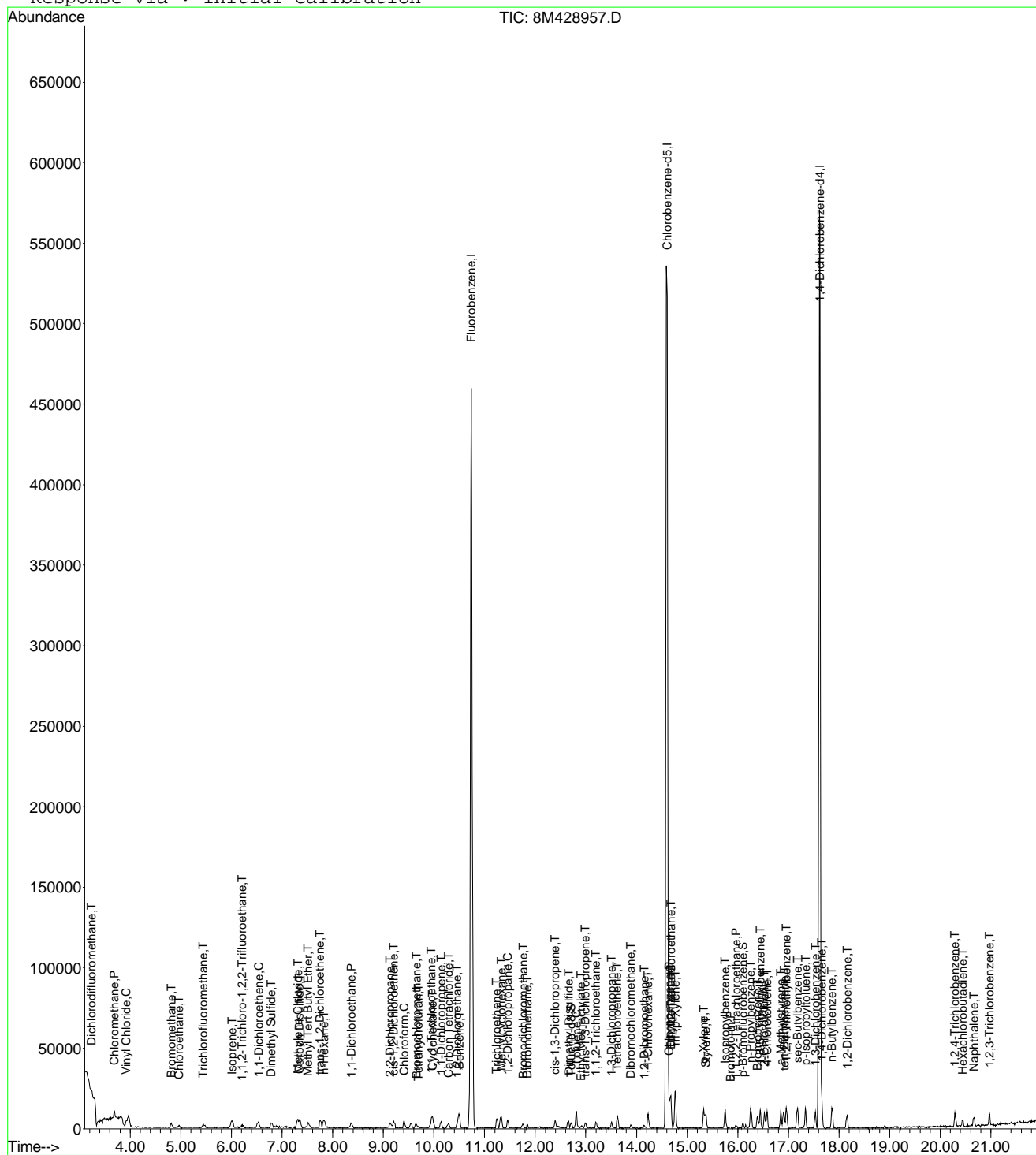
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
66) Tetrachloroethene	13.62	164	2418	0.3702	ug/L	92
67) Dibromochloromethane	13.89	129	1979	0.2878	ug/L	97
68) 1,2-Dibromoethane	14.16	107	1657	0.3149	ug/L	93
69) 1-Chlorohexane	14.23	91	3513	0.3270	ug/L	94
70) Chlorobenzene	14.65	112	7890	0.3755	ug/L	95
71) 1,1,1,2-Tetrachloroethane	14.68	131	2648	0.3427	ug/L	91
72) Ethylbenzene	14.68	106	4269	0.3755	ug/L	92
73) m-,p-Xylene	14.77	106	10707	0.7729	ug/L	97
74) o-Xylene	15.33	106	4827	0.3591	ug/L	94
75) Styrene	15.37	104	7151	0.3213	ug/L	89
76) Bromoform	15.86	173	1112	0.2608	ug/L #	27
77) Isopropylbenzene	15.75	105	12556	0.3698	ug/L	95
79) 1,1,2,2-Tetrachloroethane	15.97	83	2195	0.3636	ug/L #	80
83) n-Propylbenzene	16.26	91	15652	0.4042	ug/L	92
84) Bromobenzene	16.39	156	3007	0.3394	ug/L #	62
85) 1,3,5-Trimethylbenzene	16.44	105	10215	0.3659	ug/L	100
86) 2-Chlorotoluene	16.53	91	9099	0.3554	ug/L	95
87) 4-Chlorotoluene	16.58	91	8507	0.3803	ug/L	99
88) a-Methylstyrene	16.85	118	4857	0.3032	ug/L	98
89) tert-Butylbenzene	16.92	134	2096	0.3255	ug/L	73
90) 1,2,4-Trimethylbenzene	16.96	105	10687	0.3782	ug/L	94
91) sec-Butylbenzene	17.18	105	13726	0.3842	ug/L	97
92) p-Isopropyltoluene	17.34	119	11628	0.3797	ug/L	100
93) 1,3-Dichlorobenzene	17.53	146	6657	0.3854	ug/L	97
94) 1,4-Dichlorobenzene	17.66	146	7394	0.4227	ug/L	94
95) n-Butylbenzene	17.86	91	11247	0.4031	ug/L	92
96) 1,2-Dichlorobenzene	18.16	146	6321	0.3944	ug/L	85
98) 1,2,4-Trichlorobenzene	20.29	180	4623	0.3975	ug/L	99
99) Hexachlorobutadiene	20.45	225	2057	0.3753	ug/L #	82
100) Naphthalene	20.66	128	7298	0.3442	ug/L	95
101) 1,2,3-Trichlorobenzene	20.97	180	4025	0.3767	ug/L	93

(#) = qualifier out of range (m) = manual integration
 8M428957.D 8260WTR.M Tue Mar 05 10:54:40 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428957.D Vial: 3
 Acq On : 4 Mar 2019 13:15 Operator: EEA
 Sample : WG698192-03 0.4 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428957.D Vial: 3
 Acq On : 4 Mar 2019 13:15 Operator: EEA
 Sample : WG698192-03 0.4 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	-1.000	0.304	0.0	100	0.01
3 P	Chloromethane	-1.000	0.563	0.0	100	0.00
4 C	Vinyl Chloride	0.400	0.364	9.0	100	0.01
5 T	1,3-Butadiene	-1.000	-4.359	0.0	0	0.02
6 T	Bromomethane	-1.000	0.378	0.0	100	0.00
7 T	Chloroethane	-1.000	0.325	0.0	100	0.00
8 T	Trichlorofluoromethane	0.400	0.364	9.0	100	0.00
9 T	Diethyl ether	-1.000	0.000	0.0	0	-5.97#
10 T	Isoprene	-1.000	0.339	0.0	100	0.01
11 T	Acrolein	-1.000	0.000	0.0	0	-6.21#
12 T	1,1,2-Trichloro-1,2,2-Trifl	-1.000	0.280	0.0	100	0.00
13 T	Acetone	-1.000	0.000	0.0	0	-6.31#
14 C	1,1-Dichloroethene	0.400	0.384	4.0	100	0.01
15 T	Tert-Butyl Alcohol	-1.000	0.000	0.0	0	-6.63#
16 T	Dimethyl Sulfide	-1.000	0.286	0.0	100	0.00
17 T	Iodomethane	-1.000	0.000	0.0	0	-7.03#
18 T	Methyl acetate	-1.000	0.000	0.0	0	-7.05#
19 T	Methylene Chloride	-1.000	0.398	0.0	100	0.01
20 T	Carbon Disulfide	-1.000	0.423	0.0	100	0.00
21 T	Acrylonitrile	-1.000	0.000	0.0	0	-7.49#
22 T	Methyl Tert Butyl Ether	-1.000	0.370	0.0	100	0.00
23 T	trans-1,2-Dichloroethene	0.400	0.376	6.0	100	0.00
24 T	n-Hexane	-1.000	0.375	0.0	100	0.00
25 T	Diisopropyl ether	-1.000	0.000	0.0	0	-8.17#
26 T	Vinyl Acetate	-1.000	0.108	0.0	100	0.02
27 P	1,1-Dichloroethane	0.400	0.379	5.3	100	0.00
28 T	Ethyl-Tert-Butyl ether	-1.000	0.000	0.0	0	-8.74#
29 T	2-Butanone	-1.000	0.000	0.0	0	-8.93#
30 T	Propionitrile	-1.000	0.000	0.0	0	-9.03#
31 T	2,2-Dichloropropane	-1.000	0.424	0.0	100	0.00
32 T	cis-1,2-Dichloroethene	0.400	0.345	13.8	100	0.00
33 C	Chloroform	0.400	0.398	0.5	100	0.00
34	1-Bromopropane	-1.000	0.000	0.0	0	-9.55#
35 T	Bromochloromethane	-1.000	0.352	0.0	100	0.01
36 T	Tetrahydrofuran	-1.000	0.687	0.0	100	0.02
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.70#
38 T	1,1,1-Trichloroethane	-1.000	0.367	0.0	100	0.00
39 T	Cyclohexane	-1.000	0.383	0.0	100	0.00
40 T	1,1-Dichloropropene	0.400	0.332	17.0	100	0.00
41 T	Tert-Amyl-Methyl ether	-1.000	0.000	0.0	0	-10.24#
42 T	Carbon Tetrachloride	-1.000	0.345	0.0	100	0.00
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.33#
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	0.400	0.339	15.3	100	0.00
46 T	Benzene	0.400	0.388	3.0	100	0.00
47 T	Trichloroethene	0.400	0.348	13.0	100	0.00
48 T	Methylcyclohexane	-1.000	0.366	0.0	100	0.00
49 C	1,2-Dichloropropane	0.400	0.351	12.3	100	0.00
50 T	Bromodichloromethane	0.400	0.360	10.0	100	0.00
51 T	1,4-Dioxane	-1.000	0.000	0.0	0	-11.75#
52 T	Dibromomethane	-1.000	0.225	0.0	100	0.00
53 T	2-Chloroethyl Vinyl Ether	-1.000	0.000	0.0	0	-12.05#
54 T	4-Methyl-2-Pentanone	-1.000	0.000	0.0	0	-12.08#
55 T	cis-1,3-Dichloropropene	0.400	0.346	13.5	100	0.00
56 T	Dimethyl Disulfide	-1.000	0.261	0.0	100	0.01
57 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00

58 S	Toluene-d8	-1.000	0.178	0.0	100	0.00
59 C	Toluene	0.400	0.391	2.3	100	0.00
60 T	Ethyl Methacrylate	-1.000	0.639	0.0	100	0.00
61	Paraldehyde	-1.000	0.000	0.0	0	-13.45#
62 T	trans-1,3-Dichloropropene	-1.000	0.280	0.0	100	0.00
63 T	1,1,2-Trichloroethane	0.400	0.339	15.3	100	0.00
64 T	2-Hexanone	-1.000	0.000	0.0	0	-13.14#
65 T	1,3-Dichloropropane	0.400	0.338	15.5	100	0.00
66 T	Tetrachloroethene	0.400	0.370	7.5	100	0.00
67 T	Dibromochloromethane	-1.000	0.288	0.0	100	0.00
68 T	1,2-Dibromoethane	0.400	0.315	21.3	100	0.01
69 T	1-Chlorohexane	-1.000	0.327	0.0	100	0.00
70 P	Chlorobenzene	0.400	0.375	6.3	100	0.00
71 T	1,1,1,2-Tetrachloroethane	-1.000	0.343	0.0	100	0.00
72 C	Ethylbenzene	0.400	0.375	6.3	100	0.00
73 T	m-,p-Xylene	0.800	0.773	3.4	100	0.00
74 T	o-Xylene	0.400	0.359	10.3	100	0.00
75 T	Styrene	-1.000	0.321	0.0	100	0.00
76 P	Bromoform	-1.000	0.261	0.0	100	0.00
77 T	Isopropylbenzene	0.400	0.370	7.5	100	0.00
78 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
79 P	1,1,2,2-Tetrachloroethane	0.400	0.364	9.0	100	0.00
80 S	p-Bromofluorobenzene	-1.000	0.266	0.0	100	0.00
81 T	1,2,3-Trichloropropane	-1.000	0.000	0.0	0	-16.16#
82 T	trans-1,4-Dichloro-2-Butene	-1.000	0.000	0.0	0	-16.20#
83 T	n-Propylbenzene	0.400	0.404	-1.0	100	0.00
84 T	Bromobenzene	0.400	0.339	15.3	100	0.00
85 T	1,3,5-Trimethylbenzene	0.400	0.366	8.5	100	0.00
86 T	2-Chlorotoluene	0.400	0.355	11.3	100	0.00
87 T	4-Chlorotoluene	0.400	0.380	5.0	100	0.00
88 T	a-Methylstyrene	-1.000	0.303	0.0	100	0.00
89 T	tert-Butylbenzene	-1.000	0.325	0.0	100	0.01
90 T	1,2,4-Trimethylbenzene	0.400	0.378	5.5	100	0.00
91 T	sec-Butylbenzene	0.400	0.384	4.0	100	0.00
92 T	p-Isopropyltoluene	0.400	0.380	5.0	100	0.01
93 T	1,3-Dichlorobenzene	0.400	0.385	3.8	100	0.00
94 T	1,4-Dichlorobenzene	0.400	0.423	-5.7	100	0.00
95 T	n-Butylbenzene	0.400	0.403	-0.8	100	0.00
96 T	1,2-Dichlorobenzene	0.400	0.394	1.5	100	0.00
97 T	1,2-Dibromo-3-Chloropropane	-1.000	0.000	0.0	0	-19.15#
98 T	1,2,4-Trichlorobenzene	0.400	0.397	0.8	100	0.00
99 T	Hexachlorobutadiene	0.400	0.375	6.3	100	0.00
100 T	Naphthalene	0.400	0.344	14.0	100	0.00
101 T	1,2,3-Trichlorobenzene	0.400	0.377	5.8	100	0.00

(#) = Out of Range
8M428957.D 8260WTR.M

SPCC's out = 0 CCC's out = 0
Tue Mar 05 10:50:33 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428958.D Vial: 4
 Acq On : 4 Mar 2019 13:43 Operator: EEA
 Sample : WG698192-04 1 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:41 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	567997	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	414878	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	217135	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	0.00	111	0	0.0000	ug/L	
Spiked Amount	25.000	Range 86 - 118	Recovery	=	0.00%#	
43) 1,2-Dichloroethane-d4	0.00	65	0	0.0000	ug/L	
Spiked Amount	25.000	Range 80 - 120	Recovery	=	0.00%#	
58) Toluene-d8	12.71	98	3302	0.1562	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	0.64%#	
80) p-Bromofluorobenzene	16.10	95	2268	0.2894	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	1.16%#	

Target Compounds

						Qvalue
2) Dichlorodifluoromethane	3.23	85	8615	0.8720	ug/L	98
3) Chloromethane	3.69	50	15740	1.1078	ug/L	81
4) Vinyl Chloride	3.92	62	10943	0.9451	ug/L	97
5) 1,3-Butadiene	3.96	54	10326	Below Cal		91
6) Bromomethane	4.81	94	7076	1.0160	ug/L	96
7) Chloroethane	4.97	64	5329	0.8600	ug/L	84
8) Trichlorofluoromethane	5.44	101	10885	0.9259	ug/L	# 96
9) Diethyl ether	5.97	59	27357	4.3632	ug/L	100
10) Isoprene	6.01	67	10172	0.8657	ug/L	95
11) Acrolein	6.22	56	691	0.9611	ug/L	# 15
12) 1,1,2-Trichloro-1,2,2-Trif	6.21	101	6004	0.8865	ug/L	99
13) Acetone	6.32	43	1013	0.5790	ug/L	# 45
14) 1,1-Dichloroethene	6.53	61	11458	0.9459	ug/L	96
15) Tert-Butyl Alcohol	6.65	59	4346	7.7165	ug/L	# 87
16) Dimethyl Sulfide	6.79	62	7128	0.8169	ug/L	98
17) Iodomethane	7.06	142	1223	1.6453	ug/L	# 36
18) Methyl acetate	7.08	43	2460	0.5430	ug/L	# 65
19) Methylene Chloride	7.30	84	6798	0.8981	ug/L	95
20) Carbon Disulfide	7.35	76	22105	0.9759	ug/L	99
21) Acrylonitrile	7.51	53	2590	1.1799	ug/L	100
22) Methyl Tert Butyl Ether	7.52	73	16464	0.9138	ug/L	93
23) trans-1,2-Dichloroethene	7.75	61	10924	0.9438	ug/L	97
24) n-Hexane	7.82	57	10796	0.9504	ug/L	98
25) Diisopropyl ether	8.17	45	131662	4.5950	ug/L	99
26) Vinyl Acetate	8.36	43	6278	0.5906	ug/L	# 77
27) 1,1-Dichloroethane	8.37	63	14180	0.9477	ug/L	96
28) Ethyl-Tert-Butyl ether	8.74	59	113232	4.5595	ug/L	98
30) Propionitrile	9.04	54	664	5.3104	ug/L	# 58
31) 2,2-Dichloropropane	9.14	77	10364	0.9546	ug/L	99
32) cis-1,2-Dichloroethene	9.21	96	7715	0.9339	ug/L	90
33) Chloroform	9.41	83	12814	0.9649	ug/L	96
34) 1-Bromopropane	9.55	122	1292	0.7276	ug/L	83
35) Bromochloromethane	9.64	130	3907	0.8293	ug/L	89
36) Tetrahydrofuran	9.68	42	8257	4.7614	ug/L	89
38) 1,1,1-Trichloroethane	9.95	97	10277	0.8910	ug/L	99
39) Cyclohexane	9.98	56	14240	0.9269	ug/L	99
40) 1,1-Dichloropropene	10.14	75	9179	0.9477	ug/L	96
41) Tert-Amyl-Methyl ether	10.24	73	84665	4.4737	ug/L	98
42) Carbon Tetrachloride	10.28	117	9534	0.9096	ug/L	98
45) 1,2-Dichloroethane	10.46	62	8742	0.9624	ug/L	93

(#) = qualifier out of range (m) = manual integration
 8M428958.D 8260WTR.M Tue Mar 05 10:54:44 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428958.D Vial: 4
 Acq On : 4 Mar 2019 13:43 Operator: EEA
 Sample : WG698192-04 1 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:41 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

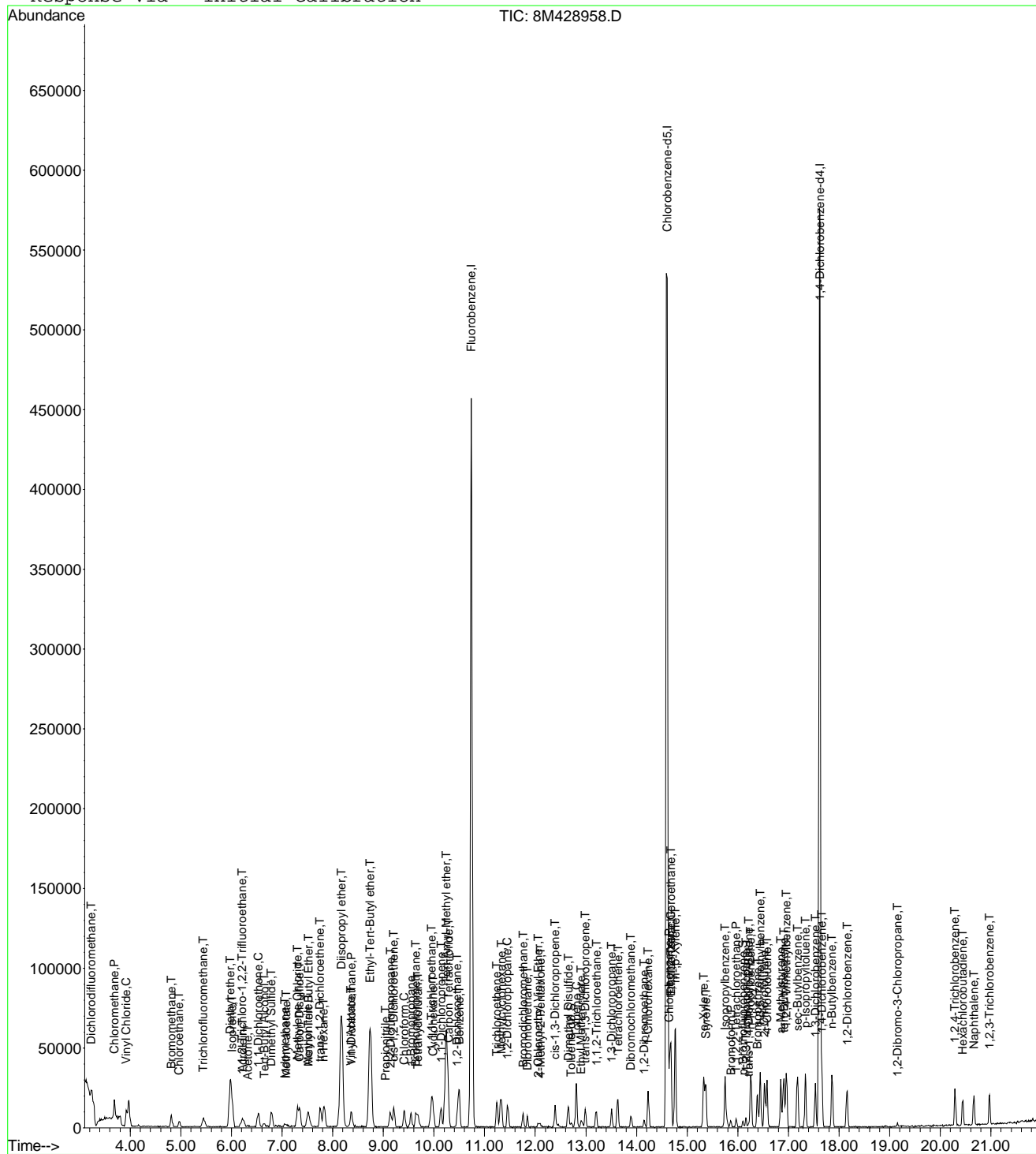
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
46) Benzene	10.50	78	28601	0.9426	ug/L	99
47) Trichloroethene	11.24	130	7731	0.9352	ug/L	97
48) Methylcyclohexane	11.33	83	11923	0.9313	ug/L	99
49) 1,2-Dichloropropane	11.45	63	7389	0.8972	ug/L	98
50) Bromodichloromethane	11.76	83	8277	0.8680	ug/L #	92
52) Dibromomethane	11.84	93	3237	0.8539	ug/L	95
53) 2-Chloroethyl Vinyl Ether	12.06	63	1864	0.5180	ug/L #	49
54) 4-Methyl-2-Pentanone	12.10	58	674	0.2977	ug/L #	41
55) cis-1,3-Dichloropropene	12.39	75	10306	0.9084	ug/L	99
56) Dimethyl Disulfide	12.65	79	4981	0.7150	ug/L	100
59) Toluene	12.82	91	29850	0.9636	ug/L	97
60) Ethyl Methacrylate	12.91	69	4388	1.0179	ug/L	85
62) trans-1,3-Dichloropropene	12.99	75	7608	0.8170	ug/L	99
63) 1,1,2-Trichloroethane	13.21	97	4956	0.9153	ug/L	99
65) 1,3-Dichloropropane	13.51	76	8573	0.9193	ug/L	94
66) Tetrachloroethene	13.63	164	6159	0.9435	ug/L	97
67) Dibromochloromethane	13.89	129	5806	0.8449	ug/L	100
68) 1,2-Dibromoethane	14.15	107	4641	0.8827	ug/L	97
69) 1-Chlorohexane	14.23	91	10107	0.9416	ug/L	97
70) Chlorobenzene	14.65	112	20039	0.9543	ug/L	82
71) 1,1,1,2-Tetrachloroethane	14.68	131	6625	0.8581	ug/L	100
72) Ethylbenzene	14.68	106	10609	0.9337	ug/L	95
73) m-,p-Xylene	14.77	106	26529	1.9163	ug/L	91
74) o-Xylene	15.33	106	12046	0.8968	ug/L	87
75) Styrene	15.37	104	19275	0.8667	ug/L	96
76) Bromoform	15.87	173	3355	0.7874	ug/L	93
77) Isopropylbenzene	15.75	105	32684	0.9633	ug/L	97
79) 1,1,2,2-Tetrachloroethane	15.97	83	5438	0.8877	ug/L #	91
81) 1,2,3-Trichloropropane	16.16	110	1456	0.7945	ug/L #	56
82) trans-1,4-Dichloro-2-Butene	16.22	53	1255	0.6060	ug/L #	49
83) n-Propylbenzene	16.26	91	38492	0.9796	ug/L	98
84) Bromobenzene	16.39	156	8281	0.9210	ug/L	95
85) 1,3,5-Trimethylbenzene	16.44	105	27707	0.9781	ug/L	96
86) 2-Chlorotoluene	16.53	91	26656	1.0259	ug/L	98
87) 4-Chlorotoluene	16.58	91	21953	0.9670	ug/L	97
88) a-Methylstyrene	16.85	118	13813	0.8498	ug/L	97
89) tert-Butylbenzene	16.91	134	5896	0.9022	ug/L	95
90) 1,2,4-Trimethylbenzene	16.96	105	26912	0.9384	ug/L	99
91) sec-Butylbenzene	17.18	105	35426	0.9772	ug/L	97
92) p-Isopropyltoluene	17.33	119	29532	0.9503	ug/L	98
93) 1,3-Dichlorobenzene	17.53	146	16643	0.9495	ug/L	97
94) 1,4-Dichlorobenzene	17.66	146	17019	0.9588	ug/L	79
95) n-Butylbenzene	17.86	91	26429	0.9334	ug/L	98
96) 1,2-Dichlorobenzene	18.16	146	15857	0.9749	ug/L	93
97) 1,2-Dibromo-3-Chloropropan	19.15	75	742	0.6720	ug/L	97
98) 1,2,4-Trichlorobenzene	20.29	180	10753	0.9111	ug/L	99
99) Hexachlorobutadiene	20.45	225	4942	0.8887	ug/L	93
100) Naphthalene	20.66	128	20225	0.9401	ug/L	98
101) 1,2,3-Trichlorobenzene	20.97	180	10148	0.9360	ug/L	96

(#) = qualifier out of range (m) = manual integration
 8M428958.D 8260WTR.M Tue Mar 05 10:54:44 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428958.D Vial: 4
 Acq On : 4 Mar 2019 13:43 Operator: EEA
 Sample : WG698192-04 1 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428958.D Vial: 4
 Acq On : 4 Mar 2019 13:43 Operator: EEA
 Sample : WG698192-04 1 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	1.000	0.872	12.8	100	0.00
3 P	Chloromethane	1.000	1.108	-10.8	100	0.00
4 C	Vinyl Chloride	1.000	0.945	5.5	100	0.01
5 T	1,3-Butadiene	-1.000	-3.263	0.0	0	0.01
6 T	Bromomethane	1.000	1.016	-1.6	100	0.00
7 T	Chloroethane	1.000	0.860	14.0	100	0.00
8 T	Trichlorofluoromethane	1.000	0.926	7.4	100	0.00
9 T	Diethyl ether	5.000	4.363	12.7	100	0.00
10 T	Isoprene	-1.000	0.866	0.0	100	0.01
11 T	Acrolein	-1.000	0.961	0.0	100	0.01
12 T	1,1,2-Trichloro-1,2,2-Trifl	1.000	0.886	11.4	100	0.00
13 T	Acetone	-1.000	0.579	0.0	100	0.01
14 C	1,1-Dichloroethene	1.000	0.946	5.4	100	0.00
15 T	Tert-Butyl Alcohol	-1.000	7.716	0.0	100	0.02
16 T	Dimethyl Sulfide	-1.000	0.817	0.0	100	0.00
17 T	Iodomethane	1.000	1.645	-64.5#	100	0.02
18 T	Methyl acetate	-1.000	0.543	0.0	100	0.02
19 T	Methylene Chloride	1.000	0.898	10.2	100	0.00
20 T	Carbon Disulfide	1.000	0.976	2.4	100	0.00
21 T	Acrylonitrile	-1.000	1.180	0.0	100	0.02
22 T	Methyl Tert Butyl Ether	1.000	0.914	8.6	100	0.01
23 T	trans-1,2-Dichloroethene	1.000	0.944	5.6	100	0.00
24 T	n-Hexane	1.000	0.950	5.0	100	0.00
25 T	Diisopropyl ether	5.000	4.595	8.1	100	0.00
26 T	Vinyl Acetate	-1.000	0.591	0.0	100	0.02
27 P	1,1-Dichloroethane	1.000	0.948	5.2	100	0.00
28 T	Ethyl-Tert-Butyl ether	5.000	4.560	8.8	100	0.00
29 T	2-Butanone	-1.000	0.000	0.0	0	-8.93#
30 T	Propionitrile	5.000	5.310	-6.2	100	0.01
31 T	2,2-Dichloropropane	1.000	0.955	4.5	100	0.00
32 T	cis-1,2-Dichloroethene	1.000	0.934	6.6	100	0.00
33 C	Chloroform	1.000	0.965	3.5	100	0.00
34	1-Bromopropane	-1.000	0.728	0.0	100	0.00
35 T	Bromochloromethane	1.000	0.829	17.1	100	0.00
36 T	Tetrahydrofuran	5.000	4.761	4.8	100	0.01
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.70#
38 T	1,1,1-Trichloroethane	1.000	0.891	10.9	100	0.01
39 T	Cyclohexane	1.000	0.927	7.3	100	0.01
40 T	1,1-Dichloropropene	1.000	0.948	5.2	100	0.00
41 T	Tert-Amyl-Methyl ether	5.000	4.474	10.5	100	0.00
42 T	Carbon Tetrachloride	1.000	0.910	9.0	100	0.00
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.33#
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	1.000	0.962	3.8	100	0.00
46 T	Benzene	1.000	0.943	5.7	100	0.00
47 T	Trichloroethene	1.000	0.935	6.5	100	0.00
48 T	Methylcyclohexane	1.000	0.931	6.9	100	0.00
49 C	1,2-Dichloropropane	1.000	0.897	10.3	100	0.00
50 T	Bromodichloromethane	1.000	0.868	13.2	100	0.00
51 T	1,4-Dioxane	-1.000	0.000	0.0	0	-11.75#
52 T	Dibromomethane	1.000	0.854	14.6	100	0.00
53 T	2-Chloroethyl Vinyl Ether	-1.000	0.518	0.0	100	0.01
54 T	4-Methyl-2-Pentanone	-1.000	0.298	0.0	100	0.02
55 T	cis-1,3-Dichloropropene	1.000	0.908	9.2	100	0.00
56 T	Dimethyl Disulfide	-1.000	0.715	0.0	100	0.00
57 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00

58 S	Toluene-d8	-1.000	0.156	0.0	100	0.00
59 C	Toluene	1.000	0.964	3.6	100	0.00
60 T	Ethyl Methacrylate	1.000	1.018	-1.8	100	0.00
61	Paraldehyde	-1.000	0.000	0.0	0	-13.45#
62 T	trans-1,3-Dichloropropene	1.000	0.817	18.3	100	0.00
63 T	1,1,2-Trichloroethane	1.000	0.915	8.5	100	0.00
64 T	2-Hexanone	-1.000	0.000	0.0	0	-13.14#
65 T	1,3-Dichloropropane	1.000	0.919	8.1	100	0.00
66 T	Tetrachloroethene	1.000	0.943	5.7	100	0.00
67 T	Dibromochloromethane	1.000	0.845	15.5	100	0.00
68 T	1,2-Dibromoethane	1.000	0.883	11.7	100	0.00
69 T	1-Chlorohexane	1.000	0.942	5.8	100	0.00
70 P	Chlorobenzene	1.000	0.954	4.6	100	0.00
71 T	1,1,1,2-Tetrachloroethane	1.000	0.858	14.2	100	0.00
72 C	Ethylbenzene	1.000	0.934	6.6	100	0.00
73 T	m-,p-Xylene	2.000	1.916	4.2	100	0.00
74 T	o-Xylene	1.000	0.897	10.3	100	0.00
75 T	Styrene	1.000	0.867	13.3	100	0.00
76 P	Bromoform	1.000	0.787	21.3	100	0.00
77 T	Isopropylbenzene	1.000	0.963	3.7	100	0.00
78 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
79 P	1,1,2,2-Tetrachloroethane	1.000	0.888	11.2	100	0.00
80 S	p-Bromofluorobenzene	-1.000	0.289	0.0	100	0.00
81 T	1,2,3-Trichloropropane	1.000	0.795	20.5	100	0.00
82 T	trans-1,4-Dichloro-2-Butene	-1.000	0.606	0.0	100	0.01
83 T	n-Propylbenzene	1.000	0.980	2.0	100	0.00
84 T	Bromobenzene	1.000	0.921	7.9	100	0.00
85 T	1,3,5-Trimethylbenzene	1.000	0.978	2.2	100	0.00
86 T	2-Chlorotoluene	1.000	1.026	-2.6	100	0.00
87 T	4-Chlorotoluene	1.000	0.967	3.3	100	0.00
88 T	a-Methylstyrene	-1.000	0.850	0.0	100	0.00
89 T	tert-Butylbenzene	1.000	0.902	9.8	100	0.00
90 T	1,2,4-Trimethylbenzene	1.000	0.938	6.2	100	0.00
91 T	sec-Butylbenzene	1.000	0.977	2.3	100	0.00
92 T	p-Isopropyltoluene	1.000	0.950	5.0	100	0.00
93 T	1,3-Dichlorobenzene	1.000	0.950	5.0	100	0.00
94 T	1,4-Dichlorobenzene	1.000	0.959	4.1	100	0.00
95 T	n-Butylbenzene	1.000	0.933	6.7	100	0.00
96 T	1,2-Dichlorobenzene	1.000	0.975	2.5	100	0.00
97 T	1,2-Dibromo-3-Chloropropane	-1.000	0.672	0.0	100	0.00
98 T	1,2,4-Trichlorobenzene	1.000	0.911	8.9	100	0.00
99 T	Hexachlorobutadiene	1.000	0.889	11.1	100	0.00
100 T	Naphthalene	1.000	0.940	6.0	100	0.00
101 T	1,2,3-Trichlorobenzene	1.000	0.936	6.4	100	0.00

(#) = Out of Range
8M428958.D 8260WTR.M

SPCC's out = 0 CCC's out = 0
Tue Mar 05 10:51:04 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428959.D Vial: 5
 Acq On : 4 Mar 2019 14:12 Operator: EEA
 Sample : WG698192-05 2 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:45 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	517395	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.59	117	373582	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.61	152	191630	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	0.00	111	0	0.0000	ug/L	
Spiked Amount	25.000	Range 86 - 118	Recovery	=	0.00%#	
43) 1,2-Dichloroethane-d4	0.00	65	0	0.0000	ug/L	
Spiked Amount	25.000	Range 80 - 120	Recovery	=	0.00%#	
58) Toluene-d8	12.71	98	2209	0.1161	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	0.48%#	
80) p-Bromofluorobenzene	16.10	95	1027	0.1485	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	0.60%#	

Target Compounds

						Qvalue
2) Dichlorodifluoromethane	3.23	85	16516	1.8352	ug/L	96
3) Chloromethane	3.69	50	29070	2.2461	ug/L	91
4) Vinyl Chloride	3.91	62	21552	2.0433	ug/L	98
5) 1,3-Butadiene	3.96	54	19915	Below Cal		93
6) Bromomethane	4.80	94	14780	2.3297	ug/L	100
7) Chloroethane	4.97	64	11143	1.9742	ug/L	97
8) Trichlorofluoromethane	5.44	101	21539	2.0113	ug/L	99
9) Diethyl ether	5.97	59	119157	20.8630	ug/L	100
10) Isoprene	6.00	67	20637	1.9282	ug/L	96
11) Acrolein	6.21	56	6115	9.3371	ug/L	91
12) 1,1,2-Trichloro-1,2,2-Trif	6.22	101	11936	1.9347	ug/L	98
13) Acetone	6.32	43	2402	1.5071	ug/L	# 45
14) 1,1-Dichloroethene	6.53	61	22437	2.0334	ug/L	98
15) Tert-Butyl Alcohol	6.64	59	22670	44.1879	ug/L	93
16) Dimethyl Sulfide	6.79	62	15368	1.9335	ug/L	100
17) Iodomethane	7.04	142	2763	1.8427	ug/L	89
18) Methyl acetate	7.07	43	5803	1.4061	ug/L	# 82
19) Methylene Chloride	7.29	84	14273	2.0701	ug/L	97
20) Carbon Disulfide	7.35	76	42463	2.0580	ug/L	96
21) Acrylonitrile	7.49	53	17921	8.9624	ug/L	100
22) Methyl Tert Butyl Ether	7.51	73	33798	2.0594	ug/L	96
23) trans-1,2-Dichloroethene	7.75	61	21304	2.0206	ug/L	100
24) n-Hexane	7.82	57	20686	1.9992	ug/L	98
25) Diisopropyl ether	8.16	45	557127	21.3454	ug/L	99
26) Vinyl Acetate	8.35	43	16460	1.7000	ug/L	# 89
27) 1,1-Dichloroethane	8.37	63	28174	2.0671	ug/L	98
28) Ethyl-Tert-Butyl ether	8.74	59	476596	21.0680	ug/L	99
29) 2-Butanone	8.94	43	3302	1.4064	ug/L	# 56
30) Propionitrile	9.04	54	11163	19.3270	ug/L	94
31) 2,2-Dichloropropane	9.13	77	20615	2.0845	ug/L	98
32) cis-1,2-Dichloroethene	9.21	96	15272	2.0295	ug/L	98
33) Chloroform	9.41	83	24223	2.0024	ug/L	99
34) 1-Bromopropane	9.54	122	2860	1.7681	ug/L	88
35) Bromochloromethane	9.64	130	8463	1.9720	ug/L	97
36) Tetrahydrofuran	9.67	42	33536	21.2299	ug/L	99
38) 1,1,1-Trichloroethane	9.94	97	21108	2.0089	ug/L	99
39) Cyclohexane	9.97	56	27878	1.9921	ug/L	99
40) 1,1-Dichloropropene	10.14	75	18198	2.0626	ug/L	94
41) Tert-Amyl-Methyl ether	10.24	73	363559	21.0894	ug/L	99
42) Carbon Tetrachloride	10.28	117	18818	1.9709	ug/L	98

(#) = qualifier out of range (m) = manual integration
 8M428959.D 8260WTR.M Tue Mar 05 10:54:47 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428959.D Vial: 5
 Acq On : 4 Mar 2019 14:12 Operator: EEA
 Sample : WG698192-05 2 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:45 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

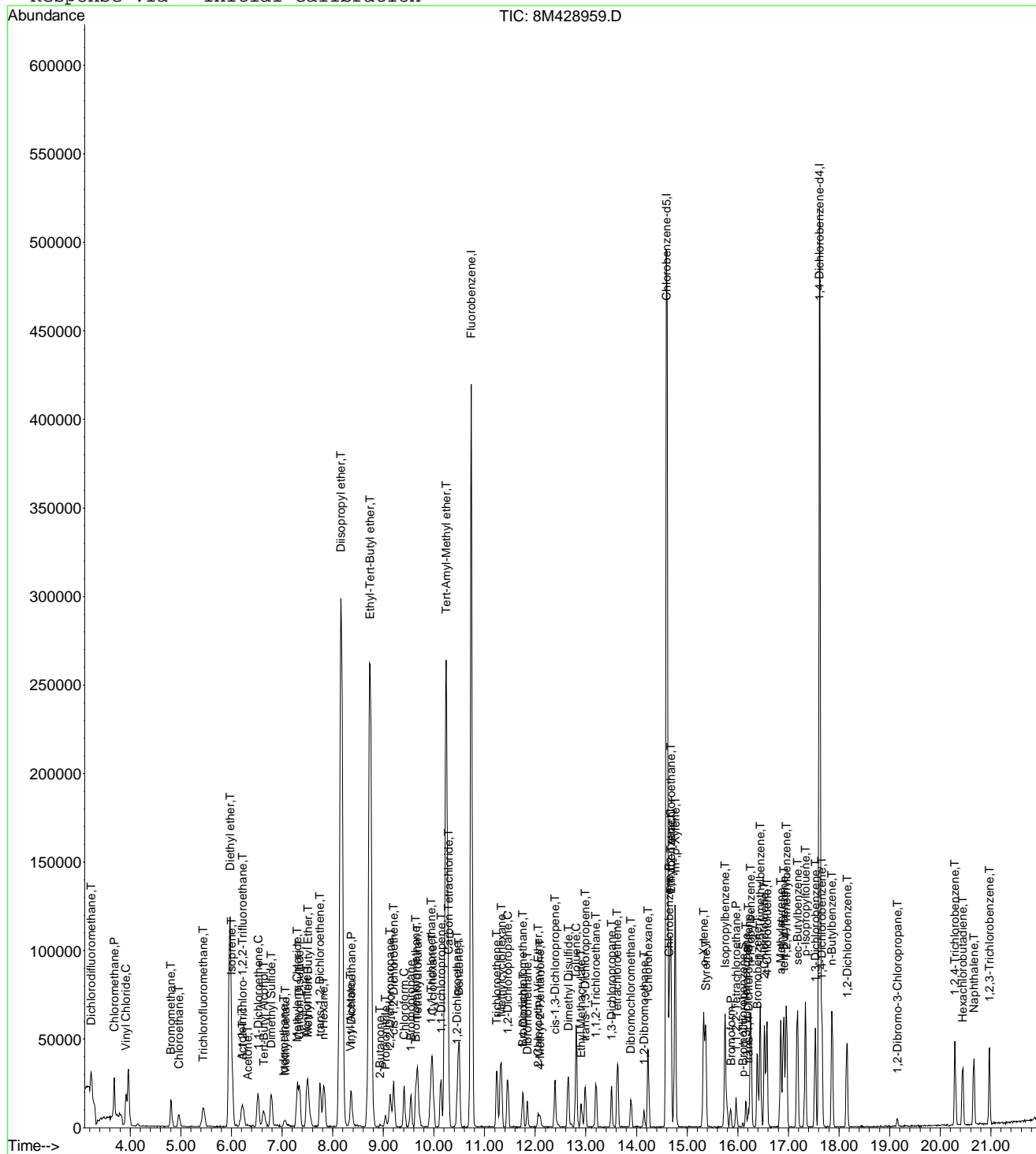
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	16949	2.0484	ug/L	96
46) Benzene	10.49	78	57876	2.0940	ug/L	98
47) Trichloroethene	11.24	130	14945	1.9847	ug/L	99
48) Methylcyclohexane	11.33	83	23479	2.0134	ug/L	99
49) 1,2-Dichloropropane	11.46	63	15773	2.1025	ug/L	95
50) Bromodichloromethane	11.75	83	17781	2.0470	ug/L	96
51) 1,4-Dioxane	11.76	88	1353	32.0520	ug/L	86
52) Dibromomethane	11.84	93	6841	1.9811	ug/L	97
53) 2-Chloroethyl Vinyl Ether	12.06	63	4465	1.3621	ug/L	85
54) 4-Methyl-2-Pentanone	12.09	58	2561	1.2417	ug/L #	64
55) cis-1,3-Dichloropropene	12.39	75	20089	1.9439	ug/L	100
56) Dimethyl Disulfide	12.65	79	10683	1.6834	ug/L	96
59) Toluene	12.80	91	59284	2.1254	ug/L	98
60) Ethyl Methacrylate	12.91	69	11491	2.0491	ug/L	99
62) trans-1,3-Dichloropropene	12.99	75	16007	1.9089	ug/L	96
63) 1,1,2-Trichloroethane	13.20	97	10122	2.0760	ug/L	98
65) 1,3-Dichloropropane	13.51	76	17551	2.0901	ug/L	100
66) Tetrachloroethene	13.62	164	12184	2.0728	ug/L	100
67) Dibromochloromethane	13.89	129	12041	1.9459	ug/L	100
68) 1,2-Dibromoethane	14.15	107	9579	2.0233	ug/L	99
69) 1-Chlorohexane	14.23	91	18690	1.9336	ug/L	96
70) Chlorobenzene	14.65	112	39126	2.0691	ug/L	91
71) 1,1,1,2-Tetrachloroethane	14.68	131	14350	2.0641	ug/L	98
72) Ethylbenzene	14.68	106	21470	2.0985	ug/L	97
73) m-,p-Xylene	14.77	106	51212	4.1082	ug/L	93
74) o-Xylene	15.33	106	24948	2.0625	ug/L	95
75) Styrene	15.37	104	39113	1.9532	ug/L	99
76) Bromoform	15.87	173	6783	1.7679	ug/L	92
77) Isopropylbenzene	15.75	105	64286	2.1042	ug/L	99
79) 1,1,2,2-Tetrachloroethane	15.97	83	11829	2.1880	ug/L	97
81) 1,2,3-Trichloropropane	16.17	110	3388	2.0949	ug/L	76
82) trans-1,4-Dichloro-2-Buten	16.21	53	3150	1.7236	ug/L #	1
83) n-Propylbenzene	16.26	91	76210	2.1975	ug/L	97
84) Bromobenzene	16.39	156	17267	2.1761	ug/L	99
85) 1,3,5-Trimethylbenzene	16.44	105	54779	2.1912	ug/L	95
86) 2-Chlorotoluene	16.53	91	45081	1.9660	ug/L	99
87) 4-Chlorotoluene	16.58	91	48478	2.4196	ug/L	89
88) a-Methylstyrene	16.85	118	27662	1.9283	ug/L	99
89) tert-Butylbenzene	16.91	134	11927	2.0680	ug/L	95
90) 1,2,4-Trimethylbenzene	16.96	105	54506	2.1536	ug/L	97
91) sec-Butylbenzene	17.18	105	69186	2.1624	ug/L	98
92) p-Isopropyltoluene	17.33	119	59081	2.1541	ug/L	99
93) 1,3-Dichlorobenzene	17.53	146	32687	2.1131	ug/L	100
94) 1,4-Dichlorobenzene	17.66	146	32560	2.0785	ug/L	89
95) n-Butylbenzene	17.86	91	51689	2.0684	ug/L	97
96) 1,2-Dichlorobenzene	18.16	146	29713	2.0699	ug/L	98
97) 1,2-Dibromo-3-Chloropropan	19.15	75	1749	1.7949	ug/L	85
98) 1,2,4-Trichlorobenzene	20.29	180	21069	2.0228	ug/L	97
99) Hexachlorobutadiene	20.45	225	10232	2.0848	ug/L	98
100) Naphthalene	20.66	128	39794	2.0958	ug/L	96
101) 1,2,3-Trichlorobenzene	20.97	180	20147	2.1056	ug/L	95

(#) = qualifier out of range (m) = manual integration
 8M428959.D 8260WTR.M Tue Mar 05 10:54:48 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428959.D Vial: 5
Acq On : 4 Mar 2019 14:12 Operator: EEA
Sample : WG698192-05 2 ug/L ICAL 8260 Inst : HPMS8
Misc : 1,1 STD92326 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428959.D Vial: 5
 Acq On : 4 Mar 2019 14:12 Operator: EEA
 Sample : WG698192-05 2 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	2.000	1.835	8.3	100	0.00
3 P	Chloromethane	2.000	2.246	-12.3	100	0.00
4 C	Vinyl Chloride	2.000	2.043	-2.2	100	0.00
5 T	1,3-Butadiene	-1.000	-1.351	0.0	0	0.01
6 T	Bromomethane	2.000	2.330	-16.5	100	0.00
7 T	Chloroethane	2.000	1.974	1.3	100	0.00
8 T	Trichlorofluoromethane	2.000	2.011	-0.6	100	0.00
9 T	Diethyl ether	20.000	20.863	-4.3	100	0.00
10 T	Isoprene	2.000	1.928	3.6	100	0.00
11 T	Acrolein	10.000	9.337	6.6	100	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	2.000	1.935	3.2	100	0.00
13 T	Acetone	-1.000	1.507	0.0	100	0.01
14 C	1,1-Dichloroethene	2.000	2.033	-1.6	100	0.00
15 T	Tert-Butyl Alcohol	40.000	44.188	-10.5	100	0.01
16 T	Dimethyl Sulfide	2.000	1.933	3.3	100	0.00
17 T	Iodomethane	2.000	1.843	7.9	100	0.00
18 T	Methyl acetate	-1.000	1.406	0.0	100	0.01
19 T	Methylene Chloride	2.000	2.070	-3.5	100	0.00
20 T	Carbon Disulfide	2.000	2.058	-2.9	100	0.00
21 T	Acrylonitrile	10.000	8.962	10.4	100	0.00
22 T	Methyl Tert Butyl Ether	2.000	2.059	-3.0	100	0.00
23 T	trans-1,2-Dichloroethene	2.000	2.021	-1.0	100	0.00
24 T	n-Hexane	2.000	1.999	0.0	100	0.00
25 T	Diisopropyl ether	20.000	21.345	-6.7	100	0.00
26 T	Vinyl Acetate	-1.000	1.700	0.0	100	0.01
27 P	1,1-Dichloroethane	2.000	2.067	-3.4	100	0.00
28 T	Ethyl-Tert-Butyl ether	20.000	21.068	-5.3	100	0.00
29 T	2-Butanone	-1.000	1.406	0.0	100	0.01
30 T	Propionitrile	20.000	19.327	3.4	100	0.01
31 T	2,2-Dichloropropane	2.000	2.084	-4.2	100	0.00
32 T	cis-1,2-Dichloroethene	2.000	2.029	-1.4	100	0.00
33 C	Chloroform	2.000	2.002	-0.1	100	0.00
34	1-Bromopropane	2.000	1.768	11.6	100	0.00
35 T	Bromochloromethane	2.000	1.972	1.4	100	0.00
36 T	Tetrahydrofuran	20.000	21.230	-6.2	100	0.00
37 S	Dibromofluoromethane	-1.000	0.000	0.0	0	-9.70#
38 T	1,1,1-Trichloroethane	2.000	2.009	-0.4	100	0.00
39 T	Cyclohexane	2.000	1.992	0.4	100	0.00
40 T	1,1-Dichloropropene	2.000	2.063	-3.2	100	0.00
41 T	Tert-Amyl-Methyl ether	20.000	21.089	-5.4	100	0.00
42 T	Carbon Tetrachloride	2.000	1.971	1.4	100	0.00
43 S	1,2-Dichloroethane-d4	-1.000	0.000	0.0	0	-10.33#
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	2.000	2.048	-2.4	100	0.00
46 T	Benzene	2.000	2.094	-4.7	100	0.00
47 T	Trichloroethene	2.000	1.985	0.7	100	0.00
48 T	Methylcyclohexane	2.000	2.013	-0.6	100	0.00
49 C	1,2-Dichloropropane	2.000	2.102	-5.1	100	0.00
50 T	Bromodichloromethane	2.000	2.047	-2.4	100	0.00
51 T	1,4-Dioxane	40.000	32.052	19.9	100	0.01
52 T	Dibromomethane	2.000	1.981	0.9	100	0.00
53 T	2-Chloroethyl Vinyl Ether	-1.000	1.362	0.0	100	0.01
54 T	4-Methyl-2-Pentanone	-1.000	1.242	0.0	100	0.01
55 T	cis-1,3-Dichloropropene	2.000	1.944	2.8	100	0.00
56 T	Dimethyl Disulfide	-1.000	1.683	0.0	100	0.00
57 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00

58 S	Toluene-d8	-1.000	0.116	0.0	100	0.00
59 C	Toluene	2.000	2.125	-6.3	100	0.00
60 T	Ethyl Methacrylate	2.000	2.049	-2.4	100	0.00
61	Paraldehyde	-1.000	0.000	0.0	0	-13.45#
62 T	trans-1,3-Dichloropropene	2.000	1.909	4.5	100	0.00
63 T	1,1,2-Trichloroethane	2.000	2.076	-3.8	100	0.00
64 T	2-Hexanone	-1.000	0.000	0.0	0	-13.14#
65 T	1,3-Dichloropropane	2.000	2.090	-4.5	100	0.00
66 T	Tetrachloroethene	2.000	2.073	-3.6	100	0.00
67 T	Dibromochloromethane	2.000	1.946	2.7	100	0.00
68 T	1,2-Dibromoethane	2.000	2.023	-1.2	100	0.00
69 T	1-Chlorohexane	2.000	1.934	3.3	100	0.00
70 P	Chlorobenzene	2.000	2.069	-3.4	100	0.00
71 T	1,1,1,2-Tetrachloroethane	2.000	2.064	-3.2	100	0.00
72 C	Ethylbenzene	2.000	2.099	-5.0	100	0.00
73 T	m-,p-Xylene	4.000	4.108	-2.7	100	0.00
74 T	o-Xylene	2.000	2.063	-3.2	100	0.00
75 T	Styrene	2.000	1.953	2.3	100	0.00
76 P	Bromoform	2.000	1.768	11.6	100	0.00
77 T	Isopropylbenzene	2.000	2.104	-5.2	100	0.00
78 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
79 P	1,1,2,2-Tetrachloroethane	2.000	2.188	-9.4	100	0.00
80 S	p-Bromofluorobenzene	-1.000	0.148	0.0	100	0.00
81 T	1,2,3-Trichloropropane	2.000	2.095	-4.8	100	0.00
82 T	trans-1,4-Dichloro-2-Butene	2.000	1.724	13.8	100	0.00
83 T	n-Propylbenzene	2.000	2.198	-9.9	100	0.00
84 T	Bromobenzene	2.000	2.176	-8.8	100	0.00
85 T	1,3,5-Trimethylbenzene	2.000	2.191	-9.5	100	0.00
86 T	2-Chlorotoluene	2.000	1.966	1.7	100	0.00
87 T	4-Chlorotoluene	2.000	2.420	-21.0	100	0.00
88 T	a-Methylstyrene	2.000	1.928	3.6	100	0.00
89 T	tert-Butylbenzene	2.000	2.068	-3.4	100	0.00
90 T	1,2,4-Trimethylbenzene	2.000	2.154	-7.7	100	0.00
91 T	sec-Butylbenzene	2.000	2.162	-8.1	100	0.00
92 T	p-Isopropyltoluene	2.000	2.154	-7.7	100	0.00
93 T	1,3-Dichlorobenzene	2.000	2.113	-5.6	100	0.00
94 T	1,4-Dichlorobenzene	2.000	2.079	-4.0	100	0.00
95 T	n-Butylbenzene	2.000	2.068	-3.4	100	0.00
96 T	1,2-Dichlorobenzene	2.000	2.070	-3.5	100	0.00
97 T	1,2-Dibromo-3-Chloropropane	2.000	1.795	10.3	100	0.00
98 T	1,2,4-Trichlorobenzene	2.000	2.023	-1.2	100	0.00
99 T	Hexachlorobutadiene	2.000	2.085	-4.2	100	0.00
100 T	Naphthalene	2.000	2.096	-4.8	100	0.00
101 T	1,2,3-Trichlorobenzene	2.000	2.106	-5.3	100	0.00

(#) = Out of Range
8M428959.D 8260WTR.M

SPCC's out = 0 CCC's out = 0
Tue Mar 05 10:51:36 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428960.D Vial: 6
 Acq On : 4 Mar 2019 14:42 Operator: EEA
 Sample : WG698192-06 5 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:49 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	527749	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	378036	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	196112	25.00	ug/L	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.70	111	25738	4.8926	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	19.56%#	
43) 1,2-Dichloroethane-d4	10.33	65	25061	4.8256	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	19.32%#	
58) Toluene-d8	12.71	98	102229	5.3088	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	21.24%#	
80) p-Bromofluorobenzene	16.10	95	36245	5.1209	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	20.48%#	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.23	85	41765	4.5498	ug/L	99
3) Chloromethane	3.69	50	67822	5.1375	ug/L	95
4) Vinyl Chloride	3.92	62	53473	4.9703	ug/L	98
5) 1,3-Butadiene	3.97	54	46618	3.3417	ug/L	95
6) Bromomethane	4.81	94	36051	5.5712	ug/L	99
7) Chloroethane	4.97	64	29010	5.0389	ug/L	98
8) Trichlorofluoromethane	5.44	101	54659	5.0039	ug/L	98
9) Diethyl ether	5.97	59	296644	50.9200	ug/L	99
10) Isoprene	6.00	67	53597	4.9095	ug/L	100
11) Acrolein	6.22	56	15443	23.1177	ug/L	100
12) 1,1,2-Trichloro-1,2,2-Trif	6.23	101	31666	5.0320	ug/L	99
13) Acetone	6.31	43	7326	4.5063	ug/L	90
14) 1,1-Dichloroethene	6.53	61	57536	5.1121	ug/L	98
15) Tert-Butyl Alcohol	6.63	59	49707	94.9870	ug/L	99
16) Dimethyl Sulfide	6.79	62	38663	4.7689	ug/L	99
17) Iodomethane	7.04	142	11611	2.8749	ug/L	95
18) Methyl acetate	7.06	43	18048	4.2873	ug/L	91
19) Methylene Chloride	7.30	84	36357	5.1696	ug/L	96
20) Carbon Disulfide	7.35	76	106758	5.0726	ug/L	99
21) Acrylonitrile	7.49	53	46771	22.9315	ug/L	100
22) Methyl Tert Butyl Ether	7.52	73	83730	5.0018	ug/L	99
23) trans-1,2-Dichloroethene	7.75	61	54727	5.0888	ug/L	100
24) n-Hexane	7.83	57	52293	4.9547	ug/L	98
25) Diisopropyl ether	8.16	45	1377409	51.7377	ug/L	100
26) Vinyl Acetate	8.35	43	45990	4.6567	ug/L	94
27) 1,1-Dichloroethane	8.37	63	70391	5.0632	ug/L	99
28) Ethyl-Tert-Butyl ether	8.74	59	1177833	51.0449	ug/L	99
29) 2-Butanone	8.94	43	10157	4.2412	ug/L #	81
30) Propionitrile	9.03	54	32733	47.1098	ug/L	97
31) 2,2-Dichloropropane	9.13	77	51691	5.1242	ug/L	99
32) cis-1,2-Dichloroethene	9.21	96	39588	5.1576	ug/L	99
33) Chloroform	9.41	83	60893	4.9351	ug/L	98
34) 1-Bromopropane	9.55	122	7921	4.8008	ug/L	96
35) Bromochloromethane	9.64	130	22117	5.0523	ug/L	99
36) Tetrahydrofuran	9.67	42	77687	48.2147	ug/L	99
38) 1,1,1-Trichloroethane	9.94	97	55149	5.1458	ug/L	99
39) Cyclohexane	9.97	56	72976	5.1123	ug/L	97
40) 1,1-Dichloropropene	10.14	75	47040	5.2270	ug/L	97
41) Tert-Amyl-Methyl ether	10.24	73	900192	51.1939	ug/L	100
42) Carbon Tetrachloride	10.28	117	49006	5.0321	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M428960.D 8260WTR.M Tue Mar 05 10:54:51 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428960.D Vial: 6
 Acq On : 4 Mar 2019 14:42 Operator: EEA
 Sample : WG698192-06 5 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:49 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

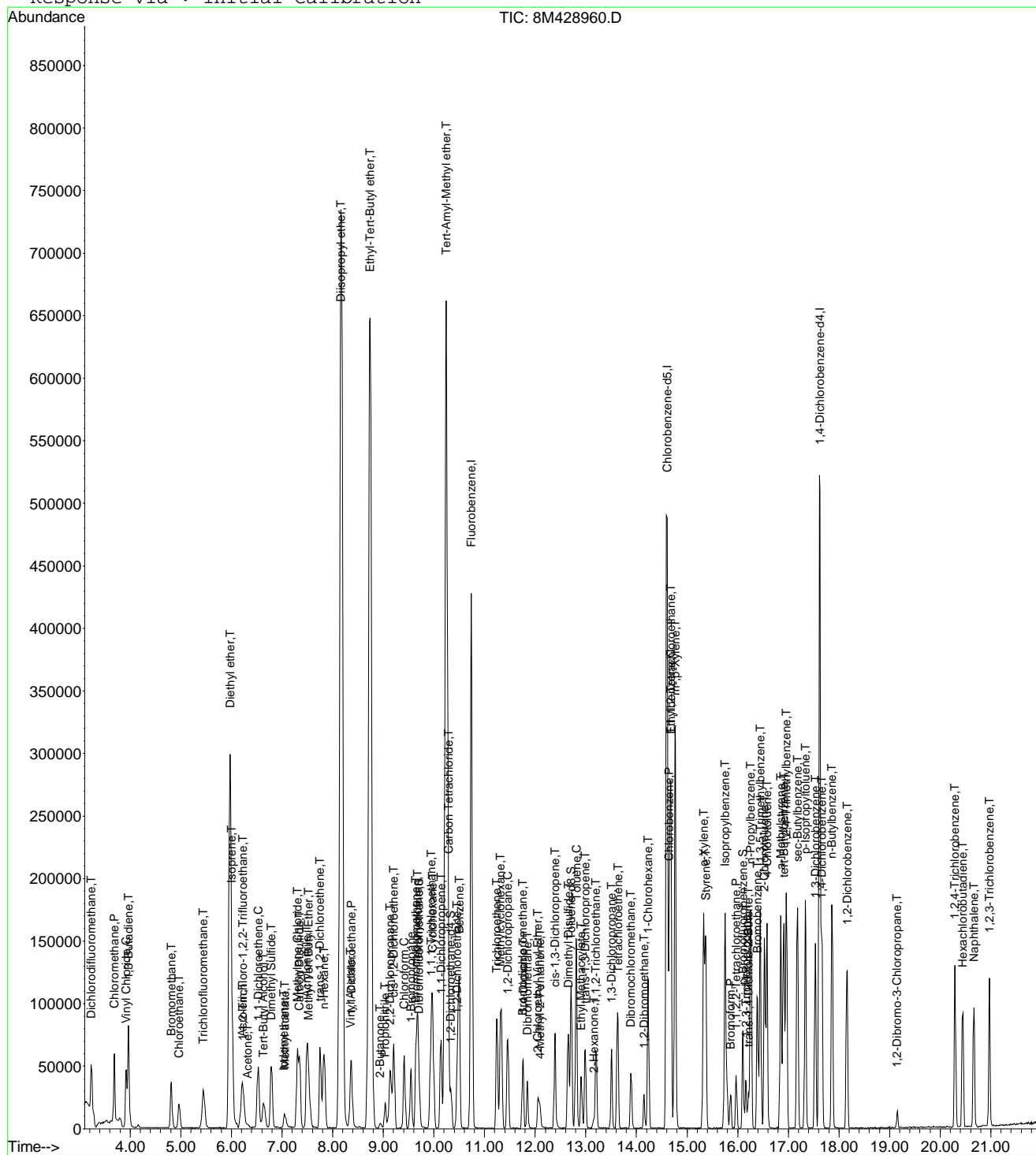
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	43201	5.1188	ug/L	99
46) Benzene	10.50	78	147053	5.2162	ug/L	99
47) Trichloroethene	11.24	130	39812	5.1834	ug/L	98
48) Methylcyclohexane	11.32	83	61933	5.2067	ug/L	98
49) 1,2-Dichloropropane	11.46	63	38788	5.0688	ug/L	98
50) Bromodichloromethane	11.75	83	45224	5.1041	ug/L	98
51) 1,4-Dioxane	11.76	88	3685	85.5833	ug/L	98
52) Dibromomethane	11.84	93	17845	5.0664	ug/L	98
53) 2-Chloroethyl Vinyl Ether	12.05	63	13492	4.0351	ug/L	94
54) 4-Methyl-2-Pentanone	12.09	58	8289	3.9402	ug/L	95
55) cis-1,3-Dichloropropene	12.39	75	52578	4.9879	ug/L	99
56) Dimethyl Disulfide	12.65	79	28739	4.4397	ug/L	94
59) Toluene	12.82	91	152840	5.4149	ug/L	98
60) Ethyl Methacrylate	12.91	69	31243	4.7002	ug/L	96
62) trans-1,3-Dichloropropene	12.99	75	43645	5.1434	ug/L	99
63) 1,1,2-Trichloroethane	13.20	97	25913	5.2522	ug/L	99
64) 2-Hexanone	13.16	58	5479	5.2779	ug/L #	58
65) 1,3-Dichloropropane	13.51	76	44680	5.2582	ug/L	97
66) Tetrachloroethene	13.63	164	31614	5.3149	ug/L	96
67) Dibromochloromethane	13.89	129	31368	5.0095	ug/L	100
68) 1,2-Dibromoethane	14.15	107	25390	5.2996	ug/L	99
69) 1-Chlorohexane	14.23	91	51086	5.2230	ug/L	100
70) Chlorobenzene	14.65	112	99963	5.2242	ug/L	98
71) 1,1,1,2-Tetrachloroethane	14.68	131	35067	4.9847	ug/L	96
72) Ethylbenzene	14.68	106	54042	5.2200	ug/L	95
73) m-,p-Xylene	14.77	106	133939	10.6179	ug/L	96
74) o-Xylene	15.33	106	63662	5.2011	ug/L	92
75) Styrene	15.37	104	102902	5.0780	ug/L	99
76) Bromoform	15.86	173	18437	4.7486	ug/L	97
77) Isopropylbenzene	15.75	105	166743	5.3934	ug/L	99
79) 1,1,2,2-Tetrachloroethane	15.97	83	29022	5.2456	ug/L	100
81) 1,2,3-Trichloropropane	16.17	110	8683	5.2462	ug/L	70
82) trans-1,4-Dichloro-2-Butene	16.21	53	9147	4.8906	ug/L #	32
83) n-Propylbenzene	16.26	91	195070	5.4964	ug/L	98
84) Bromobenzene	16.39	156	44242	5.4482	ug/L	99
85) 1,3,5-Trimethylbenzene	16.44	105	140302	5.4839	ug/L	98
86) 2-Chlorotoluene	16.53	91	135637	5.7801	ug/L	97
87) 4-Chlorotoluene	16.58	91	108219	5.2779	ug/L	98
88) a-Methylstyrene	16.85	118	75564	5.1471	ug/L	98
89) tert-Butylbenzene	16.91	134	31731	5.3760	ug/L	98
90) 1,2,4-Trimethylbenzene	16.96	105	142816	5.5138	ug/L	99
91) sec-Butylbenzene	17.18	105	181023	5.5287	ug/L	99
92) p-Isopropyltoluene	17.33	119	153347	5.4634	ug/L	99
93) 1,3-Dichlorobenzene	17.53	146	85853	5.4233	ug/L	96
94) 1,4-Dichlorobenzene	17.66	146	84516	5.2719	ug/L	97
95) n-Butylbenzene	17.86	91	137967	5.3948	ug/L	98
96) 1,2-Dichlorobenzene	18.16	146	77887	5.3019	ug/L	99
97) 1,2-Dibromo-3-Chloropropane	19.14	75	4777	4.7904	ug/L	99
98) 1,2,4-Trichlorobenzene	20.29	180	53900	5.0566	ug/L	100
99) Hexachlorobutadiene	20.45	225	25972	5.1709	ug/L	98
100) Naphthalene	20.66	128	102362	5.2679	ug/L	100
101) 1,2,3-Trichlorobenzene	20.97	180	50029	5.1092	ug/L	98

(#) = qualifier out of range (m) = manual integration
 8M428960.D 8260WTR.M Tue Mar 05 10:54:52 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428960.D Vial: 6
 Acq On : 4 Mar 2019 14:42 Operator: EEA
 Sample : WG698192-06 5 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428960.D Vial: 6
 Acq On : 4 Mar 2019 14:42 Operator: EEA
 Sample : WG698192-06 5 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	100	0.00
2 T	Dichlorodifluoromethane	5.000	4.550	9.0	100	0.00
3 P	Chloromethane	5.000	5.137	-2.7	100	0.00
4 C	Vinyl Chloride	5.000	4.970	0.6	100	0.01
5 T	1,3-Butadiene	5.000	3.342	33.2#	100	0.01
6 T	Bromomethane	5.000	5.571	-11.4	100	0.00
7 T	Chloroethane	5.000	5.039	-0.8	100	0.00
8 T	Trichlorofluoromethane	5.000	5.004	-0.1	100	0.00
9 T	Diethyl ether	50.000	50.920	-1.8	100	0.00
10 T	Isoprene	5.000	4.910	1.8	100	0.00
11 T	Acrolein	25.000	23.118	7.5	100	0.01
12 T	1,1,2-Trichloro-1,2,2-Trifl	5.000	5.032	-0.6	100	0.01
13 T	Acetone	5.000	4.506	9.9	100	0.00
14 C	1,1-Dichloroethene	5.000	5.112	-2.2	100	0.00
15 T	Tert-Butyl Alcohol	100.000	94.987	5.0	100	0.00
16 T	Dimethyl Sulfide	5.000	4.769	4.6	100	0.00
17 T	Iodomethane	5.000	2.875	42.5#	100	0.00
18 T	Methyl acetate	5.000	4.287	14.3	100	0.00
19 T	Methylene Chloride	5.000	5.170	-3.4	100	0.00
20 T	Carbon Disulfide	5.000	5.073	-1.5	100	0.00
21 T	Acrylonitrile	25.000	22.931	8.3	100	0.00
22 T	Methyl Tert Butyl Ether	5.000	5.002	-0.0	100	0.01
23 T	trans-1,2-Dichloroethene	5.000	5.089	-1.8	100	0.00
24 T	n-Hexane	5.000	4.955	0.9	100	0.00
25 T	Diisopropyl ether	50.000	51.738	-3.5	100	0.00
26 T	Vinyl Acetate	5.000	4.657	6.9	100	0.01
27 P	1,1-Dichloroethane	5.000	5.063	-1.3	100	0.00
28 T	Ethyl-Tert-Butyl ether	50.000	51.045	-2.1	100	0.00
29 T	2-Butanone	5.000	4.241	15.2	100	0.01
30 T	Propionitrile	50.000	47.110	5.8	100	0.00
31 T	2,2-Dichloropropane	5.000	5.124	-2.5	100	0.00
32 T	cis-1,2-Dichloroethene	5.000	5.158	-3.2	100	0.00
33 C	Chloroform	5.000	4.935	1.3	100	0.00
34	1-Bromopropane	5.000	4.801	4.0	100	0.00
35 T	Bromochloromethane	5.000	5.052	-1.0	100	0.00
36 T	Tetrahydrofuran	50.000	48.215	3.6	100	0.00
37 S	Dibromofluoromethane	5.000	4.893	2.1	100	0.00
38 T	1,1,1-Trichloroethane	5.000	5.146	-2.9	100	0.00
39 T	Cyclohexane	5.000	5.112	-2.2	100	0.00
40 T	1,1-Dichloropropene	5.000	5.227	-4.5	100	0.00
41 T	Tert-Amyl-Methyl ether	50.000	51.194	-2.4	100	0.00
42 T	Carbon Tetrachloride	5.000	5.032	-0.6	100	0.00
43 S	1,2-Dichloroethane-d4	5.000	4.826	3.5	100	0.00
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	5.000	5.119	-2.4	100	0.00
46 T	Benzene	5.000	5.216	-4.3	100	0.00
47 T	Trichloroethene	5.000	5.183	-3.7	100	0.00
48 T	Methylcyclohexane	5.000	5.207	-4.1	100	0.00
49 C	1,2-Dichloropropane	5.000	5.069	-1.4	100	0.00
50 T	Bromodichloromethane	5.000	5.104	-2.1	100	0.00
51 T	1,4-Dioxane	100.000	85.583	14.4	100	0.01
52 T	Dibromomethane	5.000	5.066	-1.3	100	0.00
53 T	2-Chloroethyl Vinyl Ether	5.000	4.035	19.3	100	0.00
54 T	4-Methyl-2-Pentanone	5.000	3.940	21.2	100	0.01
55 T	cis-1,3-Dichloropropene	5.000	4.988	0.2	100	0.00
56 T	Dimethyl Disulfide	5.000	4.440	11.2	100	0.00
57 I	Chlorobenzene-d5	25.000	25.000	0.0	100	0.00

58 S	Toluene-d8	5.000	5.309	-6.2	100	0.00
59 C	Toluene	5.000	5.415	-8.3	100	0.00
60 T	Ethyl Methacrylate	5.000	4.700	6.0	100	0.00
61	Paraldehyde	-1.000	0.000	0.0	0	-13.45#
62 T	trans-1,3-Dichloropropene	5.000	5.143	-2.9	100	0.00
63 T	1,1,2-Trichloroethane	5.000	5.252	-5.0	100	0.00
64 T	2-Hexanone	5.000	5.278	-5.6	100	0.01
65 T	1,3-Dichloropropane	5.000	5.258	-5.2	100	0.00
66 T	Tetrachloroethene	5.000	5.315	-6.3	100	0.00
67 T	Dibromochloromethane	5.000	5.009	-0.2	100	0.00
68 T	1,2-Dibromoethane	5.000	5.300	-6.0	100	0.00
69 T	1-Chlorohexane	5.000	5.223	-4.5	100	0.00
70 P	Chlorobenzene	5.000	5.224	-4.5	100	0.00
71 T	1,1,1,2-Tetrachloroethane	5.000	4.985	0.3	100	0.00
72 C	Ethylbenzene	5.000	5.220	-4.4	100	0.00
73 T	m-,p-Xylene	10.000	10.618	-6.2	100	0.00
74 T	o-Xylene	5.000	5.201	-4.0	100	0.00
75 T	Styrene	5.000	5.078	-1.6	100	0.00
76 P	Bromoform	5.000	4.749	5.0	100	0.00
77 T	Isopropylbenzene	5.000	5.393	-7.9	100	0.00
78 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	100	0.00
79 P	1,1,2,2-Tetrachloroethane	5.000	5.246	-4.9	100	0.00
80 S	p-Bromofluorobenzene	5.000	5.121	-2.4	100	0.00
81 T	1,2,3-Trichloropropane	5.000	5.246	-4.9	100	0.00
82 T	trans-1,4-Dichloro-2-Butene	5.000	4.891	2.2	100	0.00
83 T	n-Propylbenzene	5.000	5.496	-9.9	100	0.00
84 T	Bromobenzene	5.000	5.448	-9.0	100	0.00
85 T	1,3,5-Trimethylbenzene	5.000	5.484	-9.7	100	0.00
86 T	2-Chlorotoluene	5.000	5.780	-15.6	100	0.00
87 T	4-Chlorotoluene	5.000	5.278	-5.6	100	0.00
88 T	a-Methylstyrene	5.000	5.147	-2.9	100	0.00
89 T	tert-Butylbenzene	5.000	5.376	-7.5	100	0.00
90 T	1,2,4-Trimethylbenzene	5.000	5.514	-10.3	100	0.00
91 T	sec-Butylbenzene	5.000	5.529	-10.6	100	0.00
92 T	p-Isopropyltoluene	5.000	5.463	-9.3	100	0.00
93 T	1,3-Dichlorobenzene	5.000	5.423	-8.5	100	0.00
94 T	1,4-Dichlorobenzene	5.000	5.272	-5.4	100	0.00
95 T	n-Butylbenzene	5.000	5.395	-7.9	100	0.00
96 T	1,2-Dichlorobenzene	5.000	5.302	-6.0	100	0.00
97 T	1,2-Dibromo-3-Chloropropane	5.000	4.790	4.2	100	0.00
98 T	1,2,4-Trichlorobenzene	5.000	5.057	-1.1	100	0.00
99 T	Hexachlorobutadiene	5.000	5.171	-3.4	100	0.00
100 T	Naphthalene	5.000	5.268	-5.4	100	0.00
101 T	1,2,3-Trichlorobenzene	5.000	5.109	-2.2	100	0.00

(#) = Out of Range
8M428960.D 8260WTR.M

SPCC's out = 0 CCC's out = 0
Tue Mar 05 10:52:10 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428961.D Vial: 7
 Acq On : 4 Mar 2019 15:11 Operator: EEA
 Sample : WG698192-07 20 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:53 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.73	96	519761	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	383859	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	205732	25.00	ug/L	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.70	111	49997	9.6501	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	38.60%#	
43) 1,2-Dichloroethane-d4	10.34	65	50552	9.8836	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	39.52%#	
58) Toluene-d8	12.71	98	192225	9.8309	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	39.32%#	
80) p-Bromofluorobenzene	16.10	95	72023	9.7000	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	38.80%#	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.24	85	208518	23.0646	ug/L	99
3) Chloromethane	3.69	50	275650	21.2012	ug/L	100
4) Vinyl Chloride	3.92	62	232573	21.9498	ug/L	98
5) 1,3-Butadiene	3.96	54	166183	25.1280	ug/L	99
6) Bromomethane	4.81	94	134575	21.1162	ug/L	100
7) Chloroethane	4.97	64	121613	21.4484	ug/L	99
8) Trichlorofluoromethane	5.45	101	237529	22.0793	ug/L	100
9) Diethyl ether	5.97	59	487157	84.9074	ug/L	99
10) Isoprene	6.01	67	222399	20.6850	ug/L	100
11) Acrolein	6.21	56	26862	40.8295	ug/L	94
12) 1,1,2-Trichloro-1,2,2-Trif	6.22	101	134732	21.7392	ug/L	99
13) Acetone	6.32	43	34640	21.6351	ug/L	97
14) 1,1-Dichloroethene	6.53	61	237296	21.4077	ug/L	99
15) Tert-Butyl Alcohol	6.64	59	85003	164.9318	ug/L	99
16) Dimethyl Sulfide	6.78	62	166163	20.8103	ug/L	99
17) Iodomethane	7.04	142	117887	15.5640	ug/L	99
18) Methyl acetate	7.06	43	83477	20.1347	ug/L	99
19) Methylene Chloride	7.30	84	147627	21.3136	ug/L	100
20) Carbon Disulfide	7.34	76	444390	21.4396	ug/L	100
21) Acrylonitrile	7.49	53	85385	42.5070	ug/L	99
22) Methyl Tert Butyl Ether	7.52	73	356631	21.6317	ug/L	99
23) trans-1,2-Dichloroethene	7.75	61	227043	21.4363	ug/L	100
24) n-Hexane	7.83	57	218781	21.0477	ug/L	100
25) Diisopropyl ether	8.17	45	2247355	85.7116	ug/L	100
26) Vinyl Acetate	8.35	43	206107	21.1902	ug/L	100
27) 1,1-Dichloroethane	8.37	63	294302	21.4944	ug/L	99
28) Ethyl-Tert-Butyl ether	8.74	59	1923161	84.6268	ug/L	100
29) 2-Butanone	8.93	43	48902	20.7337	ug/L	97
30) Propionitrile	9.03	54	59321	82.9010	ug/L	99
31) 2,2-Dichloropropane	9.14	77	212343	21.3733	ug/L	99
32) cis-1,2-Dichloroethene	9.20	96	165898	21.9457	ug/L	100
33) Chloroform	9.41	83	256169	21.0803	ug/L	100
34) 1-Bromopropane	9.55	122	34230	21.0652	ug/L	100
35) Bromochloromethane	9.65	130	94314	21.8759	ug/L	99
36) Tetrahydrofuran	9.67	42	136457	85.9906	ug/L	99
38) 1,1,1-Trichloroethane	9.94	97	226839	21.4911	ug/L	99
39) Cyclohexane	9.97	56	295270	21.0029	ug/L	99
40) 1,1-Dichloropropene	10.13	75	193184	21.7961	ug/L	99
41) Tert-Amyl-Methyl ether	10.24	73	1472088	85.0043	ug/L	100
42) Carbon Tetrachloride	10.28	117	206808	21.5619	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M428961.D 8260WTR.M Tue Mar 05 10:54:55 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428961.D Vial: 7
 Acq On : 4 Mar 2019 15:11 Operator: EEA
 Sample : WG698192-07 20 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:53 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

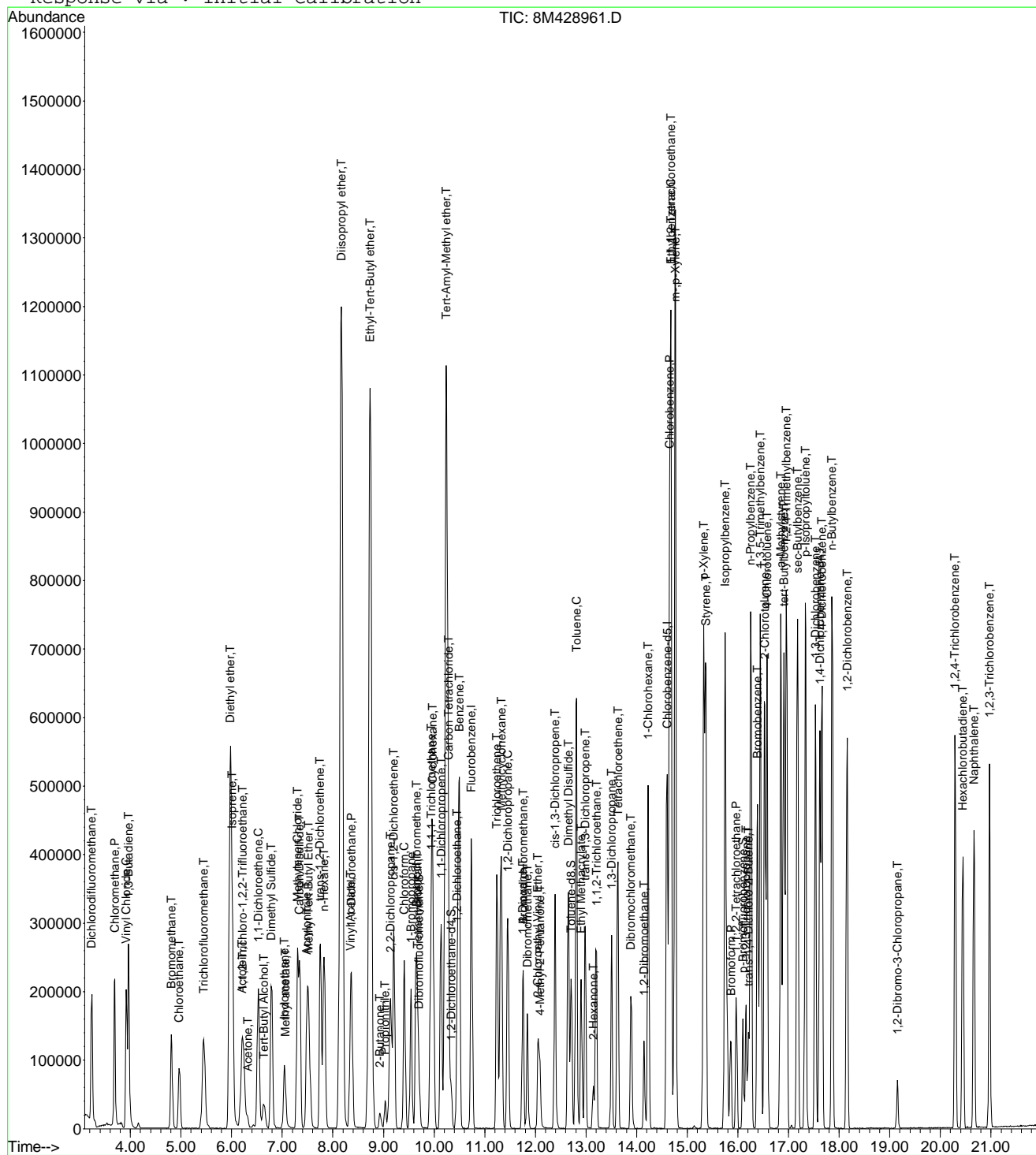
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	183714	22.1023	ug/L	100
46) Benzene	10.50	78	607917	21.8951	ug/L	99
47) Trichloroethene	11.24	130	163168	21.5703	ug/L	100
48) Methylcyclohexane	11.32	83	248513	21.2134	ug/L	100
49) 1,2-Dichloropropane	11.46	63	163424	21.6846	ug/L	100
50) Bromodichloromethane	11.76	83	192265	22.0330	ug/L	99
51) 1,4-Dioxane	11.75	88	7695	181.4613	ug/L	100
52) Dibromomethane	11.84	93	75950	21.8943	ug/L	99
53) 2-Chloroethyl Vinyl Ether	12.05	63	63845	19.3876	ug/L	99
54) 4-Methyl-2-Pentanone	12.09	58	41512	20.0362	ug/L	99
55) cis-1,3-Dichloropropene	12.39	75	231786	22.3267	ug/L	100
56) Dimethyl Disulfide	12.65	79	130210	20.4245	ug/L	99
59) Toluene	12.81	91	632121	22.0555	ug/L	99
60) Ethyl Methacrylate	12.91	69	151066	20.5854	ug/L	99
62) trans-1,3-Dichloropropene	12.99	75	192384	22.3278	ug/L	100
63) 1,1,2-Trichloroethane	13.21	97	111470	22.2505	ug/L	98
64) 2-Hexanone	13.14	58	37536	20.0876	ug/L	94
65) 1,3-Dichloropropane	13.51	76	196408	22.7638	ug/L	99
66) Tetrachloroethene	13.63	164	128648	21.3000	ug/L	99
67) Dibromochloromethane	13.89	129	138089	21.7184	ug/L	99
68) 1,2-Dibromoethane	14.15	107	109891	22.5896	ug/L	100
69) 1-Chlorohexane	14.23	91	208827	21.0265	ug/L	99
70) Chlorobenzene	14.65	112	415327	21.3761	ug/L	100
71) 1,1,1,2-Tetrachloroethane	14.67	131	154027	21.5625	ug/L	99
72) Ethylbenzene	14.67	106	228610	21.7469	ug/L	96
73) m-,p-Xylene	14.77	106	558736	43.6213	ug/L	97
74) o-Xylene	15.33	106	274667	22.0997	ug/L	97
75) Styrene	15.37	104	454662	22.0964	ug/L	100
76) Bromoform	15.86	173	86855	22.0311	ug/L	99
77) Isopropylbenzene	15.75	105	695094	22.1422	ug/L	99
79) 1,1,2,2-Tetrachloroethane	15.97	83	129068	22.2374	ug/L	99
81) 1,2,3-Trichloropropane	16.16	110	38369	22.0982	ug/L	88
82) trans-1,4-Dichloro-2-Butene	16.20	53	41016	20.9044	ug/L	# 16
83) n-Propylbenzene	16.26	91	827992	22.2389	ug/L	99
84) Bromobenzene	16.39	156	185655	21.7935	ug/L	99
85) 1,3,5-Trimethylbenzene	16.44	105	584546	21.7795	ug/L	100
86) 2-Chlorotoluene	16.54	91	553801	22.4963	ug/L	98
87) 4-Chlorotoluene	16.58	91	462934	21.5220	ug/L	99
88) a-Methylstyrene	16.85	118	319438	20.7415	ug/L	98
89) tert-Butylbenzene	16.91	134	131955	21.3110	ug/L	100
90) 1,2,4-Trimethylbenzene	16.96	105	600342	22.0941	ug/L	98
91) sec-Butylbenzene	17.18	105	761899	22.1812	ug/L	99
92) p-Isopropyltoluene	17.33	119	655213	22.2521	ug/L	99
93) 1,3-Dichlorobenzene	17.53	146	355729	21.4204	ug/L	99
94) 1,4-Dichlorobenzene	17.66	146	357635	21.2654	ug/L	99
95) n-Butylbenzene	17.86	91	592667	22.0909	ug/L	99
96) 1,2-Dichlorobenzene	18.16	146	330508	21.4463	ug/L	99
97) 1,2-Dibromo-3-Chloropropane	19.15	75	22970	21.9575	ug/L	97
98) 1,2,4-Trichlorobenzene	20.29	180	241342	21.5827	ug/L	99
99) Hexachlorobutadiene	20.44	225	113124	21.4695	ug/L	99
100) Naphthalene	20.66	128	461267	22.6284	ug/L	100
101) 1,2,3-Trichlorobenzene	20.97	180	221691	21.5813	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M428961.D 8260WTR.M Tue Mar 05 10:54:56 2019

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428961.D Vial: 7
 Acq On : 4 Mar 2019 15:11 Operator: EEA
 Sample : WG698192-07 20 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428962.D Vial: 8
 Acq On : 4 Mar 2019 15:40 Operator: EEA
 Sample : WG698192-08 50 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:57 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	533134	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	400675	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	217564	25.00	ug/L	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.70	111	139380	26.2275	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	104.92%	
43) 1,2-Dichloroethane-d4	10.33	65	137787	26.2635	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	105.04%	
58) Toluene-d8	12.71	98	528437	25.8916	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	103.56%	
80) p-Bromofluorobenzene	16.10	95	204314	26.0202	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	104.08%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.23	85	519958	56.0710	ug/L	100
3) Chloromethane	3.68	50	651136	48.8249	ug/L	100
4) Vinyl Chloride	3.91	62	579242	53.2965	ug/L	100
5) 1,3-Butadiene	3.95	54	346328	56.1865	ug/L	100
6) Bromomethane	4.81	94	300176	45.9193	ug/L	100
7) Chloroethane	4.97	64	309416	53.2015	ug/L	100
8) Trichlorofluoromethane	5.44	101	596452	54.0520	ug/L	100
9) Diethyl ether	5.97	59	603609	102.5651	ug/L	100
10) Isoprene	6.00	67	579917	52.5843	ug/L	100
11) Acrolein	6.21	56	34765	51.5164	ug/L	100
12) 1,1,2-Trichloro-1,2,2-Trif	6.22	101	340157	53.5082	ug/L	100
13) Acetone	6.31	43	87864	53.5007	ug/L	100
14) 1,1-Dichloroethene	6.53	61	596109	52.4292	ug/L	100
15) Tert-Butyl Alcohol	6.63	59	99887	188.9498	ug/L	100
16) Dimethyl Sulfide	6.79	62	430595	52.5752	ug/L	100
17) Iodomethane	7.03	142	441742	52.8466	ug/L	100
18) Methyl acetate	7.05	43	218390	51.3546	ug/L	100
19) Methylene Chloride	7.30	84	367767	51.7644	ug/L	100
20) Carbon Disulfide	7.34	76	1134616	53.3665	ug/L	100
21) Acrylonitrile	7.49	53	108702	52.7575	ug/L	100
22) Methyl Tert Butyl Ether	7.51	73	873403	51.6479	ug/L	100
23) trans-1,2-Dichloroethene	7.75	61	573205	52.7617	ug/L	100
24) n-Hexane	7.83	57	560051	52.5278	ug/L	100
25) Diisopropyl ether	8.17	45	2794825	103.9178	ug/L	100
26) Vinyl Acetate	8.34	43	511316	51.2506	ug/L	100
27) 1,1-Dichloroethane	8.37	63	736955	52.4736	ug/L	100
28) Ethyl-Tert-Butyl ether	8.74	59	2384559	102.2981	ug/L	100
29) 2-Butanone	8.93	43	127185	52.5718	ug/L	100
30) Propionitrile	9.03	54	74479	100.4636	ug/L	100
31) 2,2-Dichloropropane	9.14	77	528648	51.8762	ug/L	100
32) cis-1,2-Dichloroethene	9.20	96	414079	53.4021	ug/L	100
33) Chloroform	9.41	83	630762	50.6038	ug/L	100
34) 1-Bromopropane	9.55	122	88643	53.1827	ug/L	100
35) Bromochloromethane	9.64	130	235134	53.1708	ug/L	100
36) Tetrahydrofuran	9.67	42	165023	101.3834	ug/L	100
38) 1,1,1-Trichloroethane	9.94	97	574212	53.0371	ug/L	100
39) Cyclohexane	9.97	56	764351	53.0053	ug/L	100
40) 1,1-Dichloropropene	10.14	75	486257	53.4861	ug/L	100
41) Tert-Amyl-Methyl ether	10.24	73	1819195	102.4127	ug/L	100
42) Carbon Tetrachloride	10.28	117	522667	53.1266	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M428962.D 8260WTR.M Tue Mar 05 10:54:59 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428962.D Vial: 8
 Acq On : 4 Mar 2019 15:40 Operator: EEA
 Sample : WG698192-08 50 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:54:57 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

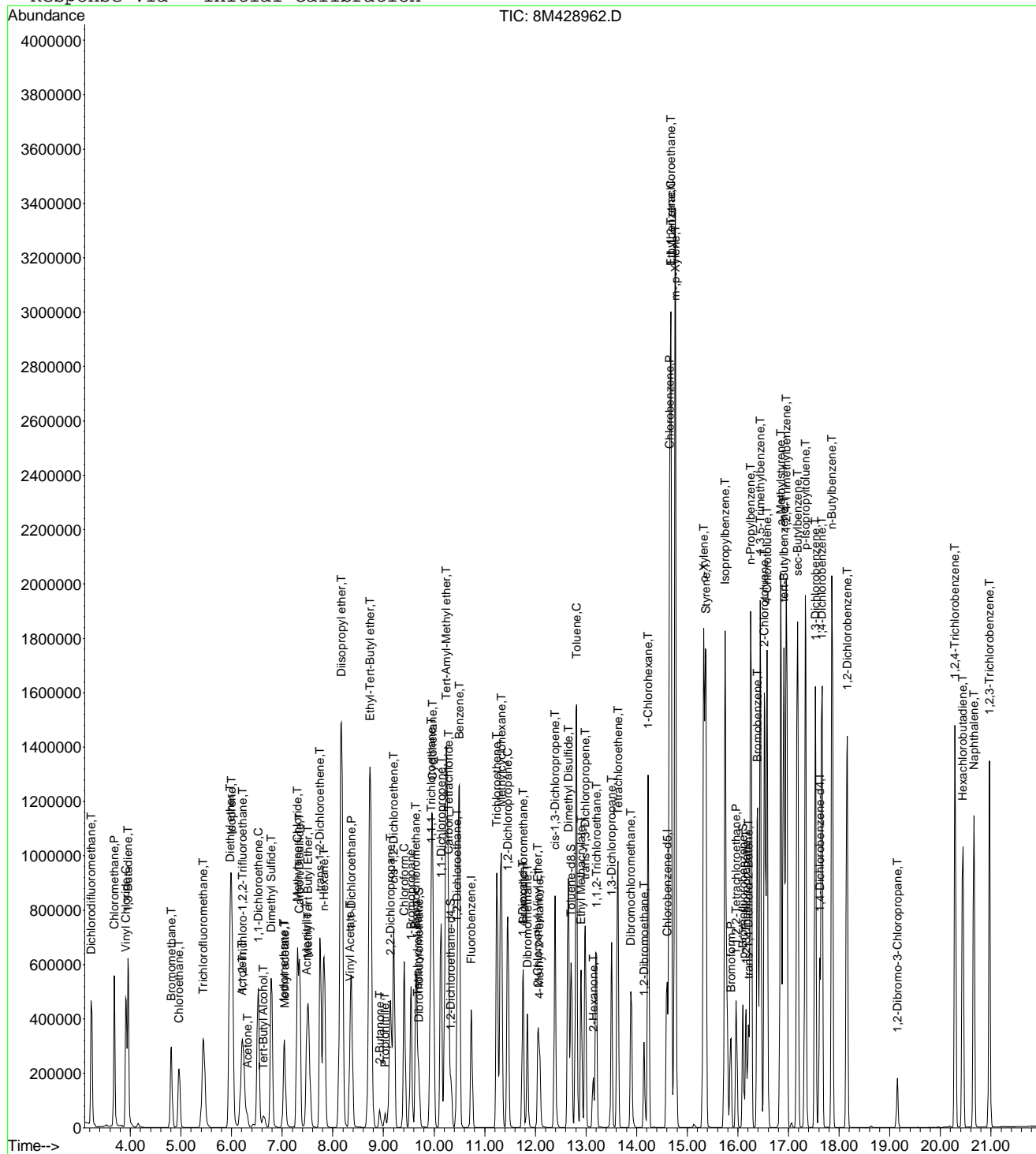
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	454650	53.3261	ug/L	100
46) Benzene	10.50	78	1499682	52.6586	ug/L	100
47) Trichloroethene	11.24	130	413618	53.3075	ug/L	100
48) Methylcyclohexane	11.32	83	630056	52.4333	ug/L	100
49) 1,2-Dichloropropane	11.46	63	416442	53.8713	ug/L	100
50) Bromodichloromethane	11.76	83	482368	53.8914	ug/L	100
51) 1,4-Dioxane	11.75	88	9416	216.4757	ug/L	100
52) Dibromomethane	11.84	93	185991	52.2711	ug/L	100
53) 2-Chloroethyl Vinyl Ether	12.05	63	174497	51.6598	ug/L	100
54) 4-Methyl-2-Pentanone	12.08	58	109795	51.6645	ug/L	100
55) cis-1,3-Dichloropropene	12.39	75	583167	54.7643	ug/L	100
56) Dimethyl Disulfide	12.65	79	345713	52.8677	ug/L	100
59) Toluene	12.81	91	1554401	51.9588	ug/L	100
60) Ethyl Methacrylate	12.91	69	395254	50.8804	ug/L	100
62) trans-1,3-Dichloropropene	12.99	75	483580	53.7681	ug/L	100
63) 1,1,2-Trichloroethane	13.21	97	277584	53.0830	ug/L	100
64) 2-Hexanone	13.14	58	103399	48.5852	ug/L	100
65) 1,3-Dichloropropane	13.51	76	470894	52.2865	ug/L	100
66) Tetrachloroethene	13.63	164	327264	51.9103	ug/L	100
67) Dibromochloromethane	13.89	129	354221	53.3730	ug/L	100
68) 1,2-Dibromoethane	14.15	107	273327	53.8280	ug/L	100
69) 1-Chlorohexane	14.23	91	544258	52.5006	ug/L	100
70) Chlorobenzene	14.65	112	1031350	50.8539	ug/L	100
71) 1,1,1,2-Tetrachloroethane	14.67	131	391985	52.5717	ug/L	100
72) Ethylbenzene	14.67	106	583481	53.1750	ug/L	100
73) m-,p-Xylene	14.77	106	1397295	104.5103	ug/L	100
74) o-Xylene	15.33	106	695543	53.6147	ug/L	100
75) Styrene	15.37	104	1156210	53.8331	ug/L	100
76) Bromoform	15.86	173	224940	54.6622	ug/L	100
77) Isopropylbenzene	15.75	105	1743408	53.2055	ug/L	100
79) 1,1,2,2-Tetrachloroethane	15.97	83	312670	50.9410	ug/L	100
81) 1,2,3-Trichloropropane	16.16	110	96184	52.3835	ug/L	100
82) trans-1,4-Dichloro-2-Butene	16.20	53	108529	52.3052	ug/L	100
83) n-Propylbenzene	16.26	91	2055567	52.2075	ug/L	100
84) Bromobenzene	16.39	156	469393	52.1041	ug/L	100
85) 1,3,5-Trimethylbenzene	16.44	105	1494686	52.6616	ug/L	100
86) 2-Chlorotoluene	16.53	91	1367937	52.5459	ug/L	100
87) 4-Chlorotoluene	16.58	91	1153673	50.7178	ug/L	100
88) a-Methylstyrene	16.85	118	875115	53.7320	ug/L	100
89) tert-Butylbenzene	16.91	134	334890	51.1440	ug/L	100
90) 1,2,4-Trimethylbenzene	16.96	105	1523293	53.0121	ug/L	100
91) sec-Butylbenzene	17.18	105	1910066	52.5837	ug/L	100
92) p-Isopropyltoluene	17.33	119	1655407	53.1628	ug/L	100
93) 1,3-Dichlorobenzene	17.53	146	902835	51.4080	ug/L	100
94) 1,4-Dichlorobenzene	17.66	146	905126	50.8929	ug/L	100
95) n-Butylbenzene	17.86	91	1510654	53.2454	ug/L	100
96) 1,2-Dichlorobenzene	18.16	146	838191	51.4315	ug/L	100
97) 1,2-Dibromo-3-Chloropropane	19.15	75	57558	52.0287	ug/L	100
98) 1,2,4-Trichlorobenzene	20.29	180	620877	52.5041	ug/L	100
99) Hexachlorobutadiene	20.44	225	294584	52.8677	ug/L	100
100) Naphthalene	20.66	128	1155751	53.6143	ug/L	100
101) 1,2,3-Trichlorobenzene	20.97	180	566747	52.1715	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M428962.D 8260WTR.M Tue Mar 05 10:55:00 2019

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428962.D Vial: 8
Acq On : 4 Mar 2019 15:40 Operator: EEA
Sample : WG698192-08 50 ug/L ICAL 8260 Inst : HPMS8
Misc : 1,1 STD92326 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428963.D Vial: 9
 Acq On : 4 Mar 2019 16:09 Operator: EEA
 Sample : WG698192-09 100 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Mar 05 10:55:01 2019

Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	578503	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	434394	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	240678	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	9.70	111	299207	51.8870	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	207.56%#	
43) 1,2-Dichloroethane-d4	10.33	65	292145	51.3184	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	205.28%#	
58) Toluene-d8	12.71	98	1114155	50.3523	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	201.40%#	
80) p-Bromofluorobenzene	16.10	95	448708	51.6568	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	206.64%#	

Target Compounds

						Qvalue
2) Dichlorodifluoromethane	3.23	85	1012567	100.6293	ug/L	99
3) Chloromethane	3.68	50	1237917	85.5444	ug/L	99
4) Vinyl Chloride	3.92	62	1139316	96.6082	ug/L	100
5) 1,3-Butadiene	3.95	54	629287	97.4416	ug/L	99
6) Bromomethane	4.80	94	599065	84.4547	ug/L	98
7) Chloroethane	4.97	64	625726	99.1508	ug/L	99
8) Trichlorofluoromethane	5.44	101	1172724	97.9406	ug/L	100
9) Diethyl ether	5.97	59	1235386	193.4538	ug/L	100
10) Isoprene	6.00	67	1161960	97.0984	ug/L	100
11) Acrolein	6.21	56	74704	102.0183	ug/L	100
12) 1,1,2-Trichloro-1,2,2-Trif	6.22	101	672607	97.5063	ug/L	99
13) Acetone	6.31	43	175252	98.3428	ug/L	95
14) 1,1-Dichloroethene	6.53	61	1185524	96.0923	ug/L	100
15) Tert-Butyl Alcohol	6.64	59	217661	379.4451	ug/L	98
16) Dimethyl Sulfide	6.79	62	867504	97.6144	ug/L	99
17) Iodomethane	7.03	142	932933	101.4253	ug/L	100
18) Methyl acetate	7.06	43	458352	99.3291	ug/L	99
19) Methylene Chloride	7.30	84	735238	95.3712	ug/L	100
20) Carbon Disulfide	7.34	76	2243994	97.2684	ug/L	99
21) Acrylonitrile	7.49	53	233053	104.2394	ug/L	97
22) Methyl Tert Butyl Ether	7.51	73	1764776	96.1742	ug/L	99
23) trans-1,2-Dichloroethene	7.75	61	1148539	97.4283	ug/L	100
24) n-Hexane	7.83	57	1114202	96.3067	ug/L	100
25) Diisopropyl ether	8.16	45	5483147	187.8866	ug/L	98
26) Vinyl Acetate	8.34	43	1086706	100.3813	ug/L	100
27) 1,1-Dichloroethane	8.37	63	1473636	96.6987	ug/L	100
28) Ethyl-Tert-Butyl ether	8.74	59	4719674	186.5960	ug/L	98
29) 2-Butanone	8.93	43	256859	97.8459	ug/L	98
30) Propionitrile	9.03	54	159010	193.3041	ug/L	98
31) 2,2-Dichloropropane	9.13	77	1032620	93.3841	ug/L	100
32) cis-1,2-Dichloroethene	9.21	96	831013	98.7674	ug/L	100
33) Chloroform	9.41	83	1261043	93.2349	ug/L	100
34) 1-Bromopropane	9.55	122	178392	98.6352	ug/L	100
35) Bromochloromethane	9.64	130	477677	99.5458	ug/L	99
36) Tetrahydrofuran	9.67	42	335892	190.1747	ug/L	98
38) 1,1,1-Trichloroethane	9.94	97	1134879	96.6024	ug/L	99
39) Cyclohexane	9.97	56	1513886	96.7499	ug/L	99
40) 1,1-Dichloropropene	10.14	75	974256	98.7595	ug/L	100
41) Tert-Amyl-Methyl ether	10.24	73	3624583	188.0457	ug/L	99
42) Carbon Tetrachloride	10.28	117	1042270	97.6334	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M428963.D 8260WTR.M Tue Mar 05 10:55:03 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428963.D Vial: 9
 Acq On : 4 Mar 2019 16:09 Operator: EEA
 Sample : WG698192-09 100 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:55:01 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

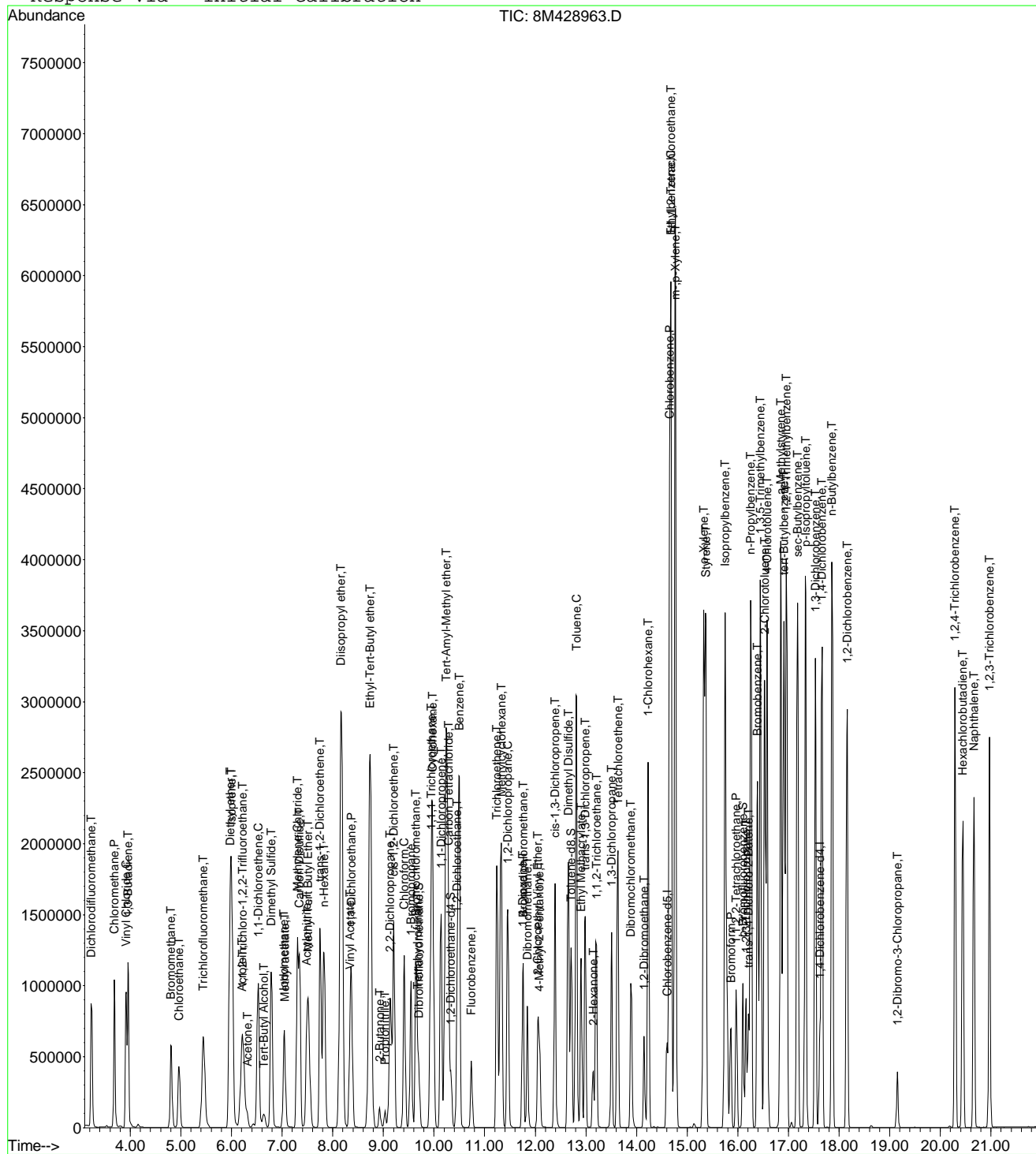
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	908506	98.2021	ug/L	100
46) Benzene	10.50	78	2934603	94.9621	ug/L	99
47) Trichloroethene	11.24	130	825611	98.0606	ug/L	99
48) Methylcyclohexane	11.33	83	1240651	95.1499	ug/L	99
49) 1,2-Dichloropropane	11.46	63	827768	98.6830	ug/L	100
50) Bromodichloromethane	11.76	83	967751	99.6404	ug/L	100
51) 1,4-Dioxane	11.75	88	20243	428.8924	ug/L	99
52) Dibromomethane	11.84	93	380355	98.5121	ug/L	100
53) 2-Chloroethyl Vinyl Ether	12.05	63	369044	100.6870	ug/L	100
54) 4-Methyl-2-Pentanone	12.08	58	234067	101.5033	ug/L	99
55) cis-1,3-Dichloropropene	12.39	75	1173352	101.5463	ug/L	100
56) Dimethyl Disulfide	12.65	79	701550	98.8700	ug/L	99
59) Toluene	12.81	91	3010430	92.8182	ug/L	98
60) Ethyl Methacrylate	12.91	69	813537	96.1671	ug/L	100
62) trans-1,3-Dichloropropene	12.99	75	977484	100.2477	ug/L	99
63) 1,1,2-Trichloroethane	13.21	97	555137	97.9197	ug/L	99
64) 2-Hexanone	13.15	58	222332	93.7044	ug/L	96
65) 1,3-Dichloropropane	13.51	76	954169	97.7237	ug/L	99
66) Tetrachloroethene	13.63	164	655070	95.8410	ug/L	99
67) Dibromochloromethane	13.89	129	723064	100.4923	ug/L	100
68) 1,2-Dibromoethane	14.15	107	557984	101.3574	ug/L	99
69) 1-Chlorohexane	14.23	91	1085750	96.6046	ug/L	100
70) Chlorobenzene	14.65	112	2043702	92.9488	ug/L	99
71) 1,1,1,2-Tetrachloroethane	14.68	131	801855	99.1943	ug/L	99
72) Ethylbenzene	14.68	106	1176858	98.9266	ug/L	94
73) m-,p-Xylene	14.77	106	2751893	189.8501	ug/L	92
74) o-Xylene	15.33	106	1415977	100.6756	ug/L	95
75) Styrene	15.37	104	2327941	99.9753	ug/L	98
76) Bromoform	15.86	173	470887	105.5470	ug/L	99
77) Isopropylbenzene	15.75	105	3379273	95.1238	ug/L	97
79) 1,1,2,2-Tetrachloroethane	15.97	83	658687	97.0087	ug/L	99
81) 1,2,3-Trichloropropane	16.16	110	197378	97.1721	ug/L	99
82) trans-1,4-Dichloro-2-Butene	16.21	53	229702	100.0724	ug/L	99
83) n-Propylbenzene	16.26	91	3941692	90.4972	ug/L	96
84) Bromobenzene	16.39	156	967544	97.0859	ug/L	99
85) 1,3,5-Trimethylbenzene	16.44	105	2952548	94.0355	ug/L	98
86) 2-Chlorotoluene	16.54	91	2680366	93.0717	ug/L	98
87) 4-Chlorotoluene	16.58	91	2307353	91.6944	ug/L	98
88) a-Methylstyrene	16.85	118	1801911	100.0119	ug/L	99
89) tert-Butylbenzene	16.91	134	692727	95.6325	ug/L	96
90) 1,2,4-Trimethylbenzene	16.96	105	3001972	94.4384	ug/L	97
91) sec-Butylbenzene	17.18	105	3734793	92.9438	ug/L	98
92) p-Isopropyltoluene	17.34	119	3247181	94.2672	ug/L	97
93) 1,3-Dichlorobenzene	17.53	146	1851935	95.3232	ug/L	100
94) 1,4-Dichlorobenzene	17.66	146	1836509	93.3451	ug/L	98
95) n-Butylbenzene	17.86	91	2982166	95.0167	ug/L	98
96) 1,2-Dichlorobenzene	18.16	146	1717023	95.2386	ug/L	99
97) 1,2-Dibromo-3-Chloropropane	19.15	75	121519	99.2961	ug/L	97
98) 1,2,4-Trichlorobenzene	20.29	180	1284312	98.1768	ug/L	100
99) Hexachlorobutadiene	20.44	225	610618	99.0607	ug/L	99
100) Naphthalene	20.66	128	2348094	98.4651	ug/L	99
101) 1,2,3-Trichlorobenzene	20.97	180	1162600	96.7443	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M428963.D 8260WTR.M Tue Mar 05 10:55:04 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428963.D Vial: 9
Acq On : 4 Mar 2019 16:09 Operator: EEA
Sample : WG698192-09 100 ug/L ICAL 8260 Inst : HPMS8
Misc : 1,1 STD92326 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428964.D Vial: 10
 Acq On : 4 Mar 2019 16:38 Operator: EEA
 Sample : WG698192-10 200 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Mar 05 10:55:05 2019

Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	565092	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	432785	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	251063	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	9.69	111	546178	96.9632	ug/L	-0.01
Spiked Amount	25.000	Range 86 - 118	Recovery	=	387.84%#	
43) 1,2-Dichloroethane-d4	10.33	65	539183	96.9611	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	387.84%#	
58) Toluene-d8	12.71	98	2011484	91.2434	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	364.96%#	
80) p-Bromofluorobenzene	16.10	95	844391	93.1883	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	372.76%#	

Target Compounds

						Qvalue
2) Dichlorodifluoromethane	3.22	85	2004123	203.8974	ug/L	98
3) Chloromethane	3.67	50	2402052	169.9296	ug/L	97
4) Vinyl Chloride	3.90	62	2214658	192.2484	ug/L	98
5) 1,3-Butadiene	3.94	54	1233022	200.4621	ug/L	99
6) Bromomethane	4.79	94	1228006	177.2298	ug/L	98
7) Chloroethane	4.95	64	1254003	203.4216	ug/L	100
8) Trichlorofluoromethane	5.43	101	2324194	198.7129	ug/L	99
10) Isoprene	5.99	67	2324648	198.8679	ug/L	100
12) 1,1,2-Trichloro-1,2,2-Trif	6.21	101	1357837	201.5142	ug/L	99
13) Acetone	6.30	43	350194	201.1753	ug/L	92
14) 1,1-Dichloroethene	6.52	61	2353107	195.2569	ug/L	99
16) Dimethyl Sulfide	6.78	62	1737568	200.1569	ug/L	98
17) Iodomethane	7.03	142	1826830	201.8012	ug/L	100
18) Methyl acetate	7.04	43	952086	211.2223	ug/L	99
19) Methylene Chloride	7.29	84	1473339	195.6494	ug/L	99
20) Carbon Disulfide	7.33	76	4313442	191.4082	ug/L	98
21) Acrylonitrile	7.50	53	35855	16.4177	ug/L #	29
22) Methyl Tert Butyl Ether	7.51	73	3512538	195.9640	ug/L	98
23) trans-1,2-Dichloroethene	7.75	61	2274275	197.5008	ug/L	99
24) n-Hexane	7.82	57	2289573	202.5971	ug/L	100
26) Vinyl Acetate	8.34	43	1985980	187.8028	ug/L	99
27) 1,1-Dichloroethane	8.36	63	2878727	193.3825	ug/L	99
29) 2-Butanone	8.92	43	538864	210.1423	ug/L	98
31) 2,2-Dichloropropane	9.13	77	2026979	187.6584	ug/L	99
32) cis-1,2-Dichloroethene	9.21	96	1649584	200.7090	ug/L	99
33) Chloroform	9.41	83	2468111	186.8097	ug/L	99
34) 1-Bromopropane	9.55	122	362532	205.2057	ug/L	100
35) Bromochloromethane	9.64	130	957734	204.3243	ug/L	99
36) Tetrahydrofuran	9.68	42	2780	1.6113	ug/L #	40
38) 1,1,1-Trichloroethane	9.94	97	2235602	194.8137	ug/L	99
39) Cyclohexane	9.97	56	3040534	198.9270	ug/L	99
40) 1,1-Dichloropropene	10.14	75	1922547	199.5121	ug/L	99
42) Carbon Tetrachloride	10.28	117	2047091	196.3097	ug/L	100
45) 1,2-Dichloroethane	10.46	62	1787189	197.7653	ug/L	98
46) Benzene	10.49	78	5433612	180.0014	ug/L	95
47) Trichloroethene	11.24	130	1663967	202.3254	ug/L	99
48) Methylcyclohexane	11.32	83	2521969	198.0090	ug/L	99
49) 1,2-Dichloropropane	11.45	63	1659991	202.5937	ug/L	100
50) Bromodichloromethane	11.75	83	1922180	202.6060	ug/L	99
51) 1,4-Dioxane	11.74	88	2147	46.5685	ug/L #	14

(#) = qualifier out of range (m) = manual integration
 8M428964.D 8260WTR.M Tue Mar 05 10:55:07 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428964.D Vial: 10
 Acq On : 4 Mar 2019 16:38 Operator: EEA
 Sample : WG698192-10 200 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:55:05 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
52) Dibromomethane	11.84	93	767150	203.4077	ug/L	99
53) 2-Chloroethyl Vinyl Ether	12.05	63	781751	218.3484	ug/L	100
54) 4-Methyl-2-Pentanone	12.08	58	492715	218.7371	ug/L	99
55) cis-1,3-Dichloropropene	12.39	75	2325494	206.0332	ug/L	98
56) Dimethyl Disulfide	12.65	79	1426255	205.7734	ug/L	97
59) Toluene	12.80	91	5454725	168.8066	ug/L	94
60) Ethyl Methacrylate	12.91	69	1691777	200.2070	ug/L	99
62) trans-1,3-Dichloropropene	12.98	75	1954014	201.1425	ug/L	98
63) 1,1,2-Trichloroethane	13.20	97	1124859	199.1497	ug/L	99
64) 2-Hexanone	13.13	58	490171	204.0804	ug/L	92
65) 1,3-Dichloropropane	13.51	76	1902968	195.6219	ug/L	99
66) Tetrachloroethene	13.63	164	1321277	194.0302	ug/L	99
67) Dibromochloromethane	13.89	129	1465222	204.3954	ug/L	99
68) 1,2-Dibromoethane	14.15	107	1139513	207.7613	ug/L	100
69) 1-Chlorohexane	14.23	91	2209339	197.3067	ug/L	99
70) Chlorobenzene	14.64	112	3906155	178.3149	ug/L	95
71) 1,1,1,2-Tetrachloroethane	14.67	131	1596855	198.2750	ug/L	100
72) Ethylbenzene	14.67	106	2320121	195.7543	ug/L	78
73) m-,p-Xylene	14.77	106	5020397	347.6395	ug/L	77
74) o-Xylene	15.33	106	2789652	199.0809	ug/L	87
75) Styrene	15.37	104	4453512	191.9708	ug/L	94
76) Bromoform	15.86	173	1003376	225.7378	ug/L	99
77) Isopropylbenzene	15.75	105	6080157	171.7879	ug/L	90
79) 1,1,2,2-Tetrachloroethane	15.97	83	1359009	191.8702	ug/L	99
81) 1,2,3-Trichloropropane	16.16	110	417199	196.8972	ug/L	100
82) trans-1,4-Dichloro-2-Buten	16.20	53	496588	207.3956	ug/L	99
83) n-Propylbenzene	16.26	91	6822994	150.1692	ug/L	87
84) Bromobenzene	16.39	156	1975669	190.0437	ug/L	98
85) 1,3,5-Trimethylbenzene	16.44	105	5450748	166.4197	ug/L	92
86) 2-Chlorotoluene	16.53	91	5047792	168.0268	ug/L	92
87) 4-Chlorotoluene	16.58	91	4250089	161.9125	ug/L	91
88) a-Methylstyrene	16.85	118	3580082	190.4868	ug/L	96
89) tert-Butylbenzene	16.91	134	1426368	188.7682	ug/L	87
90) 1,2,4-Trimethylbenzene	16.96	105	5486719	165.4659	ug/L	90
91) sec-Butylbenzene	17.18	105	6582290	157.0307	ug/L	90
92) p-Isopropyltoluene	17.33	119	5837012	162.4420	ug/L	90
93) 1,3-Dichlorobenzene	17.53	146	3620888	178.6659	ug/L	96
94) 1,4-Dichlorobenzene	17.66	146	3596351	175.2323	ug/L	95
95) n-Butylbenzene	17.86	91	5409792	165.2351	ug/L	91
96) 1,2-Dichlorobenzene	18.16	146	3368677	179.1224	ug/L	96
97) 1,2-Dibromo-3-Chloropropan	19.15	75	258649	202.6060	ug/L	96
98) 1,2,4-Trichlorobenzene	20.29	180	2623960	192.2867	ug/L	99
99) Hexachlorobutadiene	20.44	225	1253776	194.9869	ug/L	99
100) Naphthalene	20.66	128	4525002	181.9028	ug/L	96
101) 1,2,3-Trichlorobenzene	20.97	180	2400819	191.5173	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M428964.D 8260WTR.M Tue Mar 05 10:55:08 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428965.D Vial: 11
 Acq On : 4 Mar 2019 17:07 Operator: EEA
 Sample : WG698192-11 300 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Mar 05 10:55:09 2019

Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.73	96	542966	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	396623	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	212626	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	9.69	111	971	0.1794	ug/L	-0.01
Spiked Amount	25.000	Range 86 - 118	Recovery	=	0.72%#	
43) 1,2-Dichloroethane-d4	10.33	65	805	0.1507	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	0.60%#	
58) Toluene-d8	12.71	98	6199	0.3068	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	1.24%#	
80) p-Bromofluorobenzene	16.10	95	4824	0.6286	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	2.52%#	

Target Compounds

						Qvalue
2) Dichlorodifluoromethane	3.22	85	6424	0.6802	ug/L	97
3) Chloromethane	3.67	50	41635	3.0654	ug/L	99
4) Vinyl Chloride	3.90	62	2989	0.2700	ug/L	98
5) 1,3-Butadiene	3.94	54	1715174	292.4400	ug/L	99
6) Bromomethane	4.79	94	186263	27.9775	ug/L	100
8) Trichlorofluoromethane	5.42	101	2587	0.2302	ug/L	# 74
9) Diethyl ether	5.96	59	1818871	303.4656	ug/L	99
10) Isoprene	6.00	67	3378731	300.8207	ug/L	100
11) Acrolein	6.20	56	110342	160.5492	ug/L	95
12) 1,1,2-Trichloro-1,2,2-Trif	6.21	101	1873	0.2893	ug/L	99
13) Acetone	6.30	43	480528	287.2970	ug/L	93
14) 1,1-Dichloroethene	6.52	61	3073	0.2654	ug/L	97
15) Tert-Butyl Alcohol	6.63	59	329899	612.7485	ug/L	96
16) Dimethyl Sulfide	6.77	62	2528685	303.1588	ug/L	98
17) Iodomethane	7.03	142	2145401	246.3138	ug/L	99
18) Methyl acetate	7.04	43	1376364	317.7924	ug/L	98
19) Methylene Chloride	7.29	84	3022	0.4177	ug/L	99
20) Carbon Disulfide	7.33	76	5921410	273.4691	ug/L	96
21) Acrylonitrile	7.48	53	323034	153.9425	ug/L	97
22) Methyl Tert Butyl Ether	7.51	73	4004	0.2325	ug/L	# 1
23) trans-1,2-Dichloroethene	7.75	61	7116	0.6431	ug/L	97
24) n-Hexane	7.82	57	3192652	294.0200	ug/L	100
25) Diisopropyl ether	8.16	45	7633508	278.6911	ug/L	96
26) Vinyl Acetate	8.34	43	3174122	312.3901	ug/L	98
28) Ethyl-Tert-Butyl ether	8.73	59	6671315	281.0184	ug/L	96
29) 2-Butanone	8.91	43	764637	310.3389	ug/L	97
30) Propionitrile	9.03	54	238789	306.5840	ug/L	99
32) cis-1,2-Dichloroethene	9.20	96	2397	0.3035	ug/L	88
33) Chloroform	9.40	83	1936	0.1525	ug/L	100
34) 1-Bromopropane	9.55	122	522744	307.9488	ug/L	98
35) Bromochloromethane	9.64	130	978	0.2172	ug/L	# 32
36) Tetrahydrofuran	9.67	42	488537	294.7023	ug/L	97
39) Cyclohexane	9.97	56	4327622	294.6725	ug/L	99
40) 1,1-Dichloropropene	10.13	75	4234	0.4573	ug/L	# 70
41) Tert-Amyl-Methyl ether	10.24	73	5156927	285.0554	ug/L	98
42) Carbon Tetrachloride	10.27	117	1259	0.1257	ug/L	# 78
45) 1,2-Dichloroethane	10.45	62	1438	0.1656	ug/L	# 38
46) Benzene	10.49	78	6854	0.2363	ug/L	# 83
47) Trichloroethene	11.24	130	2826	0.3576	ug/L	96
48) Methylcyclohexane	11.32	83	3560427	290.9335	ug/L	98

(#) = qualifier out of range (m) = manual integration
 8M428965.D 8260WTR.M Tue Mar 05 10:55:11 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428965.D Vial: 11
 Acq On : 4 Mar 2019 17:07 Operator: EEA
 Sample : WG698192-11 300 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:55:09 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

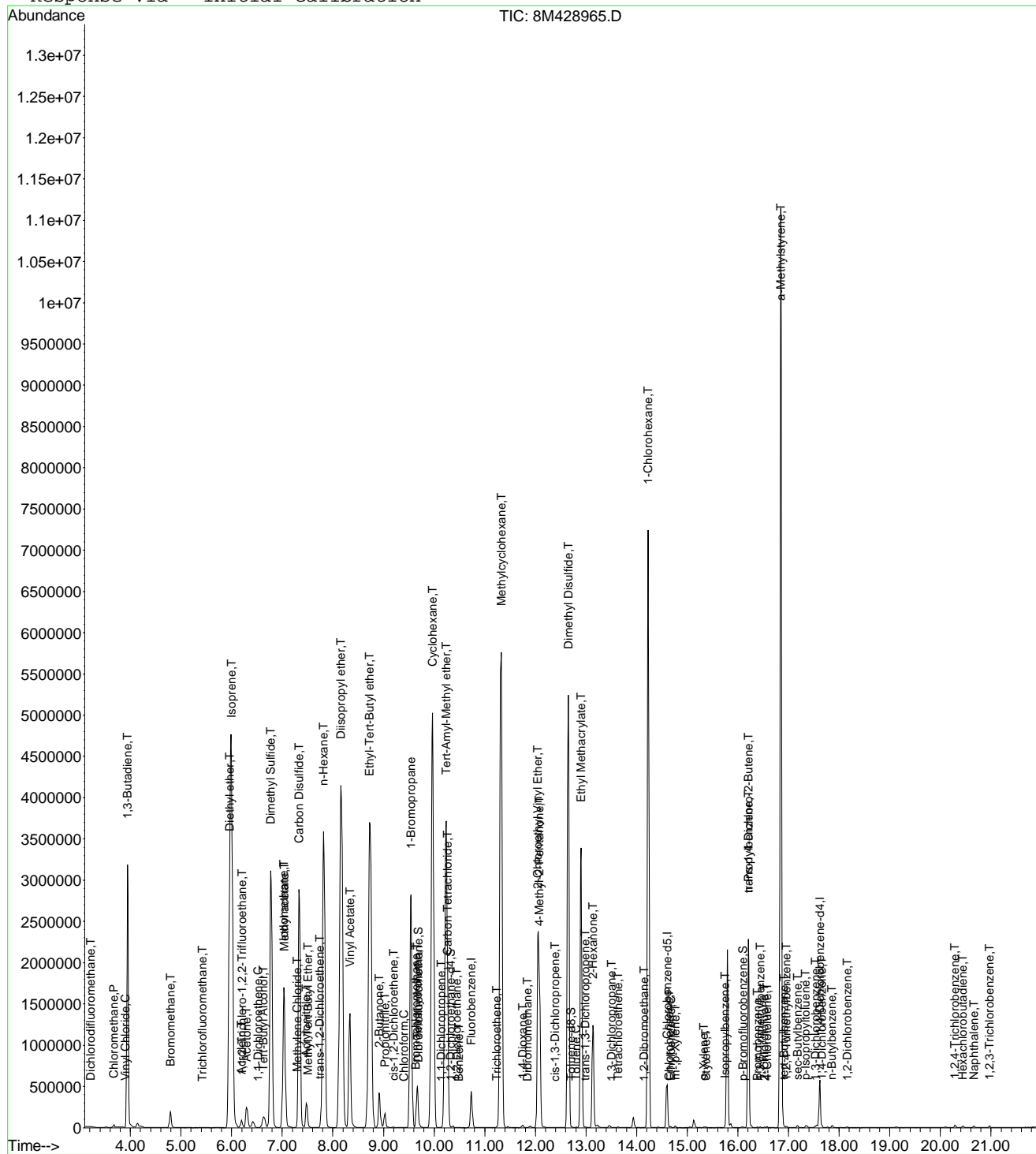
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
51) 1,4-Dioxane	11.75	88	28018	632.4753	ug/L	92
52) Dibromomethane	11.84	93	926	0.2555	ug/L	97
53) 2-Chloroethyl Vinyl Ether	12.05	63	1126775	327.5407	ug/L	99
54) 4-Methyl-2-Pentanone	12.08	58	693546	320.4412	ug/L	99
55) cis-1,3-Dichloropropene	12.39	75	2506	0.2311	ug/L	95
56) Dimethyl Disulfide	12.65	79	2029725	304.7726	ug/L	92
59) Toluene	12.80	91	6757	0.2282	ug/L	99
60) Ethyl Methacrylate	12.91	69	2343646	302.3930	ug/L	99
62) trans-1,3-Dichloropropene	12.99	75	3194	0.3588	ug/L	98
64) 2-Hexanone	13.13	58	670462	303.2643	ug/L #	47
65) 1,3-Dichloropropane	13.51	76	1201	0.1347	ug/L #	1
66) Tetrachloroethene	13.63	164	2546	0.4080	ug/L	88
68) 1,2-Dibromoethane	14.15	107	1277	0.2541	ug/L	75
69) 1-Chlorohexane	14.23	91	3057292	297.9276	ug/L	99
70) Chlorobenzene	14.64	112	6538	0.3257	ug/L	87
72) Ethylbenzene	14.67	106	3192	0.2939	ug/L	84
73) m-,p-Xylene	14.77	106	8006	0.6049	ug/L	85
74) o-Xylene	15.33	106	3229	0.2514	ug/L	79
75) Styrene	15.37	104	5587	0.2628	ug/L	91
77) Isopropylbenzene	15.75	105	23653	0.7292	ug/L	96
82) trans-1,4-Dichloro-2-Butene	16.20	53	627246	309.3197	ug/L #	71
83) n-Propylbenzene	16.20	91	160201	4.1633	ug/L #	59
84) Bromobenzene	16.39	156	2990	0.3396	ug/L	88
85) 1,3,5-Trimethylbenzene	16.44	105	9318	0.3359	ug/L	95
86) 2-Chlorotoluene	16.52	91	8631	0.3392	ug/L	94
87) 4-Chlorotoluene	16.58	91	10061	0.4526	ug/L	93
88) a-Methylstyrene	16.85	118	4498901	282.6472	ug/L	93
89) tert-Butylbenzene	16.91	134	3409	0.5327	ug/L #	1
90) 1,2,4-Trimethylbenzene	16.96	105	10435	0.3716	ug/L	84
91) sec-Butylbenzene	17.18	105	27689	0.7800	ug/L	94
92) p-Isopropyltoluene	17.33	119	15736	0.5171	ug/L	98
93) 1,3-Dichlorobenzene	17.53	146	9884	0.5759	ug/L	98
94) 1,4-Dichlorobenzene	17.66	146	10190	0.5863	ug/L	92
95) n-Butylbenzene	17.86	91	20588	0.7425	ug/L	99
96) 1,2-Dichlorobenzene	18.16	146	6432	0.4038	ug/L	99
98) 1,2,4-Trichlorobenzene	20.29	180	14061	1.2167	ug/L	98
99) Hexachlorobutadiene	20.44	225	5816	1.0680	ug/L	93
100) Naphthalene	20.66	128	17967	0.8528	ug/L	96
101) 1,2,3-Trichlorobenzene	20.97	180	10100	0.9513	ug/L	97

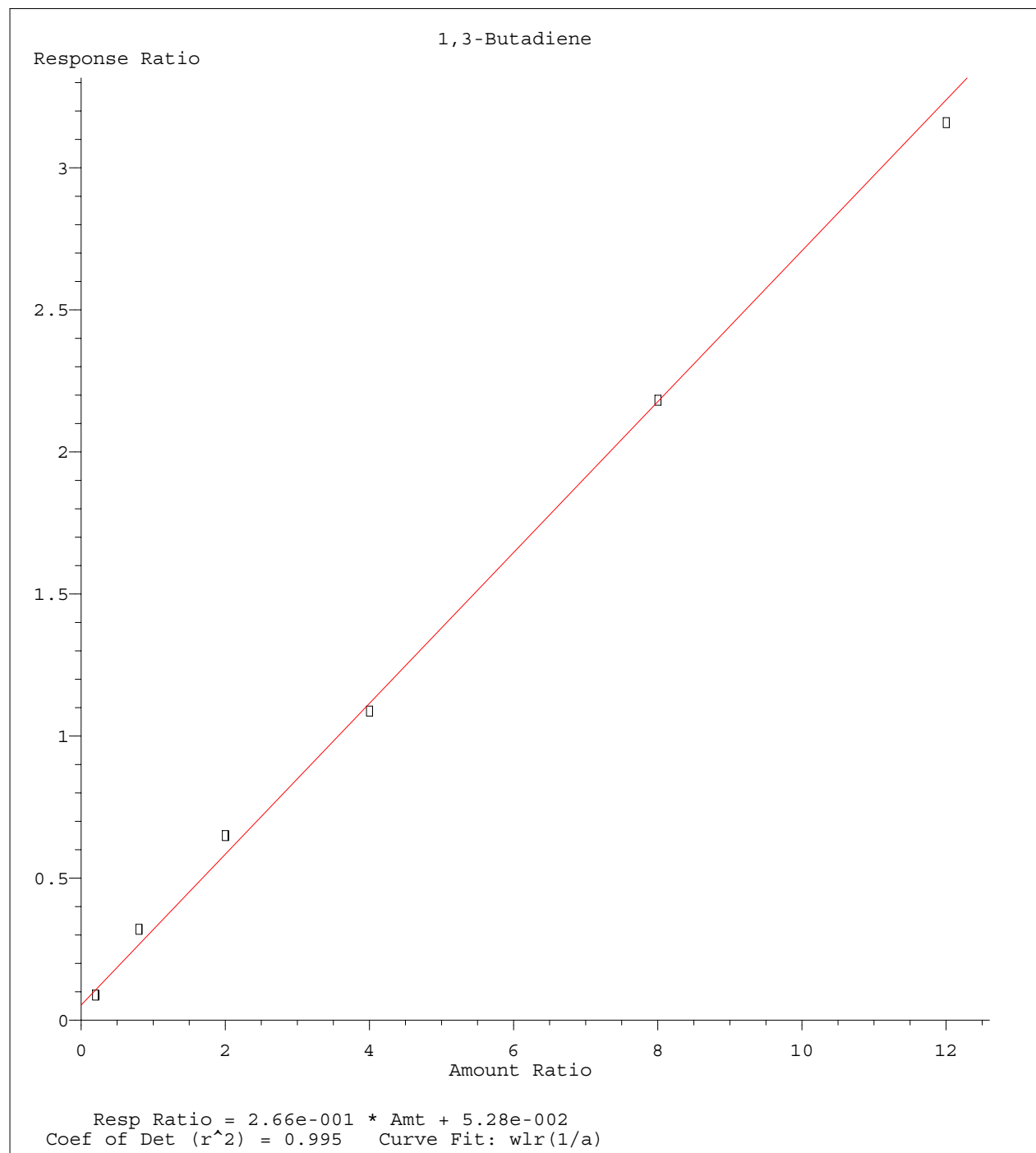
(#) = qualifier out of range (m) = manual integration
 8M428965.D 8260WTR.M Tue Mar 05 10:55:12 2019

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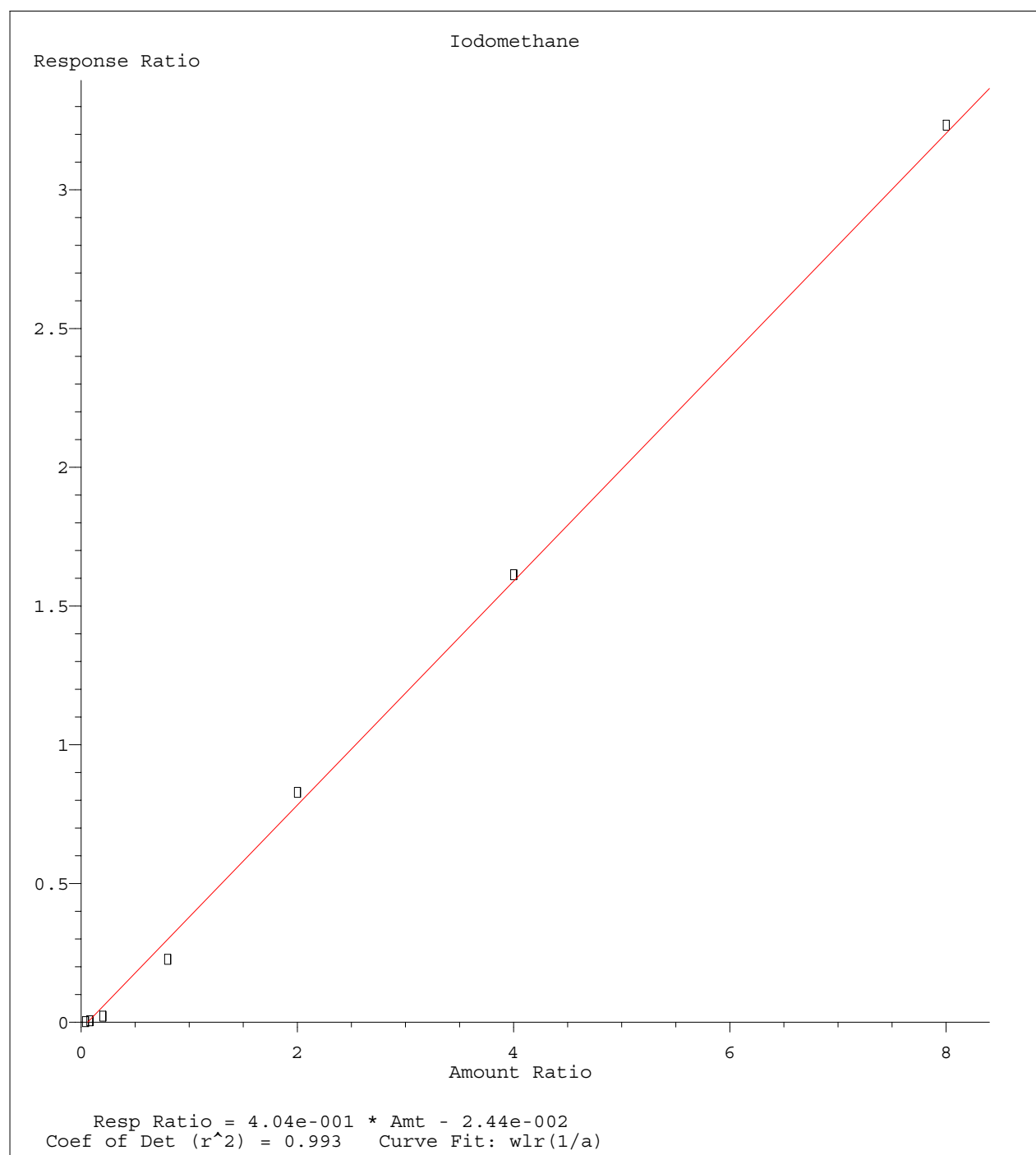
Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428965.D Vial: 11
 Acq On : 4 Mar 2019 17:07 Operator: EEA
 Sample : WG698192-11 300 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 5 10:54 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration

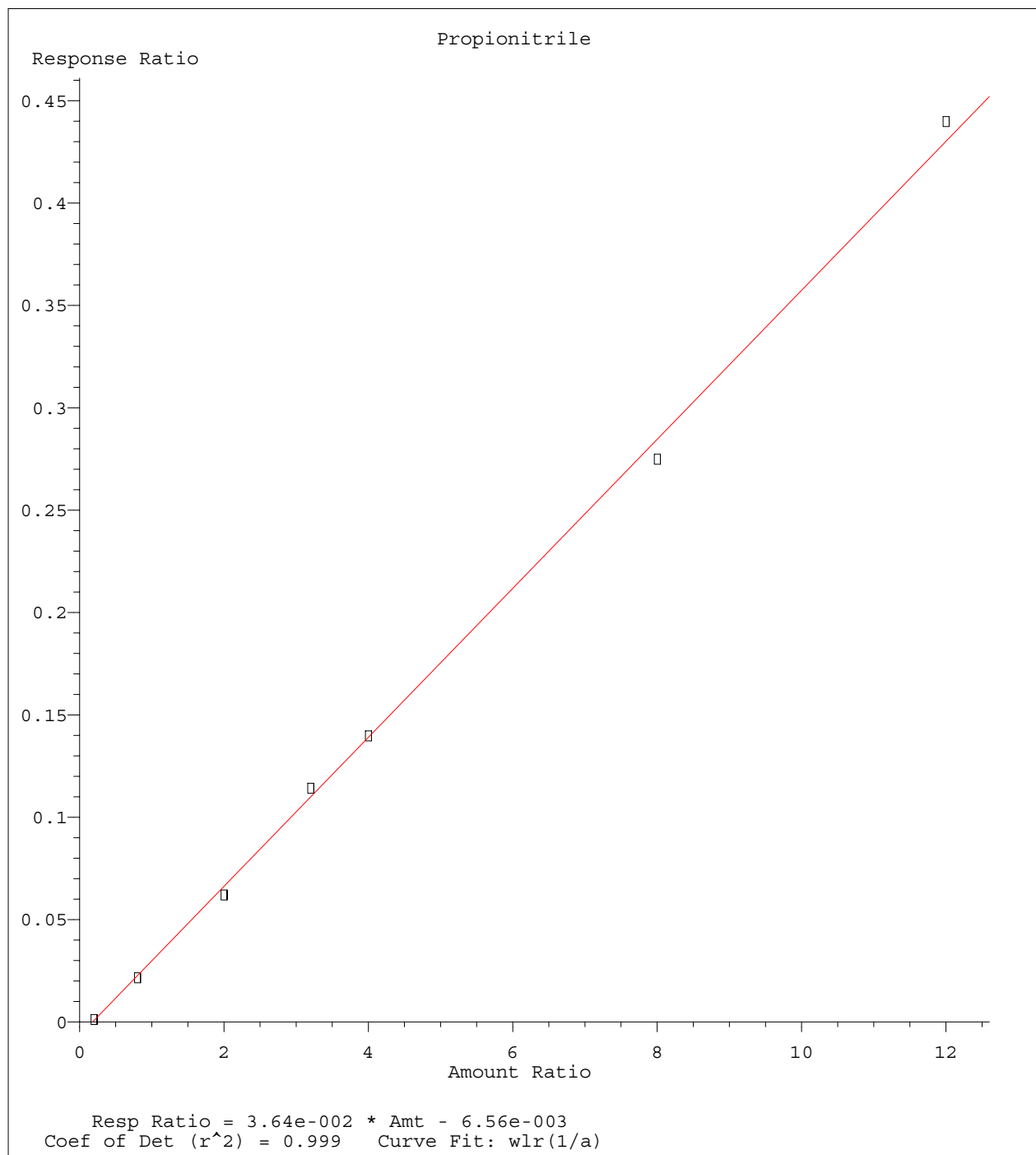




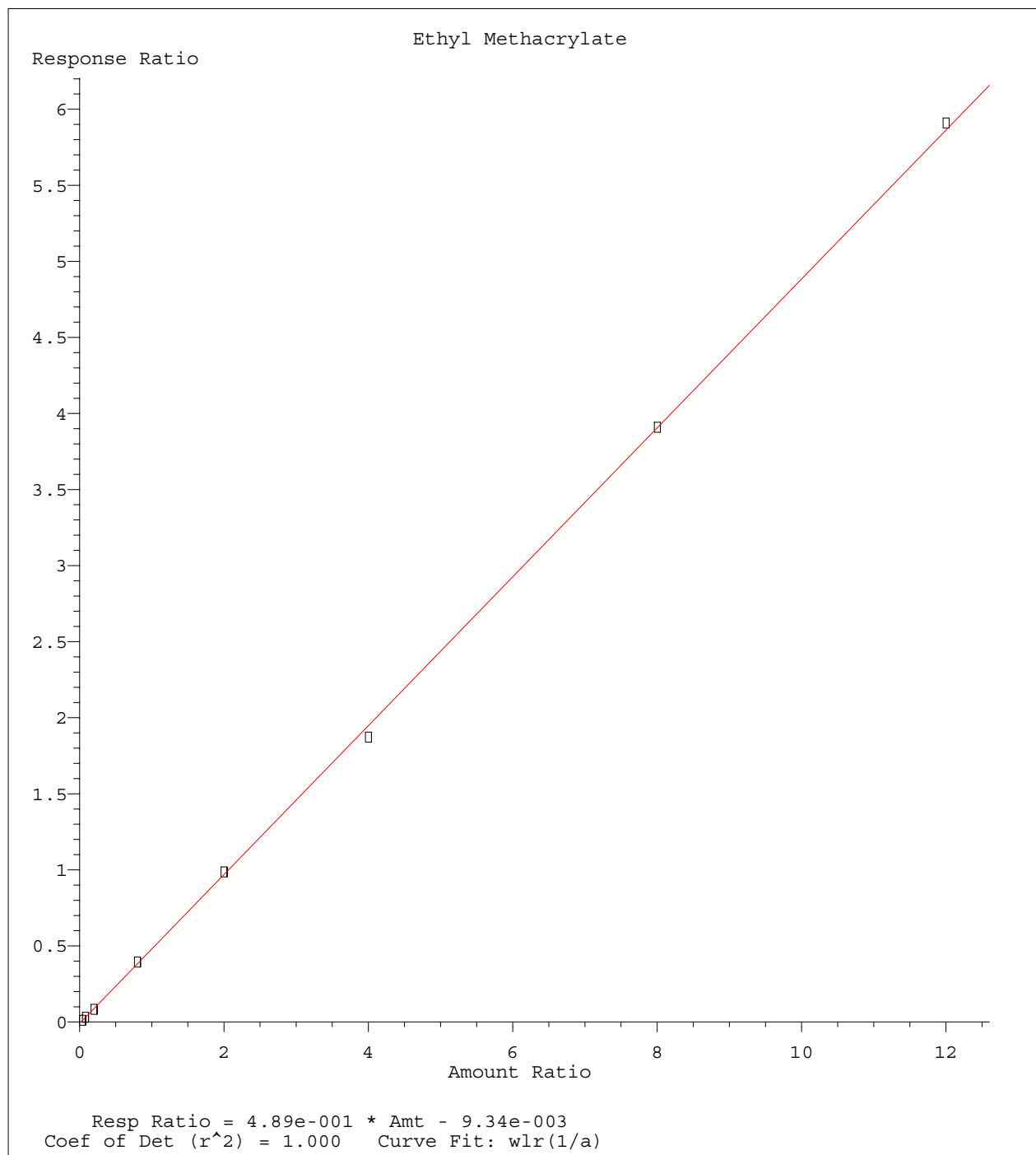
Method Name: K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M
Calibration Table Last Updated: Mon Mar 04 17:44:12 2019



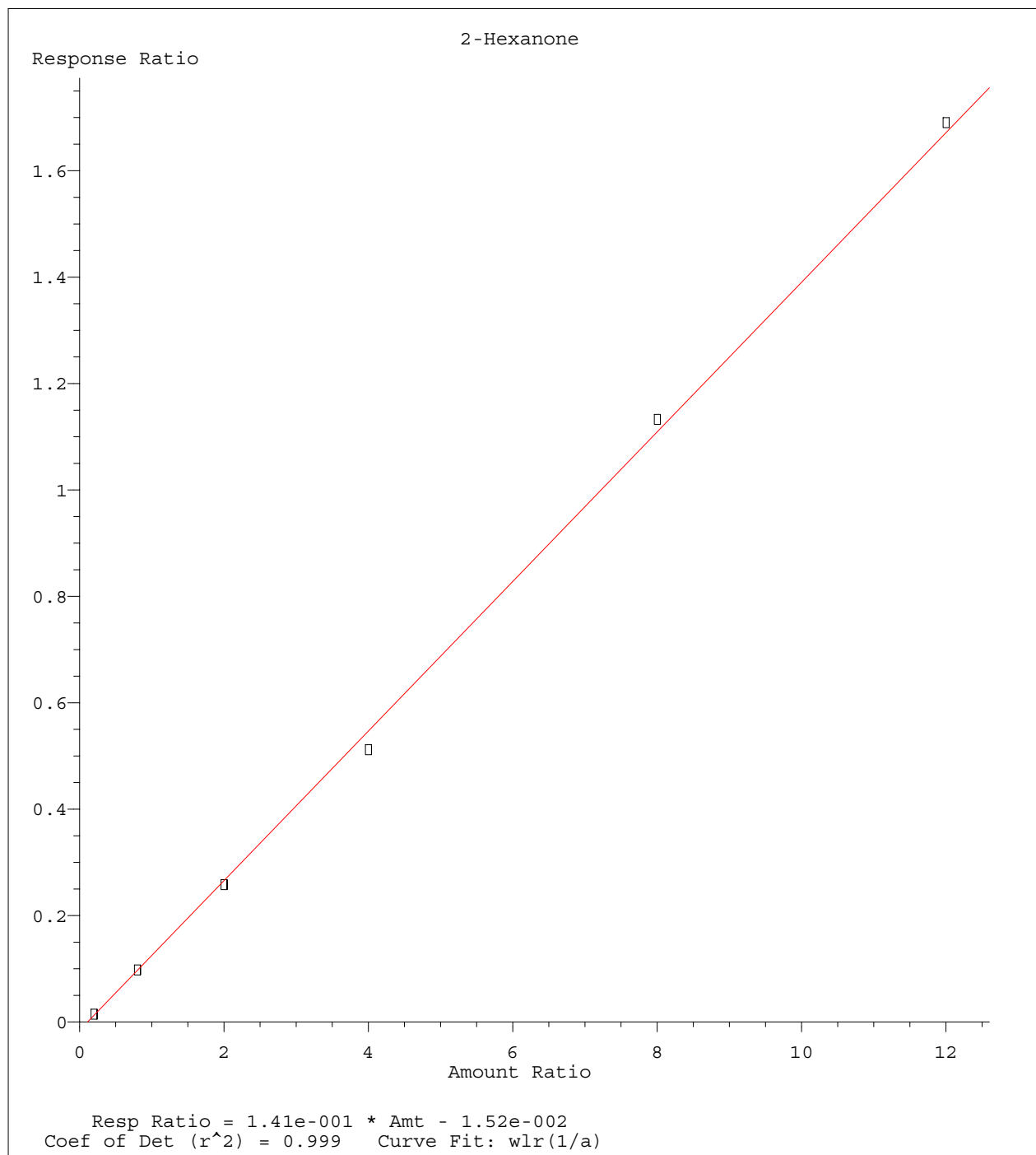
Method Name: K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M
Calibration Table Last Updated: Mon Mar 04 17:44:12 2019



Method Name: K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M
Calibration Table Last Updated: Mon Mar 04 17:44:12 2019



Method Name: K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M
Calibration Table Last Updated: Mon Mar 04 17:44:12 2019



Method Name: K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M
Calibration Table Last Updated: Mon Mar 04 17:44:12 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428967.D Vial: 13
 Acq On : 4 Mar 2019 18:06 Operator: EEA
 Sample : WG698192-12 ALT 50 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:55:31 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	511233	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	379639	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	222101	25.00	ug/L	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.70	111	130702	25.6481	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	102.60%	
43) 1,2-Dichloroethane-d4	10.33	65	128013	25.4458	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	101.80%	
58) Toluene-d8	12.71	98	502661	25.9933	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	103.96%	
80) p-Bromofluorobenzene	16.09	95	198337	24.7431	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	98.96%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.23	85	501548	56.4027	ug/L	100
3) Chloromethane	3.69	50	626433	48.9848	ug/L	100
4) Vinyl Chloride	3.91	62	466984	44.8083	ug/L	100
5) 1,3-Butadiene	3.95	54	305611	51.3080	ug/L	99
6) Bromomethane	4.79	94	229789	36.6577	ug/L	100
7) Chloroethane	4.96	64	306769	55.0060	ug/L	99
8) Trichlorofluoromethane	5.43	101	520278	49.1688	ug/L	100
9) Diethyl ether	5.97	59	569381	100.8937	ug/L	99
10) Isoprene	6.00	67	555358	52.5147	ug/L	99
11) Acrolein	6.21	56	78599	121.4613	ug/L	99
12) 1,1,2-Trichloro-1,2,2-Trif	6.21	101	327198	53.6746	ug/L	99
13) Acetone	6.31	43	73026	46.3707	ug/L	98
14) 1,1-Dichloroethene	6.52	61	562302	51.5744	ug/L	100
15) Tert-Butyl Alcohol	6.65	59	97735	192.7992	ug/L	99
16) Dimethyl Sulfide	6.79	62	452865	57.6631	ug/L	98
17) Iodomethane	7.04	142	703007	86.7080	ug/L	99
18) Methyl acetate	7.06	43	204728	50.2043	ug/L	99
19) Methylene Chloride	7.29	84	359603	52.7837	ug/L	98
20) Carbon Disulfide	7.34	76	983785	48.2544	ug/L	100
21) Acrylonitrile	7.49	53	102317	51.7859	ug/L	97
22) Methyl Tert Butyl Ether	7.51	73	786663	48.5115	ug/L	99
23) trans-1,2-Dichloroethene	7.75	61	555437	53.3164	ug/L	100
24) n-Hexane	7.82	57	484655	47.4037	ug/L	99
25) Diisopropyl ether	8.16	45	2682988	104.0331	ug/L	99
26) Vinyl Acetate	8.34	43	348376	36.4146	ug/L	100
27) 1,1-Dichloroethane	8.37	63	713238	52.9604	ug/L	100
28) Ethyl-Tert-Butyl ether	8.74	59	2309943	103.3424	ug/L	100
29) 2-Butanone	8.93	43	112492	48.4905	ug/L	100
30) Propionitrile	9.03	54	71645	100.7667	ug/L	99
31) 2,2-Dichloropropane	9.13	77	490520	50.1968	ug/L	100
32) cis-1,2-Dichloroethene	9.21	96	406093	54.6158	ug/L	99
33) Chloroform	9.41	83	617839	51.6905	ug/L	100
34) 1-Bromopropane	9.55	122	84387	52.7982	ug/L	98
35) Bromochloromethane	9.64	130	226715	53.4633	ug/L	98
36) Tetrahydrofuran	9.67	42	156031	99.9656	ug/L	100
38) 1,1,1-Trichloroethane	9.94	97	566738	54.5893	ug/L	99
39) Cyclohexane	9.97	56	604026	43.6817	ug/L	99
40) 1,1-Dichloropropene	10.14	75	490681	56.2849	ug/L	99
41) Tert-Amyl-Methyl ether	10.24	73	1773558	104.1208	ug/L	100
42) Carbon Tetrachloride	10.28	117	501816	53.1924	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M428967.D 8260WTR.M Tue Mar 05 10:55:33 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428967.D Vial: 13
 Acq On : 4 Mar 2019 18:06 Operator: EEA
 Sample : WG698192-12 ALT 50 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 05 10:55:31 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WT

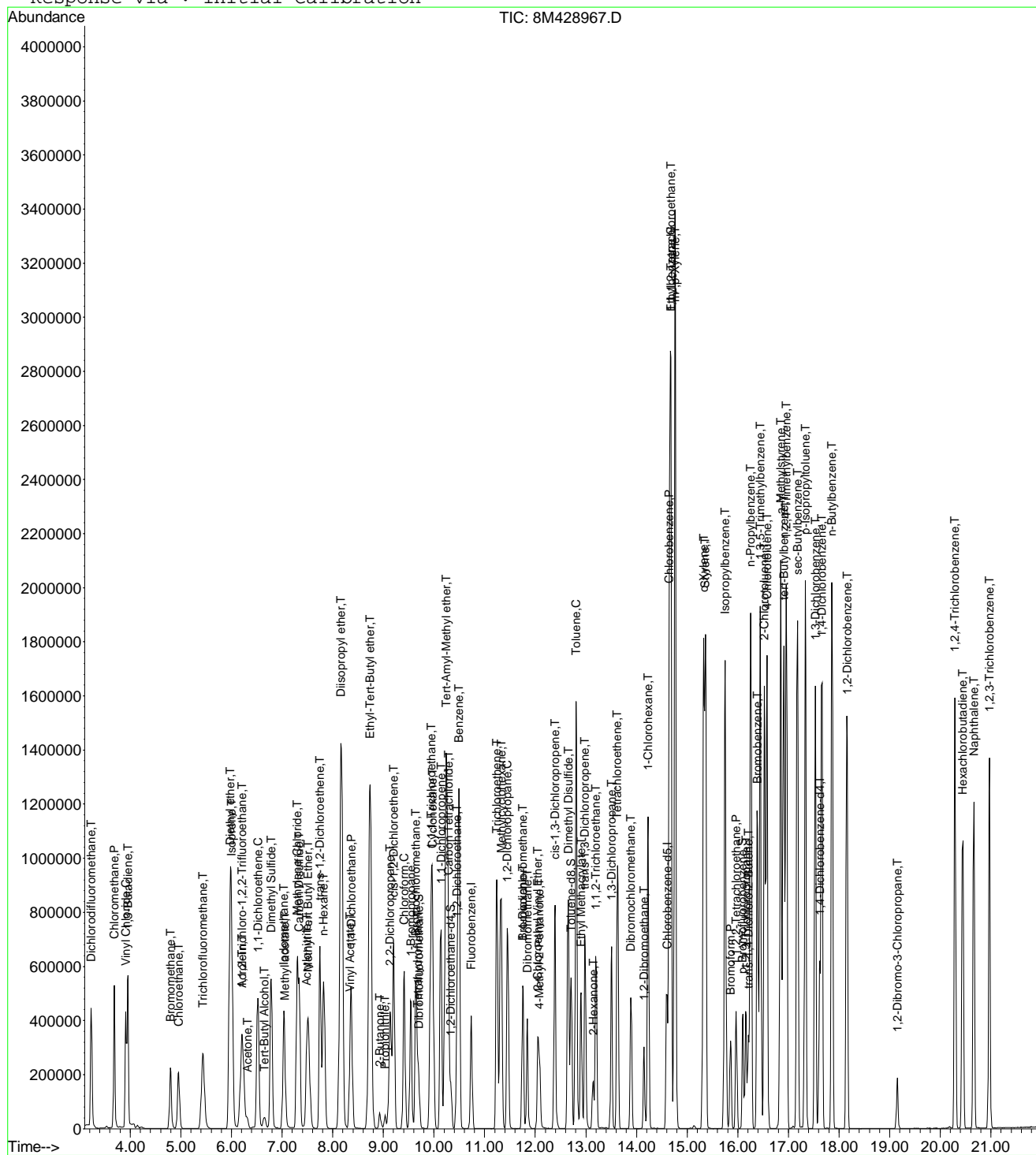
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	437454	53.5072	ug/L	100
46) Benzene	10.49	78	1473129	53.9422	ug/L	99
47) Trichloroethene	11.24	130	423874	56.9695	ug/L	99
48) Methylcyclohexane	11.33	83	548460	47.5982	ug/L	100
49) 1,2-Dichloropropane	11.45	63	403826	54.4772	ug/L	99
50) Bromodichloromethane	11.75	83	446692	52.0435	ug/L	99
51) 1,4-Dioxane	11.75	88	9939	238.2884	ug/L	97
52) Dibromomethane	11.84	93	180349	52.8568	ug/L	99
53) 2-Chloroethyl Vinyl Ether	12.05	63	167899	51.8358	ug/L	99
54) 4-Methyl-2-Pentanone	12.08	58	104920	51.4855	ug/L	99
55) cis-1,3-Dichloropropene	12.39	75	588259	57.6091	ug/L	100
56) Dimethyl Disulfide	12.65	79	322118	51.3697	ug/L	100
59) Toluene	12.80	91	1535304	54.1642	ug/L	100
60) Ethyl Methacrylate	12.91	69	370417	50.3305	ug/L	96
62) trans-1,3-Dichloropropene	12.98	75	461764	54.1873	ug/L	99
63) 1,1,2-Trichloroethane	13.20	97	264840	53.4523	ug/L	98
64) 2-Hexanone	13.15	58	97639	48.4300	ug/L	98
65) 1,3-Dichloropropane	13.51	76	475835	55.7627	ug/L	98
66) Tetrachloroethene	13.62	164	320847	53.7124	ug/L	99
67) Dibromochloromethane	13.89	129	339671	54.0166	ug/L	100
68) 1,2-Dibromoethane	14.15	107	264120	54.8969	ug/L	100
69) 1-Chlorohexane	14.23	91	512372	52.1635	ug/L	99
70) Chlorobenzene	14.65	112	1033323	53.7744	ug/L	99
71) 1,1,1,2-Tetrachloroethane	14.68	131	383135	54.2320	ug/L	99
72) Ethylbenzene	14.68	106	570055	54.8301	ug/L	100
73) m-,p-Xylene	14.76	106	1393290	109.9851	ug/L	99
74) o-Xylene	15.33	106	699921	56.9416	ug/L	99
75) Styrene	15.36	104	1173006	57.6413	ug/L	100
76) Bromoform	15.85	173	217146	55.6921	ug/L	99
77) Isopropylbenzene	15.75	105	1683333	54.2187	ug/L	100
79) 1,1,2,2-Tetrachloroethane	15.97	83	298106	47.5760	ug/L	100
81) 1,2,3-Trichloropropane	16.16	110	97497	52.0140	ug/L	80
82) trans-1,4-Dichloro-2-Butene	16.21	53	99098	46.7843	ug/L	# 19
83) n-Propylbenzene	16.26	91	2083381	51.8331	ug/L	100
84) Bromobenzene	16.39	156	471558	51.2751	ug/L	100
85) 1,3,5-Trimethylbenzene	16.44	105	1492135	51.4978	ug/L	99
86) 2-Chlorotoluene	16.53	91	1237082	46.5487	ug/L	91
87) 4-Chlorotoluene	16.58	91	1319292	56.8140	ug/L	91
88) a-Methylstyrene	16.85	118	906508	54.5225	ug/L	100
89) tert-Butylbenzene	16.91	134	342773	51.2786	ug/L	99
90) 1,2,4-Trimethylbenzene	16.96	105	1535622	52.3495	ug/L	100
91) sec-Butylbenzene	17.18	105	1921041	51.8055	ug/L	100
92) p-Isopropyltoluene	17.33	119	1678164	52.7928	ug/L	100
93) 1,3-Dichlorobenzene	17.53	146	928815	51.8070	ug/L	100
94) 1,4-Dichlorobenzene	17.66	146	951896	52.4293	ug/L	99
95) n-Butylbenzene	17.86	91	1532126	52.8991	ug/L	99
96) 1,2-Dichlorobenzene	18.16	146	901302	54.1743	ug/L	100
97) 1,2-Dibromo-3-Chloropropane	19.15	75	59487	52.6739	ug/L	98
98) 1,2,4-Trichlorobenzene	20.29	180	667869	55.3242	ug/L	100
99) Hexachlorobutadiene	20.45	225	314261	55.2469	ug/L	99
100) Naphthalene	20.66	128	1234867	56.1142	ug/L	100
101) 1,2,3-Trichlorobenzene	20.97	180	605242	54.5770	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M428967.D 8260WTR.M Tue Mar 05 10:55:33 2019

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428967.D Vial: 13
 Acq On : 4 Mar 2019 18:06 Operator: EEA
 Sample : WG698192-12 ALT 50 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 5 10:55 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428967.D Vial: 13
 Acq On : 4 Mar 2019 18:06 Operator: EEA
 Sample : WG698192-12 ALT 50 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	96	0.00
2 T	Dichlorodifluoromethane	50.000	56.403	-12.8	96	0.00
3 P	Chloromethane	50.000	48.985	2.0	96	0.00
4 C	Vinyl Chloride	50.000	44.808	10.4	81	0.00
5 T	1,3-Butadiene	50.000	51.308	-2.6	88	0.00
6 T	Bromomethane	50.000	36.658	26.7#	77	-0.02
7 T	Chloroethane	50.000	55.006	-10.0	99	0.00
8 T	Trichlorofluoromethane	50.000	49.169	1.7	87	0.00
9 T	Diethyl ether	100.000	100.894	-0.9	94	0.00
10 T	Isoprene	50.000	52.515	-5.0	96	0.00
11 T	Acrolein	50.000	121.461	-142.9#	226	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	50.000	53.675	-7.3	96	0.00
13 T	Acetone	50.000	46.371	7.3	83	0.00
14 C	1,1-Dichloroethene	50.000	51.574	-3.1	94	0.00
15 T	Tert-Butyl Alcohol	200.000	192.799	3.6	98	0.02
16 T	Dimethyl Sulfide	50.000	57.663	-15.3	105	0.00
17 T	Iodomethane	50.000	86.708	-73.4#	159	0.00
18 T	Methyl acetate	50.000	50.204	-0.4	94	0.00
19 T	Methylene Chloride	50.000	52.784	-5.6	98	0.00
20 T	Carbon Disulfide	50.000	48.254	3.5	87	0.00
21 T	Acrylonitrile	50.000	51.786	-3.6	94	0.00
22 T	Methyl Tert Butyl Ether	50.000	48.511	3.0	90	0.00
23 T	trans-1,2-Dichloroethene	50.000	53.316	-6.6	97	0.00
24 T	n-Hexane	50.000	47.404	5.2	87	0.00
25 T	Diisopropyl ether	100.000	104.033	-4.0	96	0.00
26 T	Vinyl Acetate	50.000	36.415	27.2#	68	0.00
27 P	1,1-Dichloroethane	50.000	52.960	-5.9	97	0.00
28 T	Ethyl-Tert-Butyl ether	100.000	103.342	-3.3	97	0.00
29 T	2-Butanone	50.000	48.490	3.0	88	0.00
30 T	Propionitrile	100.000	100.767	-0.8	96	0.00
31 T	2,2-Dichloropropane	50.000	50.197	-0.4	93	0.00
32 T	cis-1,2-Dichloroethene	50.000	54.616	-9.2	98	0.00
33 C	Chloroform	50.000	51.690	-3.4	98	0.00
34	1-Bromopropane	50.000	52.798	-5.6	95	0.00
35 T	Bromochloromethane	50.000	53.463	-6.9	96	0.00
36 T	Tetrahydrofuran	100.000	99.966	0.0	95	0.00
37 S	Dibromofluoromethane	25.000	25.648	-2.6	94	0.00
38 T	1,1,1-Trichloroethane	50.000	54.589	-9.2	99	0.00
39 T	Cyclohexane	50.000	43.682	12.6	79	0.00
40 T	1,1-Dichloropropene	50.000	56.285	-12.6	101	0.00
41 T	Tert-Amyl-Methyl ether	100.000	104.121	-4.1	97	0.00
42 T	Carbon Tetrachloride	50.000	53.192	-6.4	96	0.00
43 S	1,2-Dichloroethane-d4	25.000	25.446	-1.8	93	0.00
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	50.000	53.507	-7.0	96	0.00
46 T	Benzene	50.000	53.942	-7.9	98	0.00
47 T	Trichloroethene	50.000	56.970	-13.9	102	0.00
48 T	Methylcyclohexane	50.000	47.598	4.8	87	0.00
49 C	1,2-Dichloropropane	50.000	54.477	-9.0	97	0.00
50 T	Bromodichloromethane	50.000	52.044	-4.1	93	0.00
51 T	1,4-Dioxane	200.000	238.288	-19.1	106	0.00
52 T	Dibromomethane	50.000	52.857	-5.7	97	0.00
53 T	2-Chloroethyl Vinyl Ether	50.000	51.836	-3.7	96	0.00
54 T	4-Methyl-2-Pentanone	50.000	51.486	-3.0	96	0.00

(#) = Out of Range

8M428967.D 8260WTR.M

Tue Mar 05 10:56:46 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428967.D Vial: 13
 Acq On : 4 Mar 2019 18:06 Operator: EEA
 Sample : WG698192-12 ALT 50 ug/L ICAL 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	cis-1,3-Dichloropropene	50.000	57.609	-15.2	101	0.00
56 T	Dimethyl Disulfide	50.000	51.370	-2.7	93	0.00
57 I	Chlorobenzene-d5	25.000	25.000	0.0	95	0.00
58 S	Toluene-d8	25.000	25.993	-4.0	95	0.00
59 C	Toluene	50.000	54.164	-8.3	99	0.00
60 T	Ethyl Methacrylate	50.000	50.330	-0.7	94	0.00
61	Paraldehyde	-1.000	0.000	0.0	0	-13.45#
62 T	trans-1,3-Dichloropropene	50.000	54.187	-8.4	95	0.00
63 T	1,1,2-Trichloroethane	50.000	53.452	-6.9	95	0.00
64 T	2-Hexanone	50.000	48.430	3.1	94	0.00
65 T	1,3-Dichloropropane	50.000	55.763	-11.5	101	0.00
66 T	Tetrachloroethene	50.000	53.712	-7.4	98	0.00
67 T	Dibromochloromethane	50.000	54.017	-8.0	96	0.00
68 T	1,2-Dibromoethane	50.000	54.897	-9.8	97	0.00
69 T	1-Chlorohexane	50.000	52.163	-4.3	94	0.00
70 P	Chlorobenzene	50.000	53.774	-7.5	100	0.00
71 T	1,1,1,2-Tetrachloroethane	50.000	54.232	-8.5	98	0.00
72 C	Ethylbenzene	50.000	54.830	-9.7	98	0.00
73 T	m-,p-Xylene	100.000	109.985	-10.0	100	0.00
74 T	o-Xylene	50.000	56.942	-13.9	101	0.00
75 T	Styrene	50.000	57.641	-15.3	101	0.00
76 P	Bromoform	50.000	55.692	-11.4	97	0.00
77 T	Isopropylbenzene	50.000	54.219	-8.4	97	0.00
78 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	102	0.00
79 P	1,1,2,2-Tetrachloroethane	50.000	47.576	4.8	95	0.00
80 S	p-Bromofluorobenzene	25.000	24.743	1.0	97	0.00
81 T	1,2,3-Trichloropropane	50.000	52.014	-4.0	101	0.00
82 T	trans-1,4-Dichloro-2-Butene	50.000	46.784	6.4	91	0.00
83 T	n-Propylbenzene	50.000	51.833	-3.7	101	0.00
84 T	Bromobenzene	50.000	51.275	-2.5	100	0.00
85 T	1,3,5-Trimethylbenzene	50.000	51.498	-3.0	100	0.00
86 T	2-Chlorotoluene	50.000	46.549	6.9	90	0.00
87 T	4-Chlorotoluene	50.000	56.814	-13.6	114	0.00
88 T	a-Methylstyrene	50.000	54.523	-9.0	104	0.00
89 T	tert-Butylbenzene	50.000	51.279	-2.6	102	0.00
90 T	1,2,4-Trimethylbenzene	50.000	52.349	-4.7	101	0.00
91 T	sec-Butylbenzene	50.000	51.806	-3.6	101	0.00
92 T	p-Isopropyltoluene	50.000	52.793	-5.6	101	0.00
93 T	1,3-Dichlorobenzene	50.000	51.807	-3.6	103	0.00
94 T	1,4-Dichlorobenzene	50.000	52.429	-4.9	105	0.00
95 T	n-Butylbenzene	50.000	52.899	-5.8	101	0.00
96 T	1,2-Dichlorobenzene	50.000	54.174	-8.3	108	0.00
97 T	1,2-Dibromo-3-Chloropropane	50.000	52.674	-5.3	103	0.00
98 T	1,2,4-Trichlorobenzene	50.000	55.324	-10.6	108	0.00
99 T	Hexachlorobutadiene	50.000	55.247	-10.5	107	0.00
100 T	Naphthalene	50.000	56.114	-12.2	107	0.00
101 T	1,2,3-Trichlorobenzene	50.000	54.577	-9.2	107	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 8M428967.D 8260WTR.M Tue Mar 05 10:56:46 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29829.D Vial: 1
 Acq On : 12 Mar 2019 11:21 Operator: KFR
 Sample : WG699218-02 CCV 50ug/kg 8260 Inst : hpms11
 Misc : 1,1 STD92189 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 13:42:14 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3915	96	455861	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0207	117	343984	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8330	152	179518	25.0000	ug/L	0.0000

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3988	111	128745	24.4662	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	97.8648%	
43) 1,2-Dichloroethane-d4	10.0089	65	148139	23.4631	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	93.8524%	
57) Toluene-d8	12.2423	98	433016	25.2948	ug/L	-0.0103
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	101.1792%	
78) p-Bromofluorobenzene	15.4165	95	170218	24.7777	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	99.1108%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1227	85	347532	51.6060	ug/L	98
3) Chloromethane	3.5570	50	446686	34.5348	ug/L	99
4) Vinyl Chloride	3.7741	62	398360	41.2563	ug/L	99
5) 1,3-Butadiene	3.8155	54	168098	50.8291	ug/L	99
6) Bromomethane	4.6426	94	191483	51.7612	ug/L	100
7) Chloroethane	4.7977	64	179275	48.0262	ug/L	100
8) Trichlorofluoromethane	5.2734	101	457735	50.0951	ug/L	99
9) Diethyl ether	5.7904	59	437527	91.5884	ug/L	99
10) Isoprene	5.8214	67	362391	50.3539	ug/L	100
11) Acrolein	6.0178	56	35000	42.1907	ug/L	98
12) 1,1,2-Trichloro-1,2,2-Trif	6.0385	101	237628	53.0155	ug/L	97
13) Acetone	6.1109	43	67721	40.6343	ug/L	97
14) 1,1-Dichloroethene	6.3280	61	435195	49.8874	ug/L	99
15) Tert-Butyl Alcohol	6.4417	59	88246	189.8336	ug/L	98
16) Dimethyl Sulfide	6.5865	62	275244	47.7284	ug/L	99
17) Iodomethane	6.8346	142	268569	44.8924	ug/L	98
18) Methyl acetate	6.8450	43	157396	43.7837	ug/L	99
19) Methylene Chloride	7.0931	84	233883	48.1109	ug/L	97
20) Carbon Disulfide	7.1242	76	702771	52.4508	ug/L	99
21) Acrylonitrile	7.2689	53	84183	45.8203	ug/L	100
22) Methyl Tert Butyl Ether	7.2896	73	620372	45.0036	ug/L	99
23) trans-1,2-Dichloroethene	7.5274	96	244903	51.8258	ug/L	98
24) n-Hexane	7.5998	57	415691	57.1123	ug/L	99
25) Diisopropyl ether	7.9306	45	1741351	98.0258	ug/L	99
26) Vinyl Acetate	8.0961	43	380293	59.4653	ug/L	98
27) 1,1-Dichloroethane	8.1167	63	496400	48.1401	ug/L	98
28) Ethyl-Tert-Butyl ether	8.4786	59	1718516	95.7107	ug/L	99
29) 2-Butanone	8.6544	43	86859	40.9158	ug/L	97
30) Propionitrile	8.7578	54	55390	89.3043	ug/L	96
31) 2,2-Dichloropropane	8.8612	77	411617	57.7328	ug/L	100
32) cis-1,2-Dichloroethene	8.9232	96	271987	50.6984	ug/L	100
33) Chloroform	9.1197	83	448391	47.9085	ug/L	98
34) 1-Bromopropane	9.2541	122	57645	49.6081	ug/L	97
35) Bromochloromethane	9.3472	130	159466	50.6942	ug/L	97
36) Tetrahydrofuran	9.3678	42	110016	79.9917	ug/L	98
38) 1,1,1-Trichloroethane	9.6263	97	439122	51.8165	ug/L	98
39) Cyclohexane	9.6573	56	499500	49.7840	ug/L	98
40) 1,1-Dichloropropene	9.8124	75	325275	51.0491	ug/L	99
41) Carbon Tetrachloride	9.9572	117	410930	57.0223	ug/L	99
42) Tert-Amyl-Methyl ether	9.9158	73	1258790	93.9953	ug/L	98

(#) = qualifier out of range (m) = manual integration
 11M29829.D 8260WT.M Tue Mar 12 13:42:14 2019

Data File : C:\MSDCHEM\1\DATA\031219\11M29829.D Vial: 1
 Acq On : 12 Mar 2019 11:21 Operator: KFR
 Sample : WG699218-02 CCV 50ug/kg 8260 Inst : hpms11
 Misc : 1,1 STD92189 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 13:42:14 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1226	62	377048	46.3511	ug/L	100
45) Benzene	10.1537	78	928833	49.0436	ug/L	100
46) Trichloroethene	10.8671	130	265666	49.4592	ug/L	98
47) Methylcyclohexane	10.9395	83	405862	53.6358	ug/L	98
48) 1,2-Dichloropropane	11.0635	63	269795	47.9565	ug/L	99
49) 1,4-Dioxane	11.3324	88	7715	199.8606	ug/L	98
50) Bromodichloromethane	11.3530	83	342923	51.5779	ug/L	100
51) Dibromomethane	11.4358	93	148201	46.1169	ug/L	96
52) 2-Chloroethyl Vinyl Ether	11.6219	63	103727	38.1001	ug/L	98
53) 4-Methyl-2-Pentanone	11.6529	58	77324	39.8658	ug/L	97
54) cis-1,3-Dichloropropene	11.9424	75	386178	51.5370	ug/L	99
55) Dimethyl Disulfide	12.1906	79	234103	46.5621	ug/L	100
58) Toluene	12.3456	91	1018915	50.0030	ug/L	100
59) Ethyl Methacrylate	12.4284	69	265749	46.1735	ug/L	99
60) trans-1,3-Dichloropropene	12.5007	75	337594	50.6007	ug/L	99
61) 1,1,2-Trichloroethane	12.7075	97	194151	44.7564	ug/L	100
62) 2-Hexanone	12.6455	43	128963	39.9969	ug/L	97
63) 1,3-Dichloropropane	12.9970	76	320378	45.7749	ug/L	97
64) Tetrachloroethene	13.1108	164	225193	52.1009	ug/L	98
65) Dibromochloromethane	13.3589	129	254619	51.5617	ug/L	99
66) 1,2-Dibromoethane	13.5967	107	193407	46.5207	ug/L	100
67) 1-Chlorohexane	13.6691	91	365194	54.2842	ug/L	96
68) Chlorobenzene	14.0620	112	703358	49.5788	ug/L	100
69) 1,1,1,2-Tetrachloroethane	14.0930	131	290268	48.1685	ug/L	99
70) Ethylbenzene	14.0827	106	375317	52.0044	ug/L	98
71) m-,p-Xylene	14.1654	106	895859	102.8208	ug/L	96
72) o-Xylene	14.6927	106	437288	51.7275	ug/L	98
73) Styrene	14.7237	104	735525	52.6009	ug/L	99
74) Bromoform	15.1994	173	165392	46.5121	ug/L	100
75) Isopropylbenzene	15.0856	105	1106395	52.2805	ug/L	100
77) 1,1,2,2-Tetrachloroethane	15.2924	83	219666	46.9064	ug/L	99
79) 1,2,3-Trichloropropane	15.4682	110	68203	43.6169	ug/L	97
80) trans-1,4-Dichloro-2-Butene	15.5096	53	89614	45.4533	ug/L	97
81) n-Propylbenzene	15.5612	91	1356466	51.2385	ug/L	98
82) Bromobenzene	15.6853	156	315390	48.7822	ug/L	97
83) 1,3,5-Trimethylbenzene	15.7370	105	966595	51.4315	ug/L	99
84) 2-Chlorotoluene	15.8197	91	934074	49.5850	ug/L	100
85) 4-Chlorotoluene	15.8611	91	755132	48.7728	ug/L	99
86) a-Methylstyrene	16.1092	118	555925	51.9487	ug/L	99
87) tert-Butylbenzene	16.1610	134	225571	48.7309	ug/L	96
88) 1,2,4-Trimethylbenzene	16.2126	105	984397	50.7502	ug/L	99
89) sec-Butylbenzene	16.4194	105	1204516	51.6246	ug/L	99
90) p-Isopropyltoluene	16.5642	119	1073109	52.8930	ug/L	99
91) 1,3-Dichlorobenzene	16.7503	146	611172	51.2437	ug/L	99
92) 1,4-Dichlorobenzene	16.8640	146	607155	50.9789	ug/L	100
93) n-Butylbenzene	17.0501	91	1009476	55.2105	ug/L	99
94) 1,2-Dichlorobenzene	17.3293	146	567323	49.6317	ug/L	98
95) 1,2-Dibromo-3-Chloropropane	18.2495	75	41458	42.8531	ug/L	99
96) 1,2,4-Trichlorobenzene	19.3145	180	382644	51.7479	ug/L	99
97) Hexachlorobutadiene	19.4489	225	209207	56.1342	ug/L	97
98) Naphthalene	19.6557	128	724385	45.7494	ug/L	100
99) 1,2,3-Trichlorobenzene	19.9452	180	343620	49.4621	ug/L	99

(#) = qualifier out of range (m) = manual integration
 11M29829.D 8260WT.M Tue Mar 12 13:42:14 2019

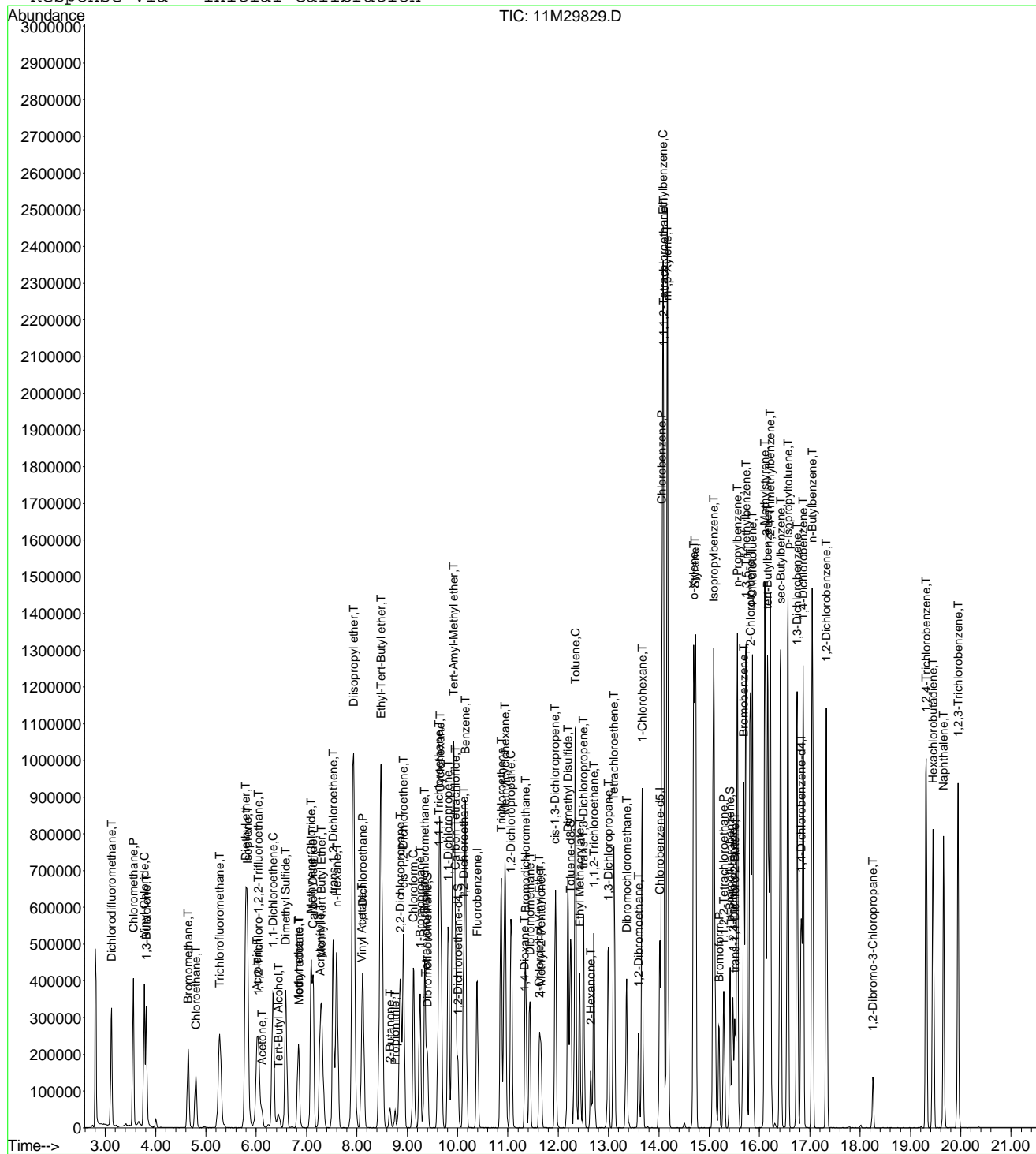
Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29829.D
Acq On : 12 Mar 2019 11:21
Sample : WG699218-02 CCV 50ug/kg 8260
Misc : 1,1 STD92189
MS Integration Params: rteint.p
Quant Time: Mar 12 13:42 2019

Vial: 1
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:13:24 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\031219\11M29829.D Vial: 1
 Acq On : 12 Mar 2019 11:21 Operator: KFR
 Sample : WG699218-02 CCV 50ug/kg 8260 Inst : hpms11
 Misc : 1,1 STD92189 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	1.000	1.000	0.0	115	0.00
2 T	Dichlorodifluoromethane	0.369	0.381	-3.3	109	0.00
3 P	Chloromethane	0.709	0.490	30.9#	83	0.00
4 C	Vinyl Chloride	0.530	0.437	17.5	93	0.00
5 T	1,3-Butadiene	0.290	0.184	36.6#	105	0.00
6 T	Bromomethane	0.203	0.210	-3.4	130	0.00
7 T	Chloroethane	0.205	0.197	3.9	106	0.00
8 T	Trichlorofluoromethane	0.501	0.502	-0.2	110	0.00
9 T	Diethyl ether	0.262	0.240	8.4	116	0.00
10 T	Isoprene	0.395	0.397	-0.5	112	0.00
11 T	Acrolein	0.045	0.038	15.6	99	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	0.246	0.261	-6.1	115	0.00
13 T	Acetone	0.091	0.074	18.7	92	-0.01
14 C	1,1-Dichloroethene	0.478	0.477	0.2	109	0.00
15 T	Tert-Butyl Alcohol	0.025	0.024	4.0	110	0.00
16 T	Dimethyl Sulfide	0.316	0.302	4.4	106	0.00
17 T	Iodomethane	0.273	0.295	-8.1	102	0.00
18 T	Methyl acetate	0.197	0.173	12.2	98	0.00
19 T	Methylene Chloride	0.267	0.257	3.7	108	0.00
20 T	Carbon Disulfide	0.735	0.771	-4.9	112	-0.01
21 T	Acrylonitrile	0.101	0.092	8.9	97	0.00
22 T	Methyl Tert Butyl Ether	0.756	0.680	10.1	100	-0.01
23 T	trans-1,2-Dichloroethene	0.259	0.269	-3.9	111	0.00
24 T	n-Hexane	0.399	0.456	-14.3	127	0.00
25 T	Diisopropyl ether	0.974	0.955	2.0	121	0.00
26 T	Vinyl Acetate	0.328	0.417	-27.1#	138	0.00
27 P	1,1-Dichloroethane	0.566	0.544	3.9	107	-0.01
28 T	Ethyl-Tert-Butyl ether	0.985	0.942	4.4	116	0.00
29 T	2-Butanone	0.116	0.095	18.1	90	0.00
30 T	Propionitrile	0.034	0.030	11.8	94	0.00
31 T	2,2-Dichloropropane	0.391	0.451	-15.3	126	0.00
32 T	cis-1,2-Dichloroethene	0.294	0.298	-1.4	109	-0.01
33 C	Chloroform	0.513	0.492	4.1	109	-0.01
34 T	1-Bromopropane	0.064	0.063	1.6	112	0.00
35 T	Bromochloromethane	0.173	0.175	-1.2	107	0.00
36 T	Tetrahydrofuran	0.075	0.060	20.0	96	0.00
37 S	Dibromofluoromethane	0.289	0.282	2.4	108	0.00
38 T	1,1,1-Trichloroethane	0.465	0.482	-3.7	112	0.00
39 T	Cyclohexane	0.550	0.548	0.4	110	0.00
40 T	1,1-Dichloropropene	0.349	0.357	-2.3	111	0.00
41 T	Carbon Tetrachloride	0.395	0.451	-14.2	118	0.00
42 T	Tert-Amyl-Methyl ether	0.734	0.690	6.0	112	0.00
43 S	1,2-Dichloroethane-d4	0.346	0.325	6.1	104	0.00
44 T	1,2-Dichloroethane	0.446	0.414	7.2	101	0.00
45 T	Benzene	1.039	1.019	1.9	108	0.00
46 T	Trichloroethene	0.295	0.291	1.4	108	0.00
47 T	Methylcyclohexane	0.415	0.445	-7.2	119	0.00
48 C	1,2-Dichloropropane	0.309	0.296	4.2	106	0.00
49 T	1,4-Dioxane	0.002	0.002	0.0	112	-0.01
50 T	Bromodichloromethane	0.365	0.376	-3.0	106	0.00
51 T	Dibromomethane	0.176	0.163	7.4	100	0.00
52 T	2-Chloroethyl Vinyl Ether	0.149	0.114	23.5	86	0.00
53 T	4-Methyl-2-Pentanone	0.106	0.085	19.8	88	0.00
54 T	cis-1,3-Dichloropropene	0.411	0.424	-3.2	108	0.00

(#) = Out of Range

11M29829.D 8260WT.M Tue Mar 12 13:42:23 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\031219\11M29829.D Vial: 1
 Acq On : 12 Mar 2019 11:21 Operator: KFR
 Sample : WG699218-02 CCV 50ug/kg 8260 Inst : hpms11
 Misc : 1,1 STD92189 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	0.251	0.257	-2.4	111	0.00
56 I	Chlorobenzene-d5	1.000	1.000	0.0	115	0.00
57 S	Toluene-d8	1.244	1.259	-1.2	112	-0.01
58 C	Toluene	1.481	1.481	0.0	108	0.00
59 T	Ethyl Methacrylate	0.418	0.386	7.7	97	0.00
60 T	trans-1,3-Dichloropropene	0.485	0.491	-1.2	105	0.00
61 T	1,1,2-Trichloroethane	0.315	0.282	10.5	100	0.00
62 T	2-Hexanone	0.234	0.187	20.1	85	0.00
63 T	1,3-Dichloropropane	0.509	0.466	8.4	100	0.00
64 T	Tetrachloroethene	0.314	0.327	-4.1	115	0.00
65 T	Dibromochloromethane	0.359	0.370	-3.1	108	0.00
66 T	1,2-Dibromoethane	0.302	0.281	7.0	100	0.00
67 T	1-Chlorohexane	0.489	0.531	-8.6	117	0.00
68 P	Chlorobenzene	1.031	1.022	0.9	108	0.00
69 T	1,1,1,2-Tetrachloroethane	0.405	0.422	-4.2	110	0.00
70 C	Ethylbenzene	0.525	0.546	-4.0	109	0.00
71 T	m-,p-Xylene	0.633	0.651	-2.8	109	0.00
72 T	o-Xylene	0.614	0.636	-3.6	109	0.00
73 T	Styrene	1.016	1.069	-5.2	107	-0.01
74 P	Bromoform	0.214	0.240	-12.1	106	0.00
75 T	Isopropylbenzene	1.538	1.608	-4.6	109	0.00
76 I	1,4-Dichlorobenzene-d4	1.000	1.000	0.0	120	0.00
77 P	1,1,2,2-Tetrachloroethane	0.652	0.612	6.1	98	0.00
78 S	p-Bromofluorobenzene	0.957	0.948	0.9	110	0.00
79 T	1,2,3-Trichloropropane	0.218	0.190	12.8	95	0.00
80 T	trans-1,4-Dichloro-2-Butene	0.275	0.250	9.1	98	0.00
81 T	n-Propylbenzene	3.687	3.778	-2.5	110	0.00
82 T	Bromobenzene	0.900	0.878	2.4	107	0.00
83 T	1,3,5-Trimethylbenzene	2.617	2.692	-2.9	110	0.00
84 T	2-Chlorotoluene	2.623	2.602	0.8	110	0.00
85 T	4-Chlorotoluene	2.156	2.103	2.5	107	0.00
86 T	a-Methylstyrene	1.490	1.548	-3.9	112	0.00
87 T	tert-Butylbenzene	0.645	0.628	2.6	112	-0.01
88 T	1,2,4-Trimethylbenzene	2.701	2.742	-1.5	109	0.00
89 T	sec-Butylbenzene	3.249	3.355	-3.3	110	0.00
90 T	p-Isopropyltoluene	2.825	2.989	-5.8	112	0.00
91 T	1,3-Dichlorobenzene	1.661	1.702	-2.5	112	0.00
92 T	1,4-Dichlorobenzene	1.659	1.691	-1.9	110	0.00
93 T	n-Butylbenzene	2.546	2.812	-10.4	114	0.00
94 T	1,2-Dichlorobenzene	1.592	1.580	0.8	108	0.00
95 T	1,2-Dibromo-3-Chloropropane	0.135	0.115	14.8	92	0.00
96 T	1,2,4-Trichlorobenzene	1.030	1.066	-3.5	111	0.00
97 T	Hexachlorobutadiene	0.519	0.583	-12.3	123	0.00
98 T	Naphthalene	2.205	2.018	8.5	95	0.00
99 T	1,2,3-Trichlorobenzene	0.967	0.957	1.0	106	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29829.D 8260WT.M Tue Mar 12 13:42:23 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29829.D Vial: 1
 Acq On : 12 Mar 2019 11:21 Operator: KFR
 Sample : WG699218-02 CCV 50ug/kg 8260 Inst : hpms11
 Misc : 1,1 STD92189 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	115	0.00
2 T	Dichlorodifluoromethane	50.000	51.606	-3.2	109	0.00
3 P	Chloromethane	50.000	34.535	30.9#	83	0.00
4 C	Vinyl Chloride	50.000	41.256	17.5	93	0.00
5 T	1,3-Butadiene	50.000	50.829	-1.7	105	0.00
6 T	Bromomethane	50.000	51.761	-3.5	130	0.00
7 T	Chloroethane	50.000	48.026	3.9	106	0.00
8 T	Trichlorofluoromethane	50.000	50.095	-0.2	110	0.00
9 T	Diethyl ether	100.000	91.588	8.4	116	0.00
10 T	Isoprene	50.000	50.354	-0.7	112	0.00
11 T	Acrolein	50.000	42.191	15.6	99	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	50.000	53.016	-6.0	115	0.00
13 T	Acetone	50.000	40.634	18.7	92	-0.01
14 C	1,1-Dichloroethene	50.000	49.887	0.2	109	0.00
15 T	Tert-Butyl Alcohol	200.000	189.834	5.1	110	0.00
16 T	Dimethyl Sulfide	50.000	47.728	4.5	106	0.00
17 T	Iodomethane	50.000	44.892	10.2	102	0.00
18 T	Methyl acetate	50.000	43.784	12.4	98	0.00
19 T	Methylene Chloride	50.000	48.111	3.8	108	0.00
20 T	Carbon Disulfide	50.000	52.451	-4.9	112	-0.01
21 T	Acrylonitrile	50.000	45.820	8.4	97	0.00
22 T	Methyl Tert Butyl Ether	50.000	45.004	10.0	100	-0.01
23 T	trans-1,2-Dichloroethene	50.000	51.826	-3.7	111	0.00
24 T	n-Hexane	50.000	57.112	-14.2	127	0.00
25 T	Diisopropyl ether	100.000	98.026	2.0	121	0.00
26 T	Vinyl Acetate	50.000	59.465	-18.9	138	0.00
27 P	1,1-Dichloroethane	50.000	48.140	3.7	107	-0.01
28 T	Ethyl-Tert-Butyl ether	100.000	95.711	4.3	116	0.00
29 T	2-Butanone	50.000	40.916	18.2	90	0.00
30 T	Propionitrile	100.000	89.304	10.7	94	0.00
31 T	2,2-Dichloropropane	50.000	57.733	-15.5	126	0.00
32 T	cis-1,2-Dichloroethene	50.000	50.698	-1.4	109	-0.01
33 C	Chloroform	50.000	47.909	4.2	109	-0.01
34 T	1-Bromopropane	50.000	49.608	0.8	112	0.00
35 T	Bromochloromethane	50.000	50.694	-1.4	107	0.00
36 T	Tetrahydrofuran	100.000	79.992	20.0	96	0.00
37 S	Dibromofluoromethane	25.000	24.466	2.1	108	0.00
38 T	1,1,1-Trichloroethane	50.000	51.816	-3.6	112	0.00
39 T	Cyclohexane	50.000	49.784	0.4	110	0.00
40 T	1,1-Dichloropropene	50.000	51.049	-2.1	111	0.00
41 T	Carbon Tetrachloride	50.000	57.022	-14.0	118	0.00
42 T	Tert-Amyl-Methyl ether	100.000	93.995	6.0	112	0.00
43 S	1,2-Dichloroethane-d4	25.000	23.463	6.1	104	0.00
44 T	1,2-Dichloroethane	50.000	46.351	7.3	101	0.00
45 T	Benzene	50.000	49.044	1.9	108	0.00
46 T	Trichloroethene	50.000	49.459	1.1	108	0.00
47 T	Methylcyclohexane	50.000	53.636	-7.3	119	0.00
48 C	1,2-Dichloropropane	50.000	47.956	4.1	106	0.00
49 T	1,4-Dioxane	200.000	199.861	0.1	112	-0.01
50 T	Bromodichloromethane	50.000	51.578	-3.2	106	0.00
51 T	Dibromomethane	50.000	46.117	7.8	100	0.00
52 T	2-Chloroethyl Vinyl Ether	50.000	38.100	23.8	86	0.00
53 T	4-Methyl-2-Pentanone	50.000	39.866	20.3	88	0.00
54 T	cis-1,3-Dichloropropene	50.000	51.537	-3.1	108	0.00

(#) = Out of Range

11M29829.D 8260WT.M Tue Mar 12 13:42:21 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\031219\11M29829.D Vial: 1
 Acq On : 12 Mar 2019 11:21 Operator: KFR
 Sample : WG699218-02 CCV 50ug/kg 8260 Inst : hpms11
 Misc : 1,1 STD92189 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	50.000	46.562	6.9	111	0.00
56 I	Chlorobenzene-d5	25.000	25.000	0.0	115	0.00
57 S	Toluene-d8	25.000	25.295	-1.2	112	-0.01
58 C	Toluene	50.000	50.003	-0.0	108	0.00
59 T	Ethyl Methacrylate	50.000	46.173	7.7	97	0.00
60 T	trans-1,3-Dichloropropene	50.000	50.601	-1.2	105	0.00
61 T	1,1,2-Trichloroethane	50.000	44.756	10.5	100	0.00
62 T	2-Hexanone	50.000	39.997	20.0	85	0.00
63 T	1,3-Dichloropropane	50.000	45.775	8.5	100	0.00
64 T	Tetrachloroethene	50.000	52.101	-4.2	115	0.00
65 T	Dibromochloromethane	50.000	51.562	-3.1	108	0.00
66 T	1,2-Dibromoethane	50.000	46.521	7.0	100	0.00
67 T	1-Chlorohexane	50.000	54.284	-8.6	117	0.00
68 P	Chlorobenzene	50.000	49.579	0.8	108	0.00
69 T	1,1,1,2-Tetrachloroethane	50.000	48.168	3.7	110	0.00
70 C	Ethylbenzene	50.000	52.004	-4.0	109	0.00
71 T	m-,p-Xylene	100.000	102.821	-2.8	109	0.00
72 T	o-Xylene	50.000	51.728	-3.5	109	0.00
73 T	Styrene	50.000	52.601	-5.2	107	-0.01
74 P	Bromoform	50.000	46.512	7.0	106	0.00
75 T	Isopropylbenzene	50.000	52.280	-4.6	109	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	120	0.00
77 P	1,1,2,2-Tetrachloroethane	50.000	46.906	6.2	98	0.00
78 S	p-Bromofluorobenzene	25.000	24.778	0.9	110	0.00
79 T	1,2,3-Trichloropropane	50.000	43.617	12.8	95	0.00
80 T	trans-1,4-Dichloro-2-Butene	50.000	45.453	9.1	98	0.00
81 T	n-Propylbenzene	50.000	51.239	-2.5	110	0.00
82 T	Bromobenzene	50.000	48.782	2.4	107	0.00
83 T	1,3,5-Trimethylbenzene	50.000	51.431	-2.9	110	0.00
84 T	2-Chlorotoluene	50.000	49.585	0.8	110	0.00
85 T	4-Chlorotoluene	50.000	48.773	2.5	107	0.00
86 T	a-Methylstyrene	50.000	51.949	-3.9	112	0.00
87 T	tert-Butylbenzene	50.000	48.731	2.5	112	-0.01
88 T	1,2,4-Trimethylbenzene	50.000	50.750	-1.5	109	0.00
89 T	sec-Butylbenzene	50.000	51.625	-3.3	110	0.00
90 T	p-Isopropyltoluene	50.000	52.893	-5.8	112	0.00
91 T	1,3-Dichlorobenzene	50.000	51.244	-2.5	112	0.00
92 T	1,4-Dichlorobenzene	50.000	50.979	-2.0	110	0.00
93 T	n-Butylbenzene	50.000	55.210	-10.4	114	0.00
94 T	1,2-Dichlorobenzene	50.000	49.632	0.7	108	0.00
95 T	1,2-Dibromo-3-Chloropropane	50.000	42.853	14.3	92	0.00
96 T	1,2,4-Trichlorobenzene	50.000	51.748	-3.5	111	0.00
97 T	Hexachlorobutadiene	50.000	56.134	-12.3	123	0.00
98 T	Naphthalene	50.000	45.749	8.5	95	0.00
99 T	1,2,3-Trichlorobenzene	50.000	49.462	1.1	106	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29829.D 8260WT.M Tue Mar 12 13:42:21 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429061.D Vial: 1
 Acq On : 11 Mar 2019 9:19 Operator: EEA
 Sample : WG699027-02 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:31:09 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	591011	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.59	117	447411	25.00	ug/L	-0.01
78) 1,4-Dichlorobenzene-d4	17.62	152	249471	25.00	ug/L	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.69	111	153213	26.0071	ug/L	-0.01
Spiked Amount	25.000	Range 86 - 118	Recovery	=	104.04%	
43) 1,2-Dichloroethane-d4	10.33	65	147296	25.3265	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	101.32%	
58) Toluene-d8	12.70	98	589200	25.8531	ug/L	-0.01
Spiked Amount	25.000	Range 88 - 110	Recovery	=	103.40%	
80) p-Bromofluorobenzene	16.09	95	233373	25.9198	ug/L	-0.01
Spiked Amount	25.000	Range 86 - 115	Recovery	=	103.68%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.22	85	484597	47.1402	ug/L	99
3) Chloromethane	3.66	50	622234	42.0886	ug/L	100
4) Vinyl Chloride	3.90	62	561164	46.5768	ug/L	100
5) 1,3-Butadiene	3.94	54	351416	51.0076	ug/L	100
6) Bromomethane	4.79	94	291549	40.2320	ug/L	99
7) Chloroethane	4.94	64	300895	46.6699	ug/L	99
8) Trichlorofluoromethane	5.43	101	579559	47.3778	ug/L	100
9) Diethyl ether	5.95	59	583015	89.3644	ug/L	98
10) Isoprene	5.99	67	570222	46.6418	ug/L	100
11) Acrolein	6.19	56	34331	45.8913	ug/L	93
12) 1,1,2-Trichloro-1,2,2-Trif	6.21	101	336896	47.8054	ug/L	100
13) Acetone	6.29	43	72788	39.9806	ug/L	99
14) 1,1-Dichloroethene	6.51	61	587248	46.5918	ug/L	100
15) Tert-Butyl Alcohol	6.62	59	89803	153.2390	ug/L	98
16) Dimethyl Sulfide	6.78	62	413363	45.5286	ug/L	98
17) Iodomethane	7.02	142	418455	45.3783	ug/L	99
18) Methyl acetate	7.03	43	219768	46.6178	ug/L	100
19) Methylene Chloride	7.29	84	362036	45.9675	ug/L	97
20) Carbon Disulfide	7.33	76	1108544	47.0341	ug/L	100
21) Acrylonitrile	7.48	53	106664	46.6987	ug/L	96
22) Methyl Tert Butyl Ether	7.50	73	830155	44.2831	ug/L	99
23) trans-1,2-Dichloroethene	7.74	61	568219	47.1808	ug/L	100
24) n-Hexane	7.81	57	611845	51.7659	ug/L	98
25) Diisopropyl ether	8.15	45	2793073	93.6825	ug/L	99
26) Vinyl Acetate	8.33	43	637737	57.6624	ug/L	99
27) 1,1-Dichloroethane	8.36	63	724898	46.5605	ug/L	100
28) Ethyl-Tert-Butyl ether	8.73	59	2297647	88.9168	ug/L	99
29) 2-Butanone	8.92	43	119054	44.3917	ug/L	98
30) Propionitrile	9.02	54	69393	85.1558	ug/L	98
31) 2,2-Dichloropropane	9.13	77	566727	50.1668	ug/L	100
32) cis-1,2-Dichloroethene	9.19	96	408738	47.5512	ug/L	98
33) Chloroform	9.40	83	626567	45.3446	ug/L	100
34) 1-Bromopropane	9.54	122	85201	46.1117	ug/L	99
35) Bromochloromethane	9.64	130	229324	46.7787	ug/L	98
36) Tetrahydrofuran	9.66	42	159421	88.3504	ug/L	98
38) 1,1,1-Trichloroethane	9.93	97	563433	46.9452	ug/L	99
39) Cyclohexane	9.96	56	767077	47.9851	ug/L	99
40) 1,1-Dichloropropene	10.13	75	482944	47.9195	ug/L	99
41) Tert-Amyl-Methyl ether	10.23	73	1730185	87.8634	ug/L	100
42) Carbon Tetrachloride	10.27	117	516382	47.3477	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M429061.D 8260WTR.M Mon Mar 11 15:31:10 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429061.D Vial: 1
 Acq On : 11 Mar 2019 9:19 Operator: EEA
 Sample : WG699027-02 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:31:09 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

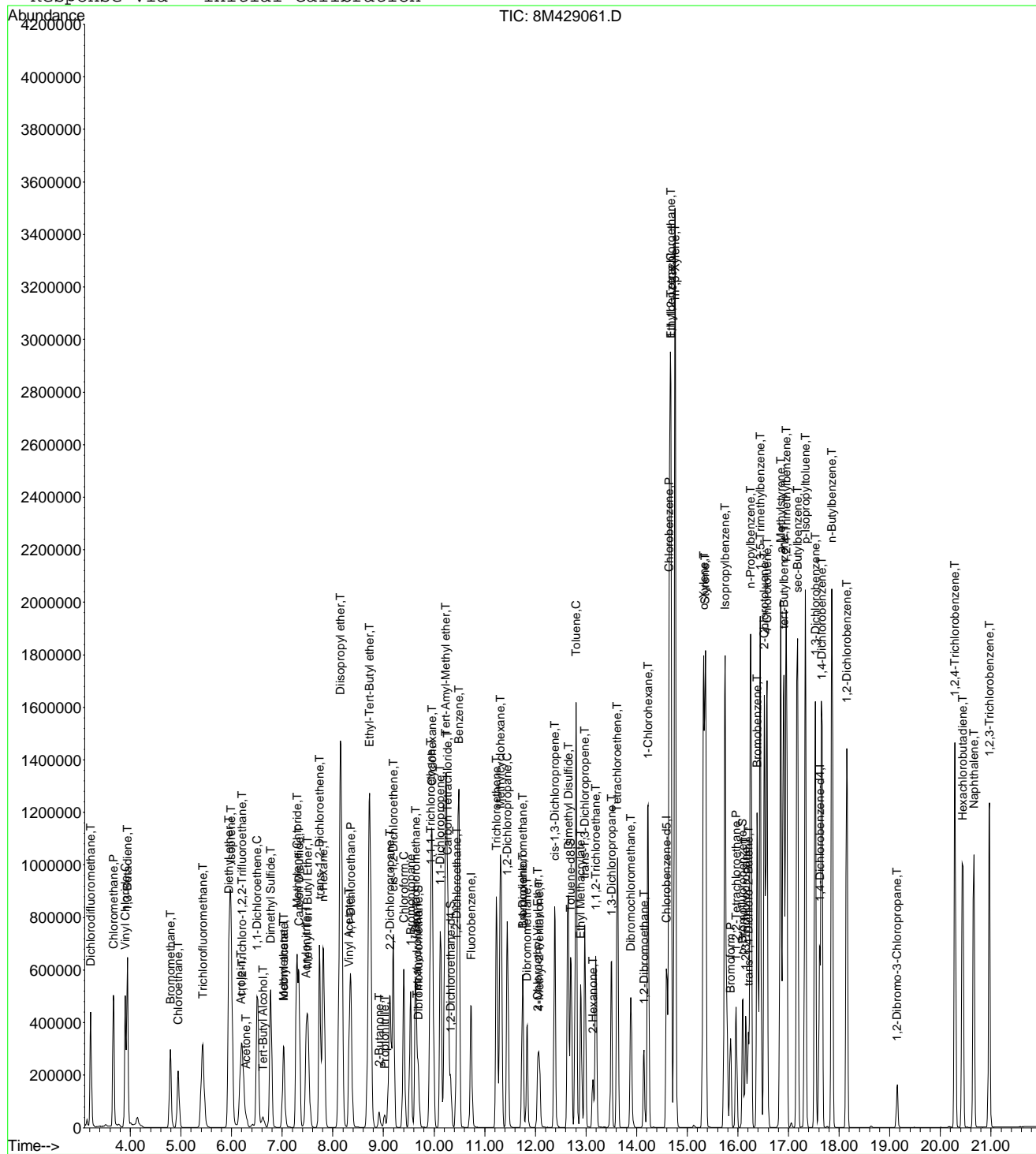
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.45	62	442131	46.7793	ug/L	99
46) Benzene	10.49	78	1485603	47.0559	ug/L	99
47) Trichloroethene	11.23	130	393302	45.7252	ug/L	99
48) Methylcyclohexane	11.31	83	640613	48.0911	ug/L	99
49) 1,2-Dichloropropane	11.45	63	411172	47.9808	ug/L	99
50) Bromodichloromethane	11.75	83	476354	48.0078	ug/L	99
51) 1,4-Dioxane	11.75	88	8839	183.3102	ug/L	93
52) Dibromomethane	11.83	93	180898	45.8611	ug/L	100
53) 2-Chloroethyl Vinyl Ether	12.05	63	115392	30.8163	ug/L	99
54) 4-Methyl-2-Pentanone	12.08	58	107553	45.6534	ug/L	99
55) cis-1,3-Dichloropropene	12.38	75	585524	49.6010	ug/L	100
56) Dimethyl Disulfide	12.65	79	331697	45.7570	ug/L	98
59) Toluene	12.80	91	1541093	46.1329	ug/L	100
60) Ethyl Methacrylate	12.90	69	370280	42.7633	ug/L	98
62) trans-1,3-Dichloropropene	12.98	75	483112	48.1049	ug/L	100
63) 1,1,2-Trichloroethane	13.20	97	268007	45.8979	ug/L	100
64) 2-Hexanone	13.13	58	98638	41.9002	ug/L	98
65) 1,3-Dichloropropane	13.51	76	460455	45.7866	ug/L	99
66) Tetrachloroethene	13.62	164	326095	46.3218	ug/L	99
67) Dibromochloromethane	13.89	129	345451	46.6143	ug/L	100
68) 1,2-Dibromoethane	14.15	107	263514	46.4745	ug/L	99
69) 1-Chlorohexane	14.23	91	541900	46.8128	ug/L	98
70) Chlorobenzene	14.64	112	1022326	45.1433	ug/L	99
71) 1,1,1,2-Tetrachloroethane	14.67	131	387635	46.5577	ug/L	99
72) Ethylbenzene	14.67	106	584603	47.7120	ug/L	100
73) m-,p-Xylene	14.76	106	1397279	93.5922	ug/L	100
74) o-Xylene	15.33	106	694695	47.9556	ug/L	99
75) Styrene	15.36	104	1146408	47.8010	ug/L	99
76) Bromoform	15.85	173	217491	47.3312	ug/L	99
77) Isopropylbenzene	15.75	105	1740410	47.5658	ug/L	100
79) 1,1,2,2-Tetrachloroethane	15.97	83	314504	44.6863	ug/L	100
81) 1,2,3-Trichloropropane	16.15	110	91112	43.2748	ug/L	97
82) trans-1,4-Dichloro-2-Butene	16.20	53	103594	43.5412	ug/L	87
83) n-Propylbenzene	16.26	91	2077693	46.0204	ug/L	100
84) Bromobenzene	16.38	156	464774	44.9929	ug/L	98
85) 1,3,5-Trimethylbenzene	16.44	105	1490512	45.7980	ug/L	100
86) 2-Chlorotoluene	16.53	91	1254981	42.0414	ug/L	99
87) 4-Chlorotoluene	16.58	91	1298396	49.7797	ug/L	99
88) a-Methylstyrene	16.85	118	859535	46.0255	ug/L	99
89) tert-Butylbenzene	16.91	134	336065	44.7593	ug/L	98
90) 1,2,4-Trimethylbenzene	16.96	105	1516412	46.0231	ug/L	100
91) sec-Butylbenzene	17.18	105	1931673	46.3771	ug/L	100
92) p-Isopropyltoluene	17.33	119	1670367	46.7824	ug/L	100
93) 1,3-Dichlorobenzene	17.53	146	915690	45.4713	ug/L	99
94) 1,4-Dichlorobenzene	17.66	146	909292	44.5880	ug/L	99
95) n-Butylbenzene	17.86	91	1537403	47.2576	ug/L	100
96) 1,2-Dichlorobenzene	18.15	146	831254	44.4823	ug/L	99
97) 1,2-Dibromo-3-Chloropropane	19.14	75	52127	41.0929	ug/L	97
98) 1,2,4-Trichlorobenzene	20.29	180	616788	45.4873	ug/L	100
99) Hexachlorobutadiene	20.44	225	300233	46.9901	ug/L	100
100) Naphthalene	20.66	128	1065041	43.0873	ug/L	100
101) 1,2,3-Trichlorobenzene	20.97	180	548043	43.9973	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M429061.D 8260WTR.M Mon Mar 11 15:31:10 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429061.D Vial: 1
 Acq On : 11 Mar 2019 9:19 Operator: EEA
 Sample : WG699027-02 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:30 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429061.D Vial: 1
 Acq On : 11 Mar 2019 9:19 Operator: EEA
 Sample : WG699027-02 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	1.000	1.000	0.0	111	0.00
2 T	Dichlorodifluoromethane	0.435	0.410	5.7	93	-0.01
3 P	Chloromethane	0.625	0.526	15.8	96	-0.02
4 C	Vinyl Chloride	0.510	0.475	6.9	97	-0.01
5 T	1,3-Butadiene	0.329	0.297	9.7	101	0.00
6 T	Bromomethane	0.307	0.247	19.5	97	-0.02
7 T	Chloroethane	0.273	0.255	6.6	97	-0.02
8 T	Trichlorofluoromethane	0.517	0.490	5.2	97	-0.01
9 T	Diethyl ether	0.276	0.247	10.5	97	-0.02
10 T	Isoprene	0.517	0.482	6.8	98	-0.01
11 T	Acrolein	0.032	0.029	9.4	99	-0.02
12 T	1,1,2-Trichloro-1,2,2-Trifl	0.298	0.285	4.4	99	-0.01
13 T	Acetone	0.077	0.062	19.5	83	-0.02
14 C	1,1-Dichloroethene	0.533	0.497	6.8	99	-0.02
15 T	Tert-Butyl Alcohol	0.025	0.019	24.0	90	-0.01
16 T	Dimethyl Sulfide	0.384	0.350	8.9	96	-0.01
17 T	Iodomethane	0.248	0.354	-42.7#	95	0.00
18 T	Methyl acetate	0.199	0.186	6.5	101	-0.02
19 T	Methylene Chloride	0.333	0.306	8.1	98	-0.01
20 T	Carbon Disulfide	0.997	0.938	5.9	98	-0.01
21 T	Acrylonitrile	0.097	0.090	7.2	98	-0.01
22 T	Methyl Tert Butyl Ether	0.793	0.702	11.5	95	-0.01
23 T	trans-1,2-Dichloroethene	0.509	0.481	5.5	99	-0.01
24 T	n-Hexane	0.500	0.518	-3.6	109	-0.02
25 T	Diisopropyl ether	1.261	1.181	6.3	100	-0.02
26 T	Vinyl Acetate	0.468	0.540	-15.4	125	-0.01
27 P	1,1-Dichloroethane	0.659	0.613	7.0	98	-0.01
28 T	Ethyl-Tert-Butyl ether	1.093	0.972	11.1	96	-0.01
29 T	2-Butanone	0.113	0.101	10.6	94	-0.01
30 T	Propionitrile	0.029	0.029	0.0	93	-0.01
31 T	2,2-Dichloropropane	0.478	0.479	-0.2	107	-0.01
32 T	cis-1,2-Dichloroethene	0.364	0.346	4.9	99	-0.01
33 C	Chloroform	0.585	0.530	9.4	99	-0.01
34	1-Bromopropane	0.078	0.072	7.7	96	-0.01
35 T	Bromochloromethane	0.207	0.194	6.3	98	0.00
36 T	Tetrahydrofuran	0.076	0.067	11.8	97	-0.01
37 S	Dibromofluoromethane	0.249	0.259	-4.0	110	-0.01
38 T	1,1,1-Trichloroethane	0.508	0.477	6.1	98	-0.01
39 T	Cyclohexane	0.676	0.649	4.0	100	-0.01
40 T	1,1-Dichloropropene	0.426	0.409	4.0	99	-0.01
41 T	Tert-Amyl-Methyl ether	0.833	0.732	12.1	95	-0.01
42 T	Carbon Tetrachloride	0.461	0.437	5.2	99	-0.01
43 S	1,2-Dichloroethane-d4	0.246	0.249	-1.2	107	0.00
44	Heptane	0.000	0.000	0.0	0#	-2.63#
45 T	1,2-Dichloroethane	0.400	0.374	6.5	97	-0.01
46 T	Benzene	1.335	1.257	5.8	99	-0.01
47 T	Trichloroethene	0.364	0.333	8.5	95	-0.01
48 T	Methylcyclohexane	0.563	0.542	3.7	102	-0.01
49 C	1,2-Dichloropropane	0.362	0.348	3.9	99	-0.01
50 T	Bromodichloromethane	0.420	0.403	4.0	99	-0.01
51 T	1,4-Dioxane	0.002	0.002	0.0	94	0.00
52 T	Dibromomethane	0.167	0.153	8.4	97	-0.01
53 T	2-Chloroethyl Vinyl Ether	0.158	0.098	38.0#	66	0.00
54 T	4-Methyl-2-Pentanone	0.100	0.091	9.0	98	0.00

(#) = Out of Range

8M429061.D 8260WTR.M

Mon Mar 11 15:31:19 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429061.D Vial: 1
 Acq On : 11 Mar 2019 9:19 Operator: EEA
 Sample : WG699027-02 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
55 T	cis-1,3-Dichloropropene	0.499	0.495	0.8	100	-0.01
56 T	Dimethyl Disulfide	0.307	0.281	8.5	96	0.00
57 I	Chlorobenzene-d5	1.000	1.000	0.0	112	-0.01
58 S	Toluene-d8	1.273	1.317	-3.5	111	-0.01
59 C	Toluene	1.867	1.722	7.8	99	-0.01
60 T	Ethyl Methacrylate	0.437	0.414	5.3	94	-0.01
61	Paraldehyde	0.000	0.000	0.0	158#	0.00
62 T	trans-1,3-Dichloropropene	0.561	0.540	3.7	100	-0.01
63 T	1,1,2-Trichloroethane	0.326	0.300	8.0	97	-0.01
64 T	2-Hexanone	0.122	0.110	9.8	95	-0.01
65 T	1,3-Dichloropropane	0.562	0.515	8.4	98	0.00
66 T	Tetrachloroethene	0.393	0.364	7.4	100	-0.01
67 T	Dibromochloromethane	0.414	0.386	6.8	98	0.00
68 T	1,2-Dibromoethane	0.317	0.294	7.3	96	0.00
69 T	1-Chlorohexane	0.647	0.606	6.3	100	0.00
70 P	Chlorobenzene	1.265	1.142	9.7	99	-0.01
71 T	1,1,1,2-Tetrachloroethane	0.465	0.433	6.9	99	0.00
72 C	Ethylbenzene	0.685	0.653	4.7	100	0.00
73 T	m-,p-Xylene	0.834	0.781	6.4	100	-0.01
74 T	o-Xylene	0.809	0.776	4.1	100	0.00
75 T	Styrene	1.340	1.281	4.4	99	-0.01
76 P	Bromoform	0.257	0.243	5.4	97	-0.01
77 T	Isopropylbenzene	2.045	1.945	4.9	100	0.00
78 I	1,4-Dichlorobenzene-d4	1.000	1.000	0.0	115	0.00
79 P	1,1,2,2-Tetrachloroethane	0.705	0.630	10.6	101	0.00
80 S	p-Bromofluorobenzene	0.902	0.935	-3.7	114	-0.01
81 T	1,2,3-Trichloropropane	0.211	0.183	13.3	95	-0.01
82 T	trans-1,4-Dichloro-2-Butene	0.238	0.208	12.6	95	0.00
83 T	n-Propylbenzene	4.524	4.164	8.0	101	0.00
84 T	Bromobenzene	1.035	0.932	10.0	99	-0.01
85 T	1,3,5-Trimethylbenzene	3.261	2.987	8.4	100	0.00
86 T	2-Chlorotoluene	2.991	2.515	15.9	92	0.00
87 T	4-Chlorotoluene	2.614	2.602	0.5	113	0.00
88 T	a-Methylstyrene	1.871	1.723	7.9	98	0.00
89 T	tert-Butylbenzene	0.752	0.674	10.4	100	0.00
90 T	1,2,4-Trimethylbenzene	3.302	3.039	8.0	100	0.00
91 T	sec-Butylbenzene	4.174	3.872	7.2	101	0.00
92 T	p-Isopropyltoluene	3.578	3.348	6.4	101	0.00
93 T	1,3-Dichlorobenzene	2.018	1.835	9.1	101	0.00
94 T	1,4-Dichlorobenzene	2.044	1.822	10.9	100	0.00
95 T	n-Butylbenzene	3.260	3.081	5.5	102	0.00
96 T	1,2-Dichlorobenzene	1.873	1.666	11.1	99	-0.01
97 T	1,2-Dibromo-3-Chloropropane	0.127	0.104	18.1	91	-0.01
98 T	1,2,4-Trichlorobenzene	1.359	1.236	9.1	99	0.00
99 T	Hexachlorobutadiene	0.640	0.602	5.9	102	0.00
100 T	Naphthalene	2.477	2.135	13.8	92	0.00
101 T	1,2,3-Trichlorobenzene	1.248	1.098	12.0	97	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 8M429061.D 8260WTR.M Mon Mar 11 15:31:19 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429061.D Vial: 1
 Acq On : 11 Mar 2019 9:19 Operator: EEA
 Sample : WG699027-02 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	111	0.00
2 T	Dichlorodifluoromethane	50.000	47.140	5.7	93	-0.01
3 P	Chloromethane	50.000	42.089	15.8	96	-0.02
4 C	Vinyl Chloride	50.000	46.577	6.8	97	-0.01
5 T	1,3-Butadiene	50.000	51.008	-2.0	101	0.00
6 T	Bromomethane	50.000	40.232	19.5	97	-0.02
7 T	Chloroethane	50.000	46.670	6.7	97	-0.02
8 T	Trichlorofluoromethane	50.000	47.378	5.2	97	-0.01
9 T	Diethyl ether	100.000	89.364	10.6	97	-0.02
10 T	Isoprene	50.000	46.642	6.7	98	-0.01
11 T	Acrolein	50.000	45.891	8.2	99	-0.02
12 T	1,1,2-Trichloro-1,2,2-Trifl	50.000	47.805	4.4	99	-0.01
13 T	Acetone	50.000	39.981	20.0	83	-0.02
14 C	1,1-Dichloroethene	50.000	46.592	6.8	99	-0.02
15 T	Tert-Butyl Alcohol	200.000	153.239	23.4	90	-0.01
16 T	Dimethyl Sulfide	50.000	45.529	8.9	96	-0.01
17 T	Iodomethane	50.000	45.378	9.2	95	0.00
18 T	Methyl acetate	50.000	46.618	6.8	101	-0.02
19 T	Methylene Chloride	50.000	45.968	8.1	98	-0.01
20 T	Carbon Disulfide	50.000	47.034	5.9	98	-0.01
21 T	Acrylonitrile	50.000	46.699	6.6	98	-0.01
22 T	Methyl Tert Butyl Ether	50.000	44.283	11.4	95	-0.01
23 T	trans-1,2-Dichloroethene	50.000	47.181	5.6	99	-0.01
24 T	n-Hexane	50.000	51.766	-3.5	109	-0.02
25 T	Diisopropyl ether	100.000	93.682	6.3	100	-0.02
26 T	Vinyl Acetate	50.000	57.662	-15.3	125	-0.01
27 P	1,1-Dichloroethane	50.000	46.560	6.9	98	-0.01
28 T	Ethyl-Tert-Butyl ether	100.000	88.917	11.1	96	-0.01
29 T	2-Butanone	50.000	44.392	11.2	94	-0.01
30 T	Propionitrile	100.000	85.156	14.8	93	-0.01
31 T	2,2-Dichloropropane	50.000	50.167	-0.3	107	-0.01
32 T	cis-1,2-Dichloroethene	50.000	47.551	4.9	99	-0.01
33 C	Chloroform	50.000	45.345	9.3	99	-0.01
34	1-Bromopropane	50.000	46.112	7.8	96	-0.01
35 T	Bromochloromethane	50.000	46.779	6.4	98	0.00
36 T	Tetrahydrofuran	100.000	88.350	11.7	97	-0.01
37 S	Dibromofluoromethane	25.000	26.007	-4.0	110	-0.01
38 T	1,1,1-Trichloroethane	50.000	46.945	6.1	98	-0.01
39 T	Cyclohexane	50.000	47.985	4.0	100	-0.01
40 T	1,1-Dichloropropene	50.000	47.920	4.2	99	-0.01
41 T	Tert-Amyl-Methyl ether	100.000	87.863	12.1	95	-0.01
42 T	Carbon Tetrachloride	50.000	47.348	5.3	99	-0.01
43 S	1,2-Dichloroethane-d4	25.000	25.327	-1.3	107	0.00
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	50.000	46.779	6.4	97	-0.01
46 T	Benzene	50.000	47.056	5.9	99	-0.01
47 T	Trichloroethene	50.000	45.725	8.5	95	-0.01
48 T	Methylcyclohexane	50.000	48.091	3.8	102	-0.01
49 C	1,2-Dichloropropane	50.000	47.981	4.0	99	-0.01
50 T	Bromodichloromethane	50.000	48.008	4.0	99	-0.01
51 T	1,4-Dioxane	200.000	183.310	8.3	94	0.00
52 T	Dibromomethane	50.000	45.861	8.3	97	-0.01
53 T	2-Chloroethyl Vinyl Ether	50.000	30.816	38.4#	66	0.00
54 T	4-Methyl-2-Pentanone	50.000	45.653	8.7	98	0.00

(#) = Out of Range

8M429061.D 8260WTR.M

Mon Mar 11 15:31:17 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429061.D Vial: 1
 Acq On : 11 Mar 2019 9:19 Operator: EEA
 Sample : WG699027-02 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	cis-1,3-Dichloropropene	50.000	49.601	0.8	100	-0.01
56 T	Dimethyl Disulfide	50.000	45.757	8.5	96	0.00
57 I	Chlorobenzene-d5	25.000	25.000	0.0	112	-0.01
58 S	Toluene-d8	25.000	25.853	-3.4	111	-0.01
59 C	Toluene	50.000	46.133	7.7	99	-0.01
60 T	Ethyl Methacrylate	50.000	42.763	14.5	94	-0.01
61	Paraldehyde	-1.000	0.000	0.0	158	0.00
62 T	trans-1,3-Dichloropropene	50.000	48.105	3.8	100	-0.01
63 T	1,1,2-Trichloroethane	50.000	45.898	8.2	97	-0.01
64 T	2-Hexanone	50.000	41.900	16.2	95	-0.01
65 T	1,3-Dichloropropane	50.000	45.787	8.4	98	0.00
66 T	Tetrachloroethene	50.000	46.322	7.4	100	-0.01
67 T	Dibromochloromethane	50.000	46.614	6.8	98	0.00
68 T	1,2-Dibromoethane	50.000	46.474	7.1	96	0.00
69 T	1-Chlorohexane	50.000	46.813	6.4	100	0.00
70 P	Chlorobenzene	50.000	45.143	9.7	99	-0.01
71 T	1,1,1,2-Tetrachloroethane	50.000	46.558	6.9	99	0.00
72 C	Ethylbenzene	50.000	47.712	4.6	100	0.00
73 T	m-,p-Xylene	100.000	93.592	6.4	100	-0.01
74 T	o-Xylene	50.000	47.956	4.1	100	0.00
75 T	Styrene	50.000	47.801	4.4	99	-0.01
76 P	Bromoform	50.000	47.331	5.3	97	-0.01
77 T	Isopropylbenzene	50.000	47.566	4.9	100	0.00
78 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	115	0.00
79 P	1,1,2,2-Tetrachloroethane	50.000	44.686	10.6	101	0.00
80 S	p-Bromofluorobenzene	25.000	25.920	-3.7	114	-0.01
81 T	1,2,3-Trichloropropane	50.000	43.275	13.5	95	-0.01
82 T	trans-1,4-Dichloro-2-Butene	50.000	43.541	12.9	95	0.00
83 T	n-Propylbenzene	50.000	46.020	8.0	101	0.00
84 T	Bromobenzene	50.000	44.993	10.0	99	-0.01
85 T	1,3,5-Trimethylbenzene	50.000	45.798	8.4	100	0.00
86 T	2-Chlorotoluene	50.000	42.041	15.9	92	0.00
87 T	4-Chlorotoluene	50.000	49.780	0.4	113	0.00
88 T	a-Methylstyrene	50.000	46.025	8.0	98	0.00
89 T	tert-Butylbenzene	50.000	44.759	10.5	100	0.00
90 T	1,2,4-Trimethylbenzene	50.000	46.023	8.0	100	0.00
91 T	sec-Butylbenzene	50.000	46.377	7.2	101	0.00
92 T	p-Isopropyltoluene	50.000	46.782	6.4	101	0.00
93 T	1,3-Dichlorobenzene	50.000	45.471	9.1	101	0.00
94 T	1,4-Dichlorobenzene	50.000	44.588	10.8	100	0.00
95 T	n-Butylbenzene	50.000	47.258	5.5	102	0.00
96 T	1,2-Dichlorobenzene	50.000	44.482	11.0	99	-0.01
97 T	1,2-Dibromo-3-Chloropropane	50.000	41.093	17.8	91	-0.01
98 T	1,2,4-Trichlorobenzene	50.000	45.487	9.0	99	0.00
99 T	Hexachlorobutadiene	50.000	46.990	6.0	102	0.00
100 T	Naphthalene	50.000	43.087	13.8	92	0.00
101 T	1,2,3-Trichlorobenzene	50.000	43.997	12.0	97	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 8M429061.D 8260WTR.M Mon Mar 11 15:31:17 2019

Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29851.D Vial: 23
 Acq On : 12 Mar 2019 22:39 Operator: KFR
 Sample : WG699219-03 QCMRL 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 13 09:04:06 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3916	96	357416	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0208	117	276741	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8331	152	144383	25.0000	ug/L	0.0000
System Monitoring Compounds						
37) Dibromofluoromethane	9.3990	111	108020	26.1818	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	104.7272%	
43) 1,2-Dichloroethane-d4	10.0090	65	123784	25.0056	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	100.0224%	
57) Toluene-d8	12.2527	98	348345	25.2930	ug/L	0.0000
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	101.1720%	
78) p-Bromofluorobenzene	15.4166	95	136387	24.6843	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	98.7372%	
Target Compounds						
					Qvalue	
2) Dichlorodifluoromethane	3.1228	85	246271	46.6420	ug/L	97
3) Chloromethane	3.5571	50	318423	31.3991	ug/L	99
4) Vinyl Chloride	3.7742	62	328529	43.3957	ug/L	99
5) 1,3-Butadiene	3.8156	54	130428	50.2688	ug/L	99
6) Bromomethane	4.6531	94	111080	38.2973	ug/L	99
7) Chloroethane	4.7979	64	134800	46.0582	ug/L	99
8) Trichlorofluoromethane	5.2735	101	341870	47.7200	ug/L	99
9) Diethyl ether	5.7904	59	362002	96.6507	ug/L	98
10) Isoprene	5.8215	67	266399	47.2114	ug/L	100
11) Acrolein	6.0179	56	23526	36.1705	ug/L	99
12) 1,1,2-Trichloro-1,2,2-Trif	6.0386	101	171549	48.8149	ug/L	100
13) Acetone	6.1213	43	58176	44.5217	ug/L	98
14) 1,1-Dichloroethene	6.3384	61	323043	47.2309	ug/L	97
15) Tert-Butyl Alcohol	6.4418	59	73220	200.8936	ug/L	98
16) Dimethyl Sulfide	6.5866	62	216845	47.9587	ug/L	98
17) Iodomethane	6.8347	142	146851	31.7598	ug/L	99
18) Methyl acetate	6.8451	43	136099	48.2873	ug/L	97
19) Methylene Chloride	7.0932	84	184356	48.3683	ug/L	95
20) Carbon Disulfide	7.1346	76	522537	49.7409	ug/L	99
21) Acrylonitrile	7.2690	53	76606	53.1808	ug/L	99
22) Methyl Tert Butyl Ether	7.3000	73	525733	48.6429	ug/L	99
23) trans-1,2-Dichloroethene	7.5275	96	184253	49.7307	ug/L	98
24) n-Hexane	7.5999	57	237576	41.6313	ug/L	98
25) Diisopropyl ether	7.9307	45	1356100	97.3653	ug/L	99
26) Vinyl Acetate	8.0962	43	93222	19.9554	ug/L	99
27) 1,1-Dichloroethane	8.1272	63	383926	47.4877	ug/L	98
28) Ethyl-Tert-Butyl ether	8.4787	59	1412530	100.3374	ug/L	99
29) 2-Butanone	8.6545	43	76503	45.9635	ug/L	99
30) Propionitrile	8.7579	54	50387	103.6138	ug/L	100
31) 2,2-Dichloropropane	8.8613	77	196328	35.1213	ug/L	99
32) cis-1,2-Dichloroethene	8.9337	96	214062	50.8914	ug/L	90
33) Chloroform	9.1301	83	349972	47.6922	ug/L	99
34) 1-Bromopropane	9.2542	122	44266	48.5869	ug/L	97
35) Bromochloromethane	9.3473	130	129082	52.3377	ug/L	95
36) Tetrahydrofuran	9.3680	42	99551	92.3194	ug/L	97
38) 1,1,1-Trichloroethane	9.6368	97	340453	51.2387	ug/L	98
39) Cyclohexane	9.6575	56	371350	47.2059	ug/L	97
40) 1,1-Dichloropropene	9.8125	75	240720	48.1846	ug/L	98
41) Carbon Tetrachloride	9.9573	117	312695	55.3422	ug/L	99
42) Tert-Amyl-Methyl ether	9.9159	73	1068169	101.7306	ug/L	100

(#) = qualifier out of range (m) = manual integration
 11M29851.D 8260WT.M Wed Mar 13 09:04:07 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\031219\11M29851.D Vial: 23
 Acq On : 12 Mar 2019 22:39 Operator: KFR
 Sample : WG699219-03 QCMRL 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 13 09:04:06 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1227	62	315829	49.5192	ug/L	99
45) Benzene	10.1538	78	737727	49.6820	ug/L	100
46) Trichloroethene	10.8672	130	240685	57.1503	ug/L	99
47) Methylcyclohexane	10.9396	83	291746	49.1745	ug/L	96
48) 1,2-Dichloropropane	11.0636	63	216673	49.1221	ug/L	100
49) 1,4-Dioxane	11.3325	88	5646	186.5480	ug/L	89
50) Bromodichloromethane	11.3531	83	276618	53.0647	ug/L	99
51) Dibromomethane	11.4359	93	128718	51.0866	ug/L	97
52) 2-Chloroethyl Vinyl Ether	11.6220	63	83391	39.0672	ug/L	100
53) 4-Methyl-2-Pentanone	11.6530	58	72320	47.5558	ug/L	96
54) cis-1,3-Dichloropropene	11.9425	75	290443	49.4369	ug/L	99
55) Dimethyl Disulfide	12.1907	79	187889	47.6064	ug/L	100
58) Toluene	12.3457	91	816217	49.7885	ug/L	99
59) Ethyl Methacrylate	12.4285	69	240005	51.8330	ug/L	98
60) trans-1,3-Dichloropropene	12.5008	75	261870	48.7880	ug/L	99
61) 1,1,2-Trichloroethane	12.7076	97	171735	49.2083	ug/L	99
62) 2-Hexanone	12.6456	43	121358	46.7836	ug/L	98
63) 1,3-Dichloropropane	12.9971	76	275860	48.9912	ug/L	98
64) Tetrachloroethene	13.1109	164	168540	48.4684	ug/L	100
65) Dibromochloromethane	13.3590	129	213691	53.7882	ug/L	100
66) 1,2-Dibromoethane	13.5968	107	168121	50.2645	ug/L	98
67) 1-Chlorohexane	13.6692	91	259215	47.8933	ug/L	97
68) Chlorobenzene	14.0621	112	560289	49.0904	ug/L	100
69) 1,1,1,2-Tetrachloroethane	14.0931	131	236697	48.8135	ug/L	99
70) Ethylbenzene	14.0828	106	297975	51.3200	ug/L	98
71) m-,p-Xylene	14.1655	106	706634	100.8092	ug/L	99
72) o-Xylene	14.6928	106	350455	51.5289	ug/L	98
73) Styrene	14.7342	104	595723	52.9548	ug/L	98
74) Bromoform	15.1995	173	146893	51.2713	ug/L	98
75) Isopropylbenzene	15.0857	105	871839	51.2071	ug/L	100
77) 1,1,2,2-Tetrachloroethane	15.2925	83	162183	43.0592	ug/L	99
79) 1,2,3-Trichloropropane	15.4683	110	62744	49.8902	ug/L	96
80) trans-1,4-Dichloro-2-Butene	15.5097	53	68235	43.0318	ug/L	86
81) n-Propylbenzene	15.5613	91	1052322	49.4229	ug/L	100
82) Bromobenzene	15.6854	156	264007	50.7716	ug/L	90
83) 1,3,5-Trimethylbenzene	15.7371	105	753573	49.8542	ug/L	100
84) 2-Chlorotoluene	15.8198	91	735799	48.5646	ug/L	99
85) 4-Chlorotoluene	15.8612	91	589716	47.3576	ug/L	99
86) a-Methylstyrene	16.1093	118	442391	51.3993	ug/L	98
87) tert-Butylbenzene	16.1610	134	176440	47.3926	ug/L	97
88) 1,2,4-Trimethylbenzene	16.2128	105	776117	49.7493	ug/L	99
89) sec-Butylbenzene	16.4195	105	934295	49.7875	ug/L	100
90) p-Isopropyltoluene	16.5643	119	821860	50.3668	ug/L	100
91) 1,3-Dichlorobenzene	16.7504	146	480449	50.0860	ug/L	99
92) 1,4-Dichlorobenzene	16.8641	146	475995	49.6919	ug/L	99
93) n-Butylbenzene	17.0502	91	735475	50.0133	ug/L	100
94) 1,2-Dichlorobenzene	17.3294	146	463676	50.4354	ug/L	99
95) 1,2-Dibromo-3-Chloropropane	18.2496	75	39128	50.2867	ug/L	95
96) 1,2,4-Trichlorobenzene	19.3146	180	303596	51.0489	ug/L	100
97) Hexachlorobutadiene	19.4490	225	151279	50.4686	ug/L	97
98) Naphthalene	19.6558	128	675701	53.0595	ug/L	99
99) 1,2,3-Trichlorobenzene	19.9453	180	295081	52.8114	ug/L	98

(#) = qualifier out of range (m) = manual integration
 11M29851.D 8260WT.M Wed Mar 13 09:04:07 2019

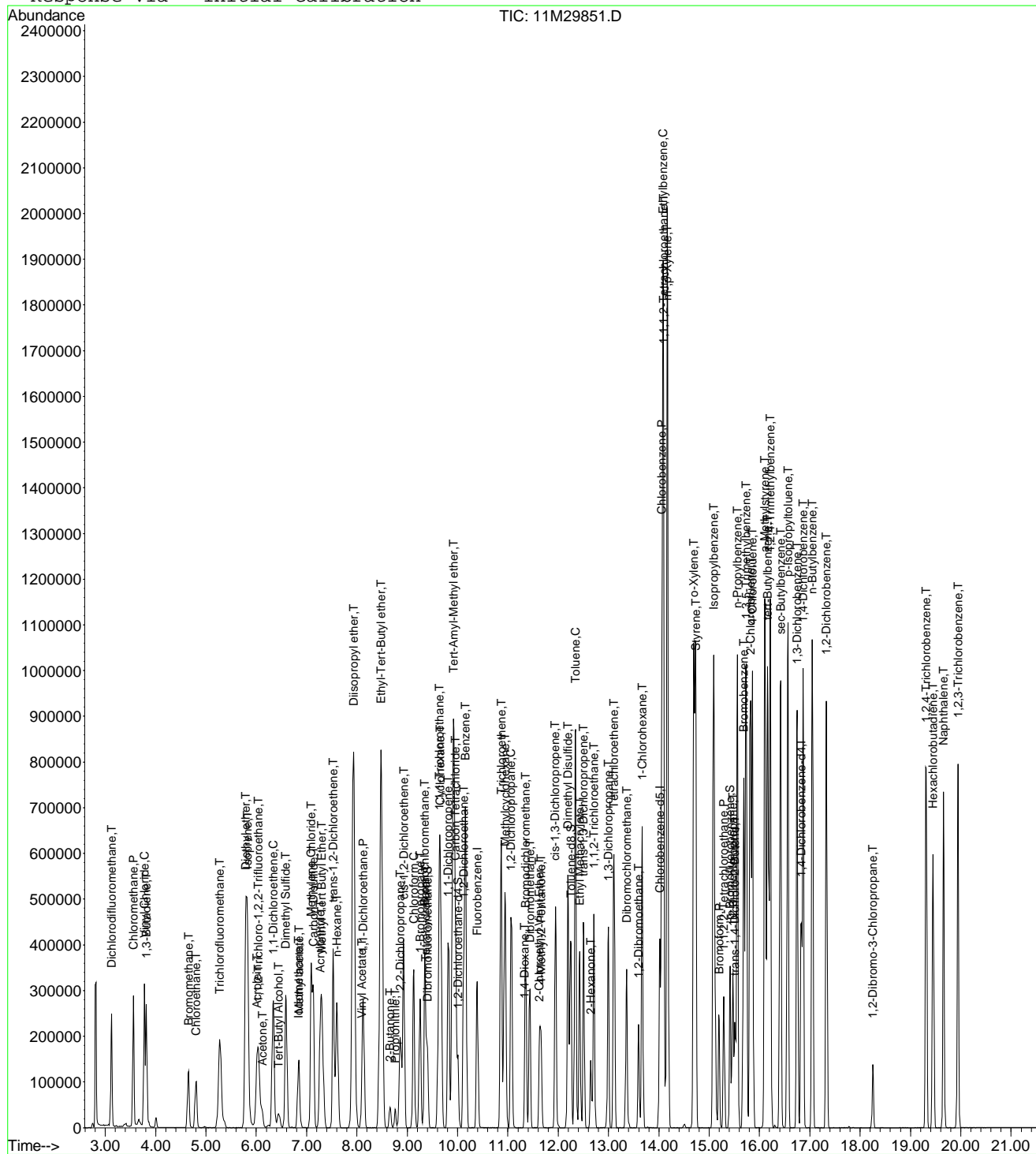
Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29851.D
Acq On : 12 Mar 2019 22:39
Sample : WG699219-03 QCMRL 50ug/L 8260
Misc : 1,1 STD92320
MS Integration Params: rteint.p
Quant Time: Mar 13 9:04 2019

Vial: 23
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:13:24 2019
Response via : Initial Calibration



Data File : C:\MSDCHEM\1\DATA\031219\11M29851.D Vial: 23
 Acq On : 12 Mar 2019 22:39 Operator: KFR
 Sample : WG699219-03 QCMRL 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	90	0.00
2 T	Dichlorodifluoromethane	50.000	46.642	6.7	77	0.00
3 P	Chloromethane	50.000	31.399	37.2#	59	0.00
4 C	Vinyl Chloride	50.000	43.396	13.2	76	0.00
5 T	1,3-Butadiene	50.000	50.269	-0.5	82	0.00
6 T	Bromomethane	50.000	38.297	23.4	76	0.01
7 T	Chloroethane	50.000	46.058	7.9	80	0.00
8 T	Trichlorofluoromethane	50.000	47.720	4.6	82	0.00
9 T	Diethyl ether	100.000	96.651	3.3	96	0.00
10 T	Isoprene	50.000	47.211	5.6	82	0.00
11 T	Acrolein	50.000	36.171	27.7#	67	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	50.000	48.815	2.4	83	0.00
13 T	Acetone	50.000	44.522	11.0	79	0.00
14 C	1,1-Dichloroethene	50.000	47.231	5.5	81	0.01
15 T	Tert-Butyl Alcohol	200.000	200.894	-0.4	91	0.00
16 T	Dimethyl Sulfide	50.000	47.959	4.1	83	0.00
17 T	Iodomethane	50.000	31.760	36.5#	56	0.00
18 T	Methyl acetate	50.000	48.287	3.4	84	0.00
19 T	Methylene Chloride	50.000	48.368	3.3	85	0.00
20 T	Carbon Disulfide	50.000	49.741	0.5	83	0.00
21 T	Acrylonitrile	50.000	53.181	-6.4	88	0.00
22 T	Methyl Tert Butyl Ether	50.000	48.643	2.7	85	0.00
23 T	trans-1,2-Dichloroethene	50.000	49.731	0.5	84	0.00
24 T	n-Hexane	50.000	41.631	16.7	73	0.00
25 T	Diisopropyl ether	100.000	97.365	2.6	94	0.00
26 T	Vinyl Acetate	50.000	19.955	60.1#	34	0.00
27 P	1,1-Dichloroethane	50.000	47.488	5.0	83	0.00
28 T	Ethyl-Tert-Butyl ether	100.000	100.337	-0.3	95	0.00
29 T	2-Butanone	50.000	45.964	8.1	79	0.00
30 T	Propionitrile	100.000	103.614	-3.6	86	0.00
31 T	2,2-Dichloropropane	50.000	35.121	29.8#	60	0.00
32 T	cis-1,2-Dichloroethene	50.000	50.891	-1.8	86	0.00
33 C	Chloroform	50.000	47.692	4.6	85	0.00
34 T	1-Bromopropane	50.000	48.587	2.8	86	0.00
35 T	Bromochloromethane	50.000	52.338	-4.7	86	0.00
36 T	Tetrahydrofuran	100.000	92.319	7.7	87	0.00
37 S	Dibromofluoromethane	25.000	26.182	-4.7	91	0.00
38 T	1,1,1-Trichloroethane	50.000	51.239	-2.5	86	0.01
39 T	Cyclohexane	50.000	47.206	5.6	82	0.00
40 T	1,1-Dichloropropene	50.000	48.185	3.6	82	0.00
41 T	Carbon Tetrachloride	50.000	55.342	-10.7	90	0.00
42 T	Tert-Amyl-Methyl ether	100.000	101.731	-1.7	95	0.00
43 S	1,2-Dichloroethane-d4	25.000	25.006	-0.0	87	0.00
44 T	1,2-Dichloroethane	50.000	49.519	1.0	85	0.00
45 T	Benzene	50.000	49.682	0.6	86	0.00
46 T	Trichloroethene	50.000	57.150	-14.3	97	0.00
47 T	Methylcyclohexane	50.000	49.174	1.7	85	0.00
48 C	1,2-Dichloropropane	50.000	49.122	1.8	85	0.00
49 T	1,4-Dioxane	200.000	186.548	6.7	82	-0.01
50 T	Bromodichloromethane	50.000	53.065	-6.1	86	0.00
51 T	Dibromomethane	50.000	51.087	-2.2	87	0.00
52 T	2-Chloroethyl Vinyl Ether	50.000	39.067	21.9	69	0.00
53 T	4-Methyl-2-Pentanone	50.000	47.556	4.9	83	0.00
54 T	cis-1,3-Dichloropropene	50.000	49.437	1.1	82	0.00

(#) = Out of Range

11M29851.D 8260WT.M Wed Mar 13 09:06:30 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\031219\11M29851.D Vial: 23
 Acq On : 12 Mar 2019 22:39 Operator: KFR
 Sample : WG699219-03 QCMRL 50ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92320 Multiplr: 1.00
 MS Integration Params: rteint.p

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	Dimethyl Disulfide	50.000	47.606	4.8	89	0.00
56 I	Chlorobenzene-d5	25.000	25.000	0.0	92	0.00
57 S	Toluene-d8	25.000	25.293	-1.2	90	0.00
58 C	Toluene	50.000	49.788	0.4	87	0.00
59 T	Ethyl Methacrylate	50.000	51.833	-3.7	88	0.00
60 T	trans-1,3-Dichloropropene	50.000	48.788	2.4	82	0.00
61 T	1,1,2-Trichloroethane	50.000	49.208	1.6	88	0.00
62 T	2-Hexanone	50.000	46.784	6.4	80	0.00
63 T	1,3-Dichloropropane	50.000	48.991	2.0	86	0.00
64 T	Tetrachloroethene	50.000	48.468	3.1	86	0.00
65 T	Dibromochloromethane	50.000	53.788	-7.6	90	0.00
66 T	1,2-Dibromoethane	50.000	50.264	-0.5	87	0.00
67 T	1-Chlorohexane	50.000	47.893	4.2	83	0.00
68 P	Chlorobenzene	50.000	49.090	1.8	86	0.00
69 T	1,1,1,2-Tetrachloroethane	50.000	48.814	2.4	90	0.00
70 C	Ethylbenzene	50.000	51.320	-2.6	86	0.00
71 T	m-,p-Xylene	100.000	100.809	-0.8	86	0.00
72 T	o-Xylene	50.000	51.529	-3.1	87	0.00
73 T	Styrene	50.000	52.955	-5.9	87	0.00
74 P	Bromoform	50.000	51.271	-2.5	94	0.00
75 T	Isopropylbenzene	50.000	51.207	-2.4	86	0.00
76 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	97	0.00
77 P	1,1,2,2-Tetrachloroethane	50.000	43.059	13.9	72	0.00
78 S	p-Bromofluorobenzene	25.000	24.684	1.3	88	0.00
79 T	1,2,3-Trichloropropane	50.000	49.890	0.2	87	0.00
80 T	trans-1,4-Dichloro-2-Butene	50.000	43.032	13.9	75	0.00
81 T	n-Propylbenzene	50.000	49.423	1.2	86	0.00
82 T	Bromobenzene	50.000	50.772	-1.5	90	0.00
83 T	1,3,5-Trimethylbenzene	50.000	49.854	0.3	86	0.00
84 T	2-Chlorotoluene	50.000	48.565	2.9	87	0.00
85 T	4-Chlorotoluene	50.000	47.358	5.3	83	0.00
86 T	a-Methylstyrene	50.000	51.399	-2.8	89	0.00
87 T	tert-Butylbenzene	50.000	47.393	5.2	87	-0.01
88 T	1,2,4-Trimethylbenzene	50.000	49.749	0.5	86	0.00
89 T	sec-Butylbenzene	50.000	49.787	0.4	86	0.00
90 T	p-Isopropyltoluene	50.000	50.367	-0.7	86	0.00
91 T	1,3-Dichlorobenzene	50.000	50.086	-0.2	88	0.00
92 T	1,4-Dichlorobenzene	50.000	49.692	0.6	87	0.00
93 T	n-Butylbenzene	50.000	50.013	-0.0	83	0.00
94 T	1,2-Dichlorobenzene	50.000	50.435	-0.9	88	0.00
95 T	1,2-Dibromo-3-Chloropropane	50.000	50.287	-0.6	87	0.00
96 T	1,2,4-Trichlorobenzene	50.000	51.049	-2.1	88	0.00
97 T	Hexachlorobutadiene	50.000	50.469	-0.9	89	0.00
98 T	Naphthalene	50.000	53.059	-6.1	89	0.00
99 T	1,2,3-Trichlorobenzene	50.000	52.811	-5.6	91	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 11M29851.D 8260WT.M Wed Mar 13 09:06:30 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429083.D Vial: 23
 Acq On : 11 Mar 2019 20:23 Operator: EEA
 Sample : WG699028-04 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00

MS Integration Params: RTEINT.P

Quant Time: Mar 12 08:50:19 2019

Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	501182	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.59	117	389505	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.61	152	219082	25.00	ug/L	0.00

System Monitoring Compounds

37) Dibromofluoromethane	9.69	111	128846	25.7910	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	103.16%	
43) 1,2-Dichloroethane-d4	10.33	65	128808	26.1173	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	104.48%	
58) Toluene-d8	12.70	98	500363	25.2191	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	100.88%	
80) p-Bromofluorobenzene	16.09	95	200405	25.3456	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	101.40%	

Target Compounds

						Qvalue
2) Dichlorodifluoromethane	3.22	85	435102	49.9117	ug/L	100
3) Chloromethane	3.68	50	605358	48.2862	ug/L	100
4) Vinyl Chloride	3.91	62	517648	50.6657	ug/L	99
5) 1,3-Butadiene	3.95	54	434413	76.6333	ug/L	97
6) Bromomethane	4.80	94	260837	42.4452	ug/L	99
7) Chloroethane	4.96	64	279019	51.0336	ug/L	99
8) Trichlorofluoromethane	5.44	101	527483	50.8494	ug/L	99
9) Diethyl ether	5.96	59	547586	98.9776	ug/L	97
10) Isoprene	6.00	67	517872	49.9521	ug/L	99
11) Acrolein	6.20	56	28001	44.1385	ug/L	99
12) 1,1,2-Trichloro-1,2,2-Trif	6.21	101	298559	49.9588	ug/L	100
13) Acetone	6.30	43	69860	45.2500	ug/L	97
14) 1,1-Dichloroethene	6.52	61	541683	50.6796	ug/L	99
15) Tert-Butyl Alcohol	6.63	59	53159	106.9683	ug/L	96
16) Dimethyl Sulfide	6.78	62	381168	49.5073	ug/L	98
17) Iodomethane	7.04	142	375792	47.9667	ug/L	99
18) Methyl acetate	7.05	43	205479	51.3990	ug/L	98
19) Methylene Chloride	7.29	84	331174	49.5856	ug/L	95
20) Carbon Disulfide	7.34	76	1007388	50.4031	ug/L	100
21) Acrylonitrile	7.48	53	98614	50.9127	ug/L	94
22) Methyl Tert Butyl Ether	7.51	73	788951	49.6283	ug/L	99
23) trans-1,2-Dichloroethene	7.75	61	525771	51.4809	ug/L	99
24) n-Hexane	7.82	57	486841	48.5724	ug/L	97
25) Diisopropyl ether	8.16	45	2628935	103.9815	ug/L	99
26) Vinyl Acetate	8.34	43	340333	36.2873	ug/L	99
27) 1,1-Dichloroethane	8.36	63	674013	51.0515	ug/L	100
28) Ethyl-Tert-Butyl ether	8.73	59	2156279	98.4023	ug/L	99
29) 2-Butanone	8.92	43	106144	46.6717	ug/L	97
30) Propionitrile	9.03	54	58096	84.1282	ug/L	99
31) 2,2-Dichloropropane	9.14	77	445551	46.5093	ug/L	100
32) cis-1,2-Dichloroethene	9.20	96	376229	51.6141	ug/L	100
33) Chloroform	9.40	83	582757	49.7332	ug/L	99
34) 1-Bromopropane	9.55	122	77110	49.2127	ug/L	97
35) Bromochloromethane	9.64	130	211667	50.9157	ug/L	96
36) Tetrahydrofuran	9.67	42	146481	95.7292	ug/L	97
38) 1,1,1-Trichloroethane	9.94	97	518442	50.9388	ug/L	100
39) Cyclohexane	9.96	56	700011	51.6383	ug/L	98
40) 1,1-Dichloropropene	10.13	75	440334	51.5226	ug/L	99
41) Tert-Amyl-Methyl ether	10.23	73	1632668	97.7717	ug/L	98
42) Carbon Tetrachloride	10.28	117	474722	51.3296	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M429083.D 8260WTR.M Tue Mar 12 08:50:19 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429083.D Vial: 23
 Acq On : 11 Mar 2019 20:23 Operator: EEA
 Sample : WG699028-04 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 08:50:19 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

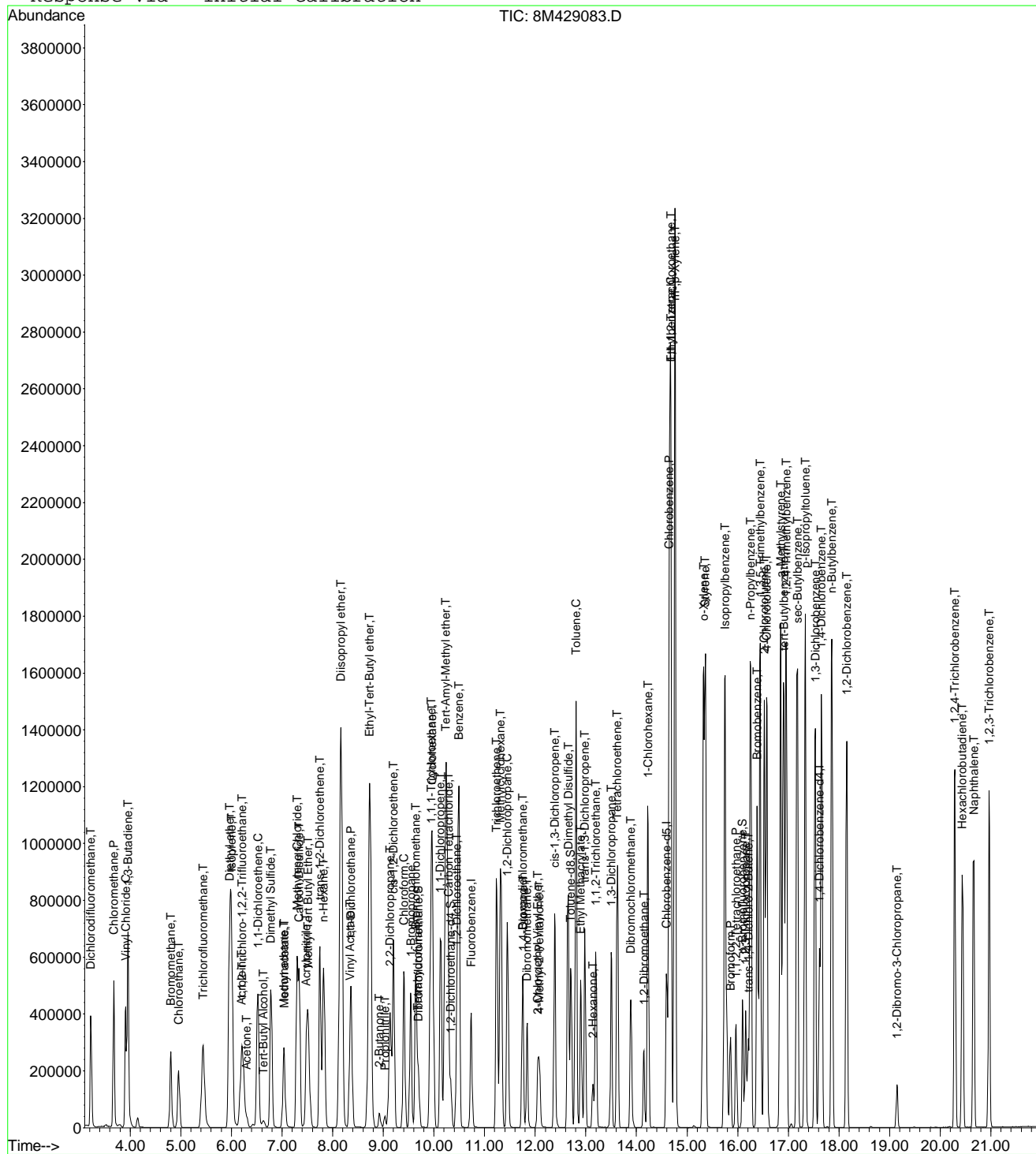
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.45	62	416841	52.0084	ug/L	99
46) Benzene	10.49	78	1370368	51.1857	ug/L	99
47) Trichloroethene	11.23	130	394861	54.1344	ug/L	99
48) Methylcyclohexane	11.32	83	559944	49.5694	ug/L	99
49) 1,2-Dichloropropane	11.45	63	376721	51.8398	ug/L	100
50) Bromodichloromethane	11.75	83	439754	52.2627	ug/L	100
51) 1,4-Dioxane	11.76	88	2790	68.2320	ug/L	91
52) Dibromomethane	11.83	93	169419	50.6492	ug/L	100
53) 2-Chloroethyl Vinyl Ether	12.05	63	101631	32.0060	ug/L	99
54) 4-Methyl-2-Pentanone	12.08	58	94950	47.5275	ug/L	98
55) cis-1,3-Dichloropropene	12.38	75	519058	51.8515	ug/L	100
56) Dimethyl Disulfide	12.65	79	305783	49.7427	ug/L	100
59) Toluene	12.81	91	1415175	48.6615	ug/L	100
60) Ethyl Methacrylate	12.90	69	352346	46.6972	ug/L	96
62) trans-1,3-Dichloropropene	12.98	75	435049	49.7592	ug/L	100
63) 1,1,2-Trichloroethane	13.20	97	251669	49.5074	ug/L	99
64) 2-Hexanone	13.14	58	85952	41.9368	ug/L	92
65) 1,3-Dichloropropane	13.50	76	434868	49.6710	ug/L	97
66) Tetrachloroethene	13.62	164	294260	48.0138	ug/L	99
67) Dibromochloromethane	13.89	129	319271	49.4864	ug/L	100
68) 1,2-Dibromoethane	14.15	107	249001	50.4435	ug/L	98
69) 1-Chlorohexane	14.23	91	479007	47.5314	ug/L	97
70) Chlorobenzene	14.65	112	938482	47.6018	ug/L	100
71) 1,1,1,2-Tetrachloroethane	14.68	131	354350	48.8871	ug/L	100
72) Ethylbenzene	14.68	106	531255	49.8039	ug/L	98
73) m-,p-Xylene	14.76	106	1269999	97.7133	ug/L	99
74) o-Xylene	15.33	106	636263	50.4517	ug/L	99
75) Styrene	15.36	104	1043462	49.9768	ug/L	99
76) Bromoform	15.86	173	201028	50.2523	ug/L	99
77) Isopropylbenzene	15.75	105	1582114	49.6678	ug/L	100
79) 1,1,2,2-Tetrachloroethane	15.97	83	254017	41.0983	ug/L	100
81) 1,2,3-Trichloropropane	16.16	110	86400	46.7290	ug/L	95
82) trans-1,4-Dichloro-2-Butene	16.21	53	87780	42.0121	ug/L	92
83) n-Propylbenzene	16.26	91	1875101	47.2941	ug/L	100
84) Bromobenzene	16.38	156	425884	46.9469	ug/L	97
85) 1,3,5-Trimethylbenzene	16.45	105	1353677	47.3631	ug/L	100
86) 2-Chlorotoluene	16.53	91	1155616	44.0825	ug/L	100
87) 4-Chlorotoluene	16.57	91	1142096	49.8610	ug/L	99
88) a-Methylstyrene	16.85	118	771772	47.0584	ug/L	98
89) tert-Butylbenzene	16.91	134	304430	46.1701	ug/L	98
90) 1,2,4-Trimethylbenzene	16.96	105	1367512	47.2610	ug/L	100
91) sec-Butylbenzene	17.18	105	1734328	47.4149	ug/L	99
92) p-Isopropyltoluene	17.33	119	1489155	47.4923	ug/L	100
93) 1,3-Dichlorobenzene	17.53	146	824584	46.6270	ug/L	100
94) 1,4-Dichlorobenzene	17.65	146	816006	45.5640	ug/L	99
95) n-Butylbenzene	17.86	91	1345813	47.1067	ug/L	100
96) 1,2-Dichlorobenzene	18.15	146	763922	46.5496	ug/L	100
97) 1,2-Dibromo-3-Chloropropane	19.14	75	49492	44.4276	ug/L	99
98) 1,2,4-Trichlorobenzene	20.29	180	557700	46.8348	ug/L	99
99) Hexachlorobutadiene	20.44	225	252665	45.0305	ug/L	99
100) Naphthalene	20.66	128	1017102	46.8555	ug/L	100
101) 1,2,3-Trichlorobenzene	20.96	180	506273	46.2817	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M429083.D 8260WTR.M Tue Mar 12 08:50:20 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429083.D Vial: 23
Acq On : 11 Mar 2019 20:23 Operator: EEA
Sample : WG699028-04 CCV 50 ug/L 8260 Inst : HPMS8
Misc : 1,1 STD92326 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 12 8:49 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429083.D Vial: 23
 Acq On : 11 Mar 2019 20:23 Operator: EEA
 Sample : WG699028-04 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	1.000	1.000	0.0	94	0.00
2 T	Dichlorodifluoromethane	0.435	0.434	0.2	84	0.00
3 P	Chloromethane	0.625	0.604	3.4	93	0.00
4 C	Vinyl Chloride	0.510	0.516	-1.2	89	0.00
5 T	1,3-Butadiene	0.329	0.433	-31.6#	125	0.00
6 T	Bromomethane	0.307	0.260	15.3	87	0.00
7 T	Chloroethane	0.273	0.278	-1.8	90	0.00
8 T	Trichlorofluoromethane	0.517	0.526	-1.7	88	0.00
9 T	Diethyl ether	0.276	0.273	1.1	91	0.00
10 T	Isoprene	0.517	0.517	0.0	89	0.00
11 T	Acrolein	0.032	0.028	12.5	81	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	0.298	0.298	0.0	88	0.00
13 T	Acetone	0.077	0.070	9.1	80	0.00
14 C	1,1-Dichloroethene	0.533	0.540	-1.3	91	0.00
15 T	Tert-Butyl Alcohol	0.025	0.013	48.0#	53	0.00
16 T	Dimethyl Sulfide	0.384	0.380	1.0	89	0.00
17 T	Iodomethane	0.248	0.375	-51.2#	85	0.00
18 T	Methyl acetate	0.199	0.205	-3.0	94	0.00
19 T	Methylene Chloride	0.333	0.330	0.9	90	0.00
20 T	Carbon Disulfide	0.997	1.005	-0.8	89	0.00
21 T	Acrylonitrile	0.097	0.098	-1.0	91	0.00
22 T	Methyl Tert Butyl Ether	0.793	0.787	0.8	90	0.00
23 T	trans-1,2-Dichloroethene	0.509	0.525	-3.1	92	0.00
24 T	n-Hexane	0.500	0.486	2.8	87	0.00
25 T	Diisopropyl ether	1.261	1.311	-4.0	94	0.00
26 T	Vinyl Acetate	0.468	0.340	27.4#	67	0.00
27 P	1,1-Dichloroethane	0.659	0.672	-2.0	91	0.00
28 T	Ethyl-Tert-Butyl ether	1.093	1.076	1.6	90	0.00
29 T	2-Butanone	0.113	0.106	6.2	83	0.00
30 T	Propionitrile	0.029	0.029	0.0	78	0.00
31 T	2,2-Dichloropropane	0.478	0.445	6.9	84	0.00
32 T	cis-1,2-Dichloroethene	0.364	0.375	-3.0	91	0.00
33 C	Chloroform	0.585	0.581	0.7	92	0.00
34	1-Bromopropane	0.078	0.077	1.3	87	0.00
35 T	Bromochloromethane	0.207	0.211	-1.9	90	0.00
36 T	Tetrahydrofuran	0.076	0.073	3.9	89	0.00
37 S	Dibromofluoromethane	0.249	0.257	-3.2	92	0.00
38 T	1,1,1-Trichloroethane	0.508	0.517	-1.8	90	0.00
39 T	Cyclohexane	0.676	0.698	-3.3	92	0.00
40 T	1,1-Dichloropropene	0.426	0.439	-3.1	91	0.00
41 T	Tert-Amyl-Methyl ether	0.833	0.814	2.3	90	0.00
42 T	Carbon Tetrachloride	0.461	0.474	-2.8	91	0.00
43 S	1,2-Dichloroethane-d4	0.246	0.257	-4.5	93	0.00
44	Heptane	0.000	0.000	0.0	0#	-2.63#
45 T	1,2-Dichloroethane	0.400	0.416	-4.0	92	0.00
46 T	Benzene	1.335	1.367	-2.4	91	0.00
47 T	Trichloroethene	0.364	0.394	-8.2	95	0.00
48 T	Methylcyclohexane	0.563	0.559	0.7	89	0.00
49 C	1,2-Dichloropropane	0.362	0.376	-3.9	90	0.00
50 T	Bromodichloromethane	0.420	0.439	-4.5	91	0.00
51 T	1,4-Dioxane	0.002	0.001	50.0#	30#	0.01
52 T	Dibromomethane	0.167	0.169	-1.2	91	0.00
53 T	2-Chloroethyl Vinyl Ether	0.158	0.101	36.1#	58	0.00
54 T	4-Methyl-2-Pentanone	0.100	0.095	5.0	86	0.00

(#) = Out of Range

8M429083.D 8260WTR.M

Tue Mar 12 08:55:13 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429083.D Vial: 23
 Acq On : 11 Mar 2019 20:23 Operator: EEA
 Sample : WG699028-04 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
55 T	cis-1,3-Dichloropropene	0.499	0.518	-3.8	89	0.00
56 T	Dimethyl Disulfide	0.307	0.305	0.7	88	0.00
57 I	Chlorobenzene-d5	1.000	1.000	0.0	97	0.00
58 S	Toluene-d8	1.273	1.285	-0.9	95	0.00
59 C	Toluene	1.867	1.817	2.7	91	0.00
60 T	Ethyl Methacrylate	0.437	0.452	-3.4	89	0.00
61	Paraldehyde	0.000	0.000	0.0	117	0.00
62 T	trans-1,3-Dichloropropene	0.561	0.558	0.5	90	0.00
63 T	1,1,2-Trichloroethane	0.326	0.323	0.9	91	0.00
64 T	2-Hexanone	0.122	0.110	9.8	83	0.00
65 T	1,3-Dichloropropane	0.562	0.558	0.7	92	0.00
66 T	Tetrachloroethene	0.393	0.378	3.8	90	0.00
67 T	Dibromochloromethane	0.414	0.410	1.0	90	0.00
68 T	1,2-Dibromoethane	0.317	0.320	-0.9	91	0.00
69 T	1-Chlorohexane	0.647	0.615	4.9	88	0.00
70 P	Chlorobenzene	1.265	1.205	4.7	91	0.00
71 T	1,1,1,2-Tetrachloroethane	0.465	0.455	2.2	90	0.00
72 C	Ethylbenzene	0.685	0.682	0.4	91	0.00
73 T	m-,p-Xylene	0.834	0.815	2.3	91	0.00
74 T	o-Xylene	0.809	0.817	-1.0	91	0.00
75 T	Styrene	1.340	1.339	0.1	90	0.00
76 P	Bromoform	0.257	0.258	-0.4	89	0.00
77 T	Isopropylbenzene	2.045	2.031	0.7	91	0.00
78 I	1,4-Dichlorobenzene-d4	1.000	1.000	0.0	101	0.00
79 P	1,1,2,2-Tetrachloroethane	0.705	0.580	17.7	81	0.00
80 S	p-Bromofluorobenzene	0.902	0.915	-1.4	98	0.00
81 T	1,2,3-Trichloropropane	0.211	0.197	6.6	90	0.00
82 T	trans-1,4-Dichloro-2-Butene	0.238	0.200	16.0	81	0.00
83 T	n-Propylbenzene	4.524	4.279	5.4	91	0.00
84 T	Bromobenzene	1.035	0.972	6.1	91	0.00
85 T	1,3,5-Trimethylbenzene	3.261	3.089	5.3	91	0.00
86 T	2-Chlorotoluene	2.991	2.637	11.8	84	0.00
87 T	4-Chlorotoluene	2.614	2.607	0.3	99	0.00
88 T	a-Methylstyrene	1.871	1.761	5.9	88	0.00
89 T	tert-Butylbenzene	0.752	0.695	7.6	91	0.00
90 T	1,2,4-Trimethylbenzene	3.302	3.121	5.5	90	0.00
91 T	sec-Butylbenzene	4.174	3.958	5.2	91	0.00
92 T	p-Isopropyltoluene	3.578	3.399	5.0	90	0.00
93 T	1,3-Dichlorobenzene	2.018	1.882	6.7	91	0.00
94 T	1,4-Dichlorobenzene	2.044	1.862	8.9	90	0.00
95 T	n-Butylbenzene	3.260	3.071	5.8	89	0.00
96 T	1,2-Dichlorobenzene	1.873	1.743	6.9	91	0.00
97 T	1,2-Dibromo-3-Chloropropane	0.127	0.113	11.0	86	0.00
98 T	1,2,4-Trichlorobenzene	1.359	1.273	6.3	90	0.00
99 T	Hexachlorobutadiene	0.640	0.577	9.8	86	0.00
100 T	Naphthalene	2.477	2.321	6.3	88	0.00
101 T	1,2,3-Trichlorobenzene	1.248	1.155	7.5	89	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 8M429083.D 8260WTR.M Tue Mar 12 08:55:13 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429083.D Vial: 23
 Acq On : 11 Mar 2019 20:23 Operator: EEA
 Sample : WG699028-04 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Fluorobenzene	25.000	25.000	0.0	94	0.00
2 T	Dichlorodifluoromethane	50.000	49.912	0.2	84	0.00
3 P	Chloromethane	50.000	48.286	3.4	93	0.00
4 C	Vinyl Chloride	50.000	50.666	-1.3	89	0.00
5 T	1,3-Butadiene	50.000	76.633	-53.3#	125	0.00
6 T	Bromomethane	50.000	42.445	15.1	87	0.00
7 T	Chloroethane	50.000	51.034	-2.1	90	0.00
8 T	Trichlorofluoromethane	50.000	50.849	-1.7	88	0.00
9 T	Diethyl ether	100.000	98.978	1.0	91	0.00
10 T	Isoprene	50.000	49.952	0.1	89	0.00
11 T	Acrolein	50.000	44.139	11.7	81	0.00
12 T	1,1,2-Trichloro-1,2,2-Trifl	50.000	49.959	0.1	88	0.00
13 T	Acetone	50.000	45.250	9.5	80	0.00
14 C	1,1-Dichloroethene	50.000	50.680	-1.4	91	0.00
15 T	Tert-Butyl Alcohol	200.000	106.968	46.5#	53	0.00
16 T	Dimethyl Sulfide	50.000	49.507	1.0	89	0.00
17 T	Iodomethane	50.000	47.967	4.1	85	0.00
18 T	Methyl acetate	50.000	51.399	-2.8	94	0.00
19 T	Methylene Chloride	50.000	49.586	0.8	90	0.00
20 T	Carbon Disulfide	50.000	50.403	-0.8	89	0.00
21 T	Acrylonitrile	50.000	50.913	-1.8	91	0.00
22 T	Methyl Tert Butyl Ether	50.000	49.628	0.7	90	0.00
23 T	trans-1,2-Dichloroethene	50.000	51.481	-3.0	92	0.00
24 T	n-Hexane	50.000	48.572	2.9	87	0.00
25 T	Diisopropyl ether	100.000	103.981	-4.0	94	0.00
26 T	Vinyl Acetate	50.000	36.287	27.4#	67	0.00
27 P	1,1-Dichloroethane	50.000	51.052	-2.1	91	0.00
28 T	Ethyl-Tert-Butyl ether	100.000	98.402	1.6	90	0.00
29 T	2-Butanone	50.000	46.672	6.7	83	0.00
30 T	Propionitrile	100.000	84.128	15.9	78	0.00
31 T	2,2-Dichloropropane	50.000	46.509	7.0	84	0.00
32 T	cis-1,2-Dichloroethene	50.000	51.614	-3.2	91	0.00
33 C	Chloroform	50.000	49.733	0.5	92	0.00
34	1-Bromopropane	50.000	49.213	1.6	87	0.00
35 T	Bromochloromethane	50.000	50.916	-1.8	90	0.00
36 T	Tetrahydrofuran	100.000	95.729	4.3	89	0.00
37 S	Dibromofluoromethane	25.000	25.791	-3.2	92	0.00
38 T	1,1,1-Trichloroethane	50.000	50.939	-1.9	90	0.00
39 T	Cyclohexane	50.000	51.638	-3.3	92	0.00
40 T	1,1-Dichloropropene	50.000	51.523	-3.0	91	0.00
41 T	Tert-Amyl-Methyl ether	100.000	97.772	2.2	90	0.00
42 T	Carbon Tetrachloride	50.000	51.330	-2.7	91	0.00
43 S	1,2-Dichloroethane-d4	25.000	26.117	-4.5	93	0.00
44	Heptane	-1.000	0.000	0.0	0	-2.63#
45 T	1,2-Dichloroethane	50.000	52.008	-4.0	92	0.00
46 T	Benzene	50.000	51.186	-2.4	91	0.00
47 T	Trichloroethene	50.000	54.134	-8.3	95	0.00
48 T	Methylcyclohexane	50.000	49.569	0.9	89	0.00
49 C	1,2-Dichloropropane	50.000	51.840	-3.7	90	0.00
50 T	Bromodichloromethane	50.000	52.263	-4.5	91	0.00
51 T	1,4-Dioxane	200.000	68.232	65.9#	30	0.01
52 T	Dibromomethane	50.000	50.649	-1.3	91	0.00
53 T	2-Chloroethyl Vinyl Ether	50.000	32.006	36.0#	58	0.00
54 T	4-Methyl-2-Pentanone	50.000	47.528	4.9	86	0.00

(#) = Out of Range

8M429083.D 8260WTR.M

Tue Mar 12 08:55:11 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429083.D Vial: 23
 Acq On : 11 Mar 2019 20:23 Operator: EEA
 Sample : WG699028-04 CCV 50 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92326 Multiplr: 1.00
 MS Integration Params: RTEINT.P

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Multiple Level Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 25% Max. Rel. Area : 150%

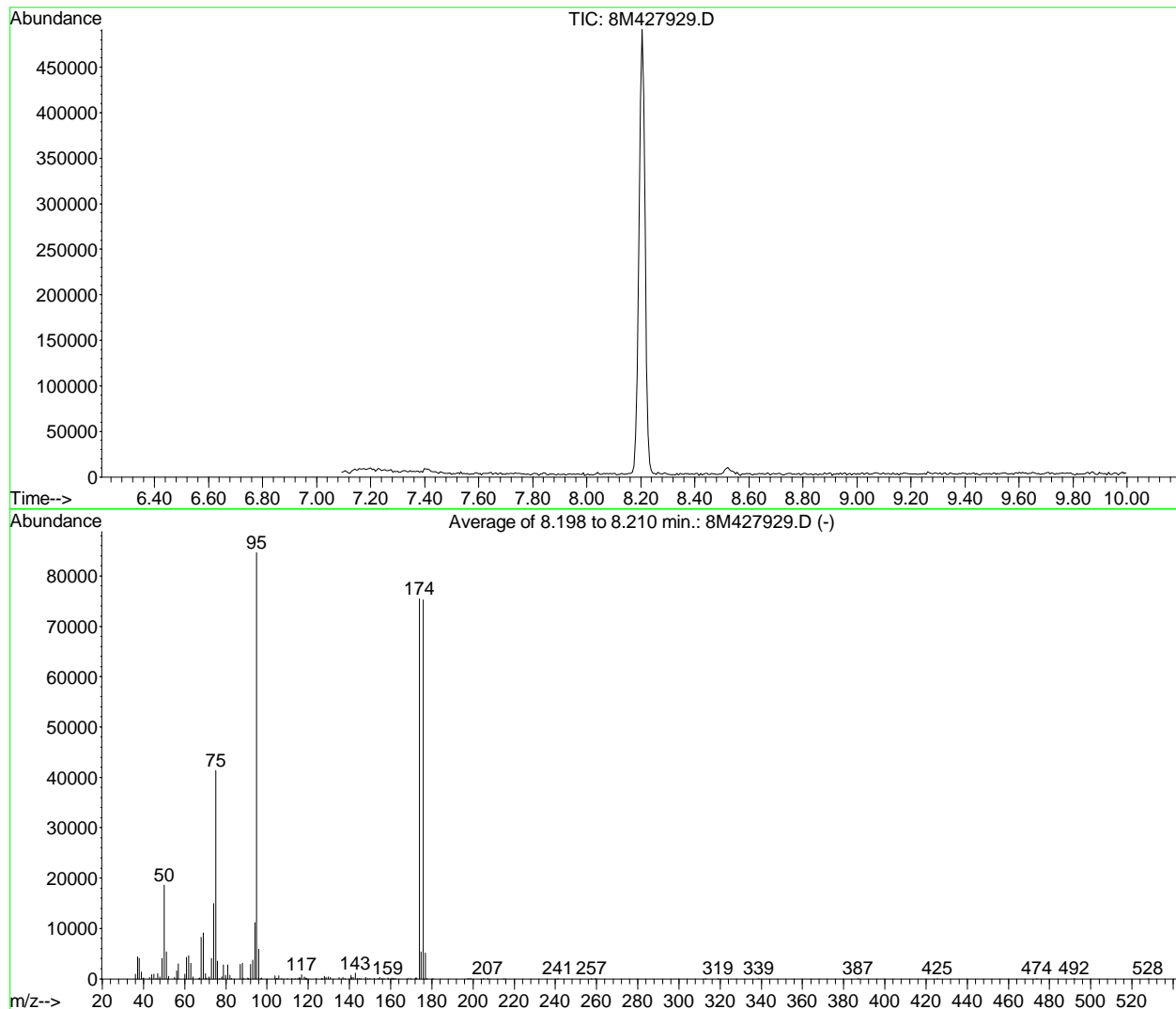
	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
55 T	cis-1,3-Dichloropropene	50.000	51.852	-3.7	89	0.00
56 T	Dimethyl Disulfide	50.000	49.743	0.5	88	0.00
57 I	Chlorobenzene-d5	25.000	25.000	0.0	97	0.00
58 S	Toluene-d8	25.000	25.219	-0.9	95	0.00
59 C	Toluene	50.000	48.662	2.7	91	0.00
60 T	Ethyl Methacrylate	50.000	46.697	6.6	89	0.00
61	Paraldehyde	-1.000	0.000	0.0	117	0.00
62 T	trans-1,3-Dichloropropene	50.000	49.759	0.5	90	0.00
63 T	1,1,2-Trichloroethane	50.000	49.507	1.0	91	0.00
64 T	2-Hexanone	50.000	41.937	16.1	83	0.00
65 T	1,3-Dichloropropane	50.000	49.671	0.7	92	0.00
66 T	Tetrachloroethene	50.000	48.014	4.0	90	0.00
67 T	Dibromochloromethane	50.000	49.486	1.0	90	0.00
68 T	1,2-Dibromoethane	50.000	50.444	-0.9	91	0.00
69 T	1-Chlorohexane	50.000	47.531	4.9	88	0.00
70 P	Chlorobenzene	50.000	47.602	4.8	91	0.00
71 T	1,1,1,2-Tetrachloroethane	50.000	48.887	2.2	90	0.00
72 C	Ethylbenzene	50.000	49.804	0.4	91	0.00
73 T	m-,p-Xylene	100.000	97.713	2.3	91	0.00
74 T	o-Xylene	50.000	50.452	-0.9	91	0.00
75 T	Styrene	50.000	49.977	0.0	90	0.00
76 P	Bromoform	50.000	50.252	-0.5	89	0.00
77 T	Isopropylbenzene	50.000	49.668	0.7	91	0.00
78 I	1,4-Dichlorobenzene-d4	25.000	25.000	0.0	101	0.00
79 P	1,1,2,2-Tetrachloroethane	50.000	41.098	17.8	81	0.00
80 S	p-Bromofluorobenzene	25.000	25.346	-1.4	98	0.00
81 T	1,2,3-Trichloropropane	50.000	46.729	6.5	90	0.00
82 T	trans-1,4-Dichloro-2-Butene	50.000	42.012	16.0	81	0.00
83 T	n-Propylbenzene	50.000	47.294	5.4	91	0.00
84 T	Bromobenzene	50.000	46.947	6.1	91	0.00
85 T	1,3,5-Trimethylbenzene	50.000	47.363	5.3	91	0.00
86 T	2-Chlorotoluene	50.000	44.083	11.8	84	0.00
87 T	4-Chlorotoluene	50.000	49.861	0.3	99	0.00
88 T	a-Methylstyrene	50.000	47.058	5.9	88	0.00
89 T	tert-Butylbenzene	50.000	46.170	7.7	91	0.00
90 T	1,2,4-Trimethylbenzene	50.000	47.261	5.5	90	0.00
91 T	sec-Butylbenzene	50.000	47.415	5.2	91	0.00
92 T	p-Isopropyltoluene	50.000	47.492	5.0	90	0.00
93 T	1,3-Dichlorobenzene	50.000	46.627	6.7	91	0.00
94 T	1,4-Dichlorobenzene	50.000	45.564	8.9	90	0.00
95 T	n-Butylbenzene	50.000	47.107	5.8	89	0.00
96 T	1,2-Dichlorobenzene	50.000	46.550	6.9	91	0.00
97 T	1,2-Dibromo-3-Chloropropane	50.000	44.428	11.1	86	0.00
98 T	1,2,4-Trichlorobenzene	50.000	46.835	6.3	90	0.00
99 T	Hexachlorobutadiene	50.000	45.030	9.9	86	0.00
100 T	Naphthalene	50.000	46.856	6.3	88	0.00
101 T	1,2,3-Trichlorobenzene	50.000	46.282	7.4	89	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 0
 8M429083.D 8260WTR.M Tue Mar 12 08:55:11 2019

Page 2

2.1.1.5 Raw QC Data

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\111218\8M427929.D Vial: 1
 Acq On : 12 Nov 2018 13:12 Operator: EEA
 Sample : WG684281-01 50ng/ BFB STD Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: rteint.p
 Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\A9FOOWT.M (RTE Integrator)
 Title : A9-FOO Water SOP:MSV01 111218 HPMS8



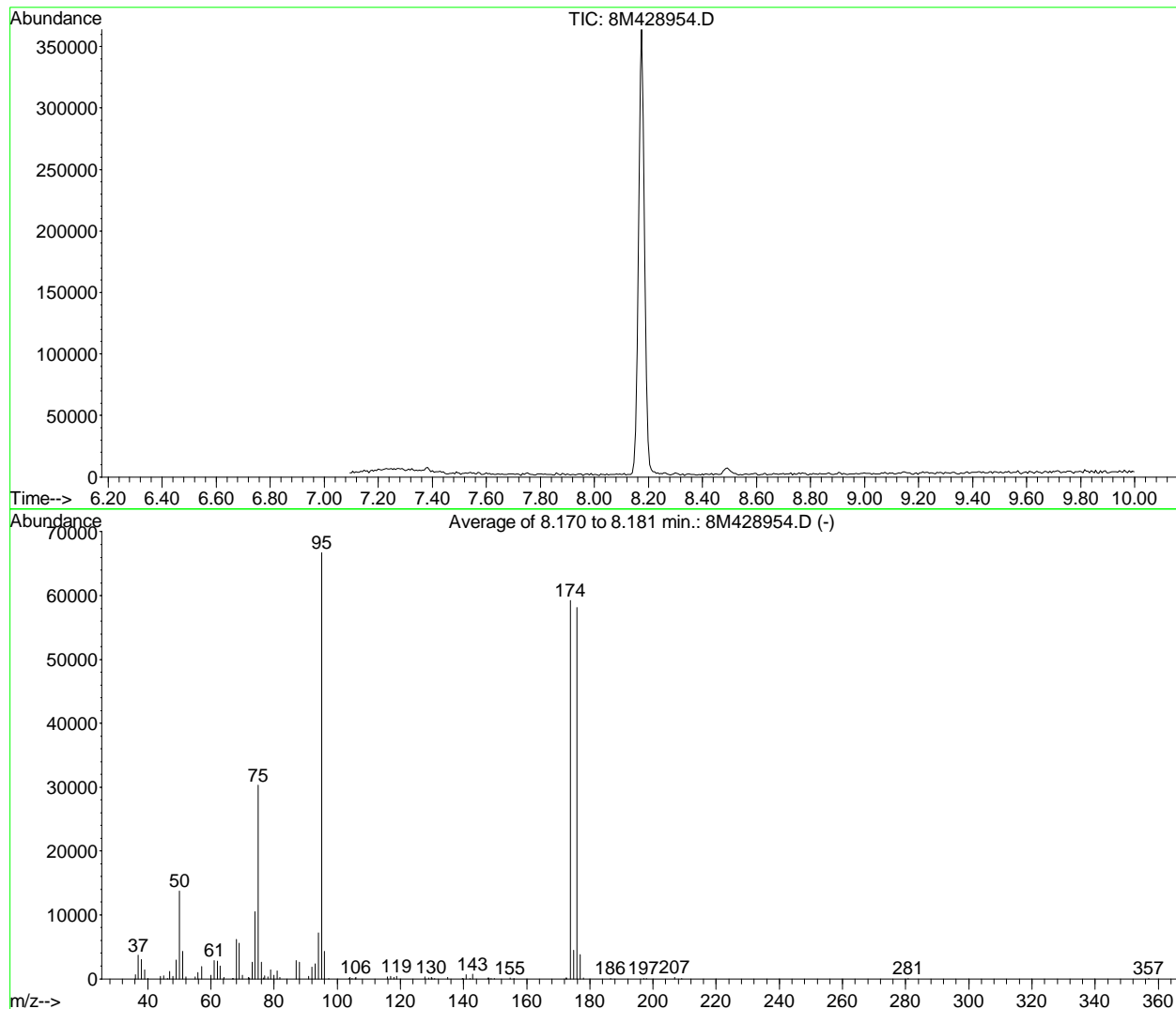
AutoFind: Scans 189, 190, 191; Background Corrected with Scan 181

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	15	40	22.0	18596	PASS
75	95	30	60	48.9	41416	PASS
95	95	100	100	100.0	84642	PASS
96	95	5	9	7.0	5906	PASS
173	174	0.00	2	0.3	193	PASS
174	95	50	100	89.2	75469	PASS
175	174	5	9	7.0	5319	PASS
176	174	95	101	99.7	75216	PASS
177	176	5	9	6.9	5164	PASS

8M427929.D A9FOOWT.M Tue Nov 13 12:15:58 2018

BFB

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\030419\8M428954.D Vial: 1
 Acq On : 4 Mar 2019 11:54 Operator: EEA
 Sample : WG698192-01 50ng BFB 8260 Inst : HPMS8
 Misc : 1,1 STD91948 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8



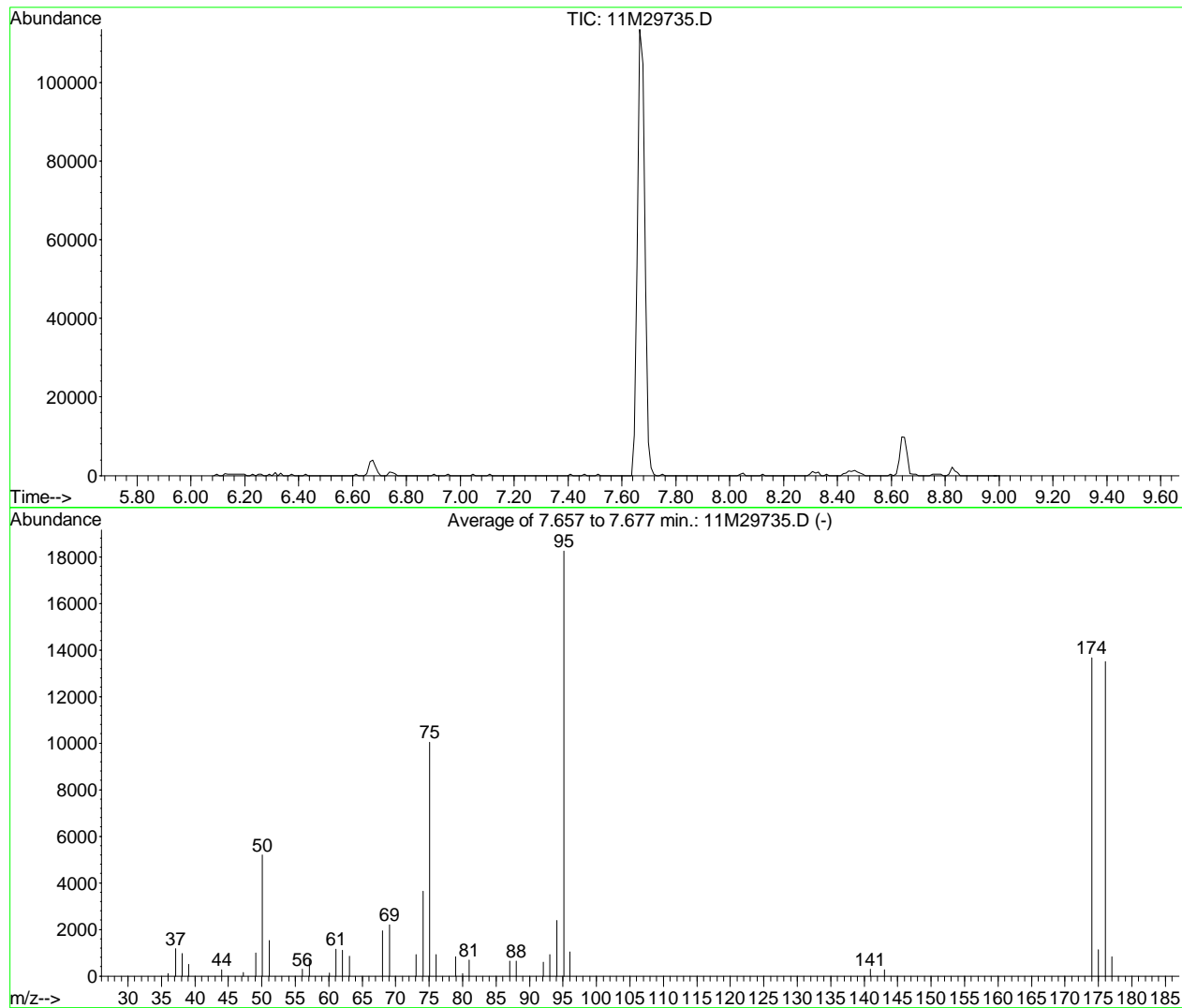
AutoFind: Scans 184, 185, 186; Background Corrected with Scan 176

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	15	40	20.6	13778	PASS
75	95	30	60	45.4	30314	PASS
95	95	100	100	100.0	66746	PASS
96	95	5	9	6.5	4307	PASS
173	174	0.00	2	0.2	137	PASS
174	95	50	100	88.7	59210	PASS
175	174	5	9	7.6	4497	PASS
176	174	95	101	98.1	58093	PASS
177	176	5	9	6.5	3763	PASS

8M428954.D 8260WTR.M Tue Mar 05 10:49:48 2019

BFB

Data File : C:\MSDCHEM\1\DATA\030519\11M29735.D Vial: 1
 Acq On : 5 Mar 2019 15:39 Operator: KFR
 Sample : WG698387-01 50ng BFB 8260 Inst : hpms11
 Misc : 1,1 STD91899 Multiplr: 1.00
 MS Integration Params: rteint.p
 Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11



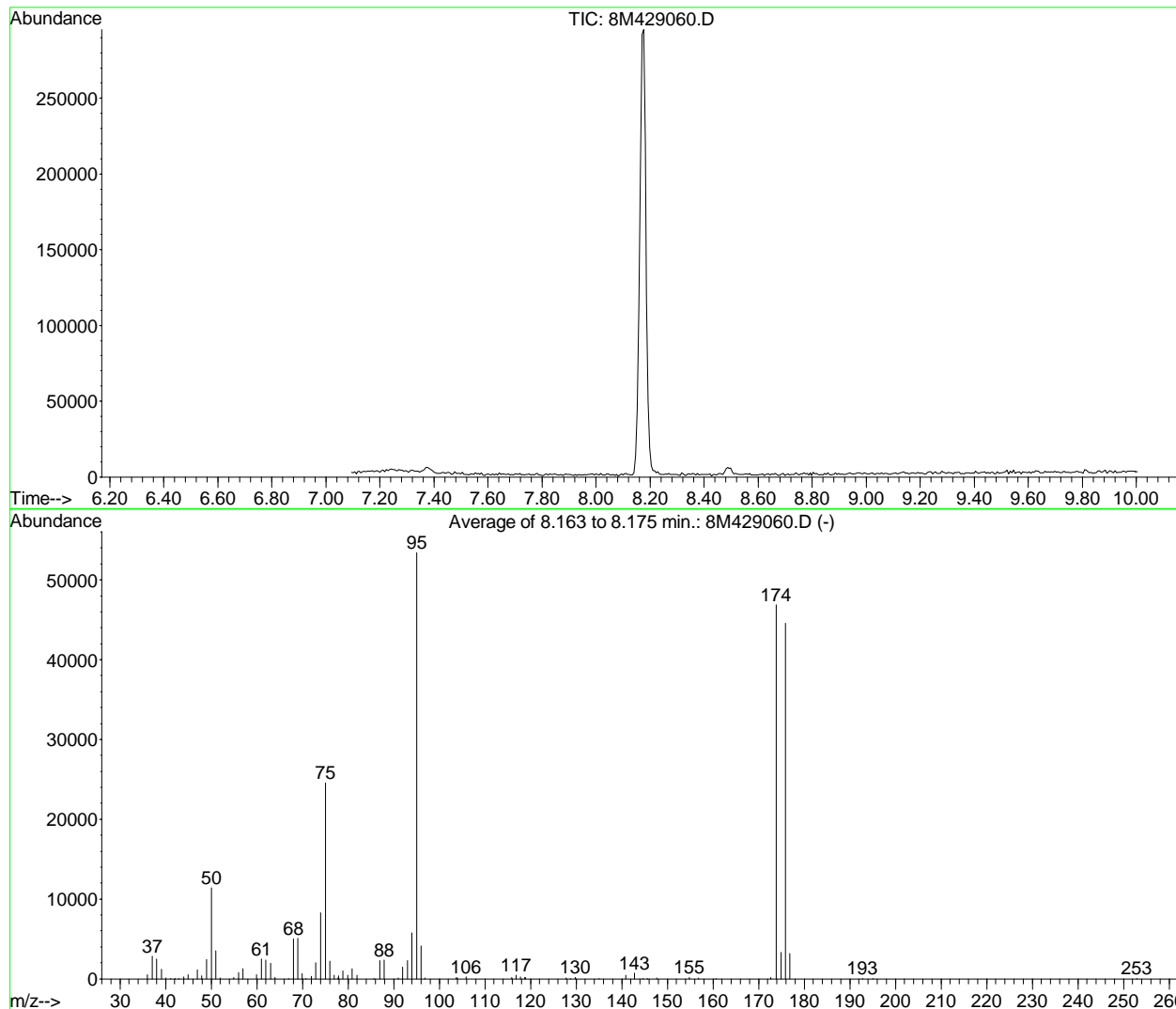
AutoFind: Scans 153, 154, 155; Background Corrected with Scan 149

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	15	40	28.5	5208	PASS
75	95	30	60	55.0	10035	PASS
95	95	100	100	100.0	18251	PASS
96	95	5	9	5.7	1034	PASS
173	174	0.00	2	0.0	0	PASS
174	95	50	100	74.8	13653	PASS
175	174	5	9	8.3	1135	PASS
176	174	95	101	98.9	13499	PASS
177	176	5	9	6.2	837	PASS

11M29735.D 8260WT.M Thu Mar 07 09:41:24 2019

BFB

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429060.D Vial: 1
 Acq On : 11 Mar 2019 8:56 Operator: EEA
 Sample : WG699027-01 50ng BFB 8260 Inst : HPMS8
 Misc : 1,1 STD92421 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8



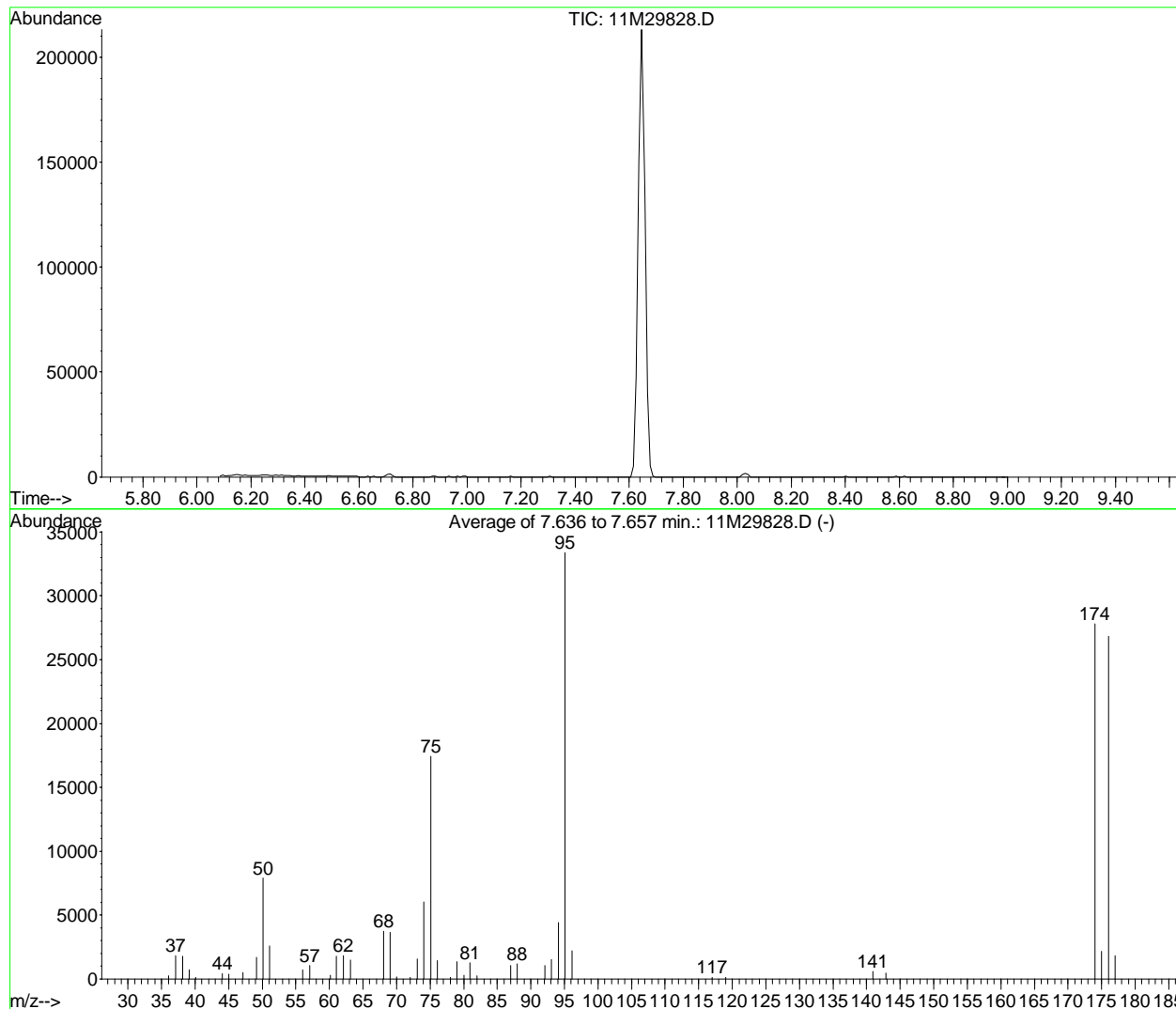
AutoFind: Scans 183, 184, 185; Background Corrected with Scan 176

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	15	40	21.3	11408	PASS
75	95	30	60	45.9	24533	PASS
95	95	100	100	100.0	53437	PASS
96	95	5	9	7.7	4118	PASS
173	174	0.00	2	0.4	184	PASS
174	95	50	100	87.8	46904	PASS
175	174	5	9	7.1	3330	PASS
176	174	95	101	95.1	44586	PASS
177	176	5	9	7.2	3188	PASS

8M429060.D 8260WTR.M Mon Mar 11 15:30:57 2019

BFB

Data File : C:\MSDCHEM\1\DATA\031219\11M29828.D Vial: 1
 Acq On : 12 Mar 2019 10:56 Operator: KFR
 Sample : WG699218-01 50ng BFB 8260 Inst : hpms11
 Misc : 1,1 STD92290 Multiplr: 1.00
 MS Integration Params: rteint.p
 Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11



AutoFind: Scans 151, 152, 153; Background Corrected with Scan 146

Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	15	40	23.7	7906	PASS
75	95	30	60	52.2	17441	PASS
95	95	100	100	100.0	33392	PASS
96	95	5	9	6.5	2173	PASS
173	174	0.00	2	0.0	0	PASS
174	95	50	100	83.3	27800	PASS
175	174	5	9	7.7	2131	PASS
176	174	95	101	96.6	26850	PASS
177	176	5	9	6.7	1801	PASS

11M29828.D 8260WT.M Tue Mar 12 13:42:38 2019

Data File : C:\MSDCHEM\1\DATA\031219\11M29831.D Vial: 3
 Acq On : 12 Mar 2019 12:31 Operator: KFR
 Sample : WG699219-01 BLANK 8260 Inst : hpms11
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 16 10:12:49 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3911	96	428800	25.0000	ug/L	0.0000
56) Chlorobenzene-d5	14.0203	117	314358	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8223	152	149164	25.0000	ug/L	-0.0107
System Monitoring Compounds						
37) Dibromofluoromethane	9.3985	111	122393	24.7269	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	98.9076%	
43) 1,2-Dichloroethane-d4	10.0085	65	142125	23.9311	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	95.7244%	
57) Toluene-d8	12.2418	98	399276	25.5219	ug/L	-0.0107
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	102.0876%	
78) p-Bromofluorobenzene	15.4161	95	151075	26.4662	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	105.8648%	
Target Compounds						
36) Tetrahydrofuran	9.3778	42	1222	0.9446	ug/L #	37

(#) = qualifier out of range (m) = manual integration
 11M29831.D 8260WT.M Sat Mar 16 10:12:50 2019

Page 1

Data File : C:\MSDCHEM\1\DATA\031219\11M29831.D

Vial: 3

Acq On : 12 Mar 2019 12:31

Operator: KFR

Sample : WG699219-01 BLANK 8260

Inst : hpms11

Misc : 1,1

Multiplr: 1.00

MS Integration Params: rteint.p

Quant Time: Mar 16 10:12 2019

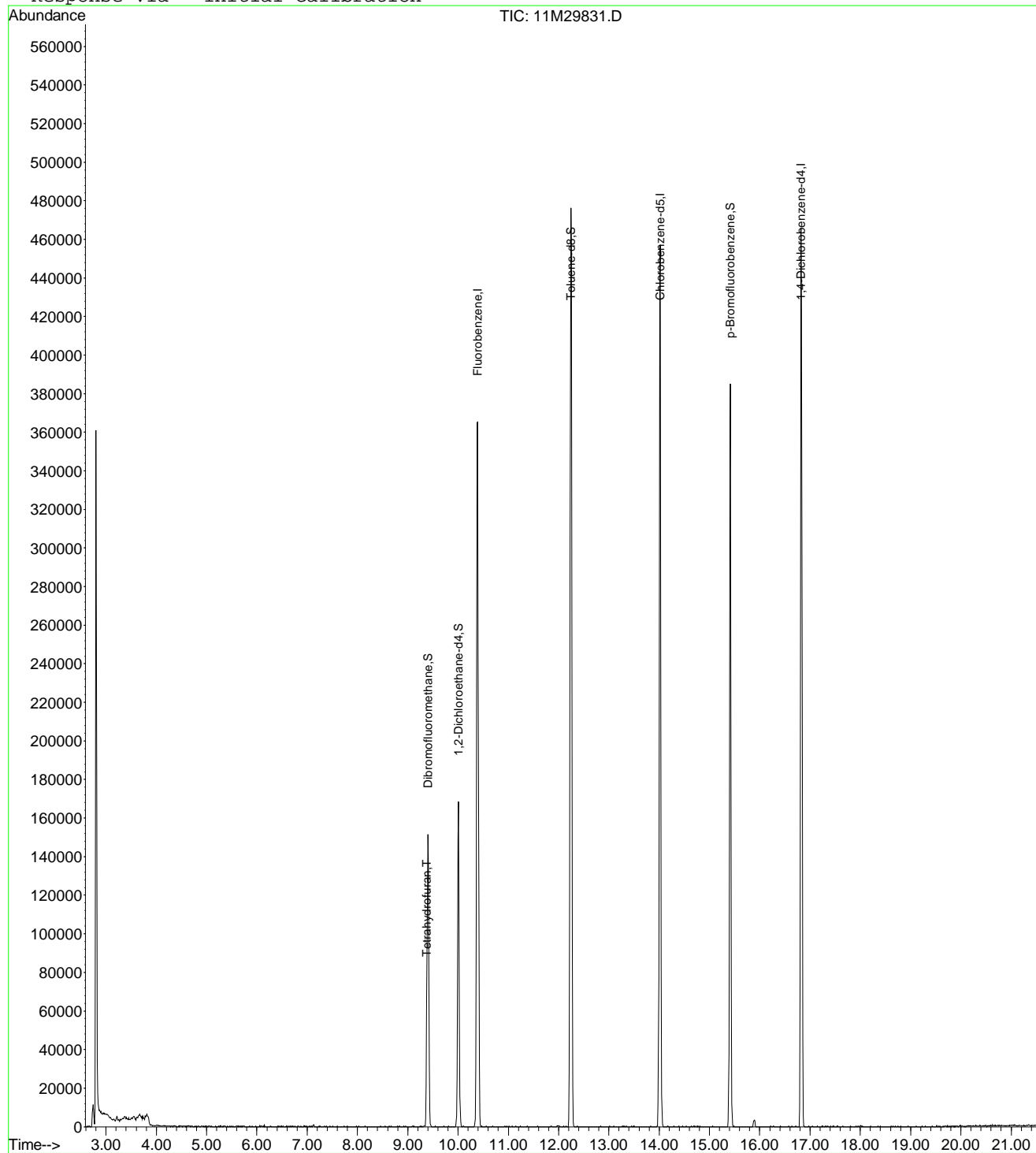
Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)

Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11

Last Update : Wed Mar 06 14:13:24 2019

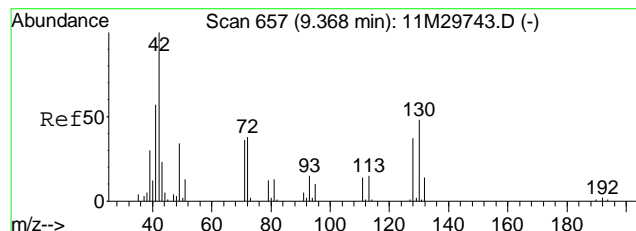
Response via : Initial Calibration



11M29831.D 8260WT.M

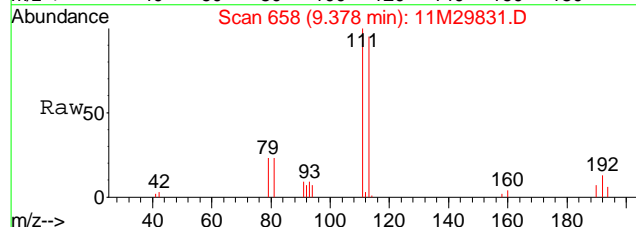
Sat Mar 16 10:12:50 2019

Page 2

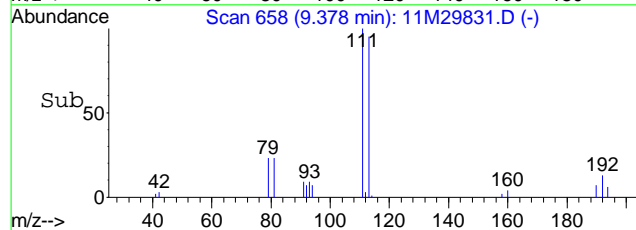
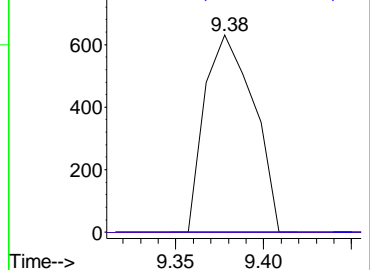


#36
 Tetrahydrofuran
 Concen: 0.9446 ug/L
 RT: 9.38 min Scan# 658
 Delta R.T. 0.01 min
 Lab File: 11M29831.D
 Acq: 12 Mar 2019 12:31

Tgt Ion	Ratio	Lower	Upper
42	100		
71	0.0	22.1	51.7#
72	0.0	22.9	53.3#



Abundance Ion 42.00 (41.70 to 42.70): 11
 Ion 71.00 (70.70 to 71.70): 11
 Ion 72.00 (71.70 to 72.70): 11



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429063.D Vial: 3
 Acq On : 11 Mar 2019 10:41 Operator: EEA
 Sample : WG699028-01 BLANK 8260 Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 09:00:08 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

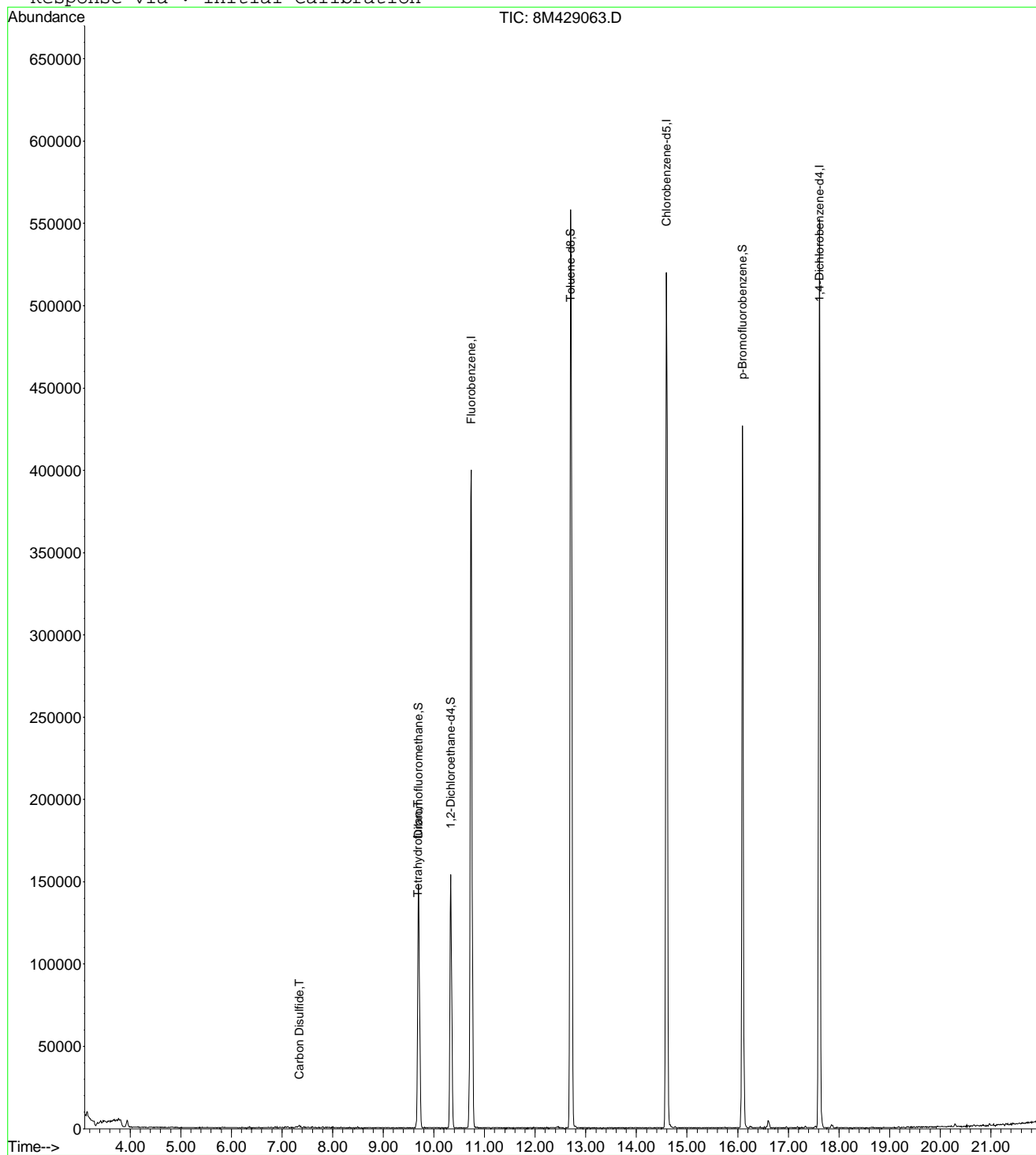
Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.74	96	508811	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.59	117	381131	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.61	152	196164	25.00	ug/L	0.00
System Monitoring Compounds						
37) Dibromofluoromethane	9.69	111	130977	25.8244	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	103.28%	
43) 1,2-Dichloroethane-d4	10.33	65	129467	25.8573	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	103.44%	
58) Toluene-d8	12.70	98	504906	26.0072	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	104.04%	
80) p-Bromofluorobenzene	16.09	95	193043	27.2669	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	109.08%	
Target Compounds						Qvalue
20) Carbon Disulfide	7.35	76	2807	0.1383	ug/L #	74
36) Tetrahydrofuran	9.68	42	1365	0.8787	ug/L #	40

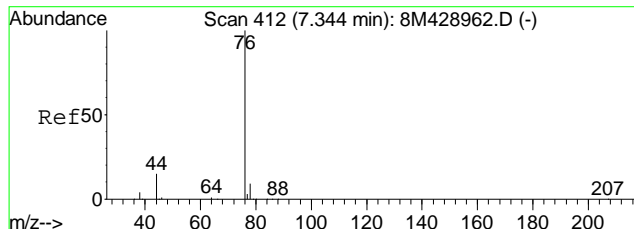
(#) = qualifier out of range (m) = manual integration
 8M429063.D 8260WTR.M Tue Mar 12 09:00:09 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429063.D Vial: 3
 Acq On : 11 Mar 2019 10:41 Operator: EEA
 Sample : WG699028-01 BLANK 8260 Inst : HPMS8
 Misc : 1,1 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 12 8:59 2019 Quant Results File: 8260WTR.RES

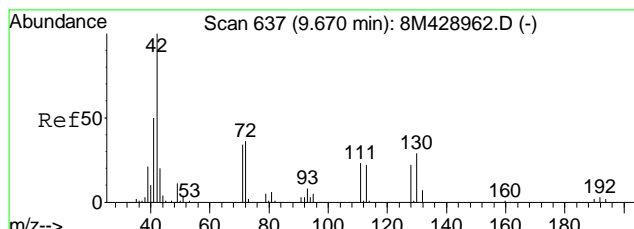
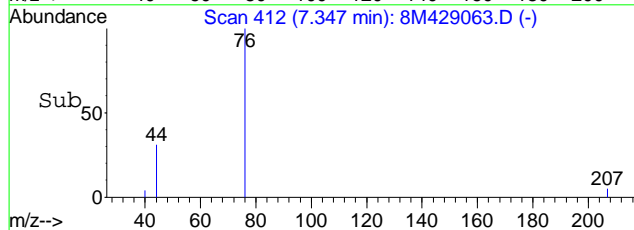
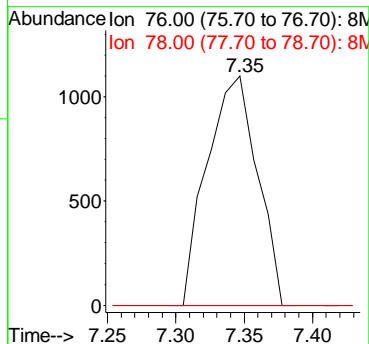
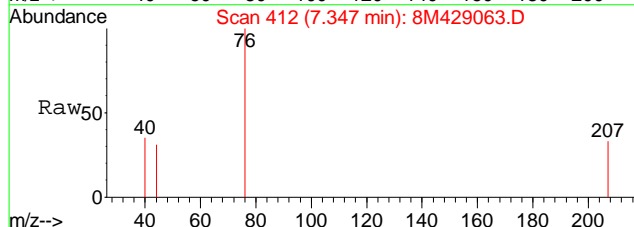
Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration





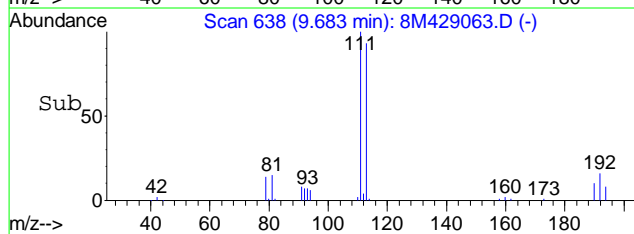
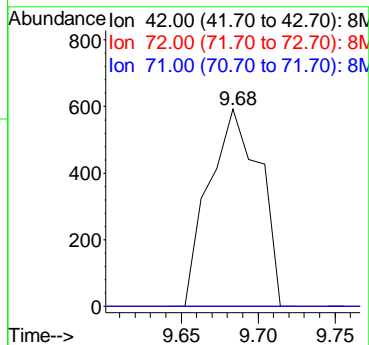
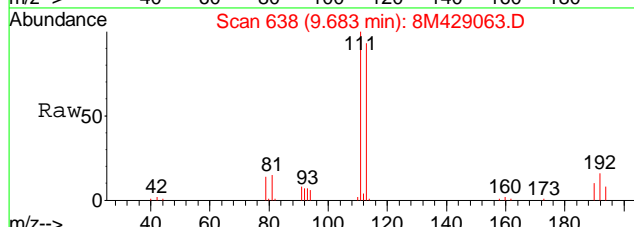
#20
 Carbon Disulfide
 Concen: 0.14 ug/L
 RT: 7.35 min Scan# 412
 Delta R.T. 0.00 min
 Lab File: 8M429063.D
 Acq: 11 Mar 2019 10:41

Tgt Ion	Ratio	Lower	Upper
76	100		
78	0.0	5.6	13.0#



#36
 Tetrahydrofuran
 Concen: 0.88 ug/L
 RT: 9.68 min Scan# 638
 Delta R.T. 0.01 min
 Lab File: 8M429063.D
 Acq: 11 Mar 2019 10:41

Tgt Ion	Ratio	Lower	Upper
42	100		
72	0.0	21.9	51.1#
71	0.0	20.2	47.2#



Data File : C:\MSDCHEM\1\DATA\031219\11M29832.D Vial: 4
 Acq On : 12 Mar 2019 13:00 Operator: KFR
 Sample : WG699219-02 LCS 20ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 13:42:01 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.3812	96	402710	25.0000	ug/L	-0.0103
56) Chlorobenzene-d5	14.0207	117	309185	25.0000	ug/L	0.0000
76) 1,4-Dichlorobenzene-d4	16.8227	152	156356	25.0000	ug/L	-0.0103

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.3989	111	117749	25.3299	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 118	Recovery	=	101.3196%	
43) 1,2-Dichloroethane-d4	10.0090	65	137384	24.6315	ug/L	0.0000
Spiked Amount	25.0000	Range 80 - 120	Recovery	=	98.5260%	
57) Toluene-d8	12.2423	98	383757	24.9404	ug/L	-0.0103
Spiked Amount	25.0000	Range 88 - 110	Recovery	=	99.7616%	
78) p-Bromofluorobenzene	15.4166	95	149808	25.0371	ug/L	0.0000
Spiked Amount	25.0000	Range 86 - 115	Recovery	=	100.1484%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.1124	85	88271	14.8376	ug/L	99
3) Chloromethane	3.5467	50	144669	12.6611	ug/L	99
4) Vinyl Chloride	3.7639	62	130239	15.2685	ug/L	100
5) 1,3-Butadiene	3.8052	54	49375	14.8118	ug/L	99
6) Bromomethane	4.6324	94	72299	22.1231	ug/L	99
7) Chloroethane	4.7875	64	60993	18.4960	ug/L	100
8) Trichlorofluoromethane	5.2631	101	148260	18.3673	ug/L	98
9) Diethyl ether	5.7801	59	407724	96.6144	ug/L	97
10) Isoprene	5.8111	67	120889	19.0144	ug/L	96
11) Acrolein	6.0075	56	63824	87.0908	ug/L	98
12) 1,1,2-Trichloro-1,2,2-Trif	6.0282	101	80948	20.4433	ug/L	98
13) Acetone	6.1109	43	21691	14.7329	ug/L	98
14) 1,1-Dichloroethene	6.3281	61	148388	19.2551	ug/L	96
15) Tert-Butyl Alcohol	6.4418	59	64261	156.4824	ug/L	99
16) Dimethyl Sulfide	6.5865	62	106646	20.9336	ug/L	97
17) Iodomethane	6.8244	142	89750	17.9109	ug/L	98
18) Methyl acetate	6.8450	43	54867	17.2771	ug/L	99
19) Methylene Chloride	7.0828	84	87965	20.4831	ug/L	89
20) Carbon Disulfide	7.1242	76	205391	17.3524	ug/L	99
21) Acrylonitrile	7.2690	53	30006	18.4877	ug/L	96
22) Methyl Tert Butyl Ether	7.2896	73	226316	18.5845	ug/L	99
23) trans-1,2-Dichloroethene	7.5275	96	87451	20.9487	ug/L	96
24) n-Hexane	7.5998	57	113265	17.6155	ug/L	99
25) Diisopropyl ether	7.9307	45	1551747	98.8815	ug/L	99
26) Vinyl Acetate	8.0961	43	141949	26.2713	ug/L	98
27) 1,1-Dichloroethane	8.1168	63	178564	19.6024	ug/L	99
28) Ethyl-Tert-Butyl ether	8.4787	59	1624371	102.4076	ug/L	99
29) 2-Butanone	8.6545	43	33989	18.1240	ug/L	99
30) Propionitrile	8.7579	54	51314	93.6519	ug/L	99
31) 2,2-Dichloropropane	8.8613	77	144234	22.9001	ug/L	100
32) cis-1,2-Dichloroethene	8.9233	96	100581	21.2228	ug/L	94
33) Chloroform	9.1197	83	165133	19.9724	ug/L	100
34) 1-Bromopropane	9.2542	122	20850	20.3113	ug/L	96
35) Bromochloromethane	9.3472	130	58985	21.2262	ug/L	96
36) Tetrahydrofuran	9.3679	42	111623	91.8719	ug/L	97
38) 1,1,1-Trichloroethane	9.6264	97	164645	21.9923	ug/L	97
39) Cyclohexane	9.6574	56	139264	15.7121	ug/L	96
40) 1,1-Dichloropropene	9.8125	75	117521	20.8782	ug/L	99
41) Carbon Tetrachloride	9.9573	117	145340	22.8298	ug/L	99
42) Tert-Amyl-Methyl ether	9.9159	73	1238293	104.6686	ug/L	99

(#) = qualifier out of range (m) = manual integration
 11M29832.D 8260WT.M Tue Mar 12 13:42:02 2019

Data File : C:\MSDCHEM\1\DATA\031219\11M29832.D Vial: 4
 Acq On : 12 Mar 2019 13:00 Operator: KFR
 Sample : WG699219-02 LCS 20ug/L 8260 Inst : hpms11
 Misc : 1,1 STD92316 Multiplr: 1.00
 MS Integration Params: rteint.p
 Quant Time: Mar 12 13:42:01 2019 Quant Results File: 8260WT.RES

Quant Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
 Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
 Last Update : Wed Mar 06 14:13:24 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
44) 1,2-Dichloroethane	10.1123	62	144963	20.1725	ug/L	99
45) Benzene	10.1537	78	341887	20.4347	ug/L	99
46) Trichloroethene	10.8671	130	95368	20.0981	ug/L	100
47) Methylcyclohexane	10.9395	83	122605	18.3411	ug/L	96
48) 1,2-Dichloropropane	11.0636	63	99560	20.0326	ug/L	97
49) 1,4-Dioxane	11.3428	88	3524	103.3397	ug/L	86
50) Bromodichloromethane	11.3531	83	122553	20.8656	ug/L	99
51) Dibromomethane	11.4255	93	57847	20.3765	ug/L	96
52) 2-Chloroethyl Vinyl Ether	11.6219	63	39012	16.2208	ug/L	99
53) 4-Methyl-2-Pentanone	11.6530	58	31500	18.3839	ug/L	97
54) cis-1,3-Dichloropropene	11.9424	75	144696	21.8589	ug/L	100
55) Dimethyl Disulfide	12.1906	79	78617	19.1926	ug/L	92
58) Toluene	12.3354	91	377129	20.5906	ug/L	98
59) Ethyl Methacrylate	12.4284	69	102550	19.8233	ug/L	98
60) trans-1,3-Dichloropropene	12.5008	75	122509	20.4291	ug/L	100
61) 1,1,2-Trichloroethane	12.7076	97	74876	19.2034	ug/L	99
62) 2-Hexanone	12.6455	43	52686	18.1792	ug/L #	73
63) 1,3-Dichloropropane	12.9971	76	128467	20.4209	ug/L	97
64) Tetrachloroethene	13.1108	164	77633	19.9828	ug/L	99
65) Dibromochloromethane	13.3590	129	93894	21.1541	ug/L	98
66) 1,2-Dibromoethane	13.5968	107	76127	20.3720	ug/L	99
67) 1-Chlorohexane	13.6692	91	115894	19.1660	ug/L	96
68) Chlorobenzene	14.0621	112	256481	20.1138	ug/L	99
69) 1,1,1,2-Tetrachloroethane	14.0931	131	101030	19.0623	ug/L	99
70) Ethylbenzene	14.0828	106	130731	20.1530	ug/L	96
71) m-,p-Xylene	14.1655	106	322539	41.1854	ug/L	94
72) o-Xylene	14.6928	106	158802	20.8992	ug/L	99
73) Styrene	14.7238	104	267340	21.2706	ug/L	99
74) Bromoform	15.1994	173	58569	18.7671	ug/L	98
75) Isopropylbenzene	15.0857	105	391966	20.6062	ug/L	100
77) 1,1,2,2-Tetrachloroethane	15.2925	83	85109	20.8659	ug/L	99
79) 1,2,3-Trichloropropane	15.4682	110	28090	20.6251	ug/L	76
80) trans-1,4-Dichloro-2-Butene	15.5096	53	31188	18.1623	ug/L	85
81) n-Propylbenzene	15.5613	91	503230	21.8247	ug/L	99
82) Bromobenzene	15.6854	156	113791	20.2076	ug/L	75
83) 1,3,5-Trimethylbenzene	15.7267	105	348221	21.2732	ug/L	99
84) 2-Chlorotoluene	15.8198	91	342570	20.8791	ug/L	99
85) 4-Chlorotoluene	15.8612	91	263859	19.5668	ug/L	100
86) a-Methylstyrene	16.1093	118	195280	20.9512	ug/L	99
87) tert-Butylbenzene	16.1610	134	80869	20.0584	ug/L	97
88) 1,2,4-Trimethylbenzene	16.2127	105	350403	20.7410	ug/L	99
89) sec-Butylbenzene	16.4092	105	435408	21.4256	ug/L	99
90) p-Isopropyltoluene	16.5642	119	380149	21.5130	ug/L	99
91) 1,3-Dichlorobenzene	16.7504	146	210729	20.2859	ug/L	98
92) 1,4-Dichlorobenzene	16.8641	146	217067	20.9256	ug/L	100
93) n-Butylbenzene	17.0502	91	337982	21.2233	ug/L	100
94) 1,2-Dichlorobenzene	17.3294	146	209056	20.9983	ug/L	98
95) 1,2-Dibromo-3-Chloropropane	18.2496	75	16737	19.8630	ug/L	95
96) 1,2,4-Trichlorobenzene	19.3146	180	135928	21.1057	ug/L	99
97) Hexachlorobutadiene	19.4490	225	70536	21.7298	ug/L	98
98) Naphthalene	19.6558	128	291099	21.1082	ug/L	100
99) 1,2,3-Trichlorobenzene	19.9453	180	126570	20.9179	ug/L	99

(#) = qualifier out of range (m) = manual integration
 11M29832.D 8260WT.M Tue Mar 12 13:42:02 2019

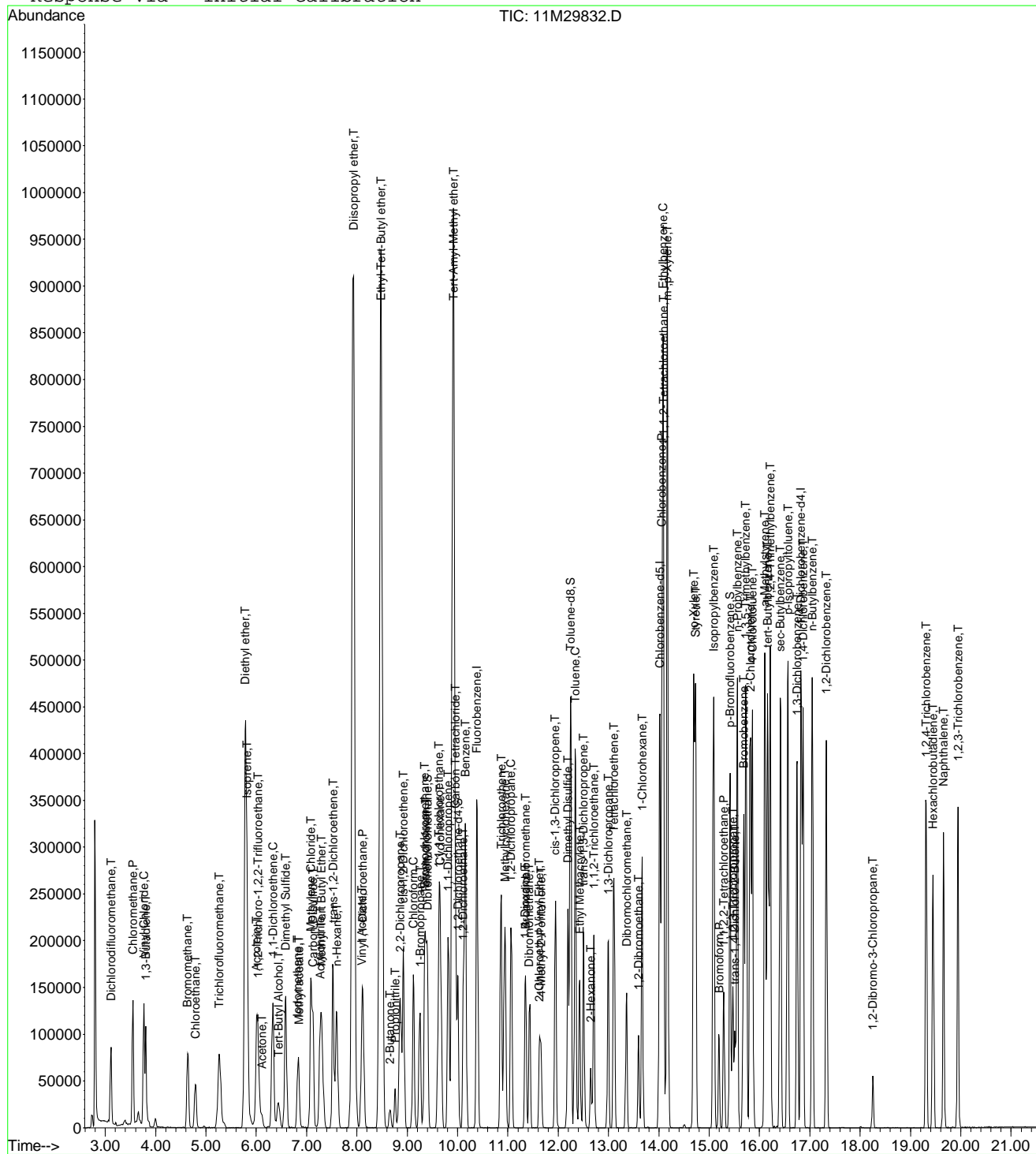
Page 2

Data File : C:\MSDCHEM\1\DATA\031219\11M29832.D
Acq On : 12 Mar 2019 13:00
Sample : WG699219-02 LCS 20ug/L 8260
Misc : 1,1 STD92316
MS Integration Params: rteint.p
Quant Time: Mar 12 13:42 2019

Vial: 4
Operator: KFR
Inst : hpms11
Multiplr: 1.00

Quant Results File: 8260WT.RES

Method : C:\MSDCHEM\1\METHODS\8260WT.M (RTE Integrator)
Title : 8260B(SOP: OVL MSV01) Water 030519 HPMS11
Last Update : Wed Mar 06 14:13:24 2019
Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429064.D Vial: 4
 Acq On : 11 Mar 2019 11:09 Operator: EEA
 Sample : WG699028-02 LCS 20 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:31:47 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.73	96	523789	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	399992	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.61	152	215609	25.00	ug/L	-0.01

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.70	111	135824	26.0143	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	104.04%	
43) 1,2-Dichloroethane-d4	10.33	65	135138	26.2181	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	104.88%	
58) Toluene-d8	12.71	98	517518	25.3999	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	101.60%	
80) p-Bromofluorobenzene	16.09	95	202468	26.0190	ug/L	-0.01
Spiked Amount	25.000	Range 86 - 115	Recovery	=	104.08%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.23	85	151396	16.6175	ug/L	100
3) Chloromethane	3.68	50	224720	17.1511	ug/L	99
4) Vinyl Chloride	3.91	62	199262	18.6614	ug/L	99
5) 1,3-Butadiene	3.95	54	135009	19.2929	ug/L	99
6) Bromomethane	4.80	94	122166	19.0217	ug/L	99
7) Chloroethane	4.96	64	115532	20.2192	ug/L	99
8) Trichlorofluoromethane	5.44	101	200977	18.5380	ug/L	100
9) Diethyl ether	5.96	59	572928	99.0886	ug/L	99
10) Isoprene	6.00	67	209562	19.3412	ug/L	99
11) Acrolein	6.20	56	83083	125.3129	ug/L	98
12) 1,1,2-Trichloro-1,2,2-Trif	6.22	101	127970	20.4894	ug/L	100
13) Acetone	6.31	43	23651	14.6581	ug/L	98
14) 1,1-Dichloroethene	6.52	61	224151	20.0663	ug/L	99
15) Tert-Butyl Alcohol	6.63	59	60801	117.0653	ug/L	99
16) Dimethyl Sulfide	6.79	62	169657	21.0845	ug/L	98
17) Iodomethane	7.03	142	149591	19.2059	ug/L	99
18) Methyl acetate	7.05	43	70693	16.9201	ug/L	98
19) Methylene Chloride	7.29	84	140627	20.1469	ug/L	98
20) Carbon Disulfide	7.34	76	365449	17.4955	ug/L	100
21) Acrylonitrile	7.49	53	35240	17.4085	ug/L	100
22) Methyl Tert Butyl Ether	7.51	73	310032	18.6606	ug/L	99
23) trans-1,2-Dichloroethene	7.75	61	218612	20.4815	ug/L	100
24) n-Hexane	7.82	57	202859	19.3658	ug/L	100
25) Diisopropyl ether	8.16	45	2748749	104.0280	ug/L	99
26) Vinyl Acetate	8.34	43	232411	23.7108	ug/L	98
27) 1,1-Dichloroethane	8.37	63	282697	20.4881	ug/L	100
28) Ethyl-Tert-Butyl ether	8.73	59	2305047	100.6513	ug/L	99
29) 2-Butanone	8.93	43	41743	17.5623	ug/L	94
30) Propionitrile	9.03	54	61071	84.5930	ug/L	98
31) 2,2-Dichloropropane	9.13	77	212405	21.2152	ug/L	99
32) cis-1,2-Dichloroethene	9.20	96	158023	20.7432	ug/L	97
33) Chloroform	9.41	83	245335	20.0335	ug/L	100
34) 1-Bromopropane	9.55	122	32497	19.8449	ug/L	99
35) Bromochloromethane	9.64	130	87672	20.1790	ug/L	98
36) Tetrahydrofuran	9.67	42	151475	94.7203	ug/L	98
38) 1,1,1-Trichloroethane	9.94	97	226239	21.2694	ug/L	100
39) Cyclohexane	9.97	56	240478	16.9739	ug/L	99
40) 1,1-Dichloropropene	10.14	75	192634	21.5669	ug/L	100
41) Tert-Amyl-Methyl ether	10.24	73	1765666	101.1726	ug/L	100
42) Carbon Tetrachloride	10.28	117	197841	20.4684	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M429064.D 8260WTR.M Mon Mar 11 15:31:48 2019

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429064.D Vial: 4
 Acq On : 11 Mar 2019 11:09 Operator: EEA
 Sample : WG699028-02 LCS 20 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:31:47 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

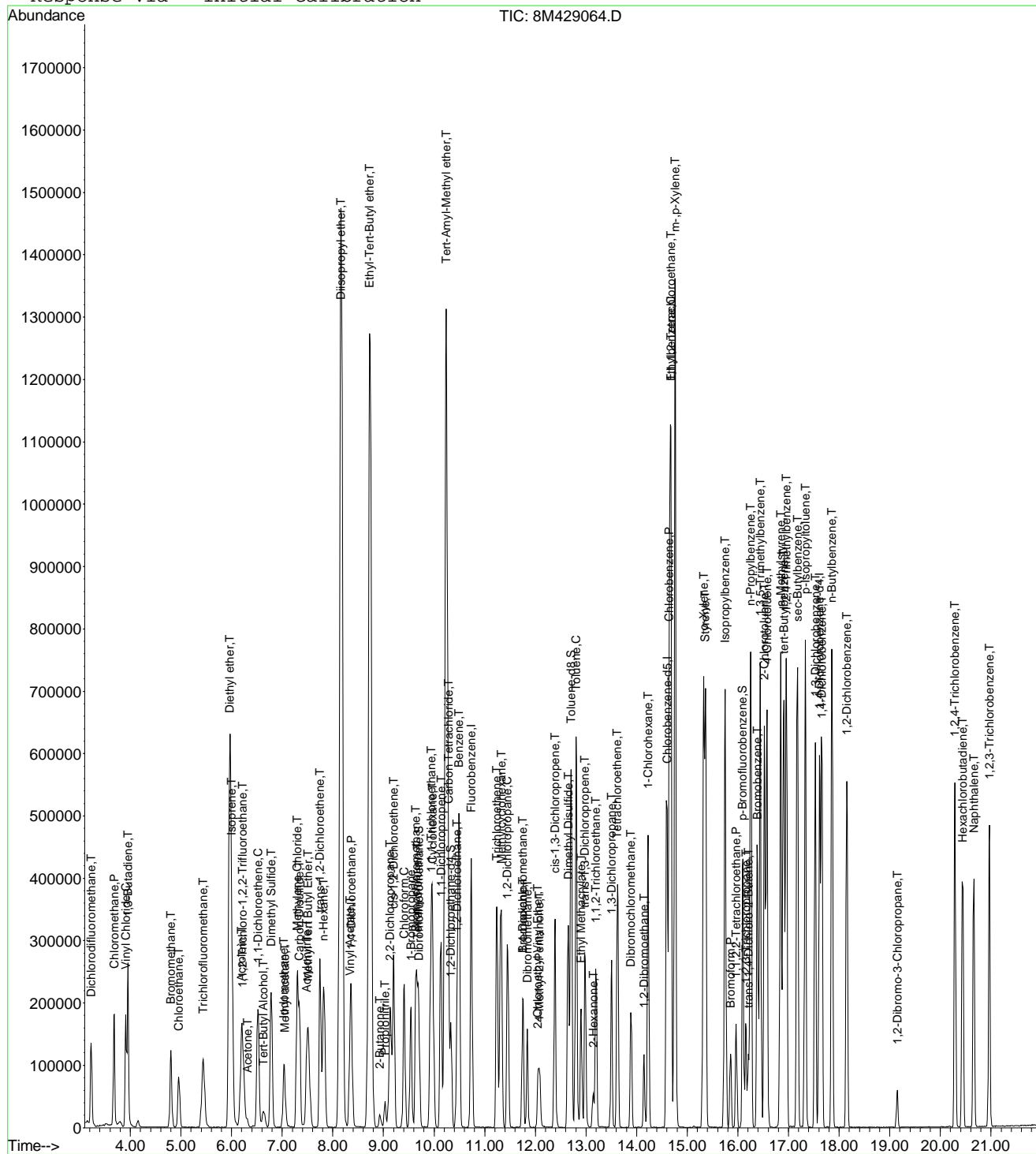
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	174114	20.7862	ug/L	100
46) Benzene	10.49	78	588082	21.0178	ug/L	99
47) Trichloroethene	11.24	130	159454	20.9172	ug/L	100
48) Methylcyclohexane	11.32	83	219888	18.6256	ug/L	99
49) 1,2-Dichloropropane	11.45	63	159717	21.0297	ug/L	98
50) Bromodichloromethane	11.75	83	176027	20.0171	ug/L	99
51) 1,4-Dioxane	11.75	88	3800	88.9214	ug/L	89
52) Dibromomethane	11.84	93	71088	20.3351	ug/L	99
53) 2-Chloroethyl Vinyl Ether	12.05	63	41068	12.3751	ug/L	99
54) 4-Methyl-2-Pentanone	12.08	58	37538	17.9788	ug/L	98
55) cis-1,3-Dichloropropene	12.39	75	231886	22.1646	ug/L	100
56) Dimethyl Disulfide	12.65	79	118828	18.4958	ug/L	92
59) Toluene	12.80	91	607977	20.3575	ug/L	99
60) Ethyl Methacrylate	12.91	69	139408	18.2852	ug/L	99
62) trans-1,3-Dichloropropene	12.98	75	183852	20.4770	ug/L	100
63) 1,1,2-Trichloroethane	13.20	97	105666	20.2413	ug/L	99
64) 2-Hexanone	13.14	58	30903	16.4379	ug/L	83
65) 1,3-Dichloropropane	13.51	76	186335	20.7253	ug/L	99
66) Tetrachloroethene	13.62	164	130129	20.6762	ug/L	98
67) Dibromochloromethane	13.89	129	131379	19.8296	ug/L	99
68) 1,2-Dibromoethane	14.15	107	102243	20.1697	ug/L	99
69) 1-Chlorohexane	14.23	91	199422	19.2696	ug/L	98
70) Chlorobenzene	14.64	112	406390	20.0725	ug/L	100
71) 1,1,1,2-Tetrachloroethane	14.67	131	145735	19.5789	ug/L	99
72) Ethylbenzene	14.67	106	224357	20.4815	ug/L	97
73) m-,p-Xylene	14.76	106	554236	41.5247	ug/L	97
74) o-Xylene	15.33	106	273803	21.1416	ug/L	98
75) Styrene	15.36	104	450889	21.0292	ug/L	99
76) Bromoform	15.85	173	78900	19.2061	ug/L	99
77) Isopropylbenzene	15.75	105	673442	20.5873	ug/L	99
79) 1,1,2,2-Tetrachloroethane	15.97	83	111817	18.3827	ug/L	98
81) 1,2,3-Trichloropropane	16.16	110	36409	20.0088	ug/L	67
82) trans-1,4-Dichloro-2-Butene	16.20	53	34280	16.6709	ug/L #	26
83) n-Propylbenzene	16.26	91	841976	21.5785	ug/L	99
84) Bromobenzene	16.39	156	178031	19.9412	ug/L	98
85) 1,3,5-Trimethylbenzene	16.44	105	581393	20.6697	ug/L	100
86) 2-Chlorotoluene	16.53	91	489690	18.9808	ug/L	92
87) 4-Chlorotoluene	16.58	91	513565	22.7821	ug/L	90
88) a-Methylstyrene	16.85	118	329328	20.4041	ug/L	100
89) tert-Butylbenzene	16.91	134	132350	20.3956	ug/L	100
90) 1,2,4-Trimethylbenzene	16.96	105	583247	20.4816	ug/L	100
91) sec-Butylbenzene	17.18	105	765412	21.2627	ug/L	99
92) p-Isopropyltoluene	17.33	119	649564	21.0497	ug/L	99
93) 1,3-Dichlorobenzene	17.53	146	347215	19.9499	ug/L	99
94) 1,4-Dichlorobenzene	17.66	146	350068	19.8619	ug/L	98
95) n-Butylbenzene	17.86	91	585078	20.8090	ug/L	99
96) 1,2-Dichlorobenzene	18.15	146	325621	20.1613	ug/L	99
97) 1,2-Dibromo-3-Chloropropane	19.15	75	19791	18.0520	ug/L	98
98) 1,2,4-Trichlorobenzene	20.29	180	236870	20.2124	ug/L	100
99) Hexachlorobutadiene	20.44	225	115509	20.9178	ug/L	99
100) Naphthalene	20.66	128	417358	19.5364	ug/L	100
101) 1,2,3-Trichlorobenzene	20.97	180	213612	19.8422	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M429064.D 8260WTR.M Mon Mar 11 15:31:48 2019

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Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429064.D Vial: 4
Acq On : 11 Mar 2019 11:09 Operator: EEA
Sample : WG699028-02 LCS 20 ug/L 8260 Inst : HPMS8
Misc : 1,1 STD92354 Multiplr: 1.00
MS Integration Params: RTEINT.P
Quant Time: Mar 11 15:31 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
Last Update : Mon Mar 04 17:44:12 2019
Response via : Initial Calibration



Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429065.D Vial: 6
 Acq On : 11 Mar 2019 11:38 Operator: EEA
 Sample : WG699028-03 LCS DUP 20 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:31:49 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

Internal Standards	R.T.	QIon	Response	Conc	Units	Dev(Min)
1) Fluorobenzene	10.73	96	524940	25.00	ug/L	0.00
57) Chlorobenzene-d5	14.60	117	402726	25.00	ug/L	0.00
78) 1,4-Dichlorobenzene-d4	17.62	152	217474	25.00	ug/L	0.00

System Monitoring Compounds	R.T.	QIon	Response	Conc	Units	Dev(Min)
37) Dibromofluoromethane	9.70	111	138268	26.4243	ug/L	0.00
Spiked Amount	25.000	Range 86 - 118	Recovery	=	105.68%	
43) 1,2-Dichloroethane-d4	10.33	65	136855	26.4930	ug/L	0.00
Spiked Amount	25.000	Range 80 - 120	Recovery	=	105.96%	
58) Toluene-d8	12.71	98	523088	25.4989	ug/L	0.00
Spiked Amount	25.000	Range 88 - 110	Recovery	=	102.00%	
80) p-Bromofluorobenzene	16.10	95	200664	25.5660	ug/L	0.00
Spiked Amount	25.000	Range 86 - 115	Recovery	=	102.28%	

Target Compounds	R.T.	QIon	Response	Conc	Units	Qvalue
2) Dichlorodifluoromethane	3.23	85	146155	16.0070	ug/L	99
3) Chloromethane	3.68	50	211116	16.0775	ug/L	100
4) Vinyl Chloride	3.91	62	191666	17.9106	ug/L	100
5) 1,3-Butadiene	3.95	54	128152	18.0099	ug/L	99
6) Bromomethane	4.80	94	112116	17.4186	ug/L	99
7) Chloroethane	4.95	64	111164	19.4121	ug/L	100
8) Trichlorofluoromethane	5.44	101	194321	17.8847	ug/L	100
9) Diethyl ether	5.97	59	558937	96.4569	ug/L	99
10) Isoprene	6.00	67	203314	18.7234	ug/L	100
11) Acrolein	6.21	56	81683	122.9311	ug/L	98
12) 1,1,2-Trichloro-1,2,2-Trif	6.22	101	123936	19.8000	ug/L	99
13) Acetone	6.31	43	24068	14.8839	ug/L	97
14) 1,1-Dichloroethene	6.53	61	213760	19.0942	ug/L	99
15) Tert-Butyl Alcohol	6.64	59	65999	126.7948	ug/L	99
16) Dimethyl Sulfide	6.78	62	167223	20.7364	ug/L	98
17) Iodomethane	7.03	142	164731	20.9540	ug/L	100
18) Methyl acetate	7.05	43	71604	17.1006	ug/L	98
19) Methylene Chloride	7.30	84	136698	19.5410	ug/L	98
20) Carbon Disulfide	7.34	76	353950	16.9078	ug/L	100
21) Acrylonitrile	7.49	53	36740	18.1097	ug/L	97
22) Methyl Tert Butyl Ether	7.51	73	306992	18.4371	ug/L	99
23) trans-1,2-Dichloroethene	7.75	61	212602	19.8748	ug/L	100
24) n-Hexane	7.83	57	190889	18.1832	ug/L	98
25) Diisopropyl ether	8.16	45	2688389	101.5206	ug/L	99
26) Vinyl Acetate	8.34	43	213747	21.7589	ug/L	99
27) 1,1-Dichloroethane	8.37	63	274055	19.8182	ug/L	99
28) Ethyl-Tert-Butyl ether	8.74	59	2271239	98.9576	ug/L	99
29) 2-Butanone	8.93	43	42830	17.9801	ug/L	94
30) Propionitrile	9.03	54	60624	83.8325	ug/L	97
31) 2,2-Dichloropropane	9.14	77	202782	20.2096	ug/L	100
32) cis-1,2-Dichloroethene	9.20	96	154451	20.2299	ug/L	99
33) Chloroform	9.41	83	239118	19.4830	ug/L	99
34) 1-Bromopropane	9.55	122	31741	19.3407	ug/L	99
35) Bromochloromethane	9.64	130	87067	19.9958	ug/L	99
36) Tetrahydrofuran	9.67	42	152088	94.8951	ug/L	98
38) 1,1,1-Trichloroethane	9.94	97	216632	20.3216	ug/L	100
39) Cyclohexane	9.97	56	230381	16.2256	ug/L	100
40) 1,1-Dichloropropene	10.13	75	189524	21.1722	ug/L	99
41) Tert-Amyl-Methyl ether	10.24	73	1728726	98.8388	ug/L	99
42) Carbon Tetrachloride	10.28	117	191924	19.8127	ug/L	99

(#) = qualifier out of range (m) = manual integration
 8M429065.D 8260WTR.M Mon Mar 11 15:31:50 2019

Page 1

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429065.D Vial: 5
 Acq On : 11 Mar 2019 11:38 Operator: EEA
 Sample : WG699028-03 LCS DUP 20 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:31:49 2019 Quant Results File: 8260WTR.RES

Quant Method : K:\ORGANICS\V...\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration
 DataAcq Meth : 8260WTR

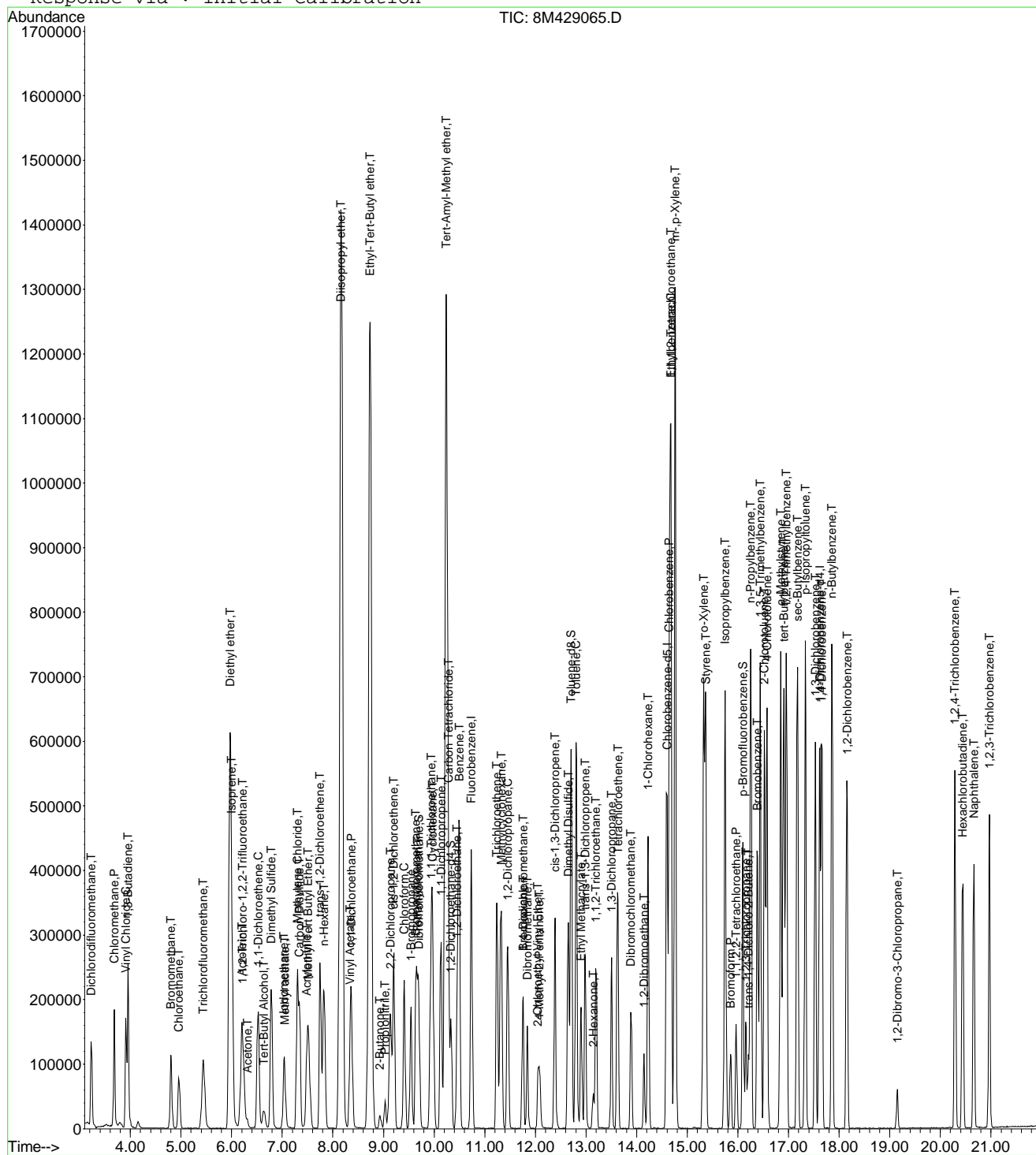
Compound	R.T.	QIon	Response	Conc	Unit	Qvalue
45) 1,2-Dichloroethane	10.46	62	170763	20.3415	ug/L	99
46) Benzene	10.50	78	569440	20.3070	ug/L	100
47) Trichloroethene	11.24	130	156275	20.4552	ug/L	100
48) Methylcyclohexane	11.32	83	212793	17.9851	ug/L	99
49) 1,2-Dichloropropane	11.46	63	155300	20.4033	ug/L	98
50) Bromodichloromethane	11.76	83	170606	19.3581	ug/L	99
51) 1,4-Dioxane	11.76	88	5284	123.3764	ug/L	94
52) Dibromomethane	11.84	93	69814	19.9269	ug/L	100
53) 2-Chloroethyl Vinyl Ether	12.05	63	42683	12.8335	ug/L	99
54) 4-Methyl-2-Pentanone	12.08	58	36454	17.4213	ug/L	99
55) cis-1,3-Dichloropropene	12.39	75	227721	21.7187	ug/L	99
56) Dimethyl Disulfide	12.65	79	117885	18.3088	ug/L	94
59) Toluene	12.80	91	594164	19.7599	ug/L	99
60) Ethyl Methacrylate	12.91	69	138560	18.0567	ug/L	99
62) trans-1,3-Dichloropropene	12.99	75	178085	19.7000	ug/L	100
63) 1,1,2-Trichloroethane	13.20	97	103204	19.6354	ug/L	98
64) 2-Hexanone	13.14	58	31330	16.5331	ug/L	85
65) 1,3-Dichloropropane	13.51	76	186262	20.5766	ug/L	100
66) Tetrachloroethene	13.63	164	123884	19.5503	ug/L	99
67) Dibromochloromethane	13.89	129	126669	18.9889	ug/L	99
68) 1,2-Dibromoethane	14.15	107	101792	19.9444	ug/L	100
69) 1-Chlorohexane	14.23	91	192038	18.4302	ug/L	99
70) Chlorobenzene	14.64	112	399541	19.6003	ug/L	99
71) 1,1,1,2-Tetrachloroethane	14.67	131	145810	19.4559	ug/L	98
72) Ethylbenzene	14.67	106	217569	19.7270	ug/L	97
73) m-,p-Xylene	14.76	106	532747	39.6437	ug/L	96
74) o-Xylene	15.33	106	265659	20.3735	ug/L	99
75) Styrene	15.37	104	436083	20.2006	ug/L	100
76) Bromoform	15.85	173	78082	18.8779	ug/L	98
77) Isopropylbenzene	15.75	105	649697	19.7265	ug/L	99
79) 1,1,2,2-Tetrachloroethane	15.97	83	109965	17.9232	ug/L	99
81) 1,2,3-Trichloropropane	16.16	110	36092	19.6645	ug/L	65
82) trans-1,4-Dichloro-2-Butene	16.20	53	33797	16.2951	ug/L #	30
83) n-Propylbenzene	16.26	91	809731	20.5742	ug/L	99
84) Bromobenzene	16.39	156	174279	19.3535	ug/L	99
85) 1,3,5-Trimethylbenzene	16.44	105	561022	19.7744	ug/L	100
86) 2-Chlorotoluene	16.52	91	540245	20.7608	ug/L	99
87) 4-Chlorotoluene	16.58	91	421664	18.5449	ug/L	100
88) a-Methylstyrene	16.85	118	316970	19.4700	ug/L	100
89) tert-Butylbenzene	16.91	134	127245	19.4408	ug/L	99
90) 1,2,4-Trimethylbenzene	16.96	105	566431	19.7205	ug/L	99
91) sec-Butylbenzene	17.18	105	740880	20.4047	ug/L	99
92) p-Isopropyltoluene	17.33	119	628854	20.2038	ug/L	99
93) 1,3-Dichlorobenzene	17.53	146	337125	19.2041	ug/L	99
94) 1,4-Dichlorobenzene	17.66	146	343541	19.3244	ug/L	99
95) n-Butylbenzene	17.86	91	570034	20.1001	ug/L	98
96) 1,2-Dichlorobenzene	18.16	146	318530	19.5531	ug/L	100
97) 1,2-Dibromo-3-Chloropropane	19.15	75	20541	18.5754	ug/L	98
98) 1,2,4-Trichlorobenzene	20.29	180	231567	19.5904	ug/L	99
99) Hexachlorobutadiene	20.44	225	110924	19.9153	ug/L	99
100) Naphthalene	20.66	128	418301	19.4126	ug/L	99
101) 1,2,3-Trichlorobenzene	20.97	180	211241	19.4537	ug/L	100

(#) = qualifier out of range (m) = manual integration
 8M429065.D 8260WTR.M Mon Mar 11 15:31:50 2019

Page 2

Data File : K:\ORGANICS\VOLATILE\HPMS8\DATA\031119\8M429065.D Vial: 5
 Acq On : 11 Mar 2019 11:38 Operator: EEA
 Sample : WG699028-03 LCS DUP 20 ug/L 8260 Inst : HPMS8
 Misc : 1,1 STD92354 Multiplr: 1.00
 MS Integration Params: RTEINT.P
 Quant Time: Mar 11 15:31 2019 Quant Results File: 8260WTR.RES

Method : K:\ORGANICS\VOLATILE\HPMS8\METHODS\8260WTR.M (RTE Integrator)
 Title : Method 8260B WTR-SOP:OVL MSV01 03-04-19 HPMS8
 Last Update : Mon Mar 04 17:44:12 2019
 Response via : Initial Calibration



2.2 General Chromatography Data

2.2.1 6850 LC/MS Data

2.2.1.1 Summary Data



Login Number: L19030638
Department: General Chromatography
Analyst: John W. Richards Jr.

METHOD

Analysis SW-846 6850

HOLDING TIMES

Sample Preparation: All holding times were met.

Sample Analysis: All holding times were met.

PREPARATION

Sample preparation proceeded normally.

CALIBRATION

Initial Calibration: For all compounds that yielded a %RSD greater than 15%, linear or higher order equations were applied. All acceptance criteria were met.

Alternate Source Standards: All acceptance criteria were met.

Continuing Calibration and Tune: All acceptance criteria were met.

BATCH QA/QC

Method Blank: All acceptance criteria were met.

Laboratory Control Sample: All acceptance criteria were met.

Matrix Spikes: Recovery out of range was observed for the following analyte: Perchlorate. Please see the applicable QC report for a detailed presentation of the failures.

Sample #	Analyte	Date	Result	Lower	Upper	Type
L19030638-03	Perchlorate	2019-03-18 20:48:40	80.5	84	119	Recovery

SAMPLES

Samples: All acceptance criteria were met.

Internal Standards: All acceptance criteria were met.

Manual Integration Reason Codes

Reason #1: Data System Fails to Select Correct Peak In some cases the chromatography system selects and integrates the 'wrong peak'. In this case the analyst must correct the selection and force the system to integrate the proper peak. Other times the system may miss the peak completely.

Reason #2: Data System Splits the Peak Incorrectly or Integrates a False Peak as a Rider Peak This phenomena is common at low concentrations where the signal:noise ratio is low. A single compound (peak) is incorrectly split into multiple peaks or integrated as a main peak with one or more rider peaks resulting in low area counts for the target compound.

Reason #3: Improperly Integrated Isomers and/or coeluting compounds. This system often fails to distinguish coeluting compounds and or isomers. The integration areas and concentrations are wrong, and they must be corrected by manual integration. Prime examples are benzo(k)fluoranthene and benzo(b)fluoranthene which are often unresolved and integrated improperly when both are present at low concentrations in standards or samples.

Reason #4: System Establishes Incorrect Baseline There are numerous situations in chromatography where the system establishes the baseline incorrectly. Some baseline errors will be obvious to the analyst and should be corrected via manual procedures.

Reason #5: Miscellaneous Other situations involving integration errors may require in-depth review and technical judgment. These cases should be brought to the attention of the laboratory management. If the form of manual integration is not clearly covered by these four cases, then review and approval by the Laboratory Director or the QA/QC Supervisor will be required.

I certify that this data package is in compliance with the terms and conditions agreed to by the client and Microbac Laboratories Inc., both technically and for completeness, except for the conditions noted above. Release of the data contained in this hard copy data package has been authorized by the Laboratory Manager or designated person, as verified by the following signature.

Narrative ID: 147535

Approved By: Eric Lawson



Certificate of Analysis

Sample #: L19030638-01	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW1	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 20:22
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 1LM.LM45229
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.542		0.400	0.200	0.100

Sample #: L19030638-02	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW1 MS	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 20:35
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 1LM.LM45230
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.735		0.400	0.200	0.100

Sample #: L19030638-03	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW1 MSD	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 20:48
Collect Date: 03/07/2019 11:30	Dilution: 1	File ID: 1LM.LM45231
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.703		0.400	0.200	0.100

Sample #: L19030638-04	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW2	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:01
Collect Date: 03/07/2019 12:30	Dilution: 1	File ID: 1LM.LM45232
Sample Tag: 01	Units: ug/L	

Certificate of Analysis

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.530		0.400	0.200	0.100

Sample #: L19030638-05	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW3	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:14
Collect Date: 03/07/2019 13:00	Dilution: 1	File ID: 1LM.LM45233
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.486		0.400	0.200	0.100

Sample #: L19030638-06	PrePrep Method: N/A	Instrument: LCMS1
Client ID: SW4	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:27
Collect Date: 03/07/2019 14:45	Dilution: 1	File ID: 1LM.LM45234
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.430		0.400	0.200	0.100

Sample #: L19030638-07	PrePrep Method: N/A	Instrument: LCMS1
Client ID: DUP01	Prep Method: 6850	Prep Date: 03/15/2019 16:30
Matrix: Water	Analytical Method: 6850	Cal Date: 03/18/2019 18:39
Workgroup #: WG699914	Analyst: JWR	Run Date: 03/18/2019 21:40
Collect Date: 03/07/2019 10:00	Dilution: 1	File ID: 1LM.LM45235
Sample Tag: 01	Units: ug/L	

Analyte	CAS #	Result	Qual	LOQ	LOD	DL
Perchlorate	14797-73-0	0.548		0.400	0.200	0.100

2.2.1.2 QC Summary Data

Example Calculation 6850 - Perchlorate**Concentration from Linear Regression****Step 1: Retrieve Curve Data From Plot, $y = mx + b$**

y = response ratio = response of analyte / response of internal standard (IS) = R_x/R_{istd}

x = amount ratio = concentration analyte/concentration internal standard (IS) = C_x / C_{istd}

m = slope from curve (1.45)

b = intercept from curve (-0.00242)

$y = 1.45x + -0.00242$

Step 2: Substitute the value for y

where $y = 12600/226000 = 0.055752$

Step 3: Solve for x

$x = (y - b)/m = 0.0040119$

Step 4: Solve for analyte concentration C_x

$C_x = (C_{is})(x) = (5 \text{ ug/L})(0.0040119) = 0.200594 \text{ ug/L}$

Example Calculation - Water:

Slope from curve, m :	1.45
Intercept from curve, b :	-0.00242
Response of analyte, R_x :	12600
Response of Internal Standard, R_{istd} :	226000
Concentration of IS, C_{istd} (ug/L):	5.00
Response Ratio:	0.05575
Amount Ratio:	0.04012
Analyte Concentration, C_x (ug/L) :	0.200594

Example Calculation - Soil:

Analyte Concentration, C_x (ug/L):	0.20059
Amount of soil extracted (g):	5.00
Final volume of extract (mL):	50.00
Percent solids (Pct wt.)	100
Concentration in soil (ug/kg):	2.005938

Microbac Laboratories Inc.
Instrument Run Log

Run Log ID: 95845

Instrument: LCMS1 Dataset: 031819_JWR.TXT
 Analyst1: JWR Analyst2: NA
 Method: 6850 SOP: HPLC06 Rev: 8

Maintenance Log ID: _____ Syringe Filter Lot#: 180728254
 Eluent ID#: _____

Workgroups: Column 1 ID: KP-RPPX250 Column 2 ID: NA
 Analytical WG699914 (waters)
 Internal STD: COA21170 Surrogate STD: NA Calibration STD WG700056 (03/18/19)
 CCV STD: STD87532 LCS STD: STD87534 MS/MSD STD: STD87534

Comments: ICAL WG700056 : Alternate Source STD87534
 Analytical Column : RPPX 5um (250x4.6mm)
 K'Prime S/N RPPX250-02115

Seq.	File ID	Sample Information	Mat	Dil	Reference	Date/Time
1	1LM.LM45214	WG700056-01 CCB	1	1		03/18/19 17:08
2	1LM.LM45215	WG700056-02 STD (0.1 ug/L)	1	1	STD87532	03/18/19 17:21
3	1LM.LM45216	WG700056-03 STD (0.2 ug/L)	1	1	STD87532	03/18/19 17:34
4	1LM.LM45217	WG700056-04 STD (0.5 ug/L)	1	1	STD87532	03/18/19 17:47
5	1LM.LM45218	WG700056-05 STD (1.0 ug/L)	1	1	STD87532	03/18/19 18:00
6	1LM.LM45219	WG700056-06 STD (2.0 ug/L)	1	1	STD87532	03/18/19 18:13
7	1LM.LM45220	WG700056-07 STD (5.0 ug/L)	1	1	STD87532	03/18/19 18:26
8	1LM.LM45221	WG700056-08 STD (10 ug/L)	1	1	STD87532	03/18/19 18:39
9	1LM.LM45222	WG700056-09 SSCV (1.0 ug/L)	1	1	STD87534	03/18/19 18:52
10	1LM.LM45223	WG699915-01 CCB	1	1		03/18/19 19:05
11	1LM.LM45224	WG699915-02 CCV (1.0ug/L)	1	1	STD87532	03/18/19 19:18
12	1LM.LM45225	WG699914-07 MRL (0.2ug/L)	1	1	STD87532	03/18/19 19:31
13	1LM.LM45226	WG699914-01 MCT (0.2ug/L)	1	1	STD87534	03/18/19 19:43
14	1LM.LM45227	WG699914-02 BLANK	1	1		03/18/19 19:56
15	1LM.LM45228	WG699914-03 LCS (0.2ug/L)	1	1	STD87534	03/18/19 20:09
16	1LM.LM45229	L19030638-01 RS	1	1		03/18/19 20:22
17	1LM.LM45230	L19030638-02 MS	1	1	STD87534	03/18/19 20:35
18	1LM.LM45231	L19030638-03 MSD	1	1	STD87534	03/18/19 20:48
19	1LM.LM45232	L19030638-04	1	1		03/18/19 21:01
20	1LM.LM45233	L19030638-05	1	1		03/18/19 21:14
21	1LM.LM45234	L19030638-06	1	1		03/18/19 21:27
22	1LM.LM45235	L19030638-07	1	1		03/18/19 21:40
23	1LM.LM45236	WG699915-03 CCV (1.0ug/L)	1	1	STD87532	03/18/19 21:53
24	1LM.LM45237	WG699914-08 MRL (0.2ug/L)	1	1	STD87532	03/18/19 22:06
25	1LM.LM45238	WG699915-04 CCB	1	1		03/18/19 22:19
26	1LM.LM45239	L19030972-01	1	1		03/18/19 22:32
27	1LM.LM45240	L19030973-01	1	1		03/18/19 22:45
28	1LM.LM45241	L19030974-01	1	1		03/18/19 22:58
29	1LM.LM45242	WG699915-05 CCV (1.0ug/L)	1	1	STD87532	03/18/19 23:10
30	1LM.LM45243	WG699914-09 MRL (0.2ug/L)	1	1	STD87532	03/18/19 23:23
31	1LM.LM45244	WG699915-06 CCB	1	1		03/18/19 23:36

Comments

Page: 1

Approved: 20-MAR-19




Microbac Laboratories Inc.
Instrument Run Log

Run Log ID:

Instrument: LCMS1 Dataset: 031819_JWR.TXT
 Analyst1: JWR Analyst2: NA
 Method: 6850 SOP: HPLC06 Rev: 8

Maintenance Log ID: _____ Syringe Filter Lot#: 180728254
 Eluent ID#: _____

Workgroups: Column 1 ID: KP-RPPX250 Column 2 ID: NA
 Analytical WG699914 (waters)
 Internal STD: COA21170 Surrogate STD: NA WG700056 (03/18/19)
 CCV STD: STD87532 LCS STD: STD87534 STD87534

Comments

Seq.	Rerun	Dil.	Reason	Analytes
18				
L19030638-03 MSD : The MSD percent recovery is below the DOD5-LCL(84%).				

Eri C. Zimm



Data Checklist

Date: 18-MAR-2019
 Analyst: JWR
 Analyst: NA
 Method: 6850
 Instrument: LCMS1
 Curve Workgroup: WG700056
 Runlog ID: 95845
 Analytical Workgroups: L19030638, L19030972, L19030973, L19030974

ANALYTICAL	
System Performance Check	X
DFTPP (GCMS)	NA
Endrin/DDT breakdown (8081/GCMS)	NA
Pentachlorophenol/benzidine tailing (GCMS)	NA
Eluent check (IC)/system pressure (HPLC)	NA
Window standard (FID)	NA
Initial Calibration	X
Average RF	NA
Linear regression or higher order curve	X
Alternate source standard (ICV) % Difference	X
Continuing Calibration (CCV)	X
% D/% Drift	X
Minimum response factors (GCMS)	X
Continuing calibration blank (CCB) (IC/LCMS)	X
Limit of quantitation verification (LOQV) (LCMS)	X
Special standards	NA
Blanks	X
TCL hits	ND
Surrogate recoveries	NA
LCS/LCSD (Laboratory Control Sample)	X
Recoveries	X
Surrogate recoveries	NA
MS/MSD/Sample duplicates	X
Recoveries	X
%RPD	X
Interference check sample (ICS) (LCMS)	MCT
Samples	X
TCL hits	X
Mass spectra (MS/HPLC)/2nd column confirmations (ECD/FID/HPLC)	X
Surrogate recoveries	NA
Internal standard areas (MS)	X
Library searches (GCMS)	NA
Calculations & correct factors	X
Compounds above calibration range	NA
Reruns	NA
Manual integrations	X
Project/client specific requirements	X
REPORTING	
Upload batch form	X
KOBRA workgroup data/forms/bench sheets	X
Case narratives	X
Check for completeness	X
Primary Reviewer	JWR
SUPERVISORY/SECONDARY REVIEW	
Check for compliance with method and project specific requirements	X
Check the completeness/accuracy of reported information	X
Data qualifiers	X
Secondary Reviewer	ECL

Primary Reviewer:
19-MAR-2019



Secondary Reviewer:
20-MAR-2019




Analytical Method:6850
Login Number:L19030638

AAB#:WG699914

Client ID	ID	Date Collected	TCLP Date	Time Held	Max Hold	Q	Extract Date	Time Held	Max Hold	Q	Run Date	Time Held	Max Hold	Q
SW1	01	03/07/19					03/15/2019	8.2	28		03/18/19	3.2	28	
SW1 MS	02	03/07/19					03/15/2019	8.2	28		03/18/19	3.2	28	
SW1 MSD	03	03/07/19					03/15/2019	8.2	28		03/18/19	3.2	28	
SW2	04	03/07/19					03/15/2019	8.2	28		03/18/19	3.2	28	
SW3	05	03/07/19					03/15/2019	8.1	28		03/18/19	3.2	28	
SW4	06	03/07/19					03/15/2019	8.1	28		03/18/19	3.2	28	
DUP01	07	03/07/19					03/15/2019	8.3	28		03/18/19	3.2	28	

* = SEE PROJECT QAPP REQUIREMENTS



METHOD BLANK SUMMARY

Login Number: L19030638
 Blank File ID: 1LM.LM45227
 Prep Date: 03/15/19 16:30
 Analyzed Date: 03/18/19 19:56
 Analyst: JWR

Work Group: WG699914
 Blank Sample ID: WG699914-02
 Instrument ID: LCMS1
 Method: 6850

This Method Blank Applies To The Following Samples:

Client ID	Lab Sample ID	Lab File ID	Time Analyzed	TAG
QCMRL	WG699914-07	1LM.LM45225	03/18/19 19:31	01
MCT	WG699914-01	1LM.LM45226	03/18/19 19:43	01
LCS	WG699914-03	1LM.LM45228	03/18/19 20:09	01
SW1	L19030638-01	1LM.LM45229	03/18/19 20:22	01
SW1 MS	L19030638-02	1LM.LM45230	03/18/19 20:35	01
SW1 MSD	L19030638-03	1LM.LM45231	03/18/19 20:48	01
SW2	L19030638-04	1LM.LM45232	03/18/19 21:01	01
SW3	L19030638-05	1LM.LM45233	03/18/19 21:14	01
SW4	L19030638-06	1LM.LM45234	03/18/19 21:27	01
DUP01	L19030638-07	1LM.LM45235	03/18/19 21:40	01
QCMRL	WG699914-08	1LM.LM45237	03/18/19 22:06	01
QCMRL	WG699914-09	1LM.LM45243	03/18/19 23:23	01

Report Name: BLANK_SUMMARY
 PDF File ID: 6354683
 Report generated 03/20/2019 16:39



Login Number: L19030638 Prep Date: 03/15/19 16:30 Sample ID: WG699914-02
 Instrument ID: LCMS1 Run Date: 03/18/19 19:56 Prep Method: 6850
 File ID: 1LM.LM45227 Analyst: JWR Method: 6850
 Workgroup (AAB#): WG699914 Matrix: Water Units: ug/L
 Contract #: _____ Cal ID: LCMS1-18-MAR-19

Analytes	DL	LOQ	Concentration	Dilution	Qualifier
Perchlorate	0.100	0.400	0.100	1	U

DL Method Detection Limit
 LOQ Reporting/Practical Quantitation Limit
 ND Analyte Not detected at or above reporting limit
 * |Analyte concentration| > 1/2 RL

Report Name: BLANK
 PDF ID: 6354684
 20-MAR-2019 16:39



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699914-03
Instrument ID: LCMS1 Run Time: 20:09 Prep Method: 6850
File ID: 1LM.LM45228 Analyst: JWR Method: 6850
Workgroup (AAB#): WG699914 Matrix: Water Units: ug/L
QC Key: DOD5 Lot#: STD87534 Cal ID: LCMS1-18-MAR-19

Analytes	Expected	Found	% Rec	LCS Limits	Q
Perchlorate	0.200	0.203	102	84 - 119	

LCS - Modified 03/06/2008
PDF File ID: 6354685
Report generated: 03/20/2019 16:39



MS/MSD REPORT

Loginum: L19030638 Cal ID: LCMS1- 18-MAR-19 Worknum: WG699914
 Instrument ID: LCMS1 Contract #: _____ Prep Method: 6850
 Parent ID: L19030638-01 File ID: ILM.LM45229 Dil: 1 Method: 6850
 Sample ID: L19030638-02 MS File ID: ILM.LM45230 Dil: 1 Matrix: Water
 Sample ID: L19030638-03 MSD File ID: ILM.LM45231 Dil: 1 Units: ug/L

Analyte	Parent	MS	MS	MS	MSD	MSD	MSD	%RPD	%Rec Limits	RPD Limit	Q
		Spiked	Found	%Rec	Spiked	Found	%Rec				
Perchlorate	0.542	0.200	0.735	96.5	0.200	0.703	80.5	4.45	84 - 119	15	*

* FAILS %REC LIMIT

FAILS RPD LIMIT

MS_MSD - Modified 03/06/2008
 PDF File ID: 6354686
 Report generated 03/20/2019 16:39



Login Number: L19030638
Analytical Method: 6850
ICAL Workgroup: WG700056

Instrument ID: LCMS1
Initial Calibration Date: 18-MAR-19 18:39
Column ID: F

Analyte	AVG RF	% RSD	LINEAR (R)	QUAD (R ²)
Perchlorate	1.377	2.02	1.00000	

R = Correlation coefficient; 0.995 minimum
R² = Coefficient of determination; 0.99 minimum

INT_CAL - Modified 03/06/2008
PDF File ID: 6354688
Report generated 03/20/2019 16:39



Login Number: L19030638
Analytical Method: 6850

Instrument ID: LCMS1
Initial Calibration Date: 18-MAR-19 18:39
Column ID: F

Analyte	WG700056-02			WG700056-03			WG700056-04		
	CONC	RESP	RF	CONC	RESP	RF	CONC	RESP	RF
Perchlorate	0.100	35100.0000	1.437	0.200	70600.0000	1.432	0.500	169000.000	1.342

INT_CAL - Modified 03/06/2008
PDF File ID: 6354688
Report generated 03/20/2019 16:39



Login Number: L19030638
 Analytical Method: 6850

Instrument ID: LCMS1
 Initial Calibration Date: 18-MAR-19 18:39
 Column ID: F

Analyte	WG700056-05			WG700056-06			WG700056-07		
	CONC	RESP	RF	CONC	RESP	RF	CONC	RESP	RF
Perchlorate	1.00	344000.000	1.354	2.00	700000.000	1.329	5.00	1760000.00	1.387

INT_CAL - Modified 03/06/2008
 PDF File ID: 6354688
 Report generated 03/20/2019 16:39



Login Number: L19030638
Analytical Method: 6850

Instrument ID: LCMS1
Initial Calibration Date: 18-MAR-19 18:39
Column ID: F

Analyte	WG700056-08		
	CONC	RESP	RF
Perchlorate	10.0	3360000.00	1.360

INT_CAL - Modified 03/06/2008
PDF File ID: 6354688
Report generated 03/20/2019 16:39



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG700056-09
 Instrument ID: LCMS1 Run Time: 18:52 Method: 6850
 File ID: 1LM.LM45222 Analyst: JWR QC Key: DOD5
 ICal Workgroup: WG700056 Cal ID: LCMS1 - 18-MAR-19

Analyte	Expected	Found	Units	RF	%D	UCL	Q
Perchlorate	1.00	0.999	ug/L	1.37	0.100	15	

* Exceeds %D Limit



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699915-01
Instrument ID: LCMS1 Run Time: 19:05 Method: 6850
File ID: LLM.LM45223 Analyst: JWR Units: ug/L
Workgroup (AAB#): WG699914 Cal ID: LCMS1 - 18-MAR-19
Matrix: WATER QAPP: DOD5

Analytes	MDL	RDL	Concentration	Qualifier
Perchlorate	0.100	0.400	0.100	U

U = Result is less than MDL.
F = Result is between MDL and RL.
* = Result is above RL.



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699915-04
Instrument ID: LCMS1 Run Time: 22:19 Method: 6850
File ID: LLM.LM45238 Analyst: JWR Units: ug/L
Workgroup (AAB#): WG699914 Cal ID: LCMS1 - 18-MAR-19
Matrix: WATER QAPP: DOD5

Analytes	MDL	RDL	Concentration	Qualifier
Perchlorate	0.100	0.400	0.100	U

U = Result is less than MDL.
F = Result is between MDL and RL.
* = Result is above RL.



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699915-06
Instrument ID: LCMS1 Run Time: 23:36 Method: 6850
File ID: 1LM.LM45244 Analyst: JWR Units: ug/L
Workgroup (AAB#): WG699914 Cal ID: LCMS1 - 18-MAR-19
Matrix: WATER QAPP: DOD5

Analytes	MDL	RDL	Concentration	Qualifier
Perchlorate	0.100	0.400	0.100	U

U = Result is less than MDL.
F = Result is between MDL and RL.
* = Result is above RL.



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699915-02
 Instrument ID: LCMS1 Run Time: 19:18 Method: 6850
 File ID: 1LM.LM45224 Analyst: JWR QC Key: DOD5
 Workgroup (AAB#): WG699914 Cal ID: LCMS1 - 18-MAR-19
 Matrix: WATER

Analyte	Expected	Found	UNITS	RF	%D	UCL	Q
Perchlorate	1.00	1.00	ug/L	1.37	0	15	

* Exceeds %D Criteria

CCV - Modified 03/05/2008
 PDF File ID: 6354690
 Report generated 03/20/2019 16:40



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699915-03
Instrument ID: LCMS1 Run Time: 21:53 Method: 6850
File ID: 1LM.LM45236 Analyst: JWR QC Key: DOD5
Workgroup (AAB#): WG699914 Cal ID: LCMS1 - 18-MAR-19
Matrix: WATER

Analyte	Expected	Found	UNITS	RF	%D	UCL	Q
Perchlorate	1.00	0.997	ug/L	1.36	0.300	15	

* Exceeds %D Criteria



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699915-05
 Instrument ID: LCMS1 Run Time: 23:10 Method: 6850
 File ID: 1LM.LM45242 Analyst: JWR QC Key: DOD5
 Workgroup (AAB#): WG699914 Cal ID: LCMS1 - 18-MAR-19
 Matrix: WATER

Analyte	Expected	Found	UNITS	RF	%D	UCL	Q
Perchlorate	1.00	0.984	ug/L	1.35	1.60	15	

* Exceeds %D Criteria



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699914-07
 Instrument ID: LCMS1 Run Time: 19:31 Prep Method: 6850
 File ID: 1LM.LM45225 Analyst: JWR Method: 6850
 Workgroup (AAB#): WG699914 Matrix: Water Units: ug/L
 Contract #: _____ Cal ID: LCMS1-18-MAR-19

Analytes	Expected	Found	% Rec	Limits	Q
Perchlorate	0.200	0.205	103	70 - 130	



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699914-08
Instrument ID: LCMS1 Run Time: 22:06 Prep Method: 6850
File ID: 1LM.LM45237 Analyst: JWR Method: 6850
Workgroup (AAB#): WG699914 Matrix: Water Units: ug/L
Contract #: _____ Cal ID: LCMS1-18-MAR-19

Analytes	Expected	Found	% Rec	Limits	Q
Perchlorate	0.200	0.212	106	70 - 130	



Login Number: L19030638 Run Date: 03/18/2019 Sample ID: WG699914-09
 Instrument ID: LCMS1 Run Time: 23:23 Prep Method: 6850
 File ID: 1LM.LM45243 Analyst: JWR Method: 6850
 Workgroup (AAB#): WG699914 Matrix: Water Units: ug/L
 Contract #: _____ Cal ID: LCMS1-18-MAR-19

Analytes	Expected	Found	% Rec	Limits	Q
Perchlorate	0.200	0.208	104	70 - 130	



Login Number: L19030638
Instrument ID: LCMS1
Workgroup (AAB#): WG699914

ICAL CCV Number: WG700056-05
CAL ID: LCMS1-18-MAR-19
Matrix: WATER

Sample Number	Dilution	Tag	IS-1
WG700056	NA	NA	1260000
Upper Limit	NA	NA	1890000
Lower Limit	NA	NA	630000
<u>L19030638-05</u>	1.00	01	1410000
L19030638-06	1.00	01	1390000
L19030638-07	1.00	01	1410000
WG699914-02	1.00	01	1420000
WG699914-03	1.00	01	1420000

IS-1 - 018LP

Underline = Response outside limits



Login Number: L19030638
Instrument ID: LCMS1
Workgroup (AAB#): WG699914

ICAL CCV Number: WG700056-05
CAL ID: LCMS1-18-MAR-19
Matrix: WATER

Sample Number	Dilution	Tag	IS-1
WG700056	NA	NA	1260000
Upper Limit	NA	NA	1890000
Lower Limit	NA	NA	630000
<u>L19030638-01</u>	1.00	01	1440000
<u>L19030638-02</u>	1.00	01	1370000

IS-1 - 018LP

Underline = Response outside limits



Login Number: L19030638
Instrument ID: LCMS1
Workgroup (AAB#): WG699914

ICAL CCV Number: WG700056-05
CAL ID: LCMS1-18-MAR-19
Matrix: WATER

Sample Number	Dilution	Tag	IS-1
WG700056	NA	NA	1260000
Upper Limit	NA	NA	1890000
Lower Limit	NA	NA	630000
<u>L19030638-03</u>	1.00	01	1410000
<u>L19030638-04</u>	1.00	01	1380000

IS-1 - 018LP

Underline = Response outside limits



Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: L19030638-01
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45229
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 20:22	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	214000	72000	2.97	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: L19030638-02
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45230
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 20:35	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	276000	86300	3.20	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: L19030638-03
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45231
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 20:48	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	271000	92300	2.94	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: L19030638-04
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45232
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 21:01	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	201000	64500	3.12	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638
Instrument: LCMS1
Analyst: JWR
Worknum: WG699914

Prep Method: 6850
Prep Date: 03/15/2019 16:30
Anal Method: 6850
Analysis Date: 03/18/2019 21:14

Samplenum: L19030638-05
File ID: 1LM.LM45233
Matrix: Water
Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	189000	62500	3.02	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: L19030638-06
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45234
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 21:27	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	165000	52300	3.15	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638
Instrument: LCMS1
Analyst: JWR
Worknum: WG699914

Prep Method: 6850
Prep Date: 03/15/2019 16:30
Anal Method: 6850
Analysis Date: 03/18/2019 21:40

Samplenum: L19030638-07
File ID: 1LM.LM45235
Matrix: Water
Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	212000	68100	3.11	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638
Instrument: LCMS1
Analyst: JWR
Worknum: WG699914

Prep Method: 6850
Prep Date: 03/15/2019 16:30
Anal Method: 6850
Analysis Date: 03/18/2019 19:43

Samplenum: WG699914-01
File ID: 1LM.LM45226
Matrix: Water
Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	79900	24800	3.22	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: WG699914-02
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45227
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 19:56	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	936	0.000	0.000	2.3	3.8	*

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: WG699914-03
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45228
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 20:09	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	80200	24500	3.27	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638
Instrument: LCMS1
Analyst: JWR
Worknum: WG699914

Prep Method: 6850
Prep Date: 03/15/2019 16:30
Anal Method: 6850
Analysis Date: 03/18/2019 19:31

Samplenum: WG699914-07
File ID: 1LM.LM45225
Matrix: Water
Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	80800	25000	3.23	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: WG699914-08
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45237
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 22:06	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	92400	31900	2.90	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: 6850	Samplenum: WG699914-09
Instrument: LCMS1	Prep Date: 03/15/2019 16:30	File ID: 1LM.LM45243
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 23:23	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	91800	27400	3.35	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG699915-01
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45223
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 19:05	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	0.000	0.000	0.000	2.3	3.8	*

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG699915-02
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45224
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 19:18	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	379000	119000	3.18	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG699915-03
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45236
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 21:53	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	426000	136000	3.13	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG699915-04
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45238
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 22:19	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	773	978	0.790	2.3	3.8	*

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG699915-05
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45242
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 23:10	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	436000	139000	3.14	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG699915-06
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45244
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 23:36	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	0.000	535	0.000	2.3	3.8	*

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-02
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45215
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 17:21	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	35100	11700	3.00	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-03
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45216
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 17:34	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	70600	21300	3.31	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-04
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45217
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 17:47	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	169000	55400	3.05	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-05
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45218
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 18:00	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	344000	109000	3.16	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-06
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45219
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 18:13	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	700000	228000	3.07	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-07
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45220
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 18:26	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	1760000	550000	3.20	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-08
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45221
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 18:39	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	3360000	1060000	3.17	2.3	3.8	

Perchlorate Ion Ratios
Microbac Laboratories Inc.



Login #: L19030638	Prep Method: _____	Samplenum: WG700056-09
Instrument: LCMS1	Prep Date: _____	File ID: 1LM.LM45222
Analyst: JWR	Anal Method: 6850	Matrix: Water
Worknum: WG699914	Analysis Date: 03/18/2019 18:52	Units: ug/L

Analyte	Res #1	Res #2	Ratio	Lower	Upper	Q
PERCHLORATE	373000	122000	3.06	2.3	3.8	

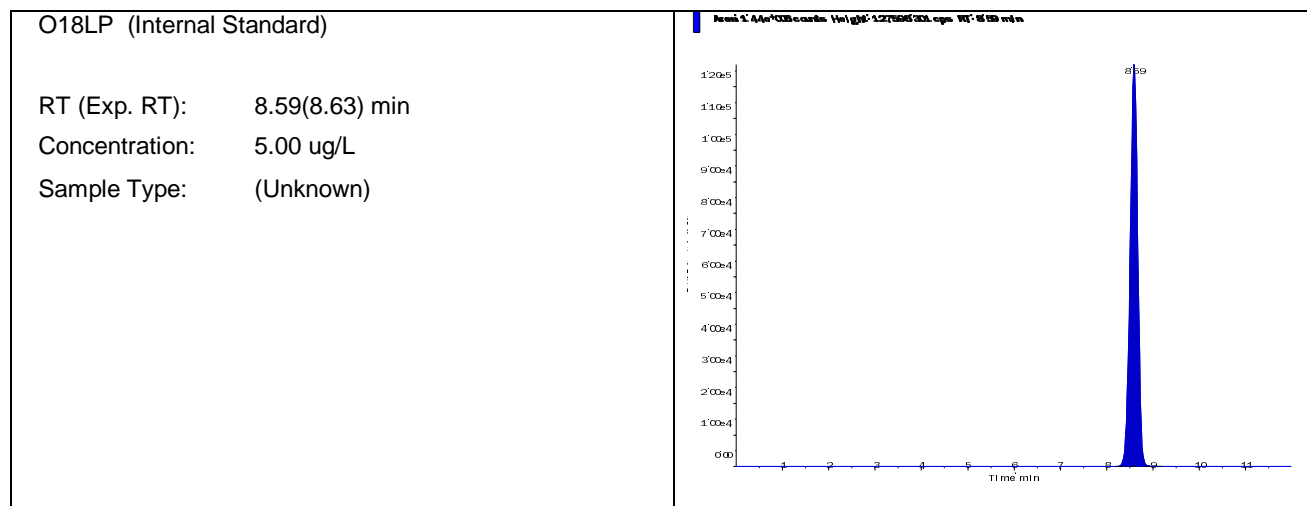
2.2.1.3 Sample Data

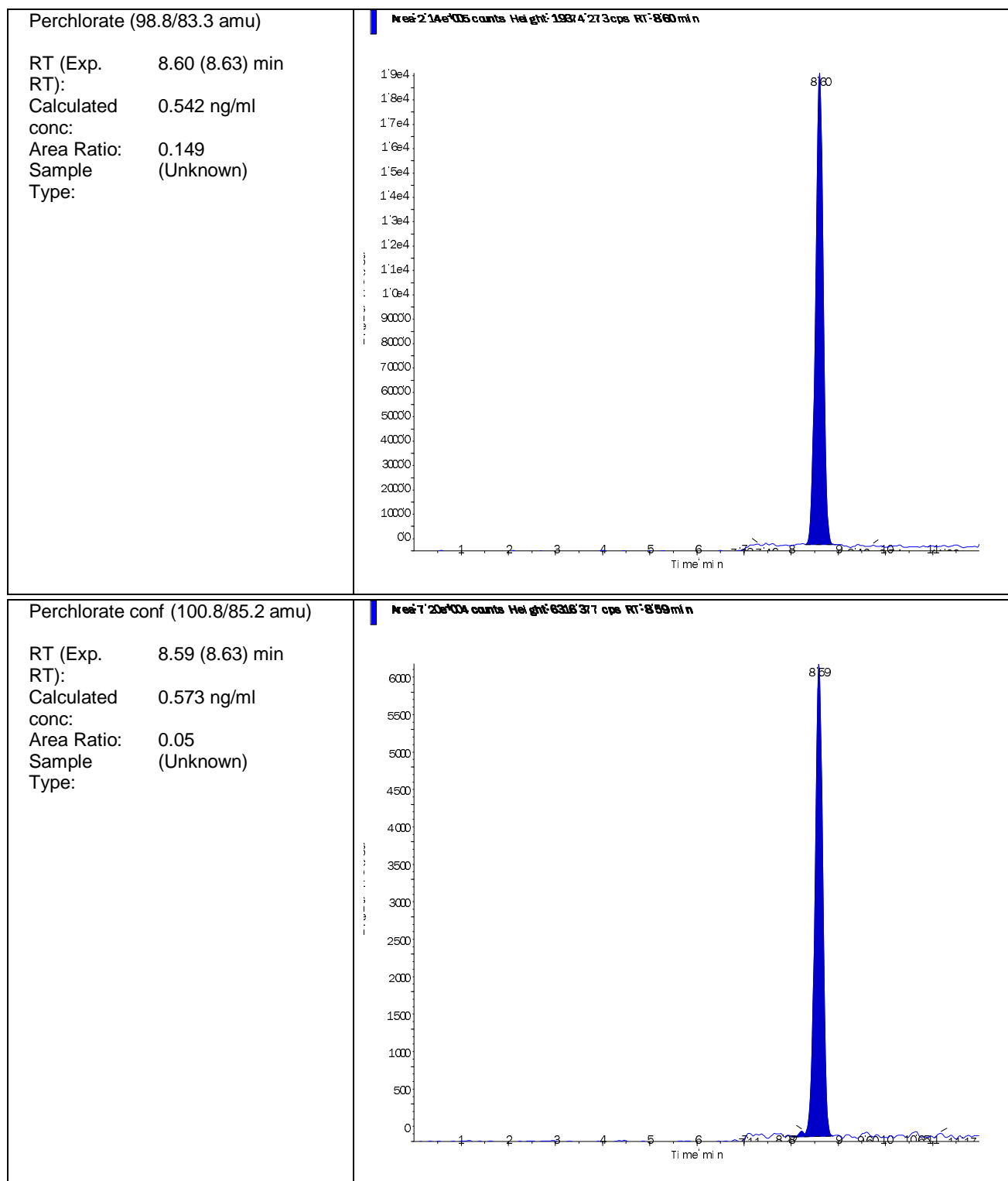
Data File	LM45229.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 8:22:48 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	L19030638-01 RS	Injection Vial	13.00
Data File	LM45229.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 8:22:48 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-04	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

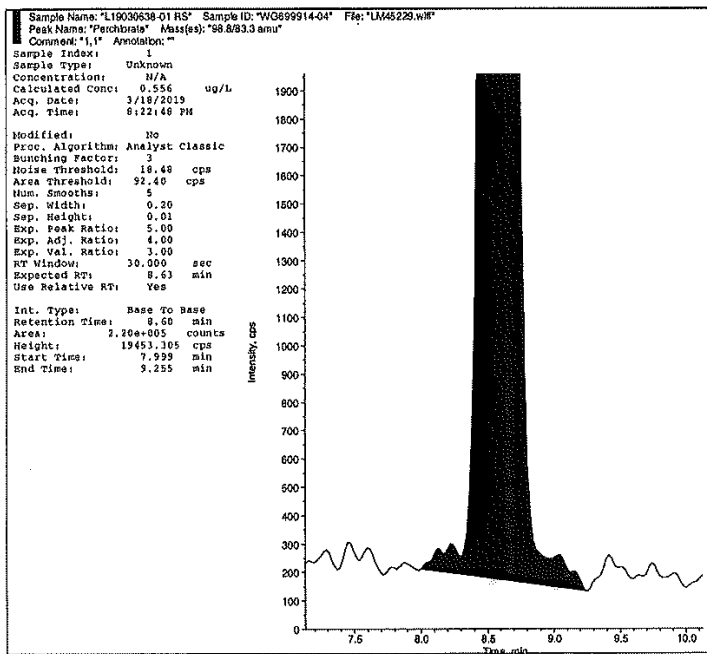
Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.440e+06	8.59	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	2.140e+05	8.60	N/A	0.542
Perchlorate conf	7.200e+04	8.59	N/A	0.573



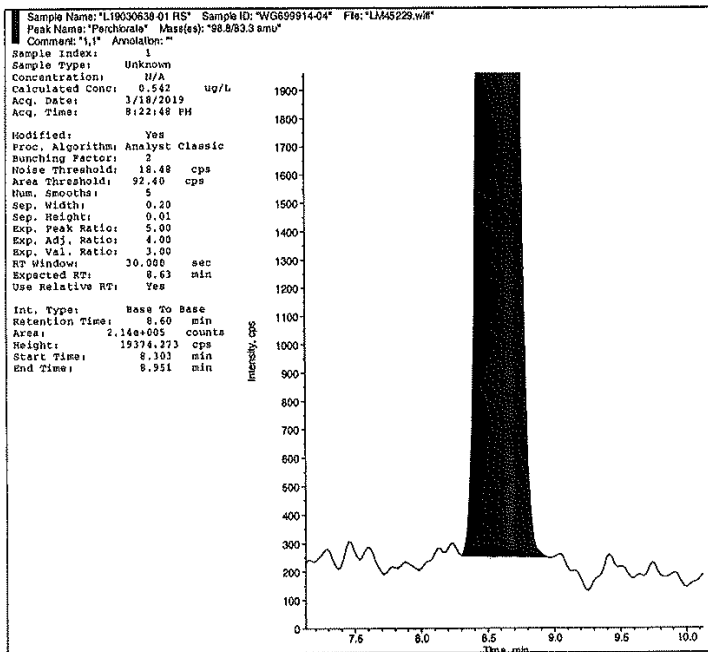


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Printing Date: Tuesday, March 19, 2019



Collected by: N/A
Electronic Signature: no
Operator: Administrator

Printing Time: 10:55:08 AM
 Printing Date: Tuesday, March 19, 2019



#4
 JWR/03/19/19

✓ EC2 3-20-19

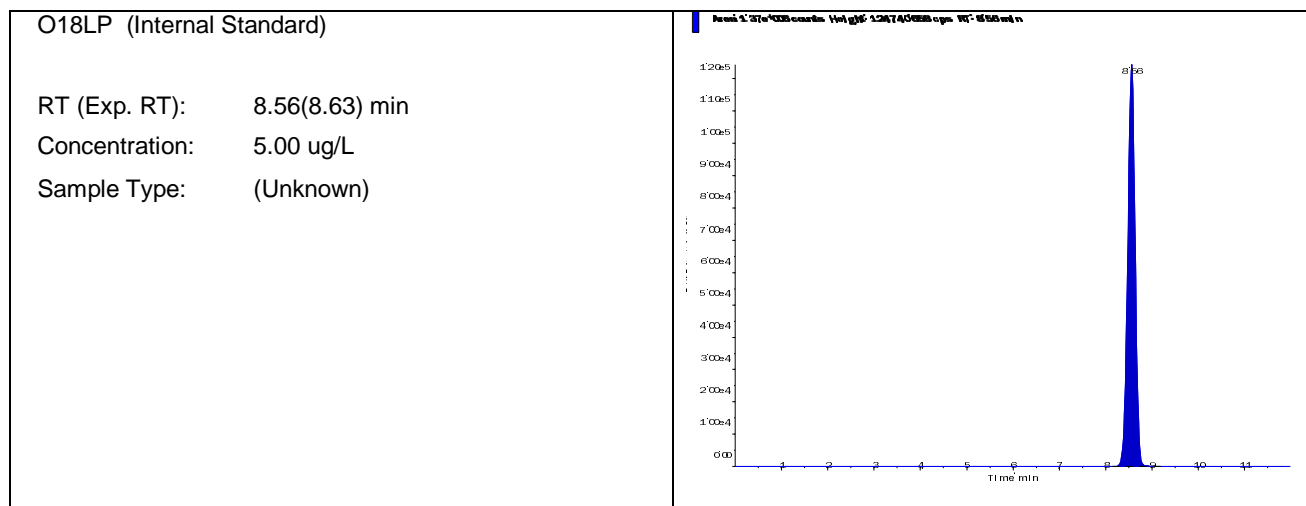
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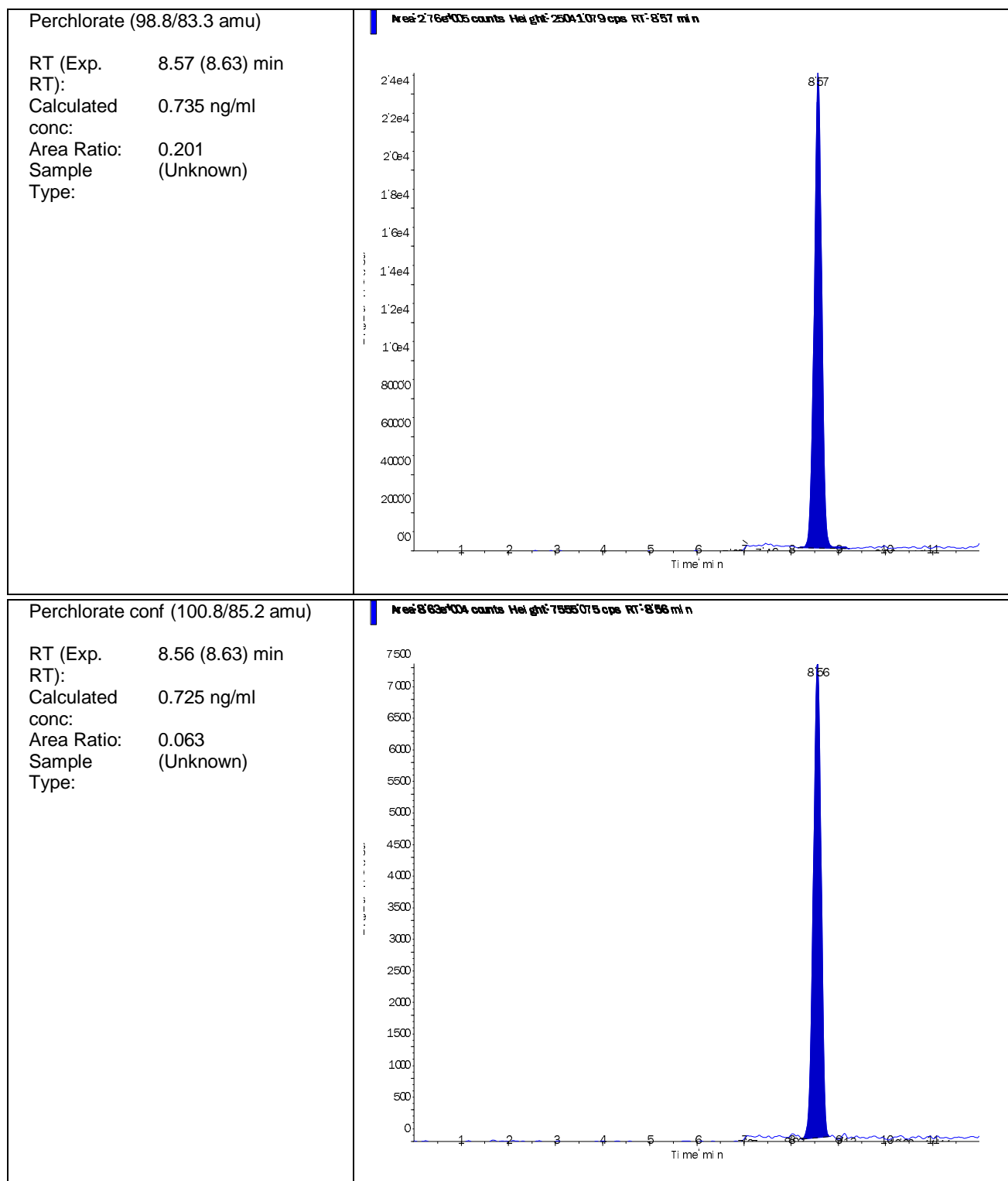
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Acquisition Date	3/18/2019 8:35:44 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	L19030638-02 MS	Injection Vial	14.00
Data File	LM45230.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 8:35:44 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-05	Dilution Factor	1.00
Sample Comment	1,1 STD87534	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.370e+06	8.56	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	2.760e+05	8.57	N/A	0.735
Perchlorate conf	8.630e+04	8.56	N/A	0.725



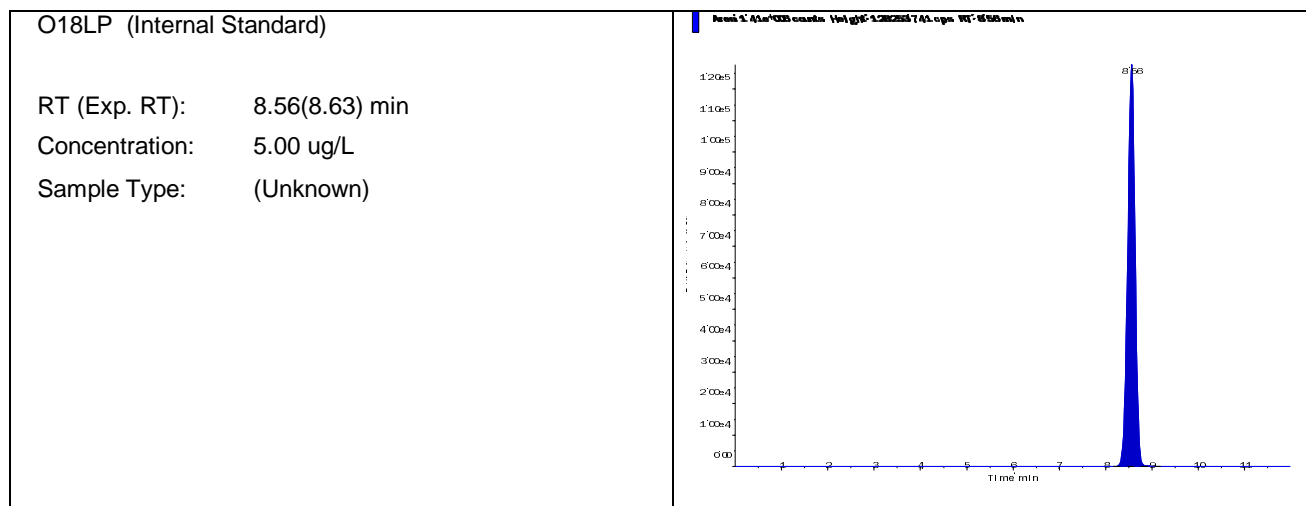


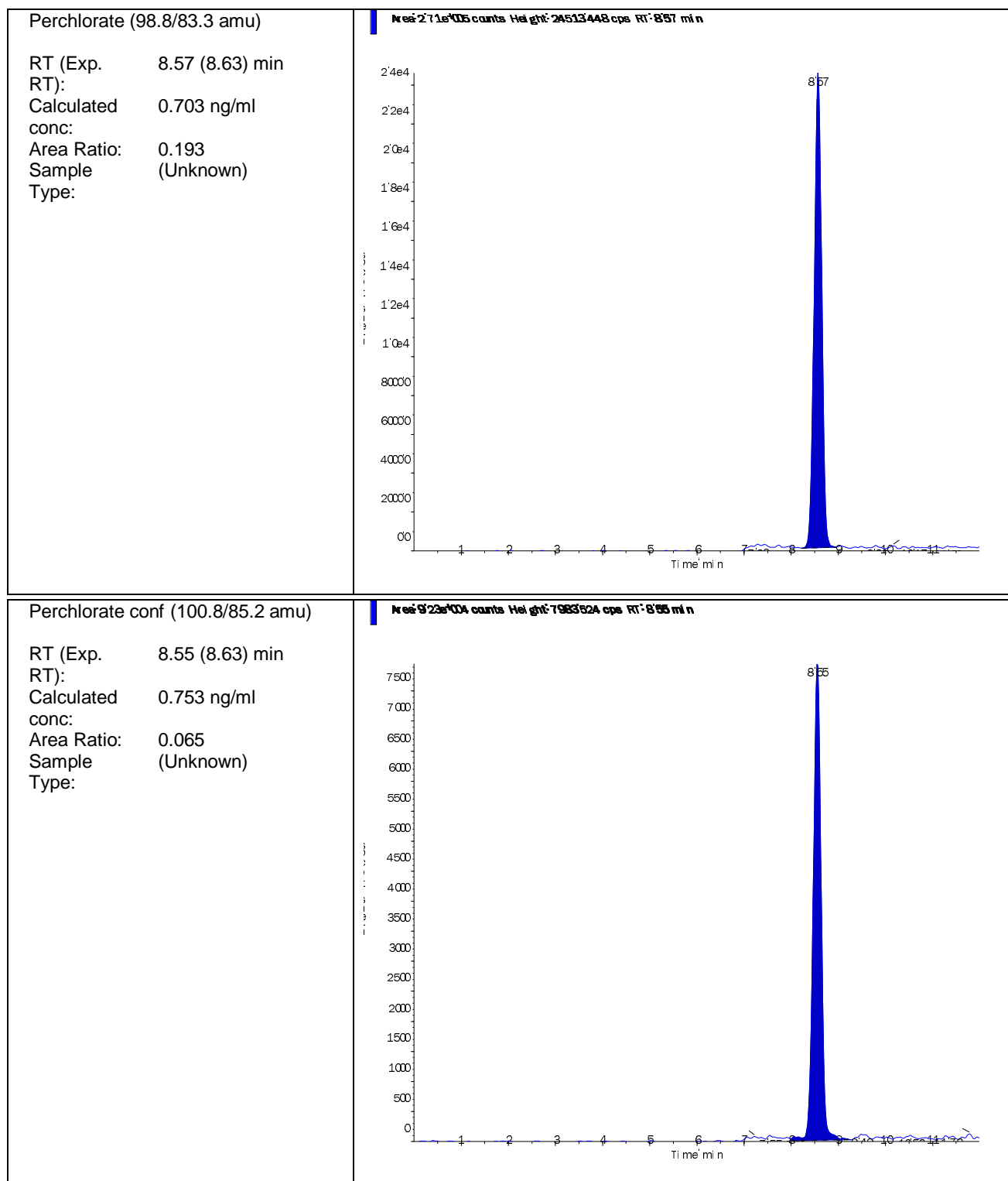
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Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	L19030638-03 MSD	Injection Vial	15.00
Data File	LM45231.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 8:48:40 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-06	Dilution Factor	1.00
Sample Comment	1,1 STD87534	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.410e+06	8.56	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	2.710e+05	8.57	N/A	0.703
Perchlorate conf	9.230e+04	8.55	N/A	0.753





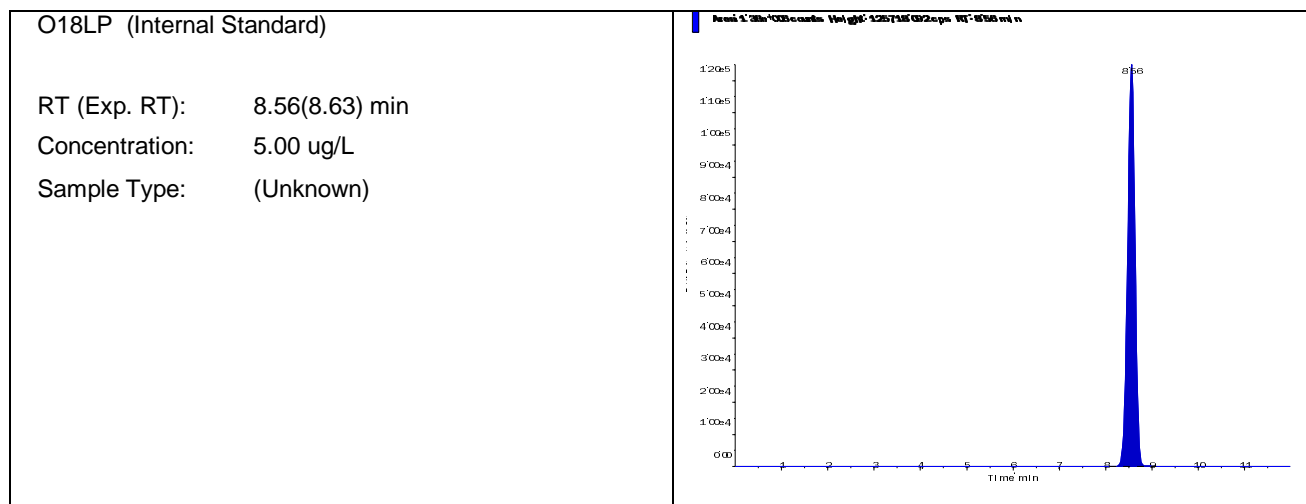
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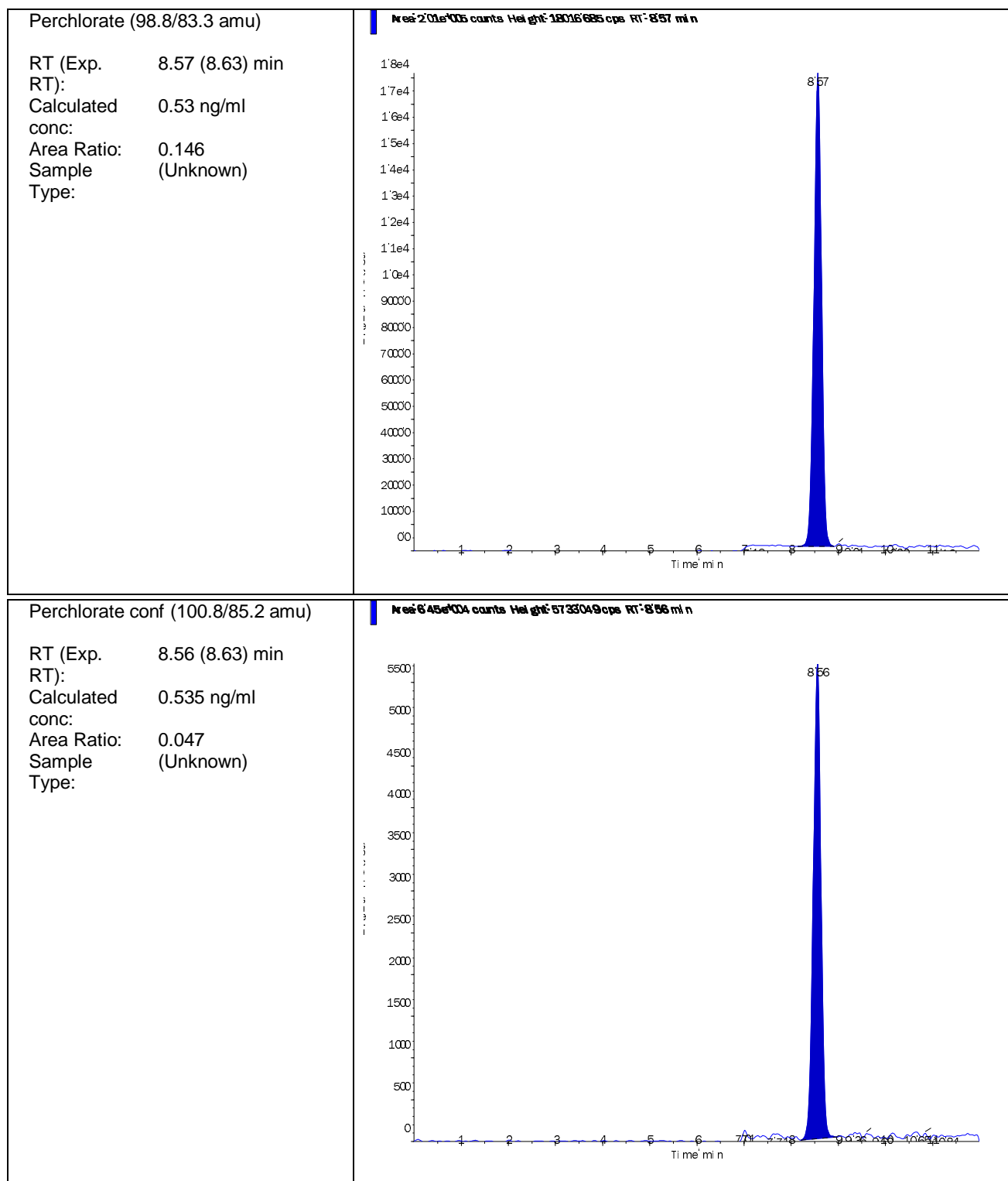
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Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	L19030638-04	Injection Vial	16.00
Data File	LM45232.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 9:01:36 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	L19030638-04	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.380e+06	8.56	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	2.010e+05	8.57	N/A	0.53
Perchlorate conf	6.450e+04	8.56	N/A	0.535





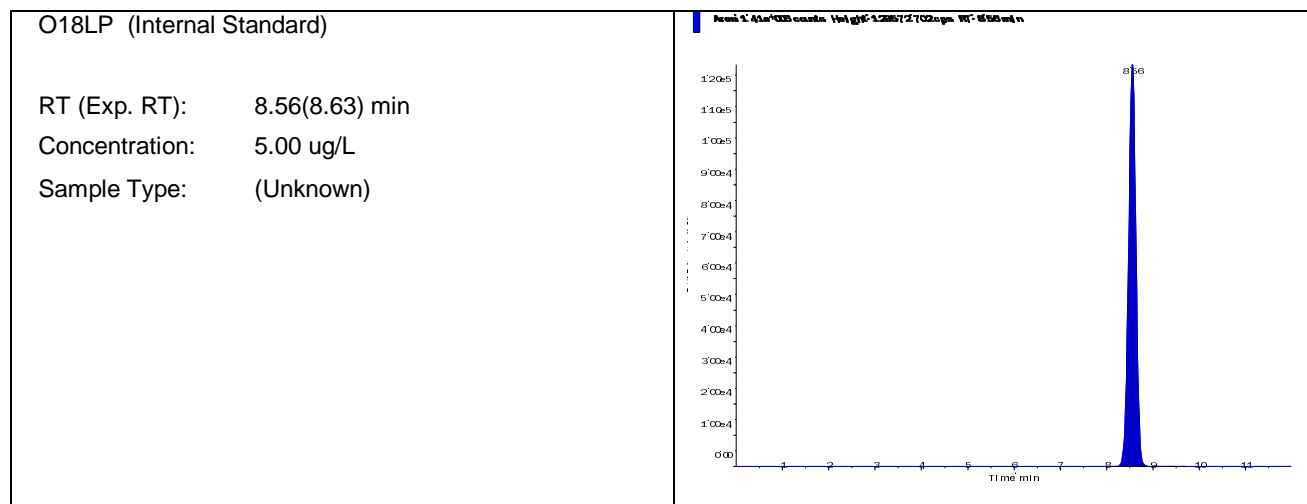
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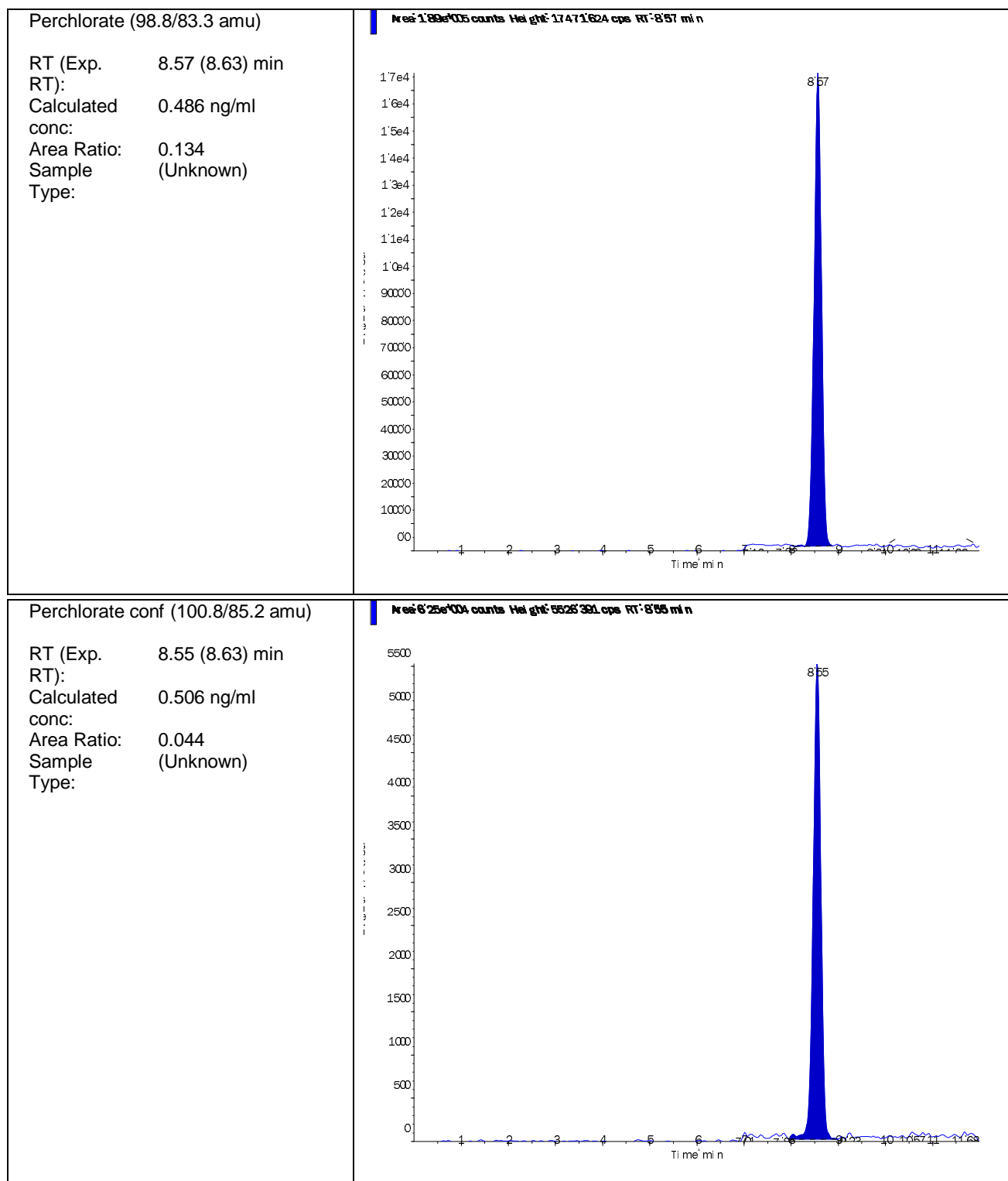
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Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	L19030638-05	Injection Vial	17.00
Data File	LM45233.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 9:14:31 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	L19030638-05	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.410e+06	8.56	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	1.890e+05	8.57	N/A	0.486
Perchlorate conf	6.250e+04	8.55	N/A	0.506



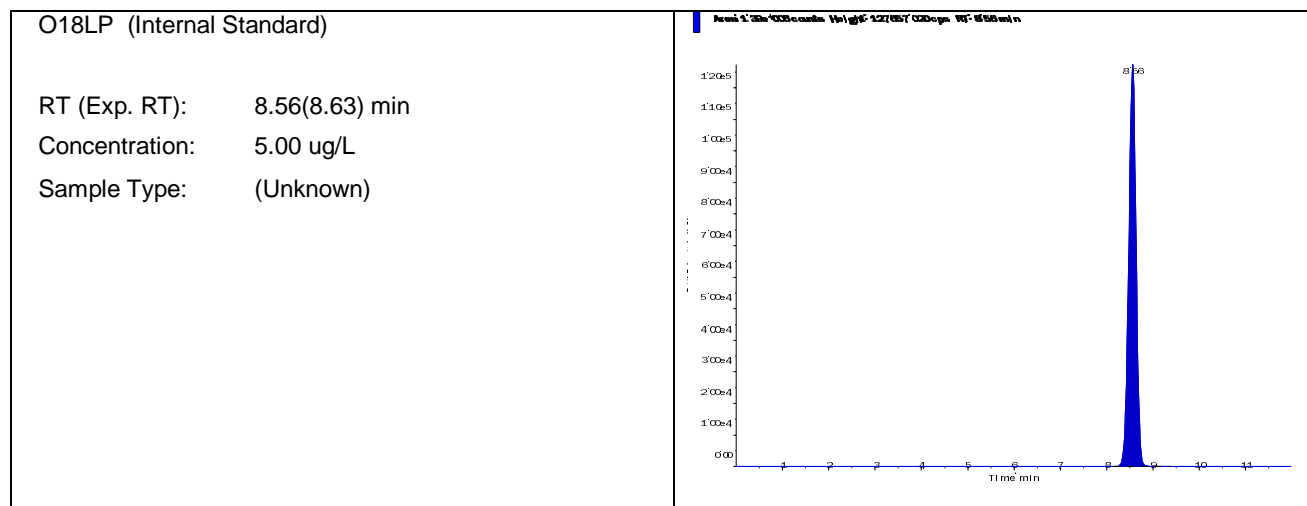


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Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

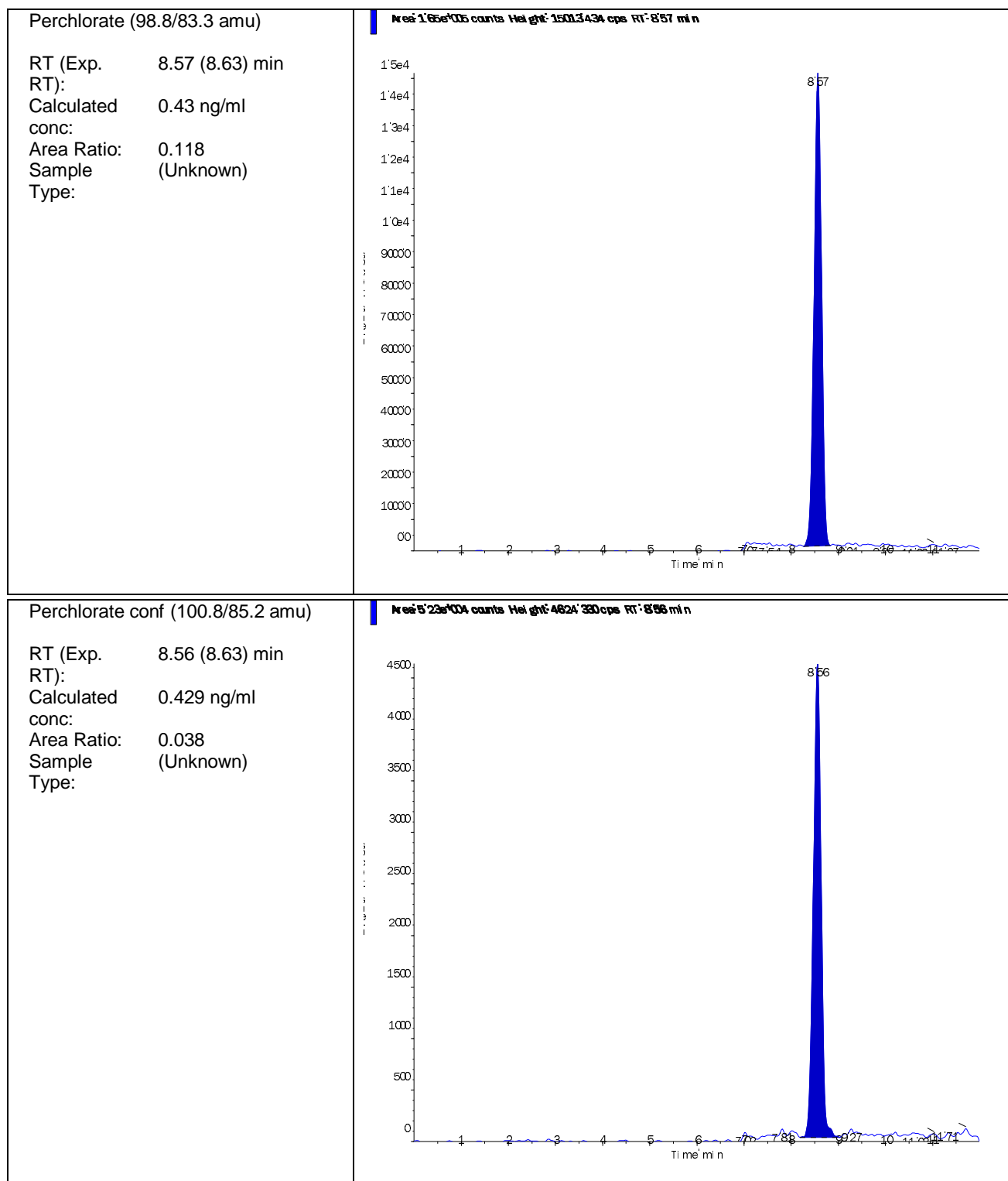
Sample Name	L19030638-06	Injection Vial	18.00
Data File	LM45234.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 9:27:26 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	L19030638-06	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.390e+06	8.56	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	1.650e+05	8.57	N/A	0.43
Perchlorate conf	5.230e+04	8.56	N/A	0.429



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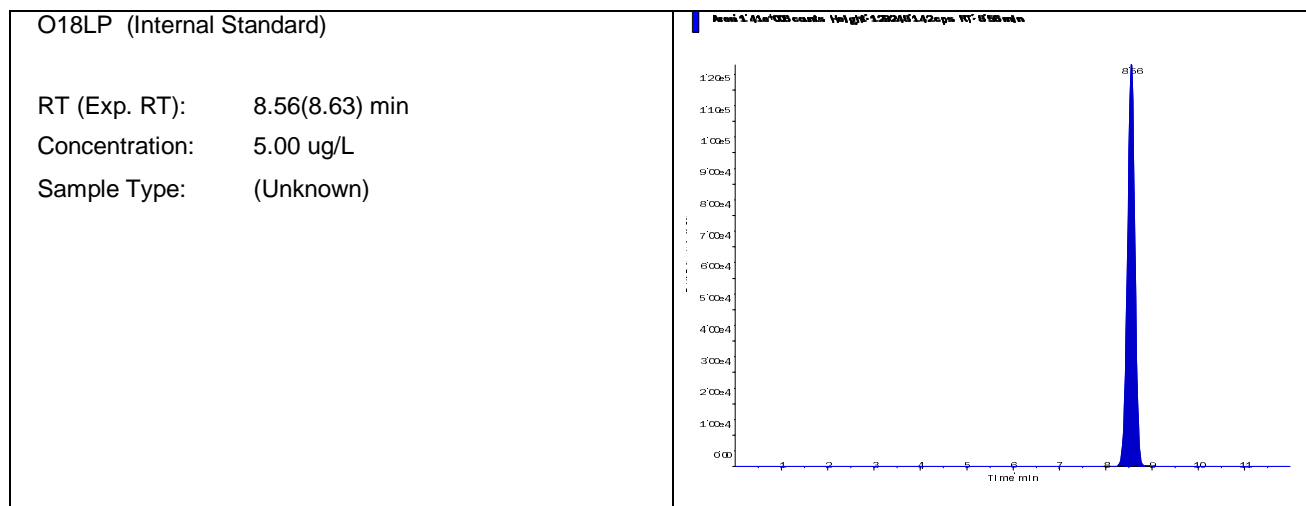


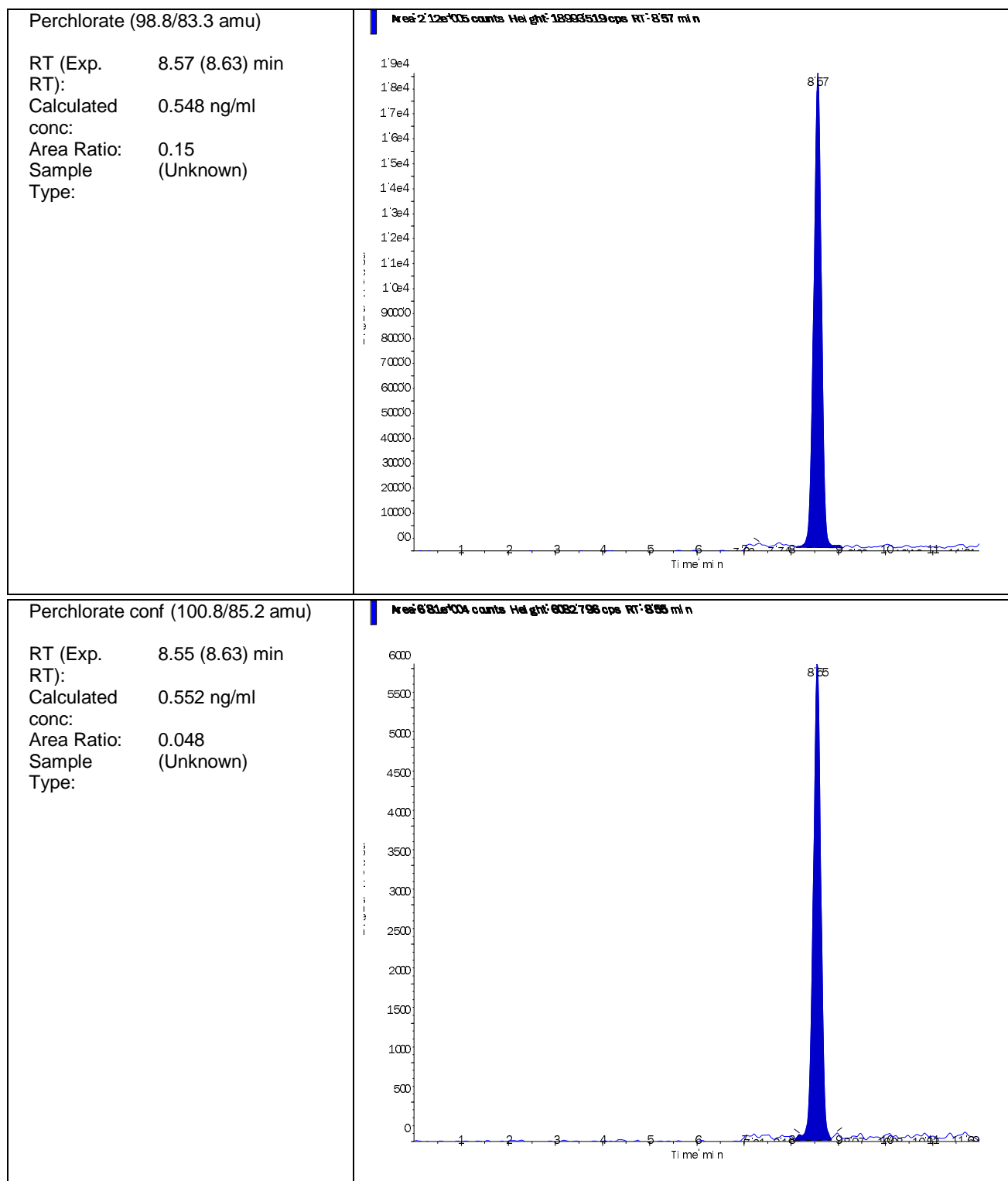
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Acquisition Date	3/18/2019 9:40:26 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	L19030638-07	Injection Vial	19.00
Data File	LM45235.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 9:40:26 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	L19030638-07	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.410e+06	8.56	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	2.120e+05	8.57	N/A	0.548
Perchlorate conf	6.810e+04	8.55	N/A	0.552





s.dataFile Page 2 of 2

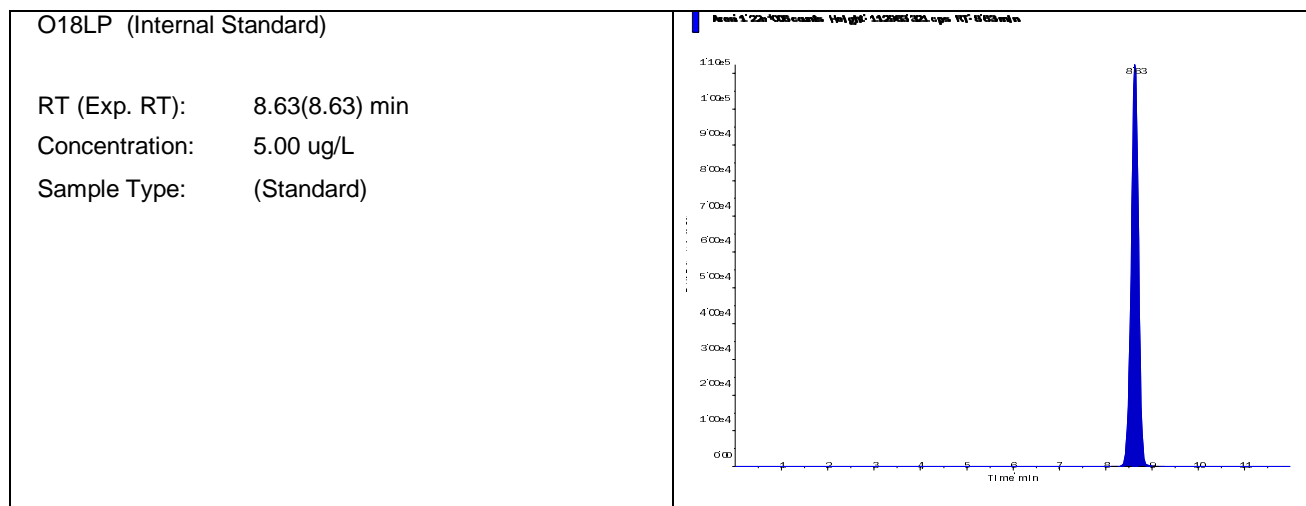
2.2.1.4 Standards Data

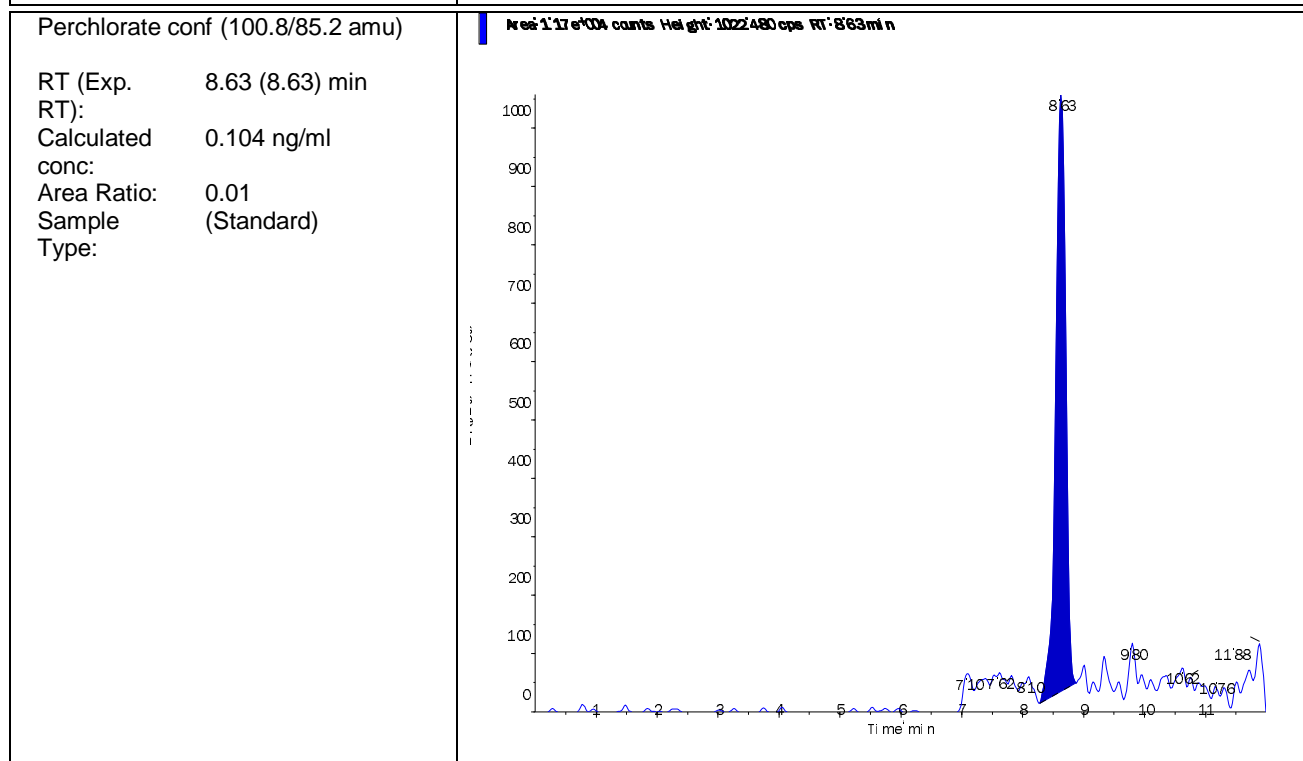
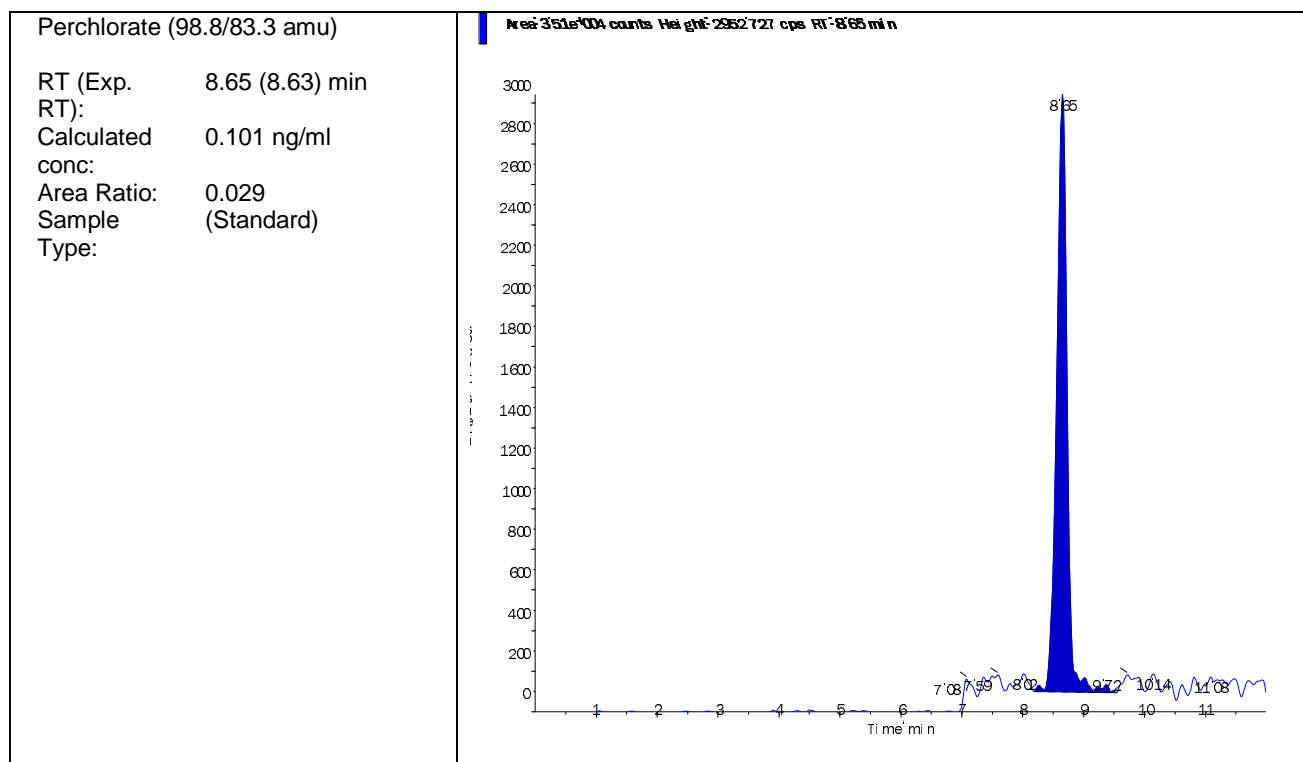
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Acquisition Date	3/18/2019 5:21:42 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-02 STD (0.1 ug/L)	Injection Vial	2.00
Data File	LM45215.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 5:21:42 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Standard
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-02	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.220e+06	8.63	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	3.510e+04	8.65	0.10	0.101
Perchlorate conf	1.170e+04	8.63	0.10	0.104



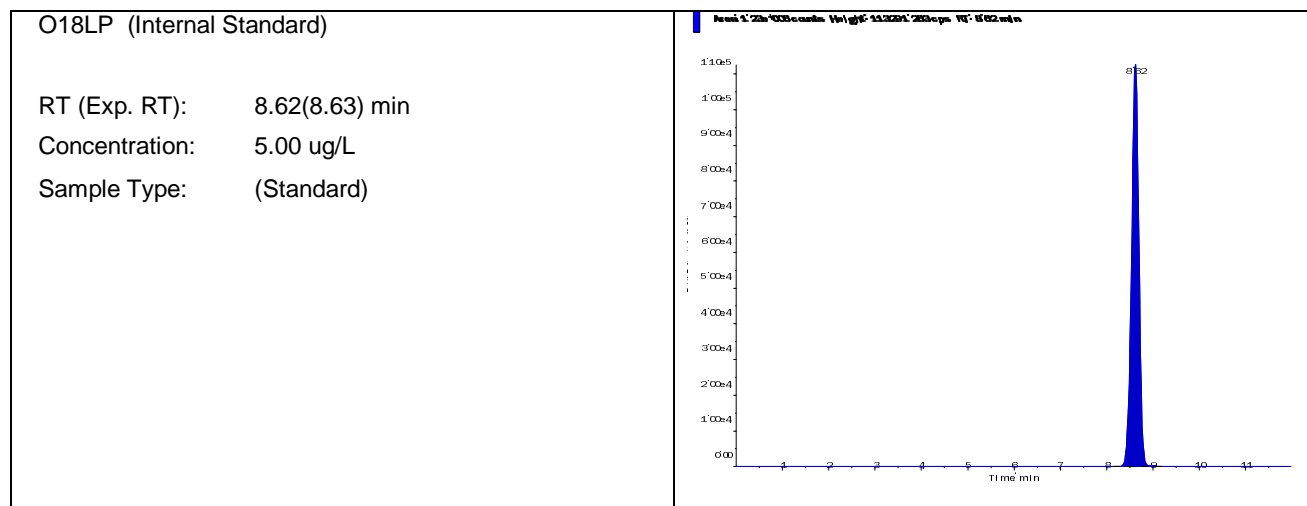


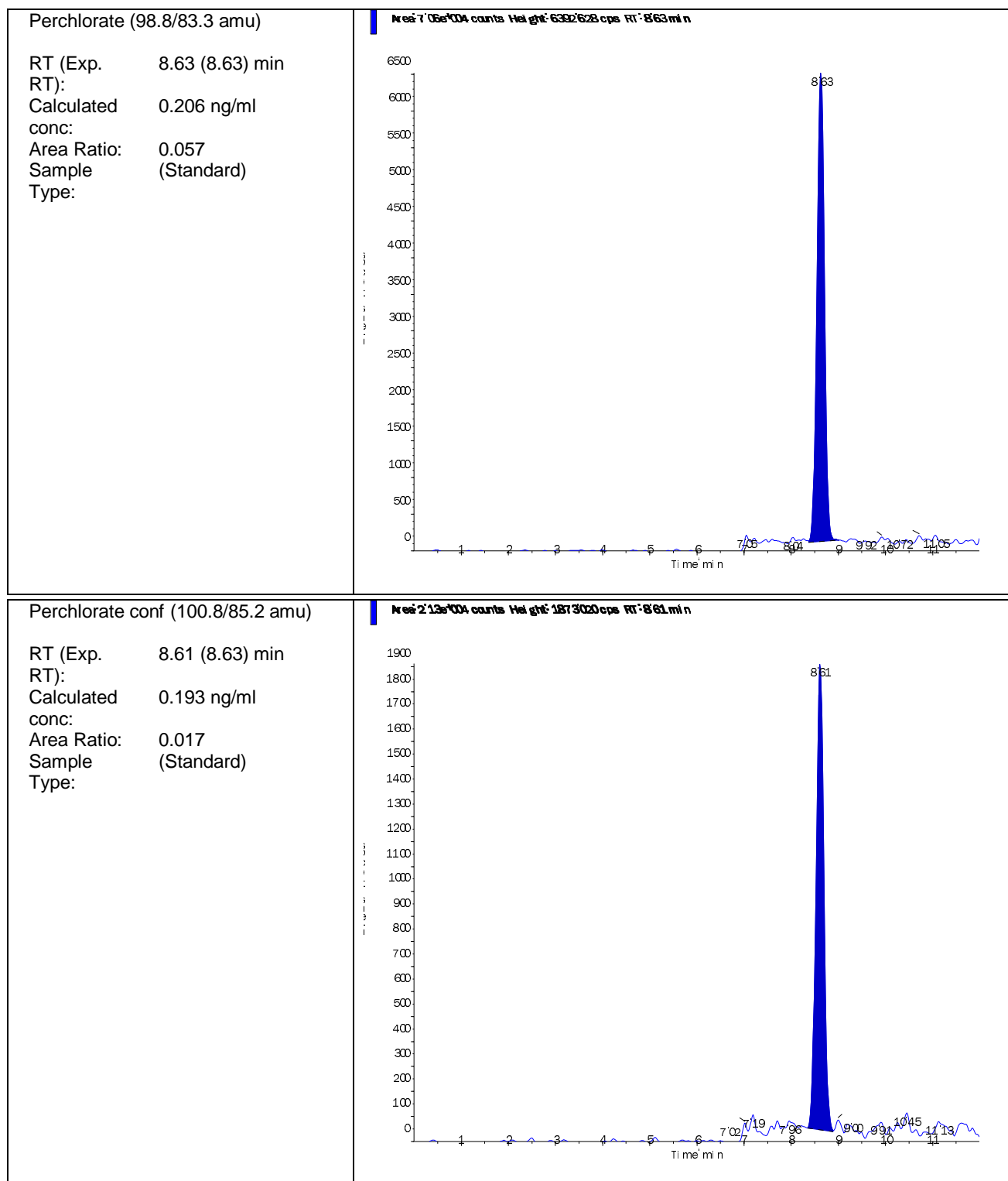
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Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-03 STD (0.2 ug/L)	Injection Vial	3.00
Data File	LM45216.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 5:34:37 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Standard
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-03	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.230e+06	8.62	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	7.060e+04	8.63	0.20	0.206
Perchlorate conf	2.130e+04	8.61	0.20	0.193



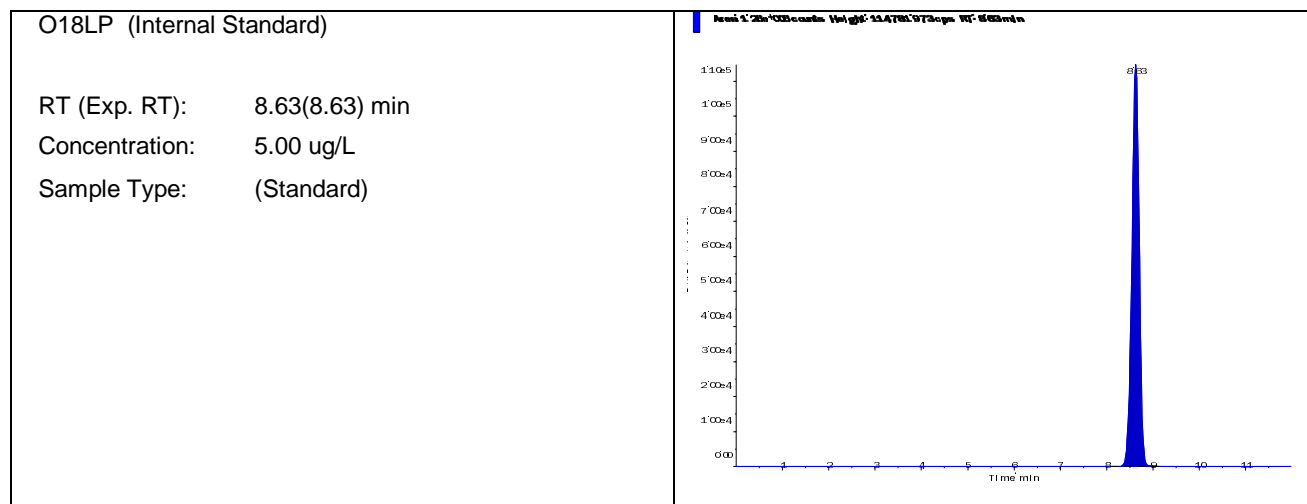


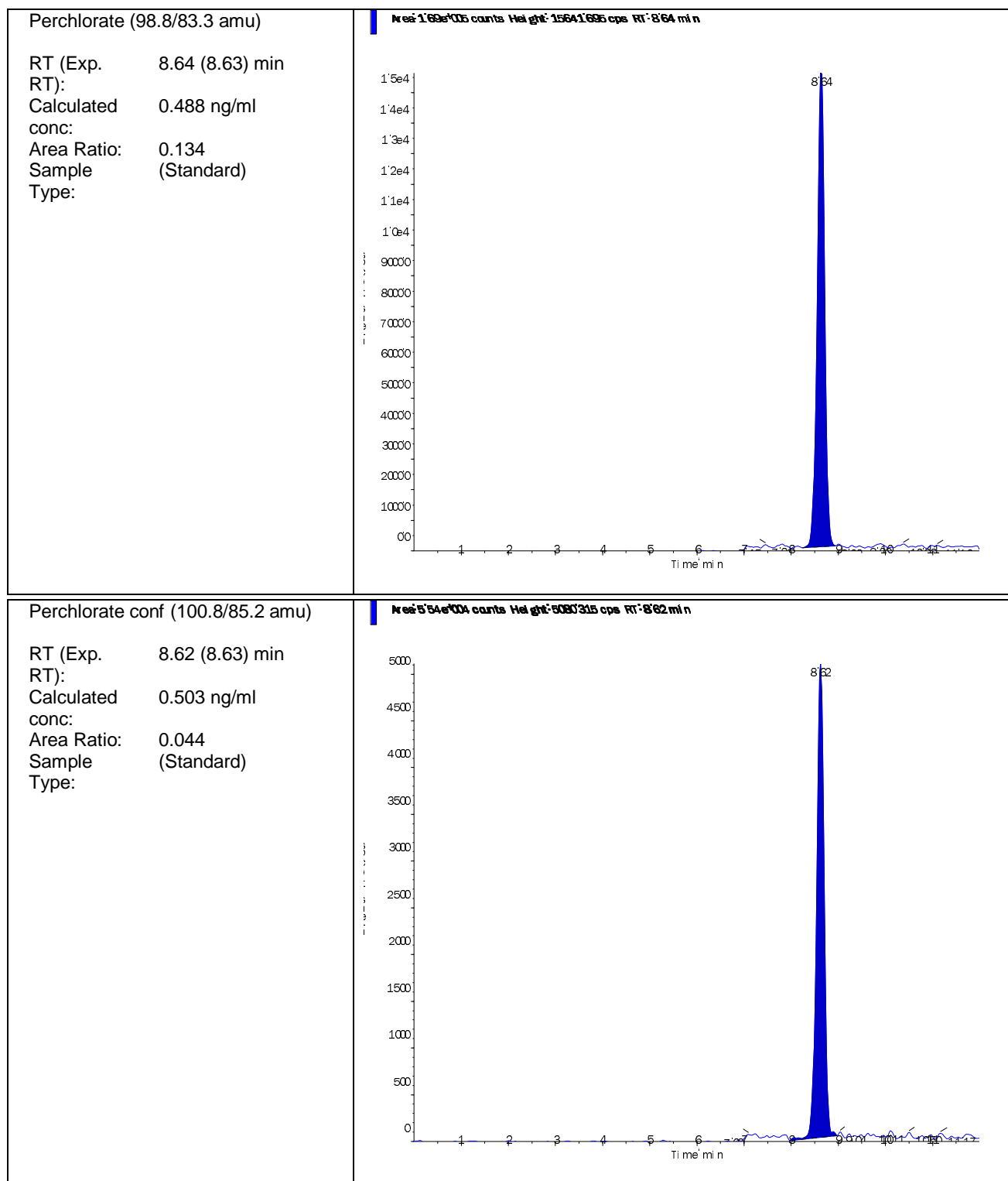
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Acquisition Date	3/18/2019 5:47:35 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-04 STD (0.5 ug/L)	Injection Vial	4.00
Data File	LM45217.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 5:47:35 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Standard
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-04	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.260e+06	8.63	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	1.690e+05	8.64	0.50	0.488
Perchlorate conf	5.540e+04	8.62	0.50	0.503





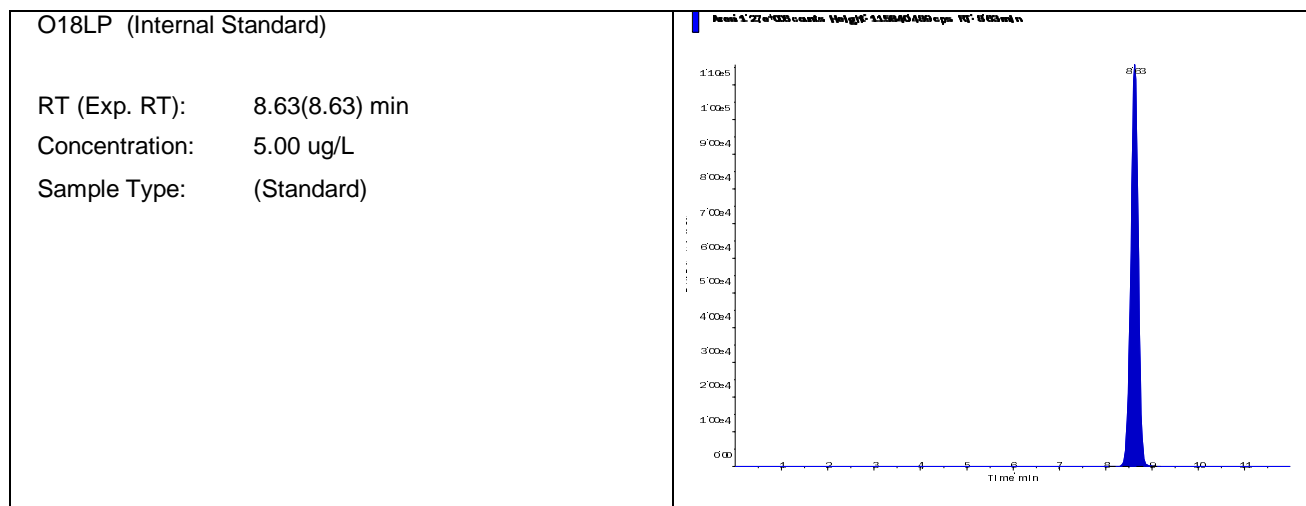
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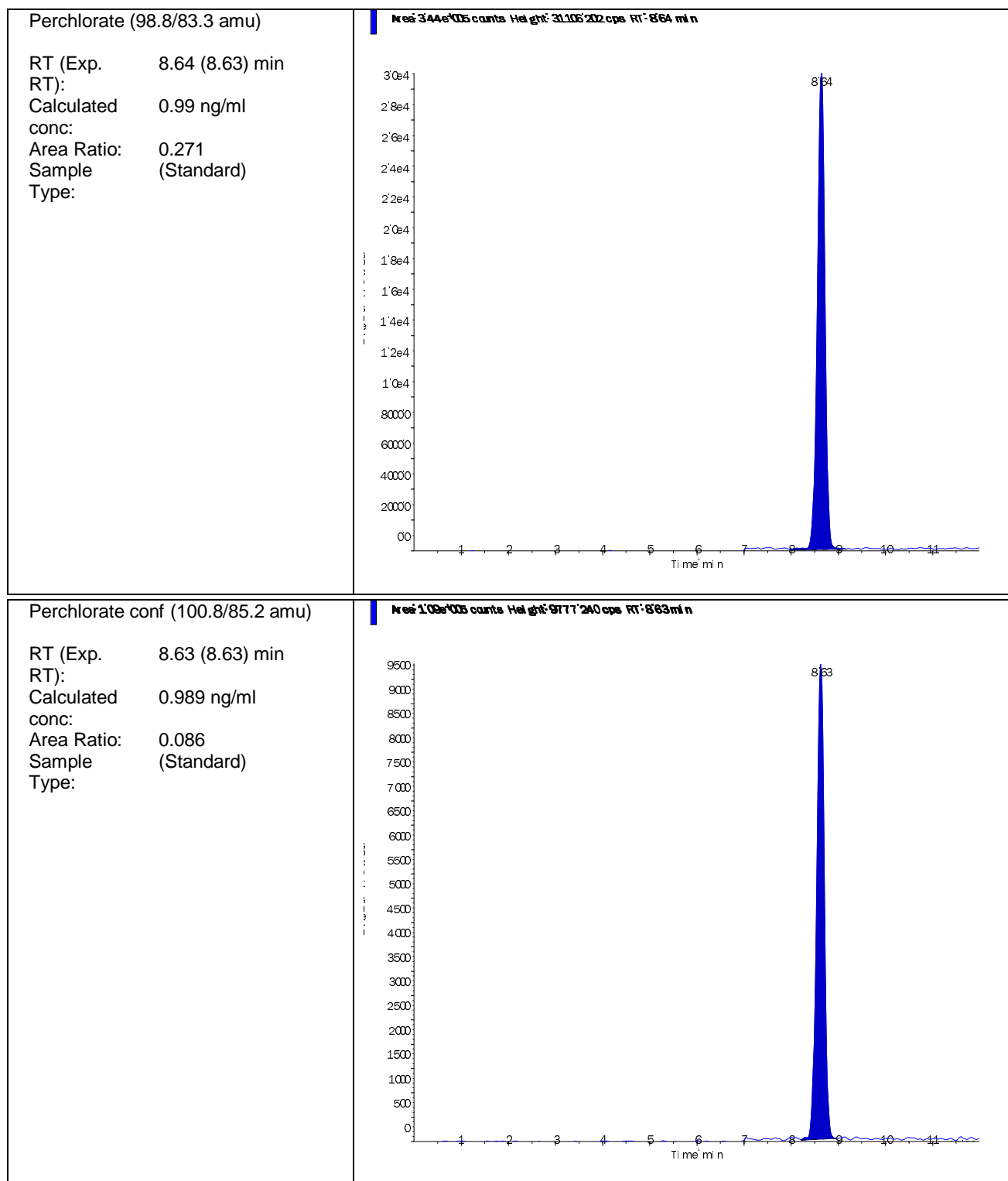
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Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-05 STD (1.0 ug/L)	Injection Vial	5.00
Data File	LM45218.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 6:00:31 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Standard
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-05	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.270e+06	8.63	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	3.440e+05	8.64	1.00	0.99
Perchlorate conf	1.090e+05	8.63	1.00	0.989





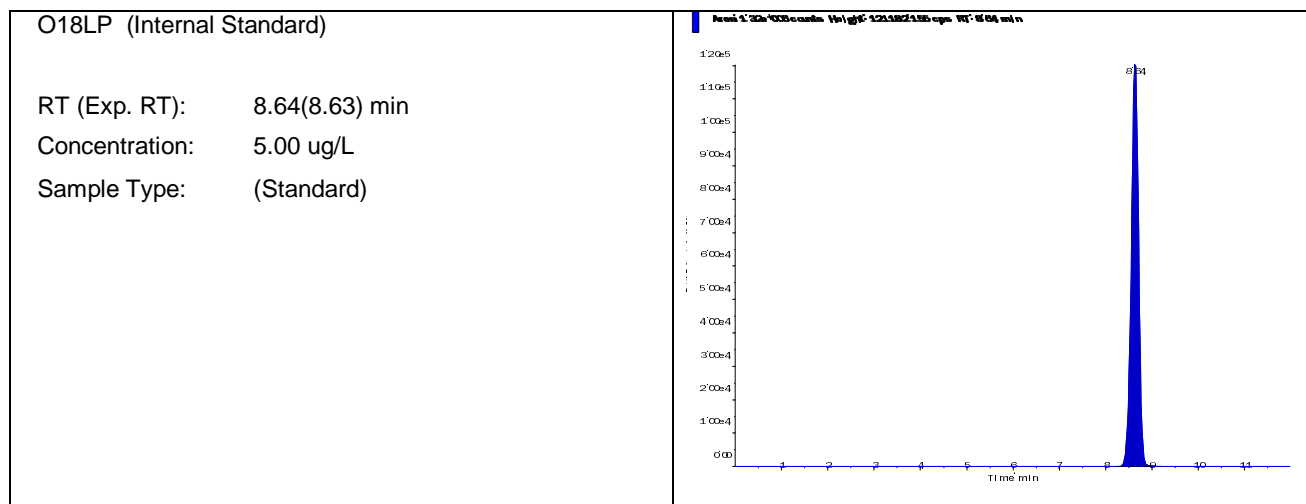
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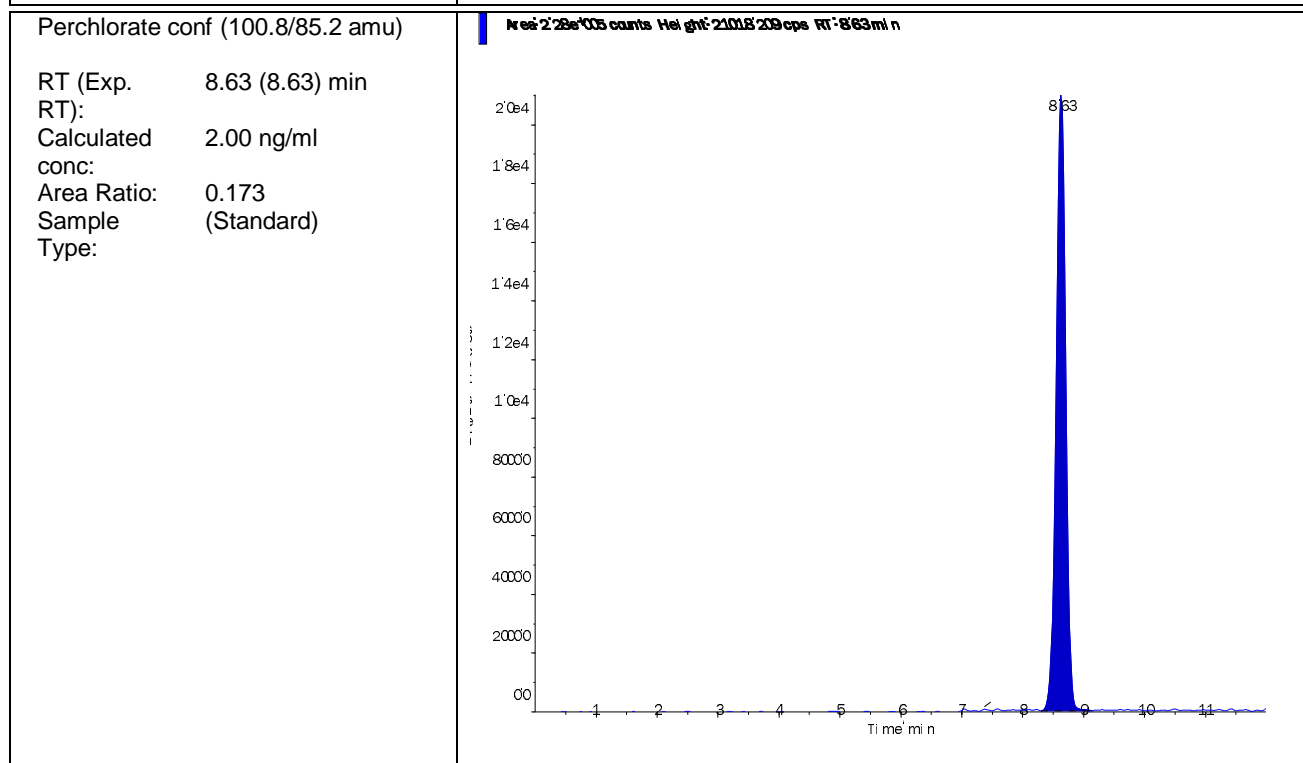
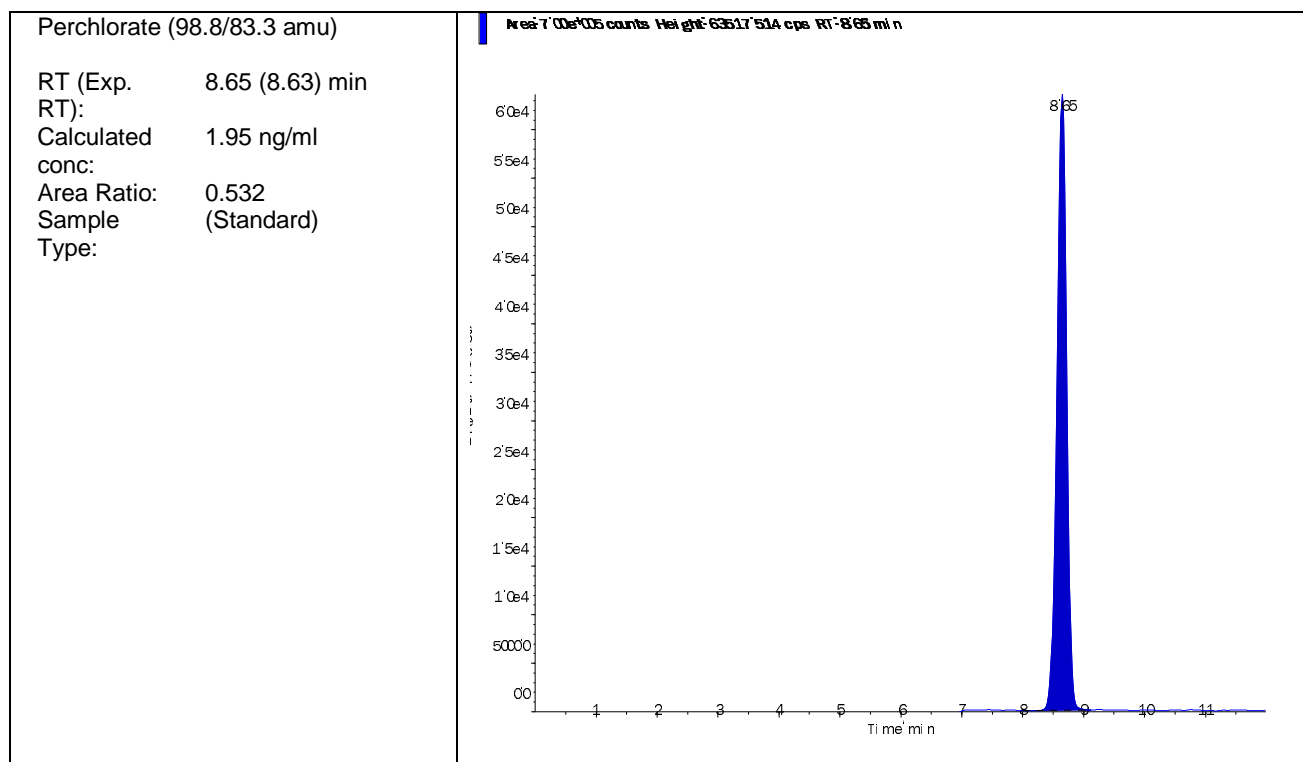
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Acquisition Date	3/18/2019 6:13:24 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-06 STD (2.0 ug/L)	Injection Vial	6.00
Data File	LM45219.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 6:13:24 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Standard
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-06	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.320e+06	8.64	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	7.000e+05	8.65	2.00	1.95
Perchlorate conf	2.280e+05	8.63	2.00	2.00



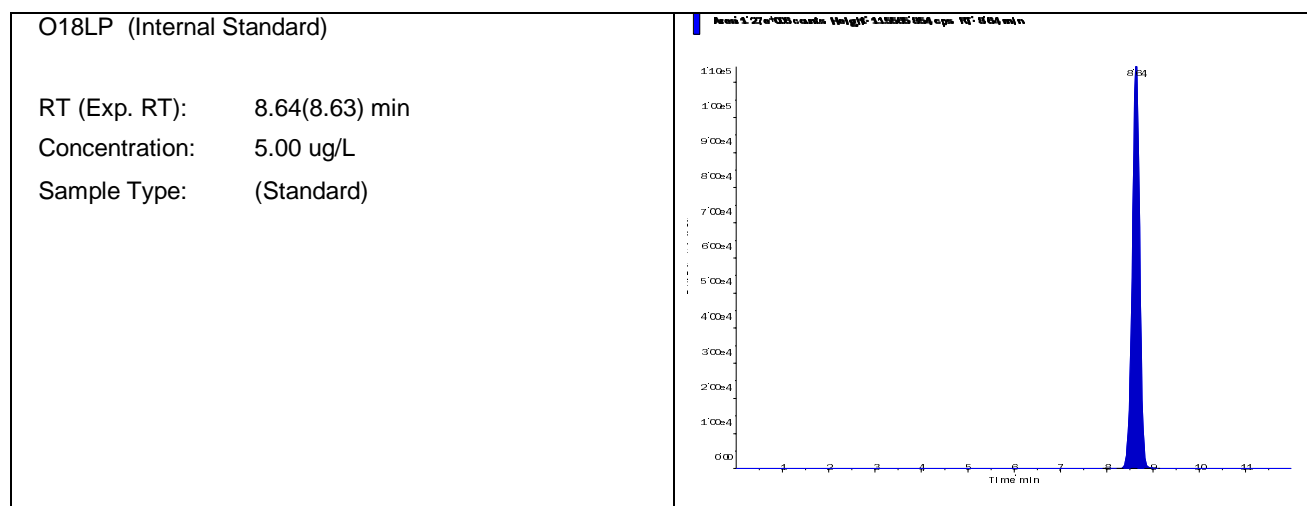


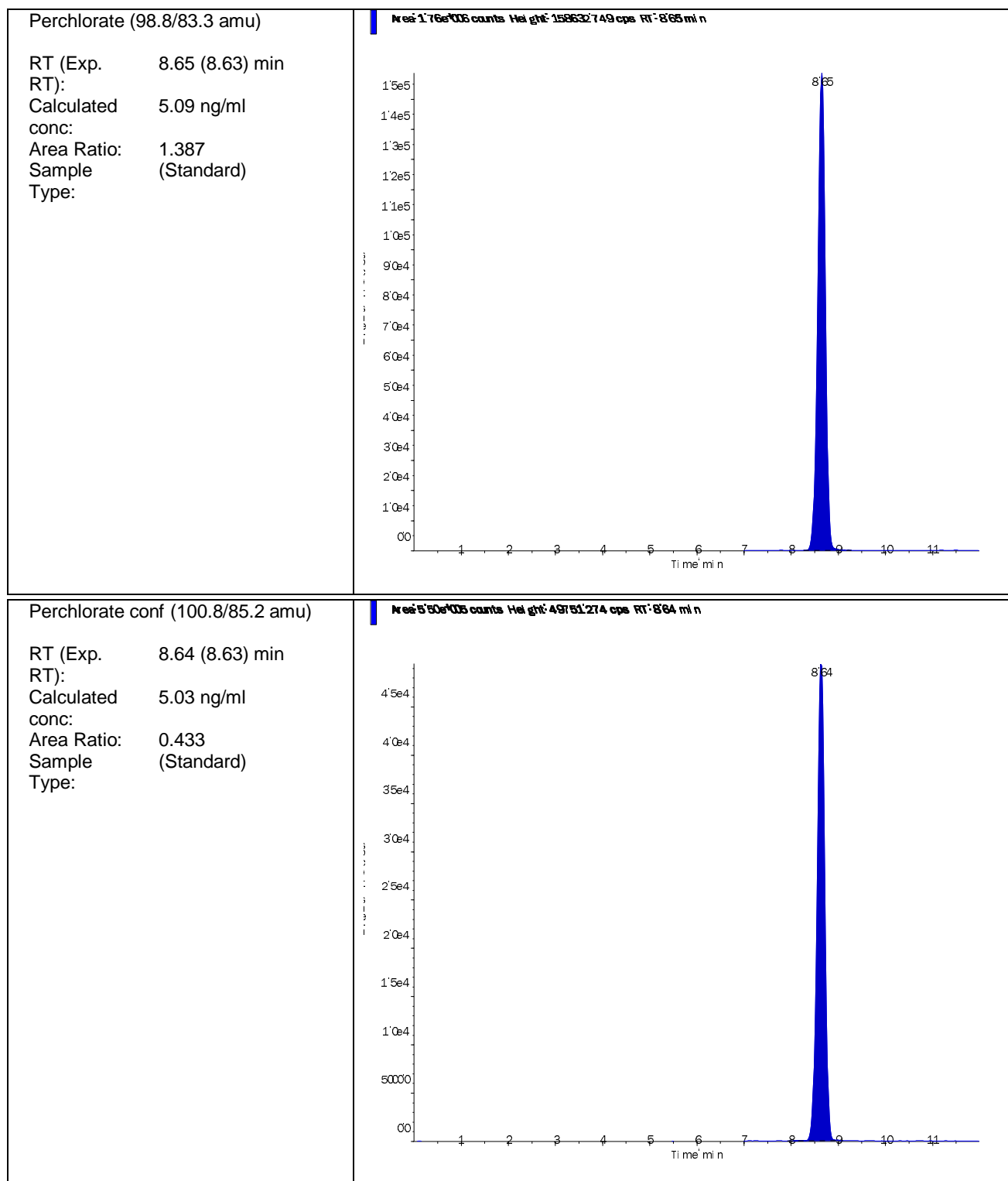
Data File	LM45220.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 6:26:23 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-07 STD (5.0 ug/L)	Injection Vial	7.00
Data File	LM45220.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 6:26:23 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Standard
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-07	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.270e+06	8.64	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	1.760e+06	8.65	5.00	5.09
Perchlorate conf	5.500e+05	8.64	5.00	5.03





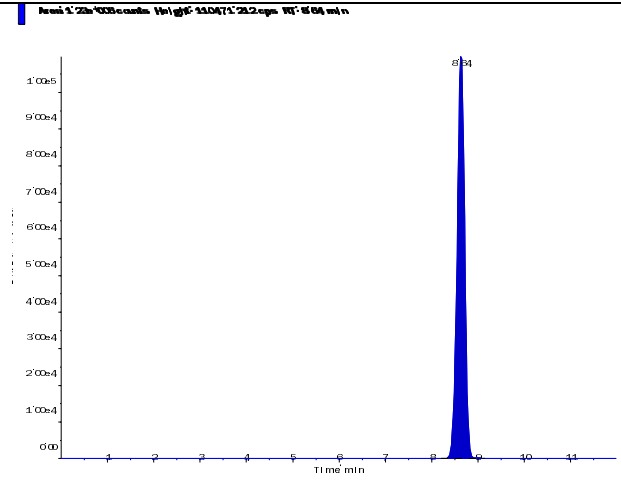
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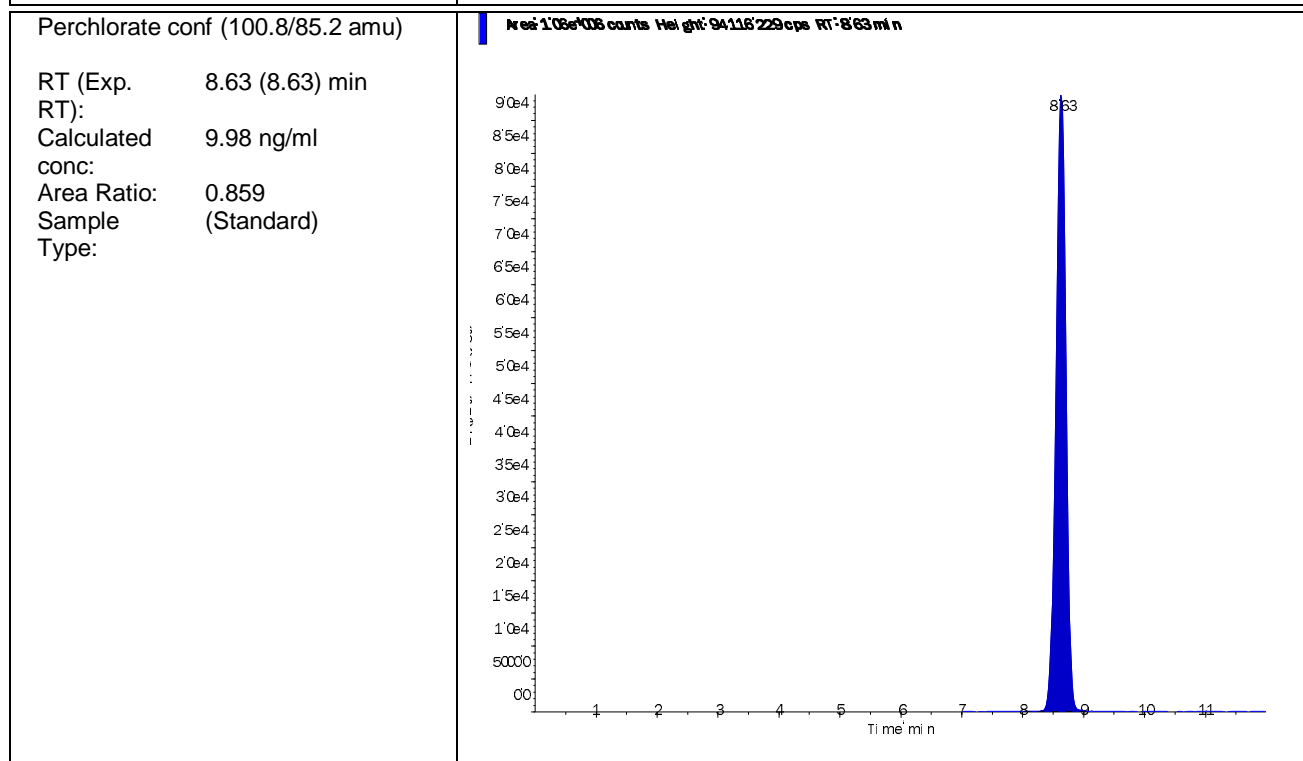
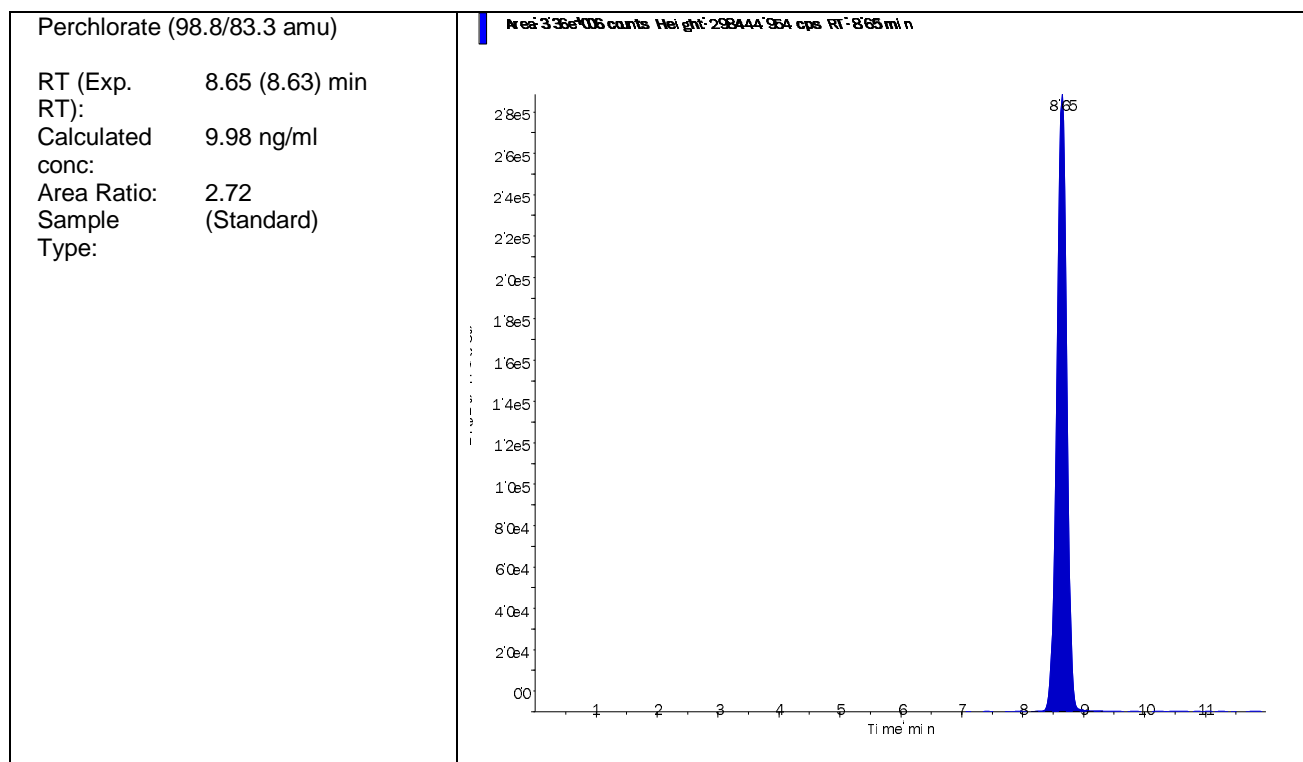
Data File	LM45221.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 6:39:19 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-08 STD (10 ug/L)	Injection Vial	8.00
Data File	LM45221.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 6:39:19 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Standard
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-08	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.230e+06	8.64	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	3.360e+06	8.65	10.00	9.98
Perchlorate conf	1.060e+06	8.63	10.00	9.98

O18LP (Internal Standard)	
RT (Exp. RT):	8.64(8.63) min
Concentration:	5.00 ug/L
Sample Type:	(Standard)



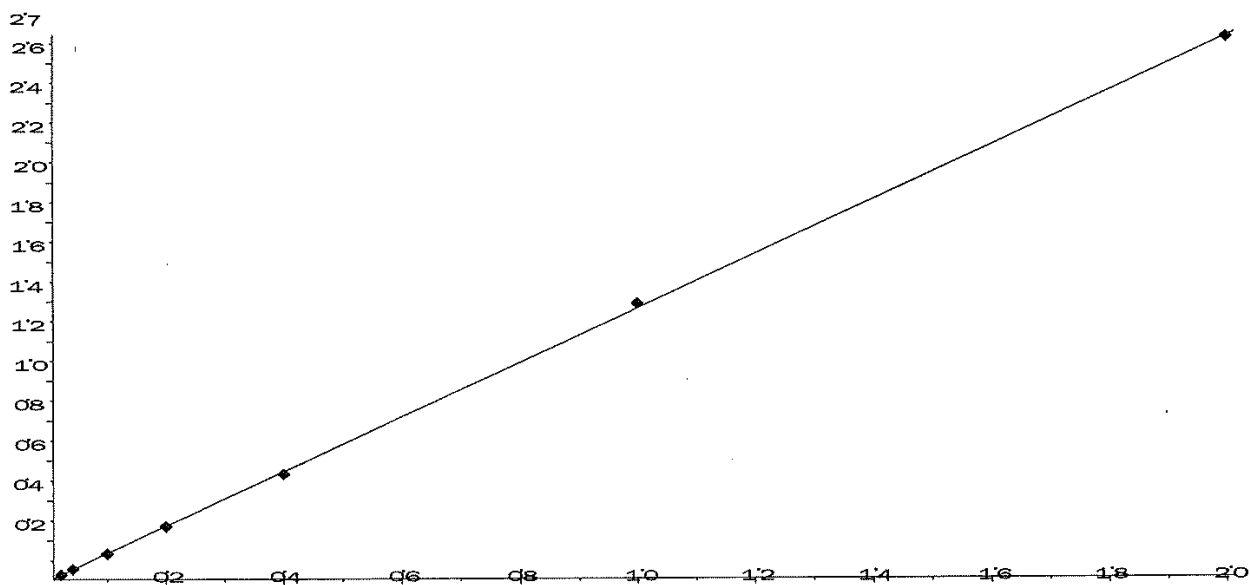
Analyte Name: Perchlorate
Internal Standard: O18LP

Data File	LM45214.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 5:08:52 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Regression Equation: $y = 1.36x + 0.00113$ ($r = 0.9999$)

Expected Concentration	Number of Values	Mean Calculated Concentration	% Accuracy	Std. Deviation	%CV
0.10	1	0.10	101.3	N/A	N/A
0.20	1	0.21	103.1	N/A	N/A
0.50	1	0.49	97.7	N/A	N/A
1.00	1	0.99	99.0	N/A	N/A
2.00	1	1.95	97.4	N/A	N/A
5.00	1	5.09	101.7	N/A	N/A
10.00	1	9.98	99.8	N/A	N/A

$y = 1.36x + 0.00113$ ($r = 0.9999$)



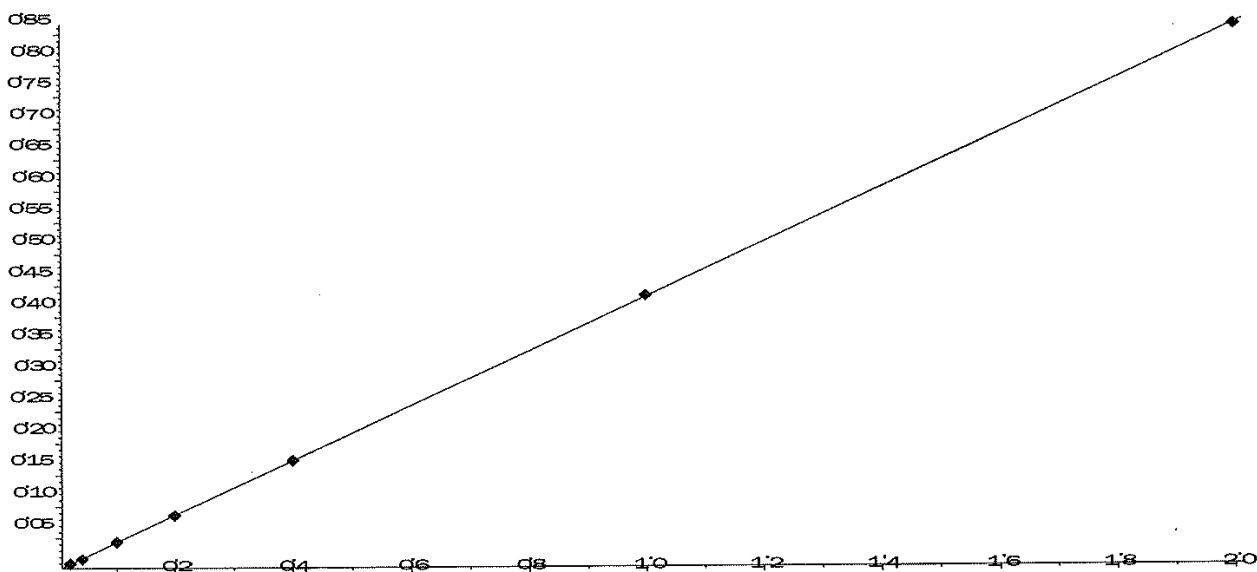
Analyte Name: Perchlorate conf
Internal Standard: O18LP

Data File	LM45214.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 5:08:52 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Regression Equation: $y = 0.43x + 0.000678$ ($r = 1.0000$)

Expected Concentration	Number of Values	Mean Calculated Concentration	% Accuracy	Std. Deviation	%CV
0.10	1	0.10	103.6	N/A	N/A
0.20	1	0.19	96.4	N/A	N/A
0.50	1	0.50	100.6	N/A	N/A
1.00	1	0.99	98.9	N/A	N/A
2.00	1	2.00	100.2	N/A	N/A
5.00	1	5.03	100.5	N/A	N/A
10.00	1	9.98	99.8	N/A	N/A

$$y = 0.43x + 0.000678 \quad (r = 1.0000)$$

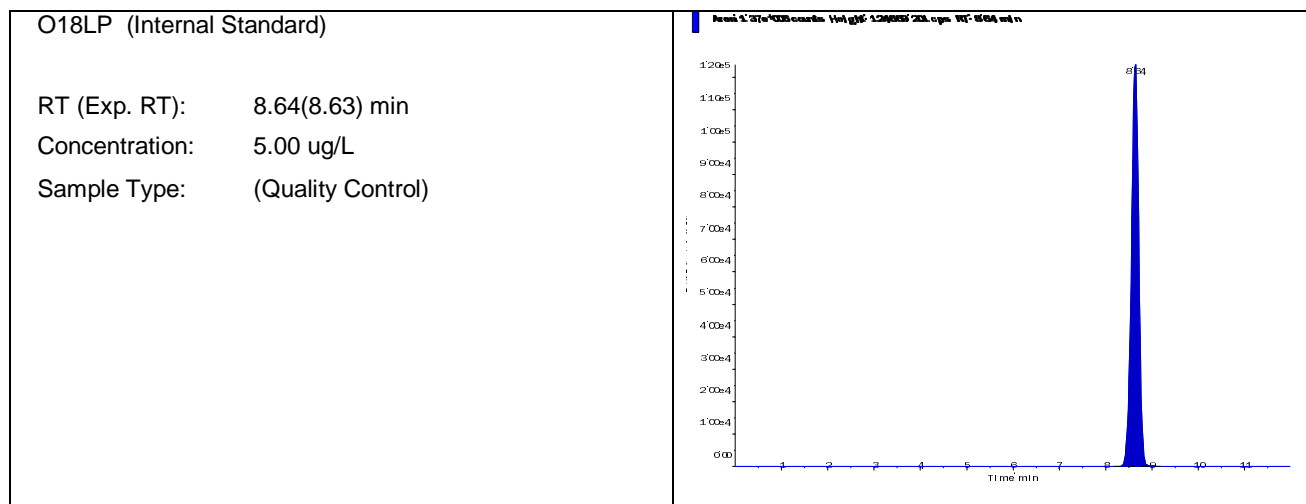


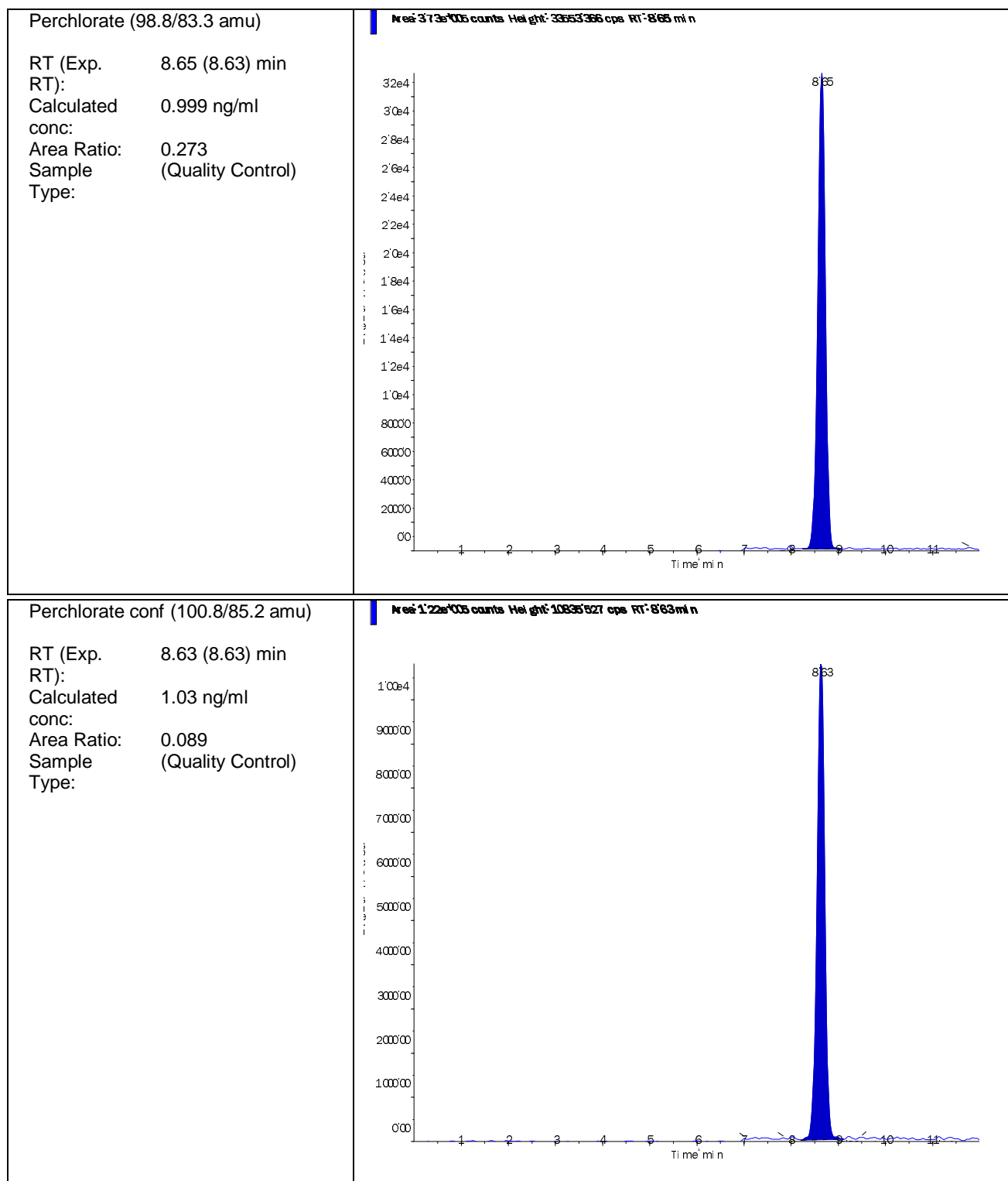
Data File	LM45222.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 6:52:15 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG700056-09 SSCV (1.0 ug/L)	Injection Vial	9.00
Data File	LM45222.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 6:52:15 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Quality Control
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG700056-09	Dilution Factor	1.00
Sample Comment	1,1 STD87534	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.370e+06	8.64	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	3.730e+05	8.65	1.00	0.999
Perchlorate conf	1.220e+05	8.63	1.00	1.03



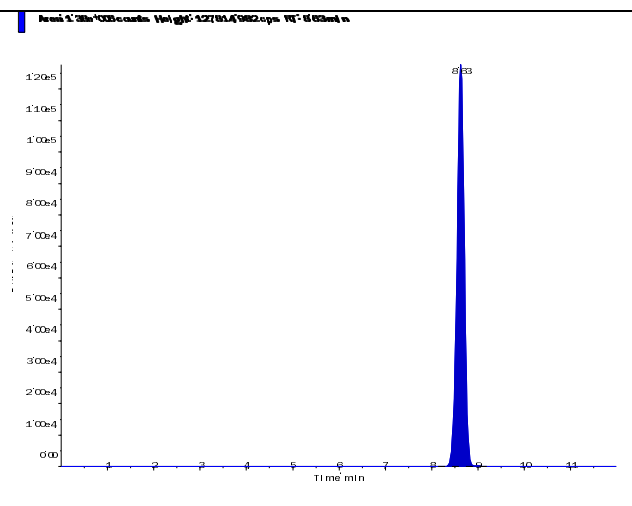


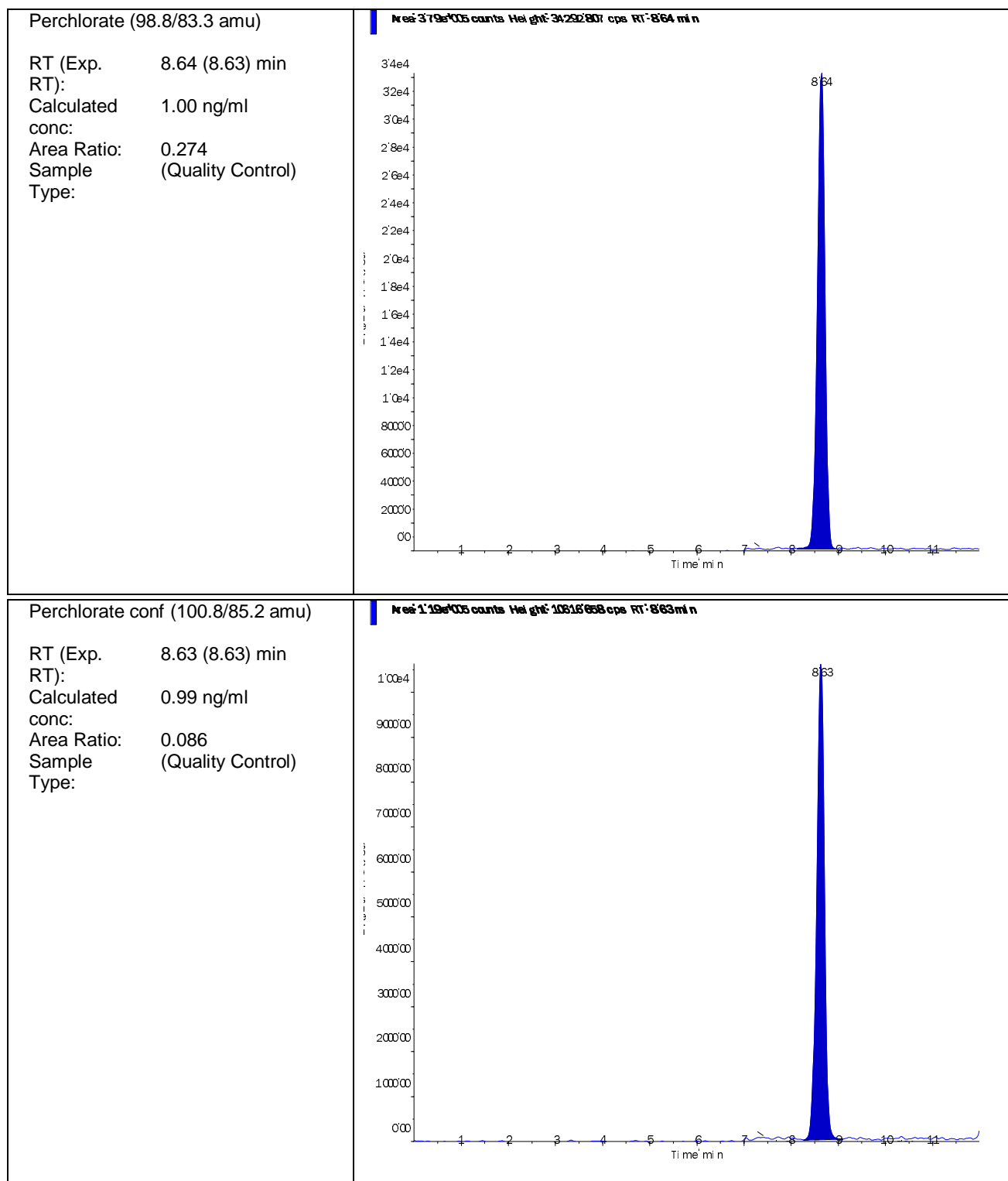
Data File	LM45224.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 7:18:09 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699915-02 CCV (1.0ug/L)	Injection Vial	5.00
Data File	LM45224.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 7:18:09 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Quality Control
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699915-02	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.380e+06	8.63	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	3.790e+05	8.64	1.00	1.00
Perchlorate conf	1.190e+05	8.63	1.00	0.99

O18LP (Internal Standard)	
RT (Exp. RT):	8.63(8.63) min
Concentration:	5.00 ug/L
Sample Type:	(Quality Control)

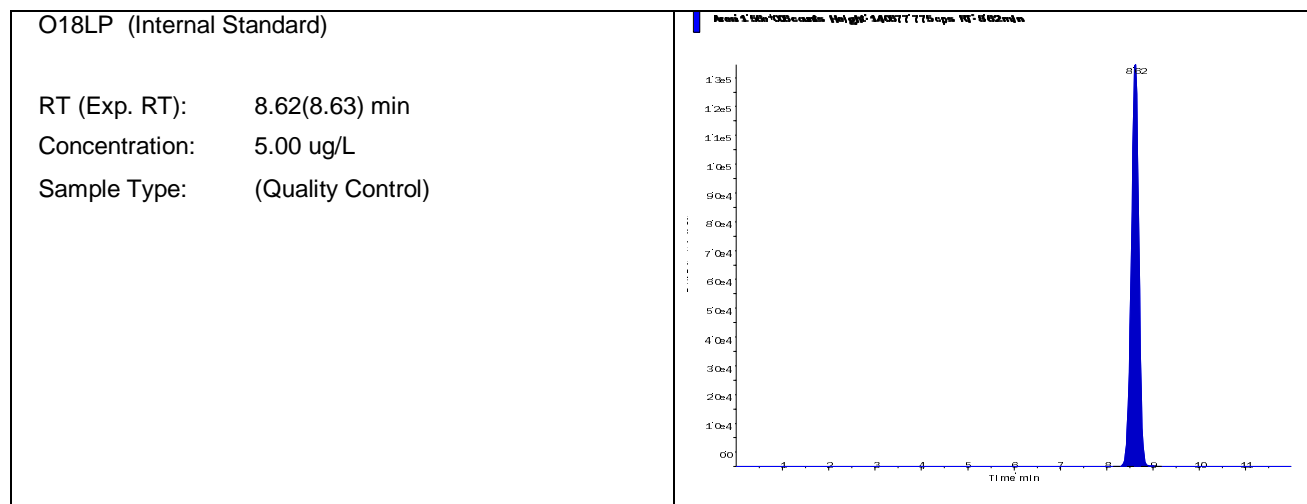


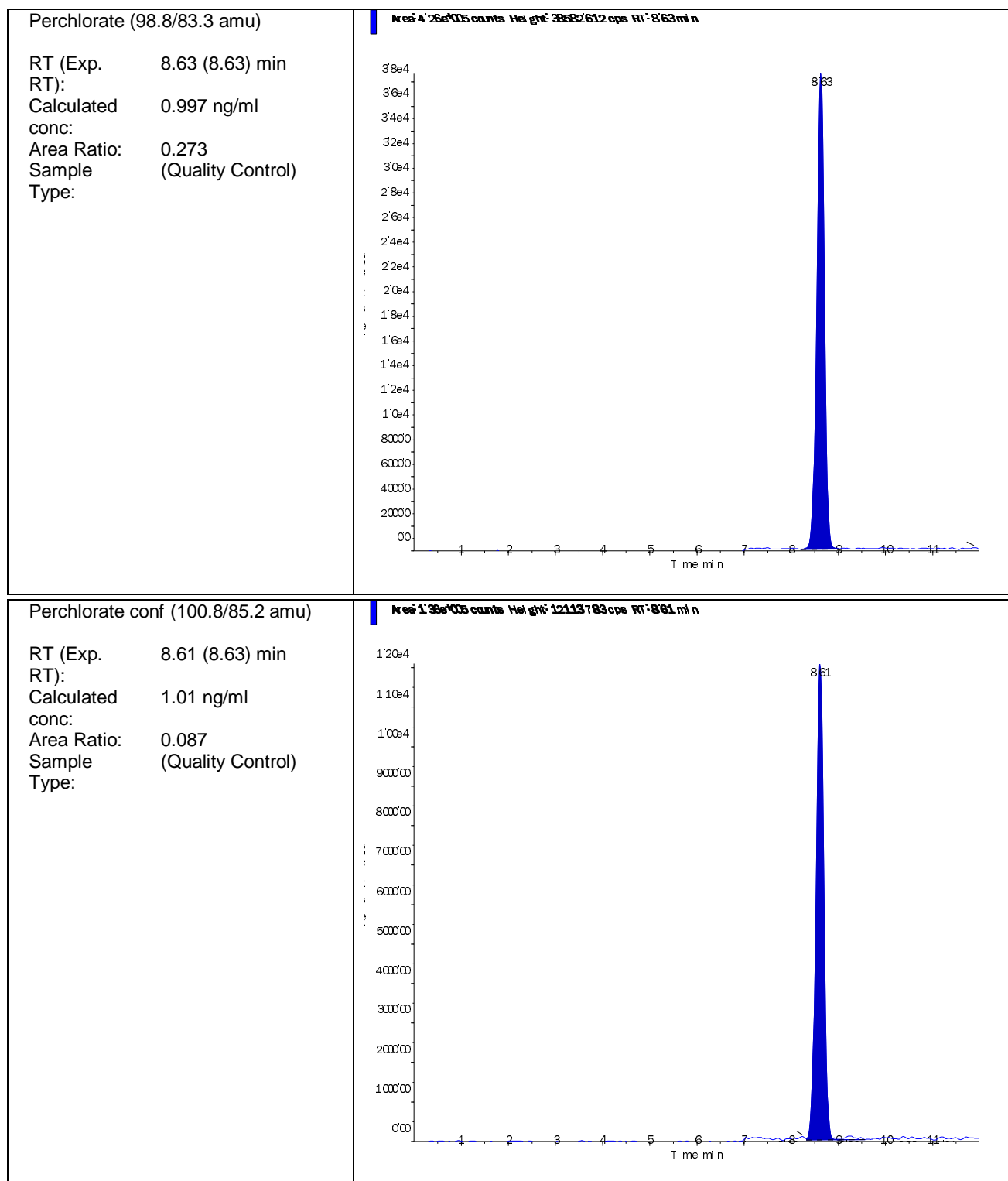
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Acquisition Date	3/18/2019 9:53:21 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699915-03 CCV (1.0ug/L)	Injection Vial	5.00
Data File	LM45236.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 9:53:21 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Quality Control
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699915-03	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.560e+06	8.62	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	4.260e+05	8.63	1.00	0.997
Perchlorate conf	1.360e+05	8.61	1.00	1.01





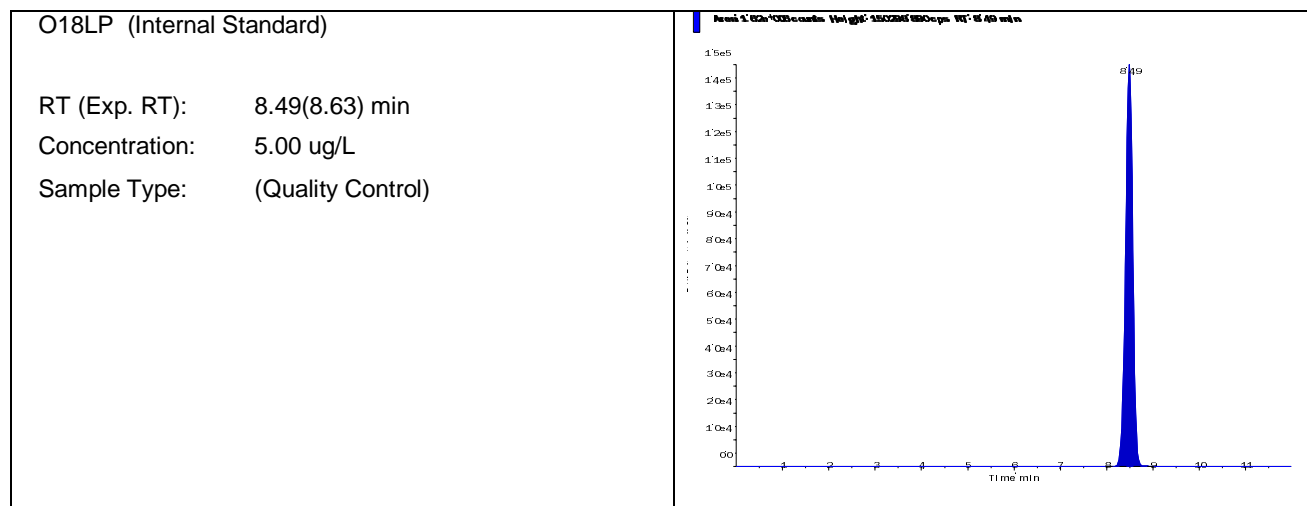
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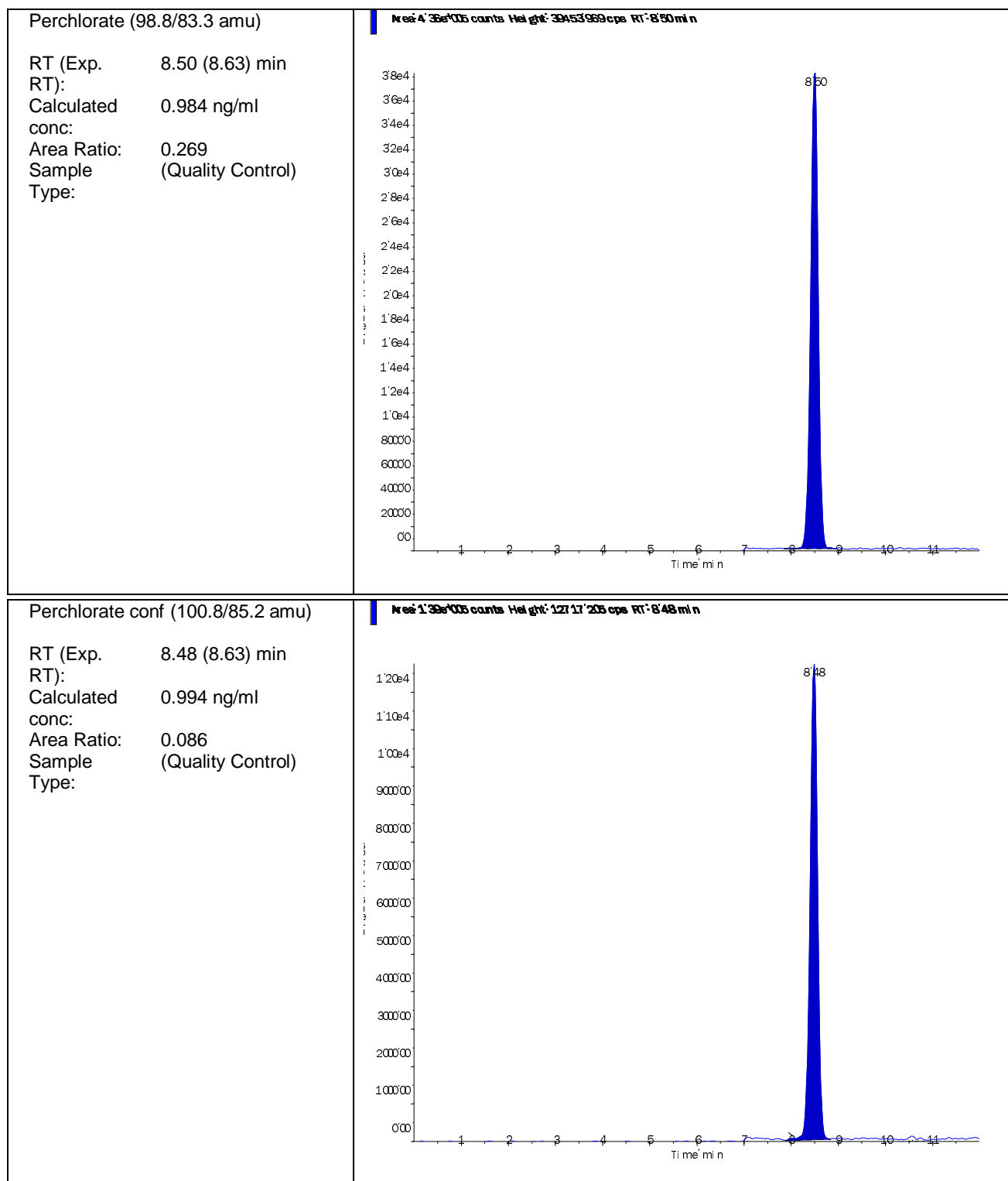
Data File	LM45242.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 11:10:55 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699915-05 CCV (1.0ug/L)	Injection Vial	5.00
Data File	LM45242.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 11:10:55 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Quality Control
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699915-05	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.620e+06	8.49	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	4.360e+05	8.50	1.00	0.984
Perchlorate conf	1.390e+05	8.48	1.00	0.994





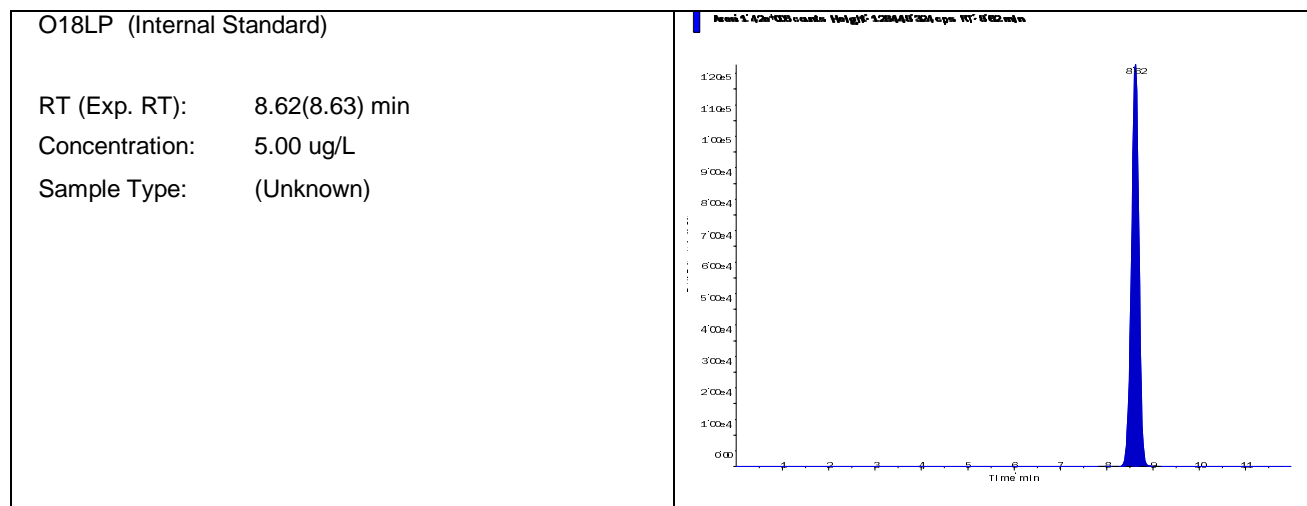
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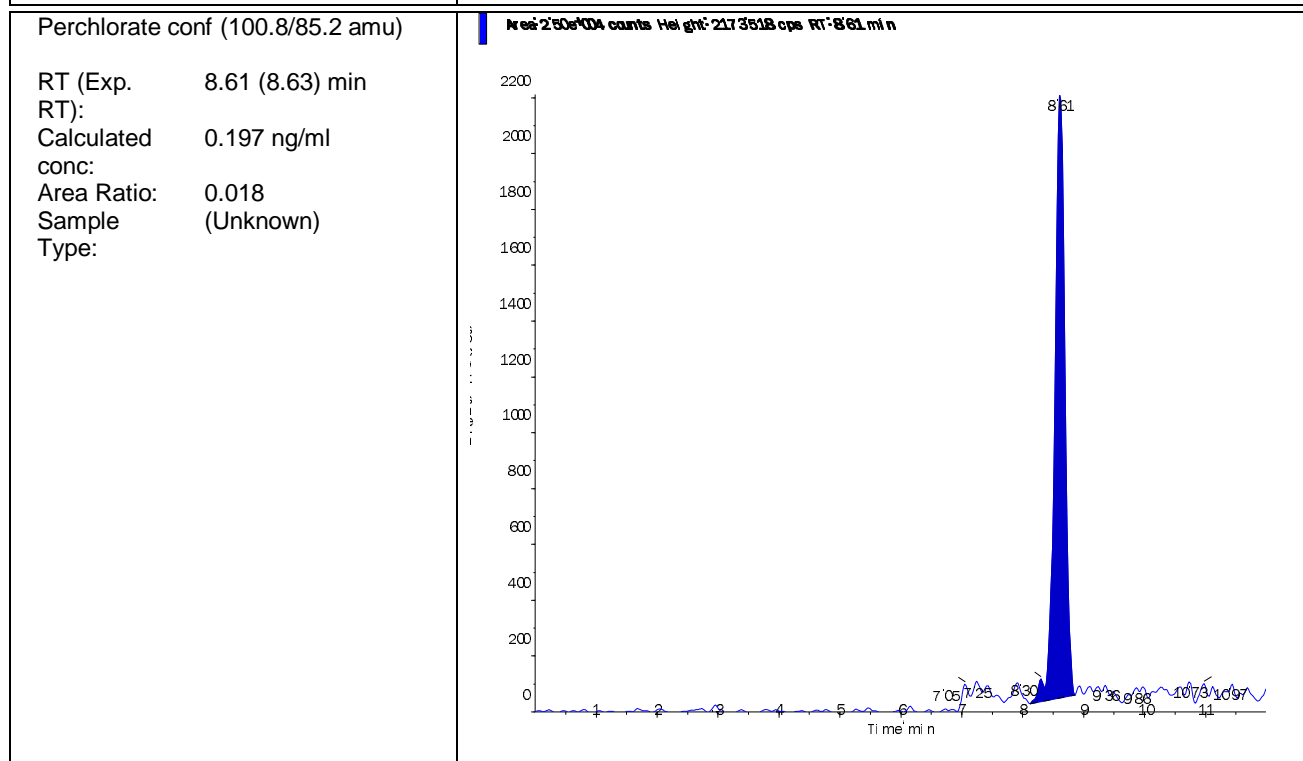
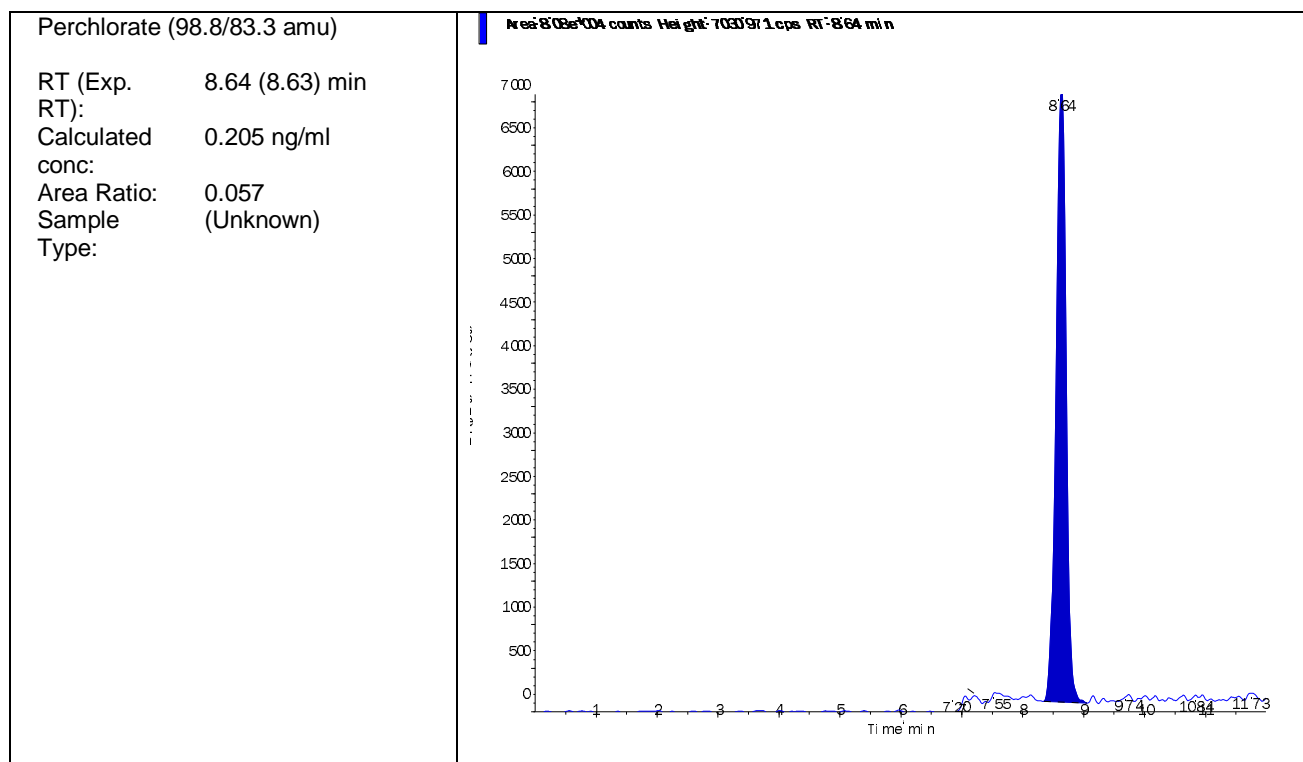
Data File	LM45225.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 7:31:05 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699914-07 MRL (0.2ug/L)	Injection Vial	3.00
Data File	LM45225.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 7:31:05 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-07	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.420e+06	8.62	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	8.080e+04	8.64	N/A	0.205
Perchlorate conf	2.500e+04	8.61	N/A	0.197



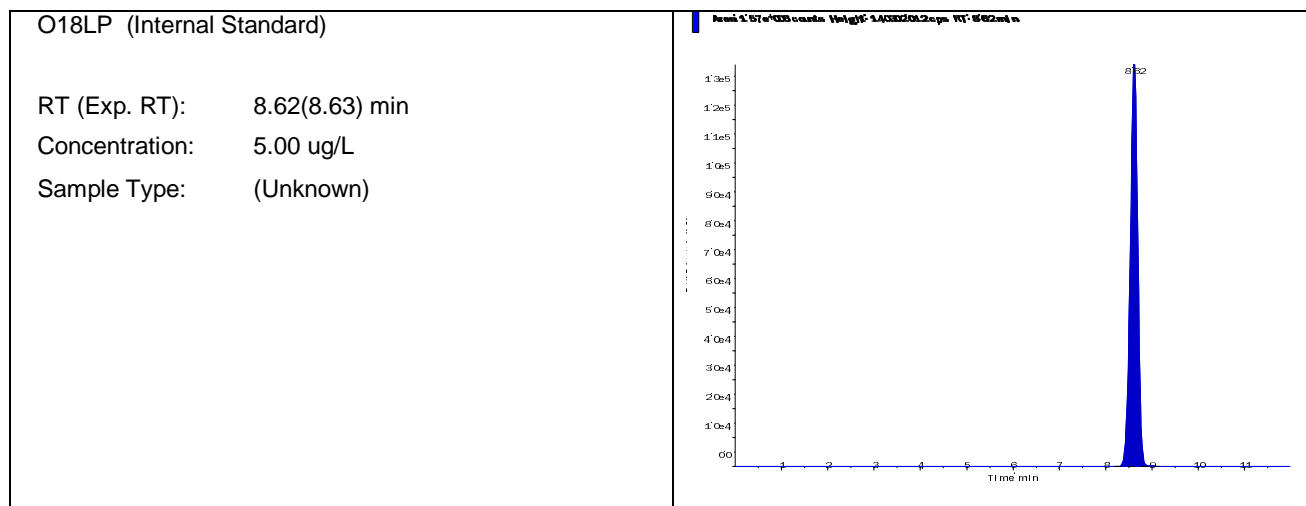


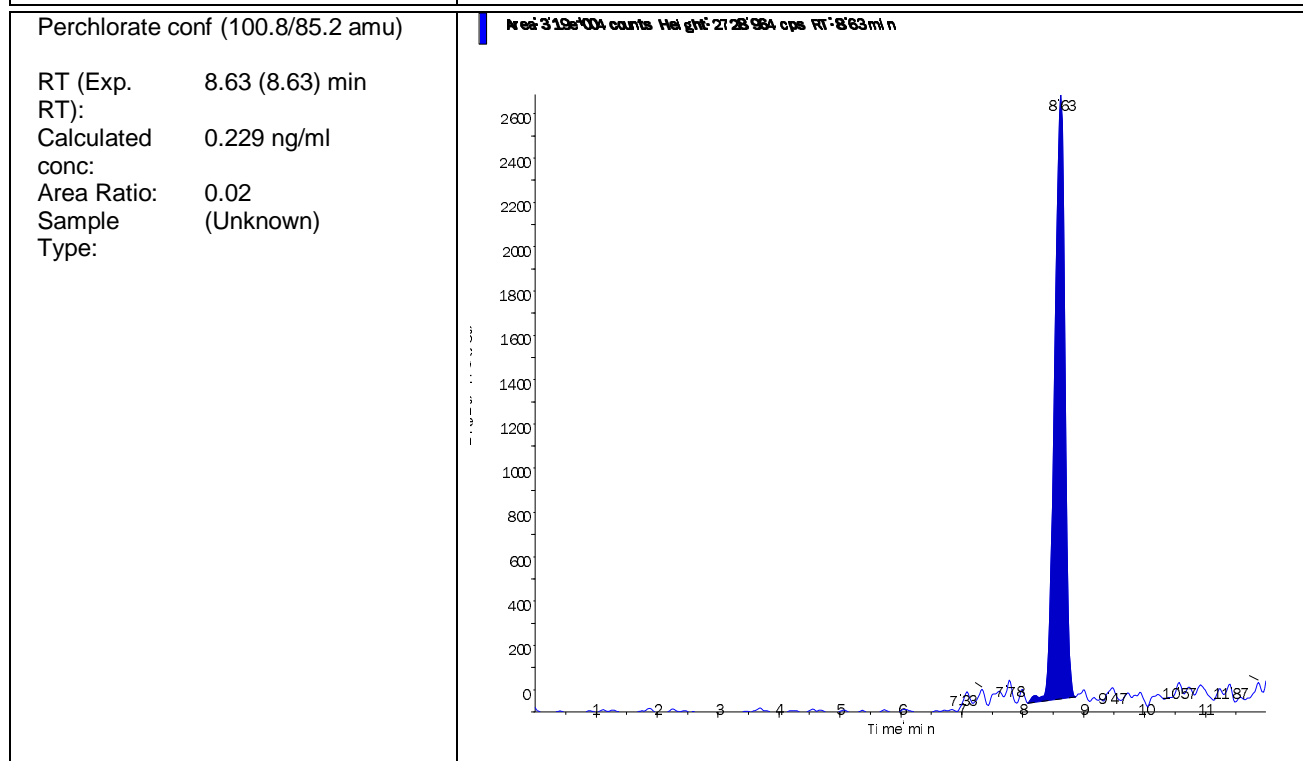
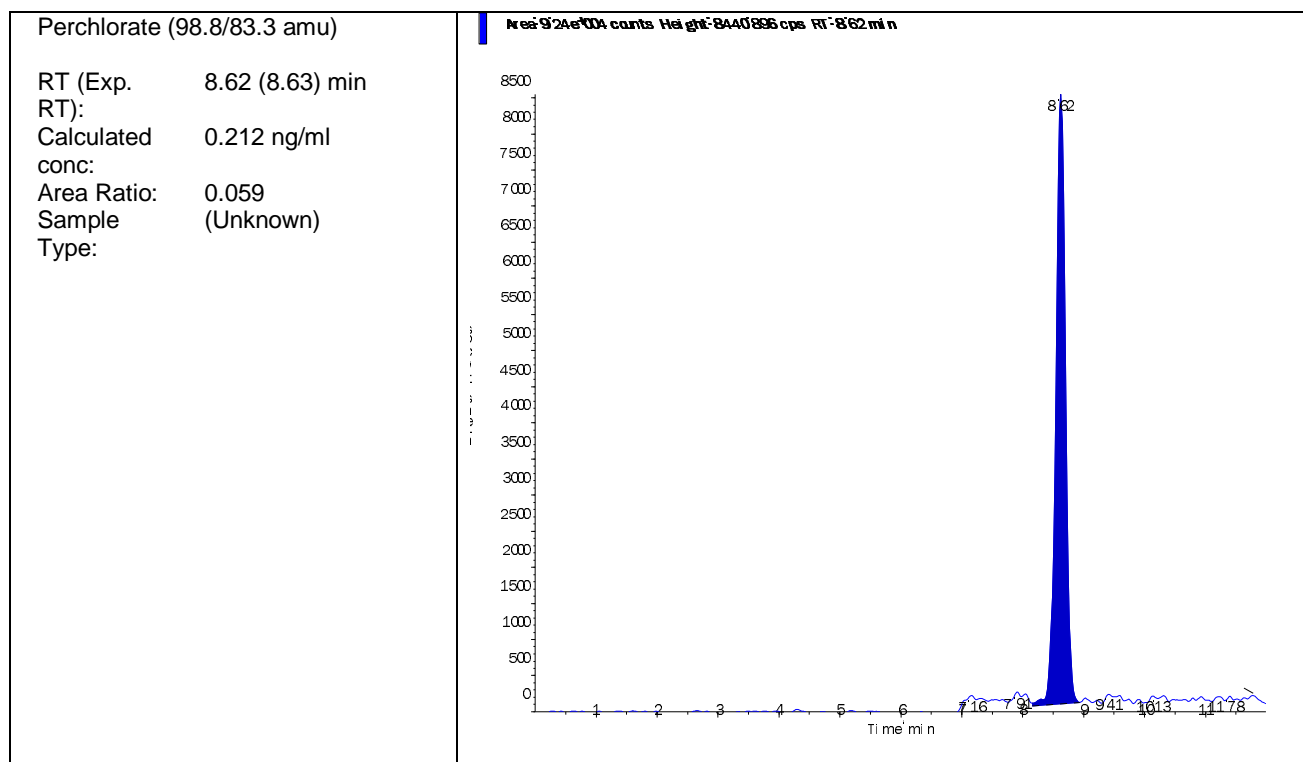
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Acquisition Date	3/18/2019 10:06:16 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699914-08 MRL (0.2ug/L)	Injection Vial	3.00
Data File	LM45237.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 10:06:16 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-08	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.570e+06	8.62	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	9.240e+04	8.62	N/A	0.212
Perchlorate conf	3.190e+04	8.63	N/A	0.229



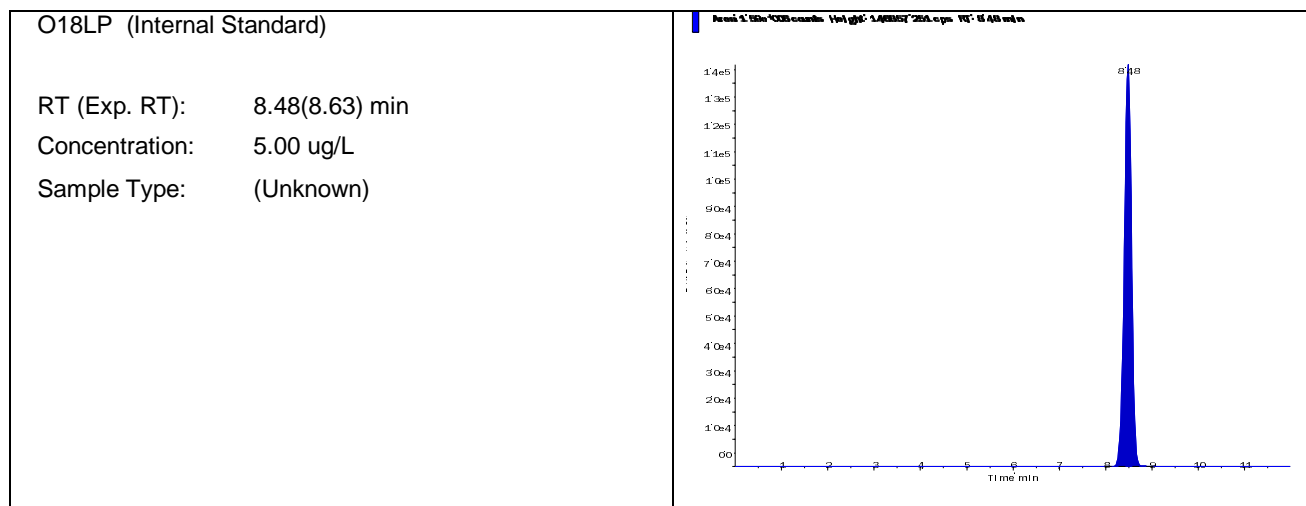


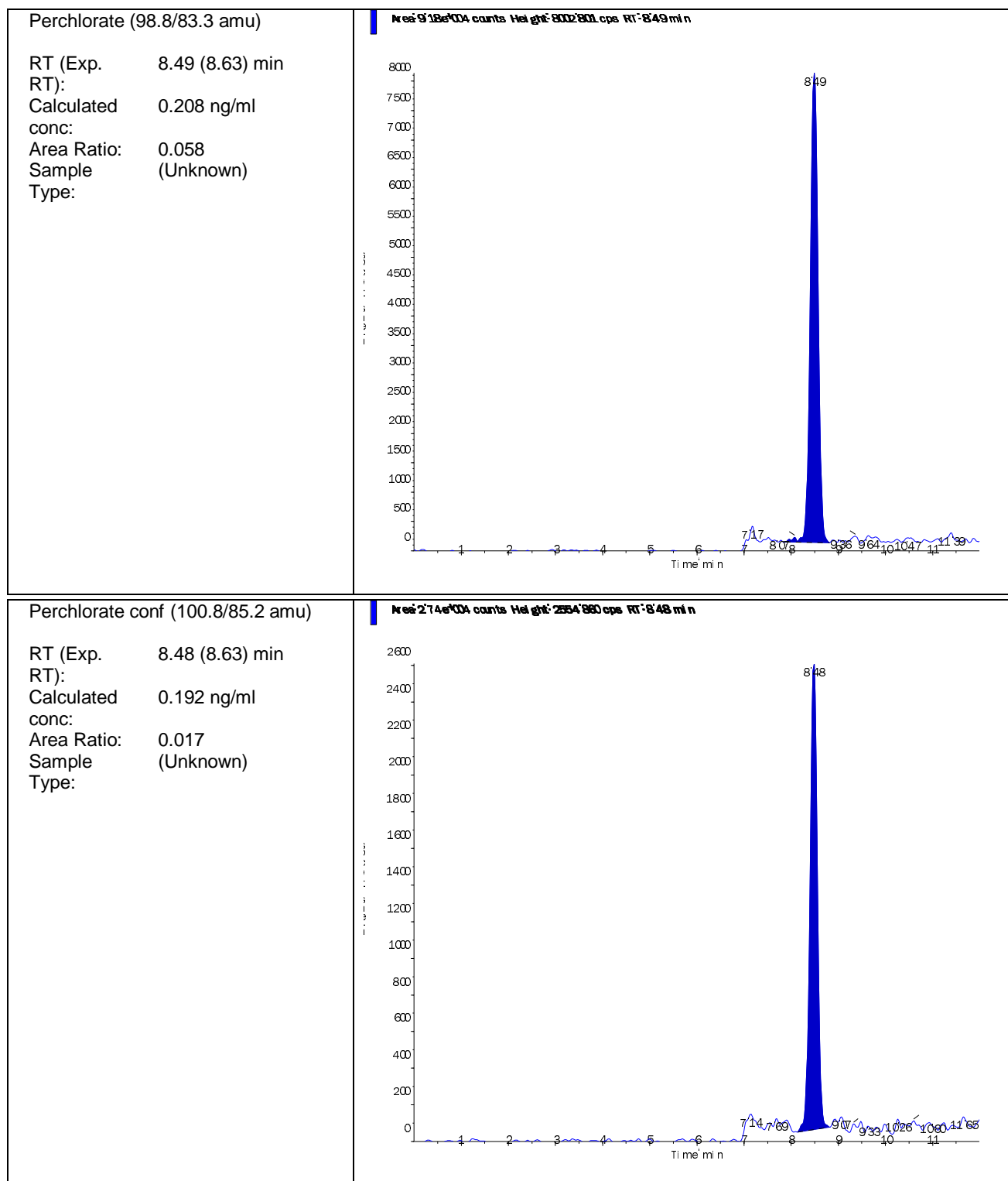
Data File	LM45243.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 11:23:52 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699914-09 MRL (0.2ug/L)	Injection Vial	3.00
Data File	LM45243.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 11:23:52 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-09	Dilution Factor	1.00
Sample Comment	1,1 STD87532	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.590e+06	8.48	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	9.180e+04	8.49	N/A	0.208
Perchlorate conf	2.740e+04	8.48	N/A	0.192



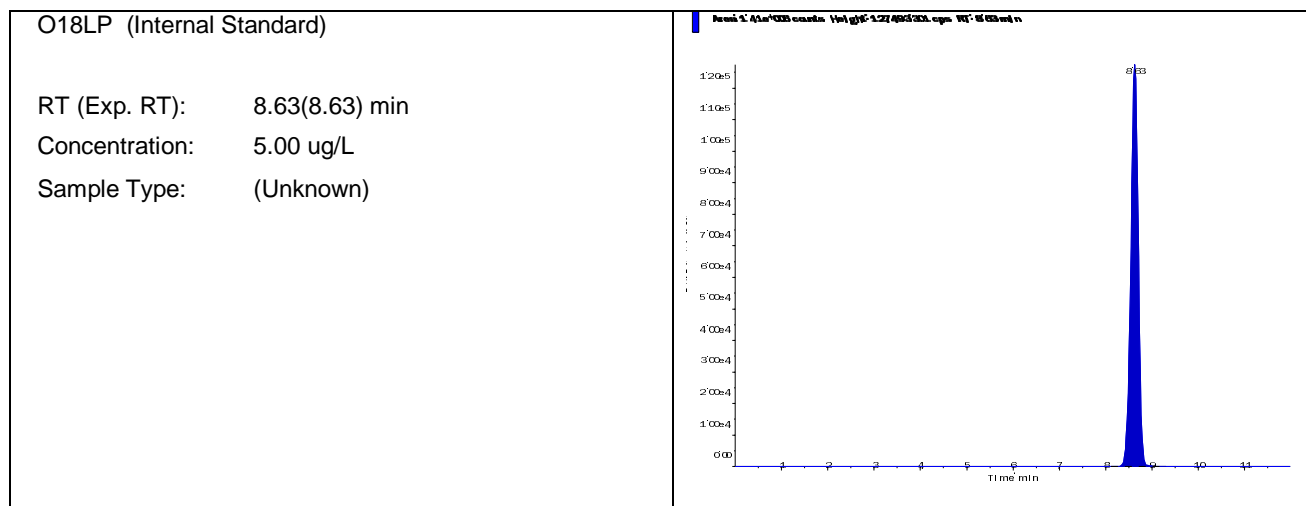


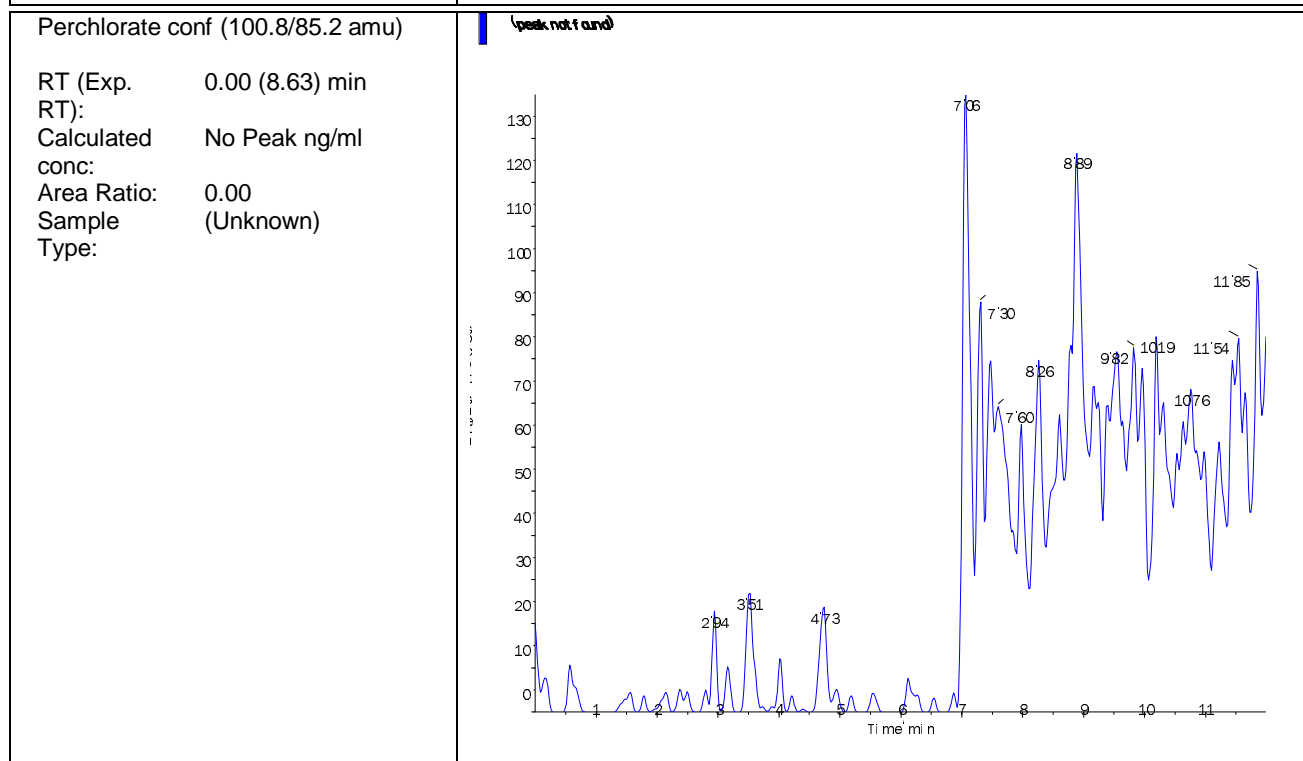
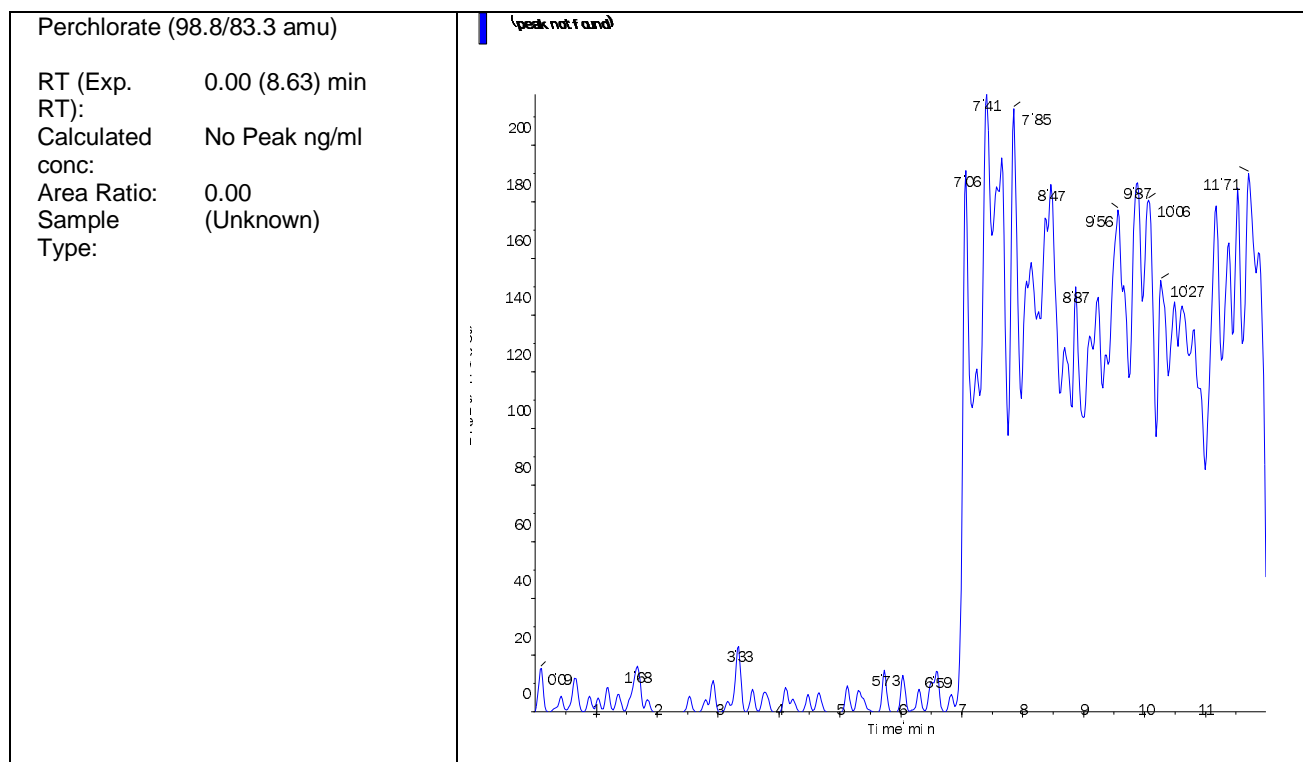
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Acquisition Date	3/18/2019 7:05:13 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699915-01 CCB	Injection Vial	1.00
Data File	LM45223.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 7:05:13 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699915-01	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.410e+06	8.63	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	0.000e+00	0.00	N/A	No Peak
Perchlorate conf	0.000e+00	0.00	N/A	No Peak



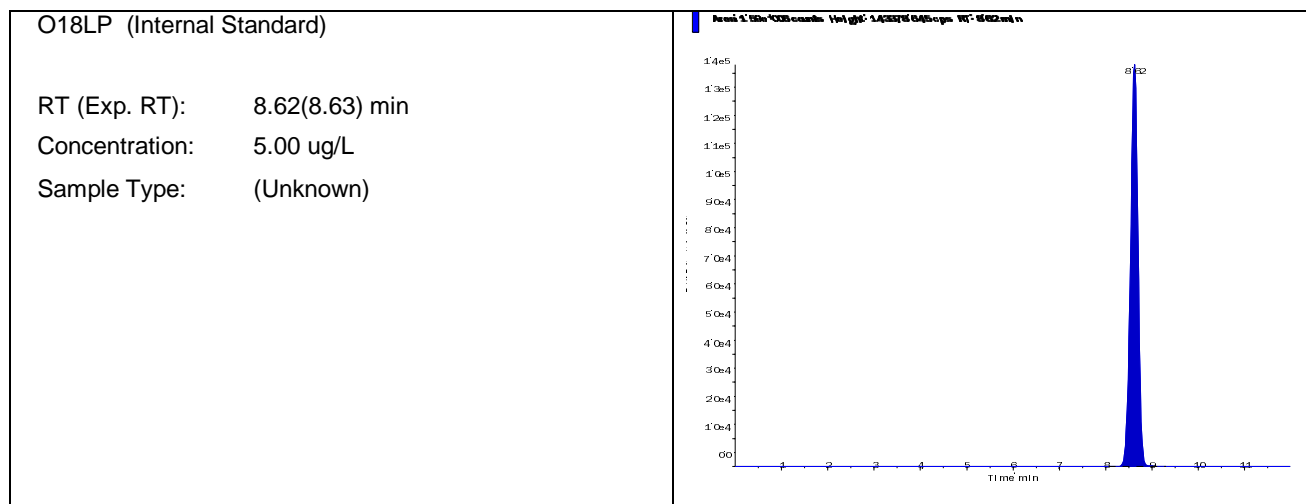


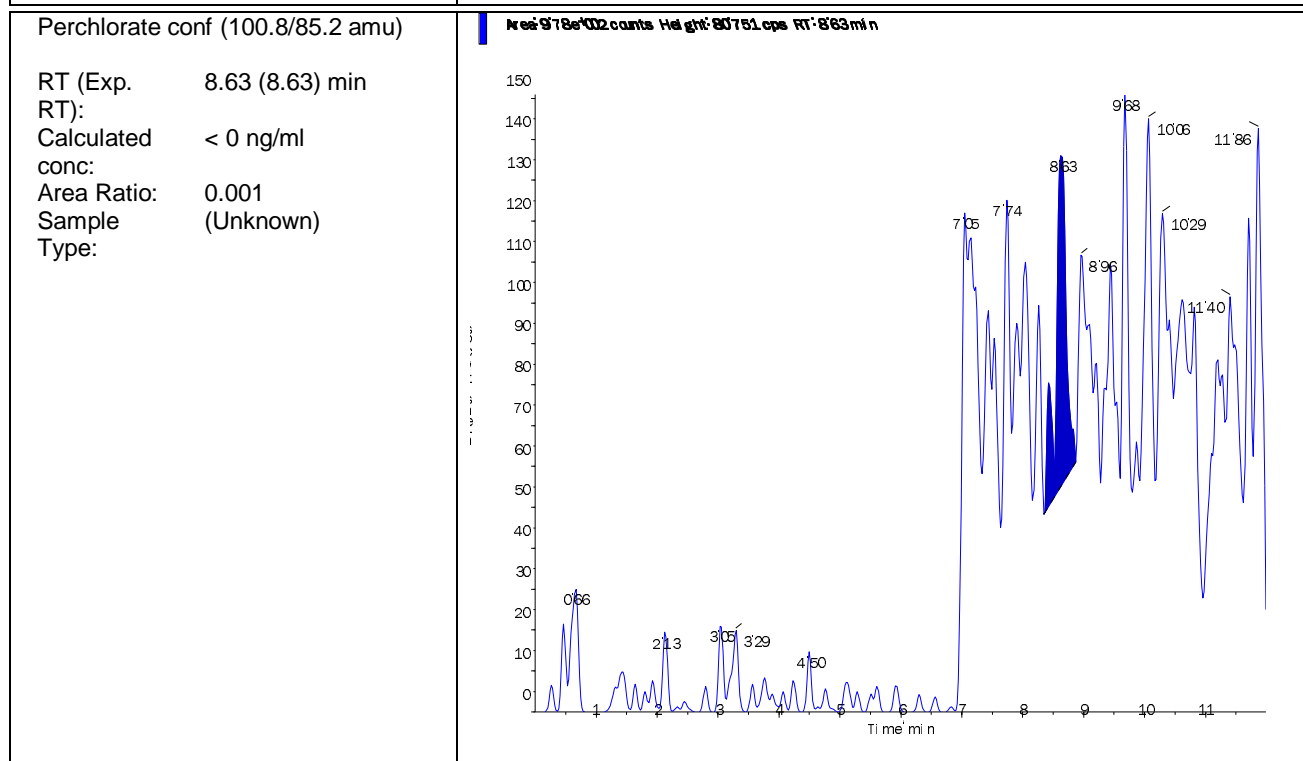
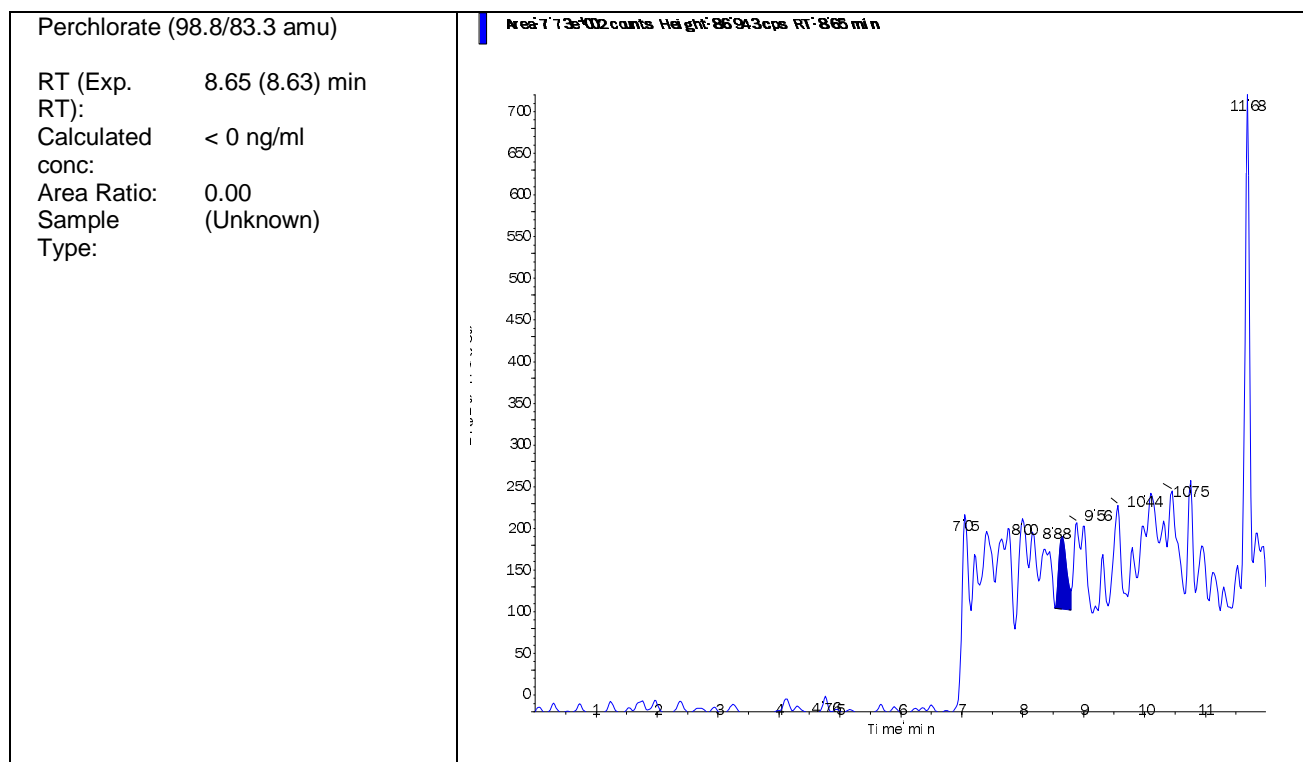
Data File	LM45238.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 10:19:13 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699915-04 CCB	Injection Vial	1.00
Data File	LM45238.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 10:19:13 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699915-04	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.590e+06	8.62	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	7.730e+02	8.65	N/A	< 0
Perchlorate conf	9.780e+02	8.63	N/A	< 0



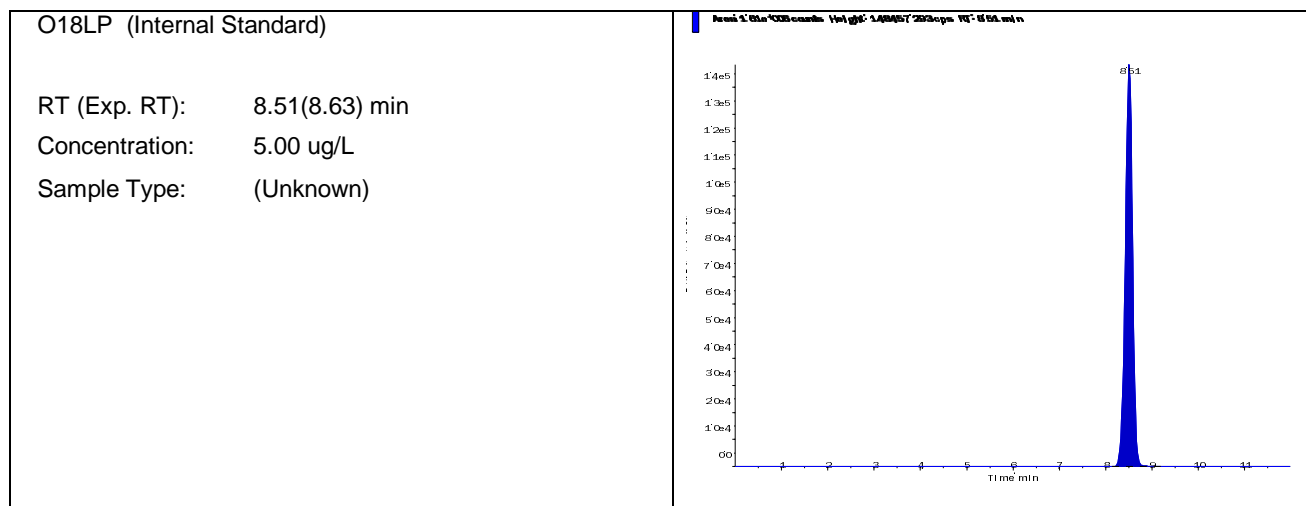


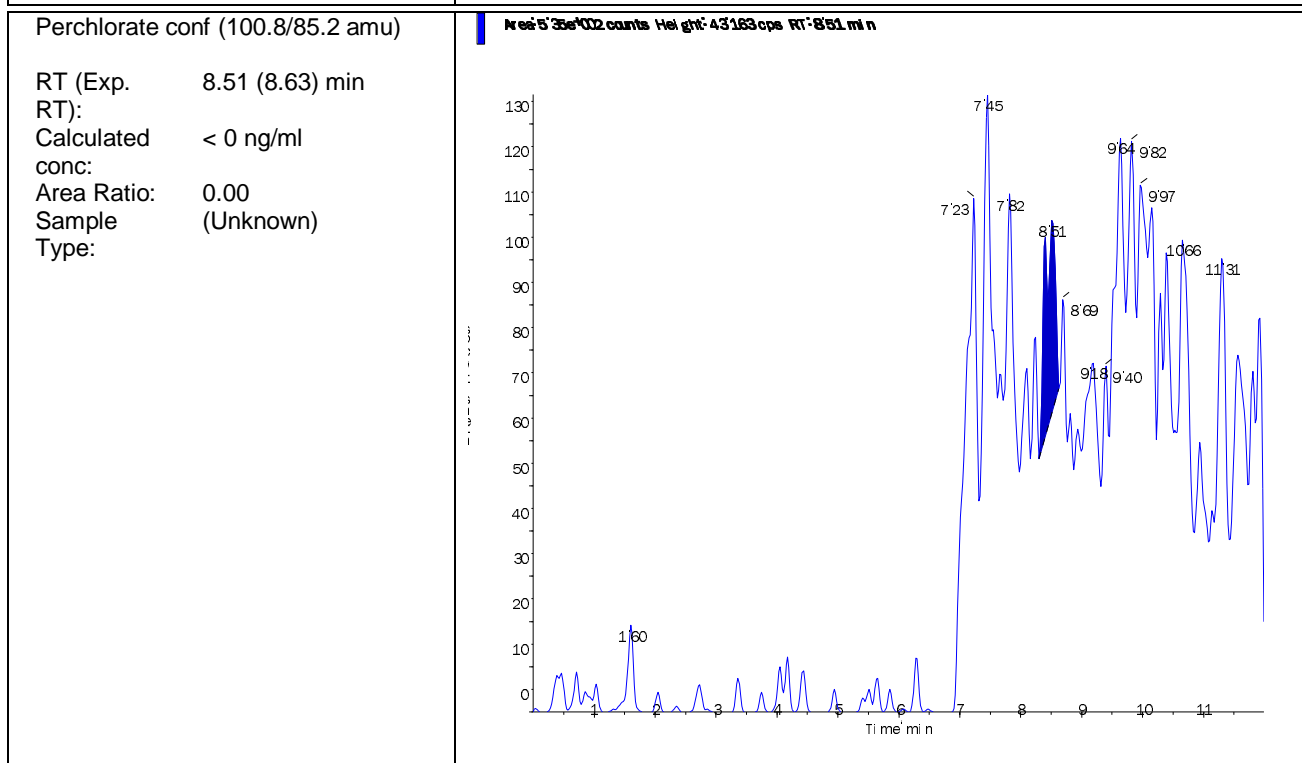
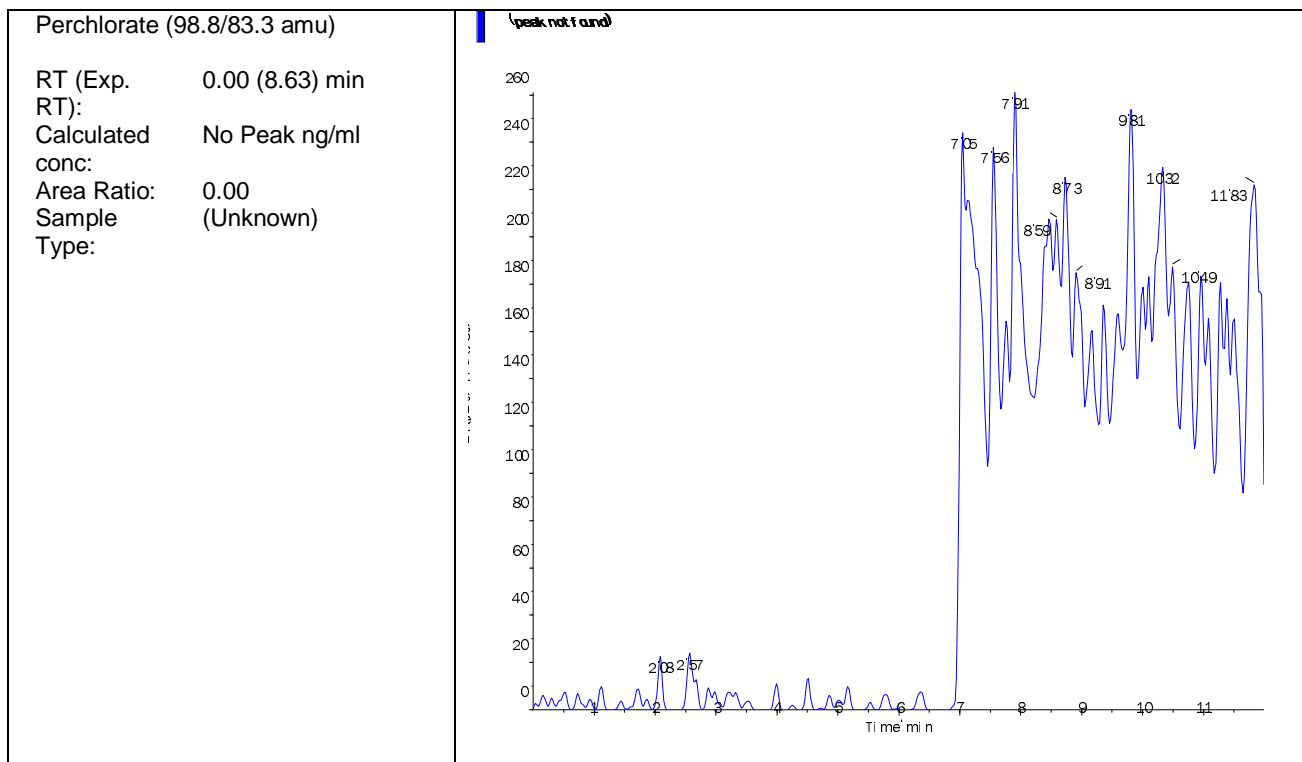
Data File	LM45244.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 11:36:47 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699915-06 CCB	Injection Vial	1.00
Data File	LM45244.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 11:36:47 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699915-06	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.610e+06	8.51	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	0.000e+00	0.00	N/A	No Peak
Perchlorate conf	5.350e+02	8.51	N/A	< 0





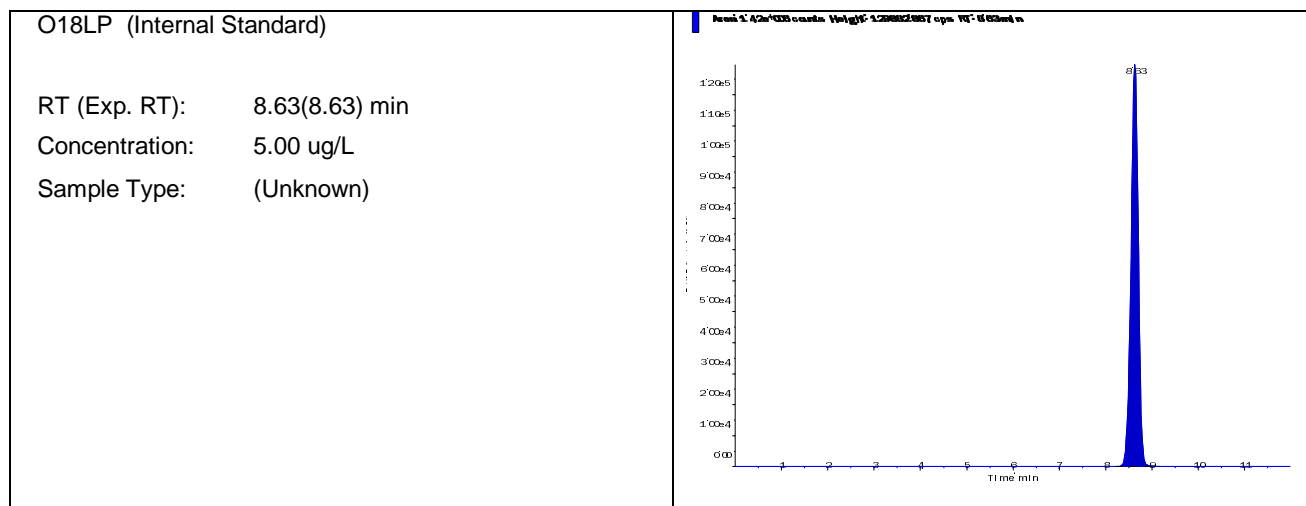
2.2.1.5 Raw QC Data

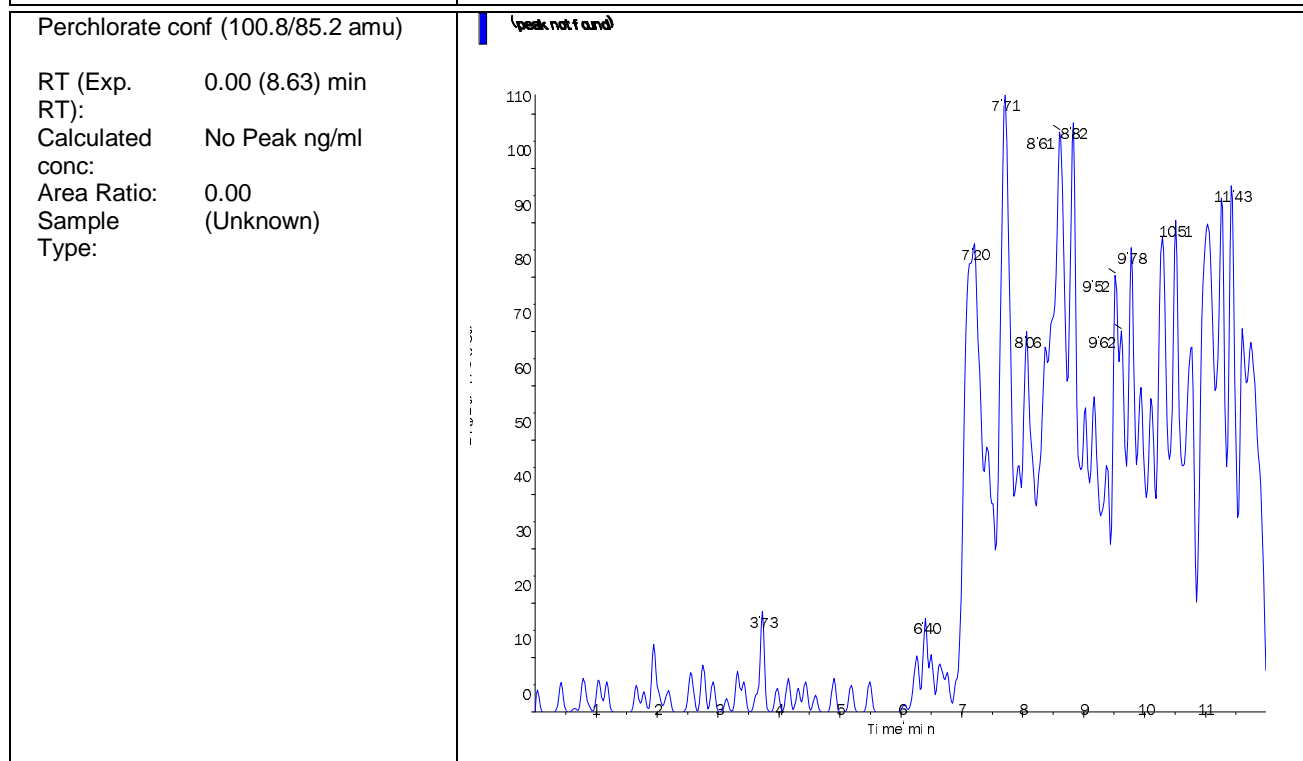
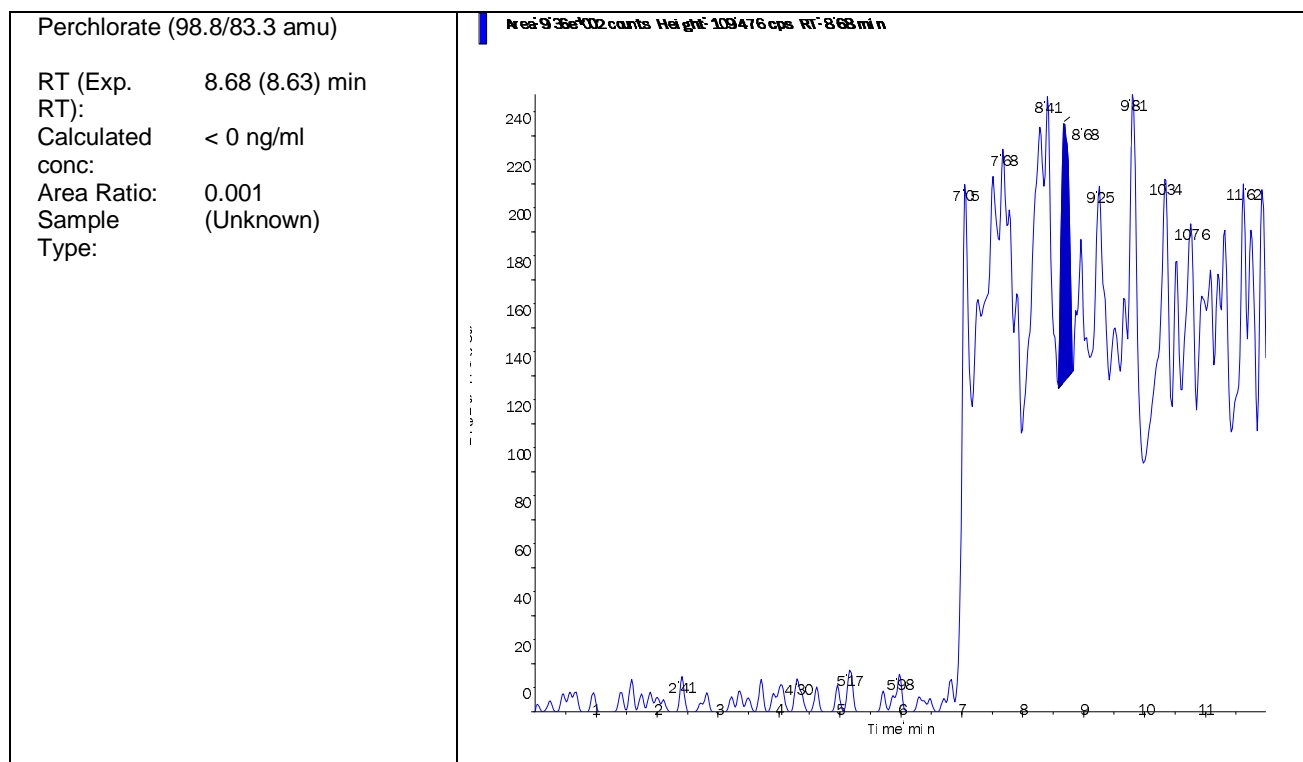
Data File	LM45227.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 7:56:57 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699914-02 BLANK	Injection Vial	11.00
Data File	LM45227.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 7:56:57 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-02	Dilution Factor	1.00
Sample Comment	11.00	Weight to Volume	0.00

Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.420e+06	8.63	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	9.360e+02	8.68	N/A	< 0
Perchlorate conf	0.000e+00	0.00	N/A	No Peak



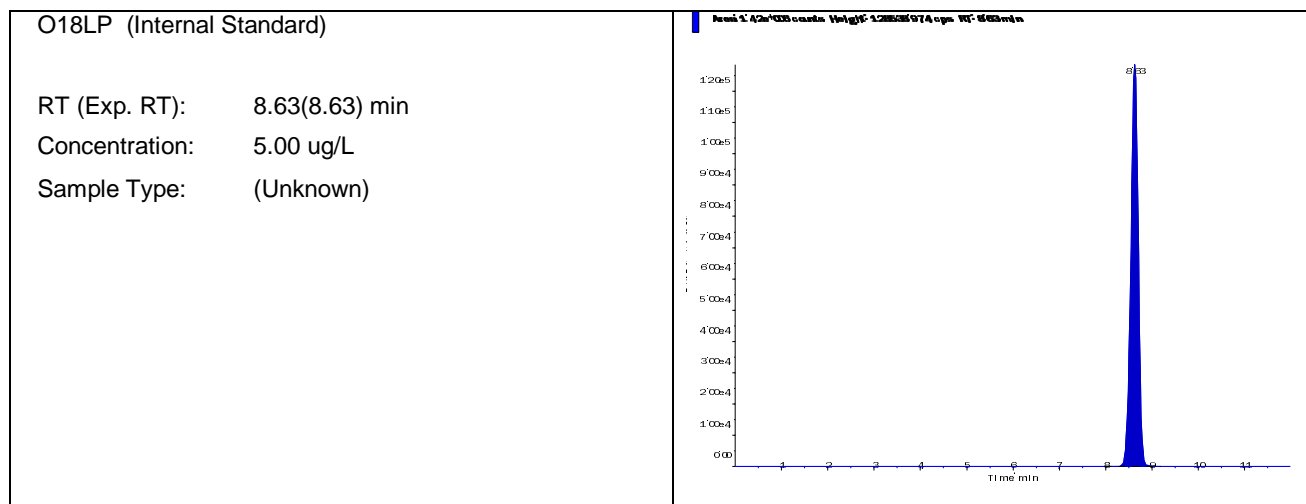


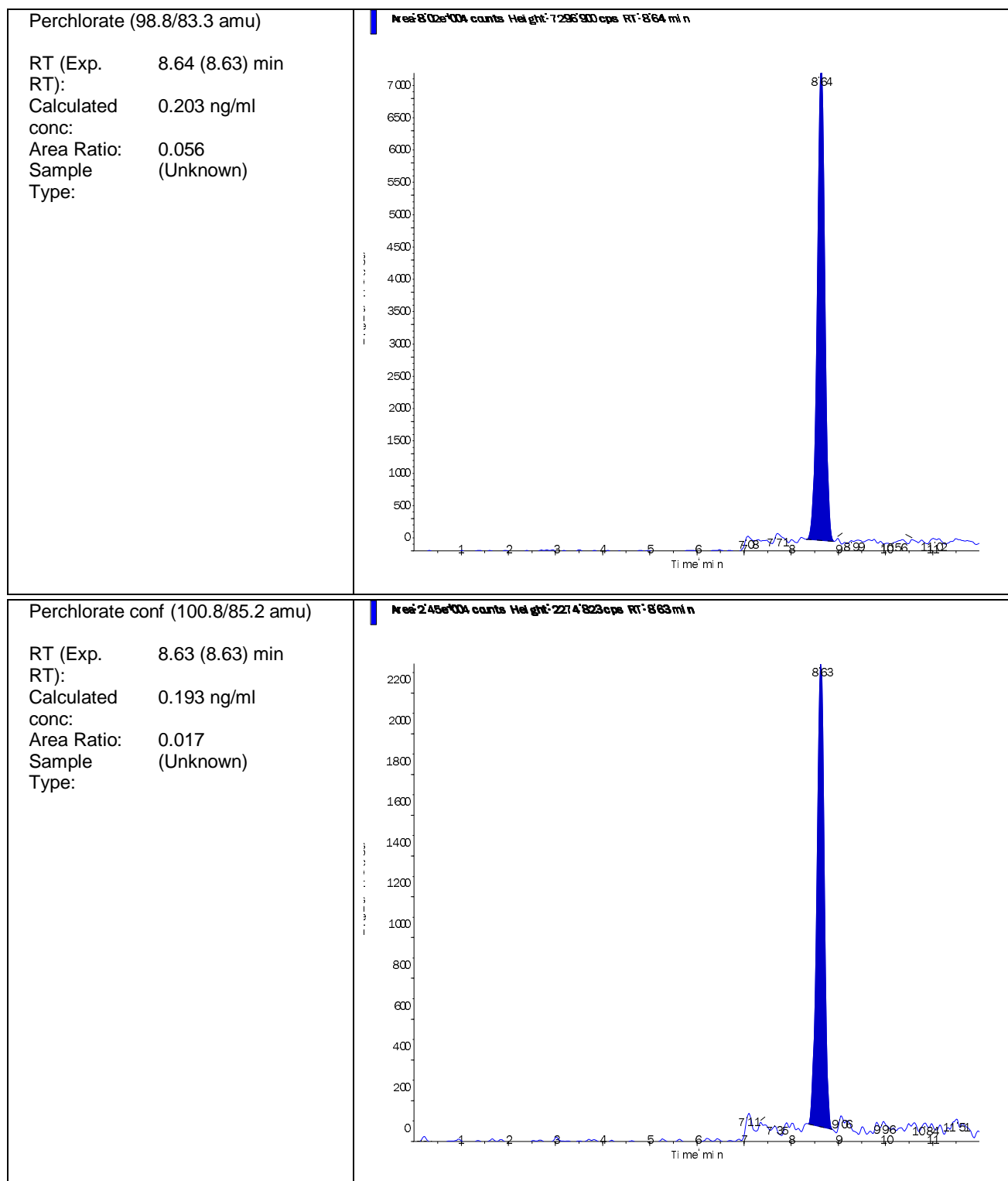
Data File	LM45228.wiff	Result Table	031819_JWR.rdb
Acquisition Date	3/18/2019 8:09:53 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Instrument Name	API 4000
Project	Perchlorate\2009_07_22		

Sample Name	WG699914-03 LCS (0.2ug/L)	Injection Vial	12.00
Data File	LM45228.wiff	Injection Volume	10.00
Acquisition Date	3/18/2019 8:09:53 PM	Algorithm Used	Analyst Classic
Acquisition Method	062911.dam	Sample Type	Unknown
Instrument Name	API 4000	Result Table	031819_JWR.rdb
Sample ID	WG699914-03	Dilution Factor	1.00
Sample Comment	1,1 STD87534	Weight to Volume	0.00

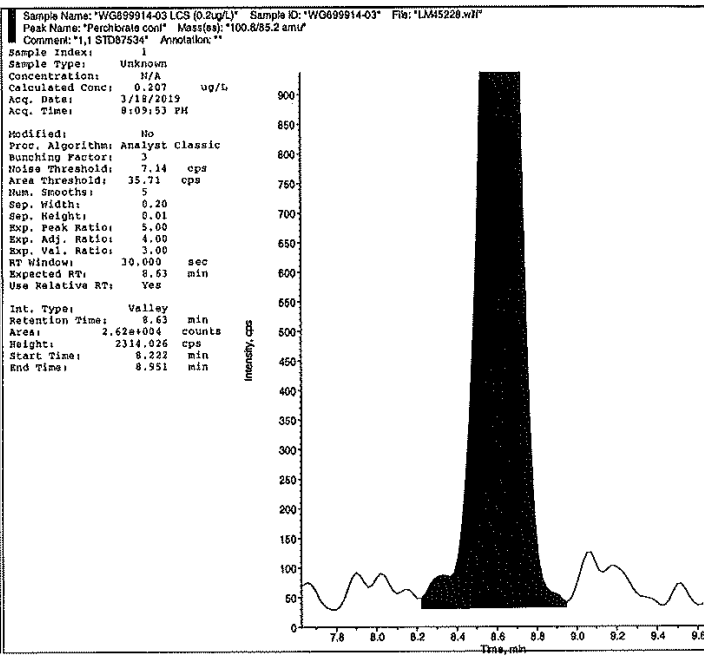
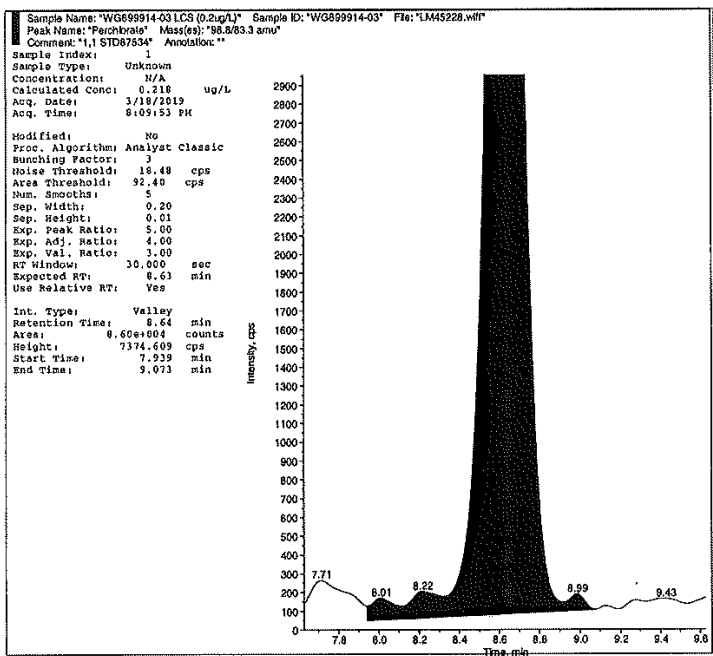
Internal Standard	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
O18LP	1.420e+06	8.63	5.00	-

Target Analyte	Area (cps)	RT (min)	Target conc. (ug/L)	Calc. Conc. (ug/L)
Perchlorate	8.020e+04	8.64	N/A	0.203
Perchlorate conf	2.450e+04	8.63	N/A	0.193



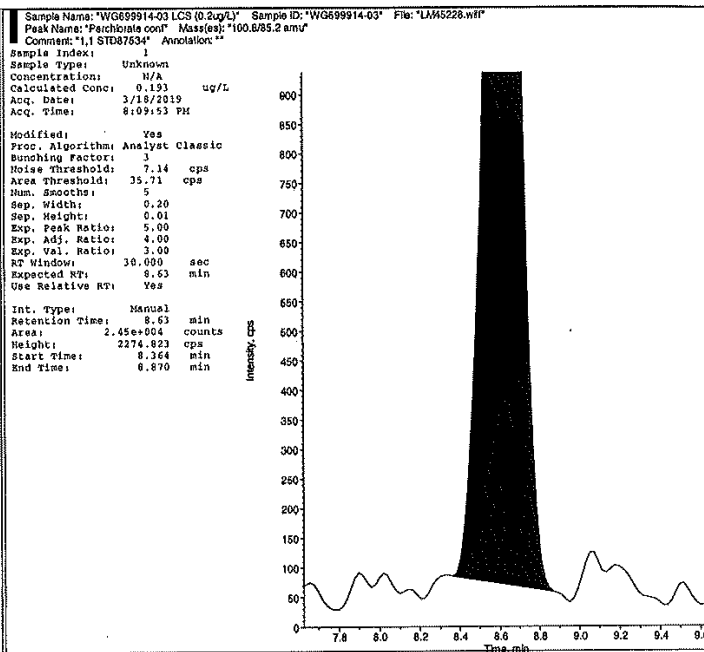
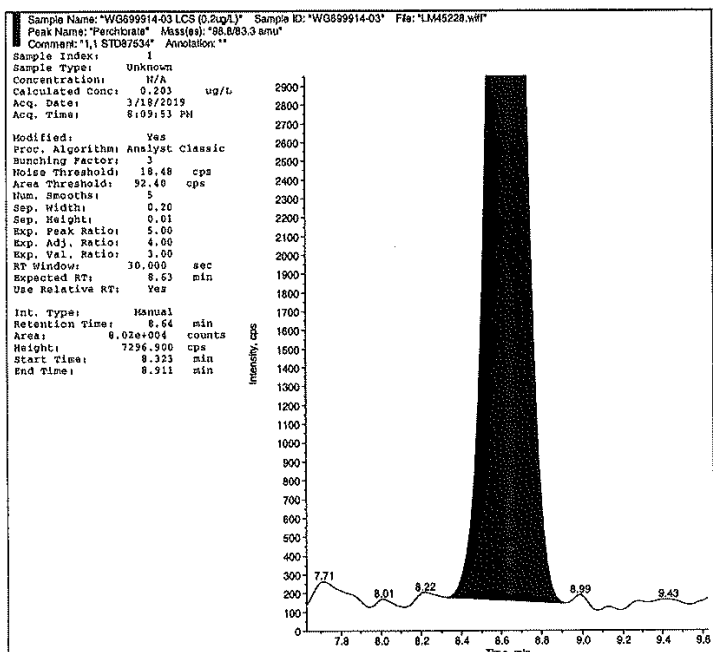


Printing Time: 10:41:19 AM
 Printing Date: Tuesday, March 19, 2019



Collected by: N/A
 Electronic Signature: no
 Operator: Administrator

Printing Time: 10:44:10 AM
 Printing Date: Tuesday, March 19, 2019



#4
 JWR/03/19/19
 JEC 3-20-19

Collected by: N/A
 Electronic Signature: no
 Operator: Administrator

3.0 Attachments

Microbac Laboratories Inc.
Ohio Valley Division Analyst List
March 25, 2019

001 - BIO-CHEM TESTING WVDEP 220	002 - REIC Consultants, Inc. WVDEP 060
003 - Sturm Environmental	004 - MICROBAC PITTSBURGH
005 - ES LABORATORIES	006 - ALCOSAN LABORATORIES
007 - ALS LABORATORIES	008 - BENCHMARK LABORATORIES
010 - MICROBAC CHICAGOLAND	AC - AMBER R. CARMICHAEL
ACG - ALEX C. GEDON	ADC - ANTHONY D. CANTER
ADG - APRIL D. GREENE	ADW - ALICIA D. WALKER
ALS - ADRIANE L. STEED	APH - ANDREW P. HOUT
AT - Asa R. Timmons	ATK - ALEX T. KLINTWORTH
AWE - ANDREW W. ESSIG	AZH - AFTER HOURS
BLG - BRENDA L. GREENWALT	BRG - BRENDA R. GREGORY
CAS - Craig A. Smith	CEB - CHAD E. BARNES
CLC - CHRYS L. CRAWFORD	COR - Corporate IT
CPD - CHAD P. DAVIS	CSH - CHRIS S. HILL
DIH - DEANNA I. HESSON	DLB - DAVID L. BUMGARNER
DLP - DOROTHY L. PAYNE	DSM - DAVID S. MOSSOR
ECL - ERIC C. LAWSON	EEA - EMILY E. ALLEN
EGS - EMILY G. SHILLING	EPT - ETHAN P. TIDD
ERP - ERIN R. PORTER	JAO - Jeff A. Ogle
JDH - JUSTIN D. HESSON	JDS - JARED D. SMITH
JDW - JAMES D. WRIGHT	JKP - JACQUELINE K. PARSONS
JLR - JIMMY L. RUSH	JRH - Justin R. Hill
JST - JOSHUA S. TAYLOR	JTP - JOSHUA T. PEMBERTON
JWR - JOHN W. RICHARDS	JYH - JI Y. HU
KAK - KATHY A. KIRBY	KEB - KATIE E. BARNES
KEH - Katelyn E. Hoover	KFR - KARISSA F. REYNOLDS
KHR - KIM H. RHODES	KKB - KERRI K. BUCK
KMC - KAYLA M. CHEVALIER	KMG - KALEN M. GANDOR
KRA - KATHY R. ALBERTSON	KRP - KATHY R. PARSONS
KWD - Kurtis W. Decker	LLS - LARRY L. STEPHENS
LSB - LESLIE S. BUCINA	LSJ - LAURA S. JONES
MAP - MARLA A. PORTER	MES - MARY E. SCHILLING
MMB - MAREN M. BEERY	MRT - MICHELLE R. TAYLOR
PDM - PIERCE D. MORRIS	PIT - MICROBAC WARRENDALE
RLB - BOB BUCHANAN	RNM - Rene N. Miller
RNP - RICK N. PETTY	SAV - SARAH A. VANDENBERG
SCB - SARAH C. BOGOLIN	SLM - STEPHANIE L. MOSSBURG
TB - TODD BOYLE	TMM - TAMMY M. MORRIS
VC - VICKI COLLIER	XXX - UNAVAILABLE OR SUBCONTRACT
ZTB - ZACH T. BARNES	

List of Valid Qualifiers

March 25, 2019

Qualkey: DOD

Qualifier	Description
*	Surrogate or spike compound out of range
+	Correlation coefficient for the MSA is less than 0.995
<	Result is less than the associated numerical value.
>	Greater than
>,H1	Result is greater than the associated numerical value. Sample analysis performed past holding time.
A	See the report narrative
B	The reported result is associated with a contaminated method blank.
B,H1	Analyte present in method blank. Sample analysis performed past holding time.
B1	Target analyte detected in method blank at or above the method reporting limit
B3	Target analyte detected in calibration blank at or above the method reporting limit
B4	The BOD unseeded dilution water blank exceeded 0.2 mg/L
B4,A	The BOD unseeded dilution water blank exceeded 0.2 mg/L. See the report narrative.
B4,F	The BOD unseeded dilution water blank exceeded 0.2 mg/L. Analyte present below the reporting limit (RL).
B4,F,A	The BOD unseeded dilution water blank exceeded 0.2 mg/L. Analyte present below the reporting limit (RL). See the report
B4,ND	The BOD unseeded dilution water blank exceeded 0.2 mg/L. Not detected at or above the reporting limit..
B4,ND,A	The BOD unseeded dilution water blank exceeded 0.2 mg/L. Not detected at or above the reporting limit. See the report na
C	Confirmed by GC/MS
CG	Confluent growth
CT1	Cooler temperature at sample receipt exceeded regulatory limit.
DL	Surrogate or spike compound was diluted out.
E	Estimated concentration due to sample matrix interference
E,CT1	Estimated results. The cooler temperature at receipt exceeded regulatory guidelines for requested testing.
EDL	Elevated sample reporting limits, presence of non-target analytes
EMPC	Estimated Maximum Possible Concentration
F, S	Estimated result below quantitation limit; method of standard additions(MSA)
F,CT1	Estimated value; the analyte concentration was less than the RL/LOQ. The cooler temperature at receipt exceeded regula
FL	Free Liquid
FP1	Did not ignite.
H1	Sample analysis performed past holding time.
H1,A	Sample analysis performed past holding time. See the report narrative
H1,CT1	Sample analysis performed past holding time. The cooler temperature at receipt exceeded regulatory guidelines for reque
I	Semiquantitative result (out of instrument calibration range)
J	Concentrations >40% difference between the two GC columns
J	Estimated concentration; sample matrix interference.
J	Estimated value ; the analyte concentration was greater than the highest standard
J	Estimated value ; the analyte concentration was less than the LOQ.
J	The reported result is an estimated value.
J,A	Present below nominal reporting limit. See case narrative.
J,B	Analyte detected in both the method blank and sample above the MDL.
J,CT1	Estimated value ; the analyte concentration was less than the LOQ. Cooler temperature at sample receipt exceeded regu
J,H1	Estimated value ; the analyte concentration was less than the LOQ. Sample analysis performed past holding time.
J,H1	The reported result is an estimated value. Sample was analyzed past holding time.
J,P	Estimate; columns don't agree to within 40%
J,S	Estimated concentration; analyzed by method of standard addition (MSA)
JB	The reported result is an estimated value. The reported result is also associated with a contaminated method blank.
JQ	Found between MDL and RL, but has quality control issue.
JQ	The reported result is an estimated value and one or more quality control criteria failed. See narrative.
L	Sample reporting limits elevated due to matrix interference
L1	The associated blank spike (LCS) recovery was above the laboratory acceptance limits.
L2	The associated blank spike (LCS) recovery was below the laboratory acceptance limits.
M	Matrix effect; the concentration is an estimate due to matrix effect.
N	Nontarget analyte; the analyte is a tentatively identified compound (TIC) by GC/MS
NA	Not applicable
ND	Not detected at or above the reporting limit (RL)
ND, B	Not detected at or above the reporting limit (RL). Analyte present in method blank.
ND, CT1	Analyte was not detected. The concentration is below the reported LOD. The cooler temperature at receipt exceeded reg
ND, L	Not detected; sample reporting limit (RL) elevated due to interference
ND, S	Not detected; analyzed by method of standard addition (MSA)
ND,A	Not detected. See report narrative.
ND,H1	Not detected; Sample analysis performed past holding time.
ND,H1,CT1	Not detected; Sample analysis performed past holding time. The cooler temperature at receipt exceeded regulatory guide
NF	Not found by library search
NFL	No free liquid
NI	Non-ignitable
NR	Analyte is not required to be analyzed
NS	Not spiked
P	Concentrations >40% difference between the two GC columns



List of Valid Qualifiers

March 25, 2019

Qualkey: DOD

Q	One or more quality control criteria failed. See narrative.
Q,H1	One or more quality control criteria failed. Sample analyzed past holding time. See narrative.
QNS	Quantity of sample not sufficient to perform analysis
RA	Reanalysis confirms reported results
RE	Reanalysis confirms sample matrix interference
S	Analyzed by method of standard addition (MSA)
SMI	Sample matrix interference on surrogate
SP	Reported results are for spike compounds only
T5	Laboratory not licensed for this parameter
TIC	Library Search Compound
TNTC	Too numerous to count
TNTC, B	Too numerous to count. Analyte present in method blank.
TNTC,CT1	Too numerous to count. The cooler temperature at receipt exceeded regulatory guidelines for requested testing.
TNTC,H1	Too numerous to count. Sample analysis performed past holding time.
U	Analyte was not detected. The concentration is below the reported LOD.
U,CT1	Analyte was not detected. The concentration is below the reported LOD. Cooler temperature at sample receipt exceeded
U,H1	Not detected; Sample analysis performed past holding time.
UJ	Undetected; the MDL and RL are estimated due to quality control discrepancies.
UQ	Analyte was not detected. The concentration is below the reported LOD. One or more quality control criteria fail. See nar
W	Post-digestion spike for furnace AA out of control limits
X	Exceeds regulatory limit
X, S	Exceeds regulatory limit; method of standard additions (MSA)
Z	Cannot be resolved from isomer - see below



Microbac Laboratories Inc.

Internal Chain of Custody Report

Login: L19030638

Account: 2886

Project: 2886.016

Samples: 8

Due Date: 22-MAR-2019

Samplenum **Container ID** **Products**
L19030638-01 307542 826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 3

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Samplenum **Container ID** **Products**
L19030638-01 307543 6850

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	W1	11-MAR-2019 14:01	BRG		
2	ANALYZ	W1	SEM	15-MAR-2019 14:03	JWR	BRG	
3	STORE	SEM	A2	20-MAR-2019 15:15	BRG	CAS	

Samplenum **Container ID** **Products**
L19030638-02 307544 826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 3

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

A1 - Sample Archive (COLD)
A2 - Sample Archive (AMBIENT)
F1 - Volatiles Freezer in Login
V1 - Volatiles Refrigerator in Login
W1 - Walkin Cooler in Login



Microbac Laboratories Inc.

Internal Chain of Custody Report

Login: L19030638

Account: 2886

Project: 2886.016

Samples: 8

Due Date: 22-MAR-2019

Samplenum **Container ID** **Products**
L19030638-02 307545 6850

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	W1	11-MAR-2019 14:01	BRG		
2	ANALYZ	W1	SEM	15-MAR-2019 14:03	JWR	BRG	
3	STORE	SEM	A2	20-MAR-2019 15:15	BRG	CAS	

Samplenum **Container ID** **Products**
L19030638-03 307546 826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 3

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Samplenum **Container ID** **Products**
L19030638-03 307547 6850

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	W1	11-MAR-2019 14:01	BRG		
2	ANALYZ	W1	SEM	15-MAR-2019 14:03	JWR	BRG	
3	STORE	SEM	A2	20-MAR-2019 15:15	BRG	CAS	

A1 - Sample Archive (COLD)
A2 - Sample Archive (AMBIENT)
F1 - Volatiles Freezer in Login
V1 - Volatiles Refrigerator in Login
W1 - Walkin Cooler in Login



Microbac Laboratories Inc.

Internal Chain of Custody Report

Login: L19030638

Account: 2886

Project: 2886.016

Samples: 8

Due Date: 22-MAR-2019

Samplenum **Container ID** **Products**
L19030638-04 307548 826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 3

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Samplenum **Container ID** **Products**
L19030638-04 307549 6850

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	W1	11-MAR-2019 14:01	BRG		
2	ANALYZ	W1	SEM	15-MAR-2019 14:03	JWR	BRG	
3	STORE	SEM	A2	20-MAR-2019 15:15	BRG	CAS	

Samplenum **Container ID** **Products**
L19030638-05 307550 826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:18	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 3

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

A1 - Sample Archive (COLD)
A2 - Sample Archive (AMBIENT)
F1 - Volatiles Freezer in Login
V1 - Volatiles Refrigerator in Login
W1 - Walkin Cooler in Login



Microbac Laboratories Inc.

Internal Chain of Custody Report

Login: L19030638

Account: 2886

Project: 2886.016

Samples: 8

Due Date: 22-MAR-2019

Samplenum **Container ID** **Products**
L19030638-05 307551 6850

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	W1	11-MAR-2019 14:01	BRG		
2	ANALYZ	W1	SEM	15-MAR-2019 14:03	JWR	BRG	
3	STORE	SEM	A2	20-MAR-2019 15:14	BRG	CAS	

Samplenum **Container ID** **Products**
L19030638-06 307552 826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Bottle: 3

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:19	EEA	BRG	

Samplenum **Container ID** **Products**
L19030638-06 307553 6850

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	W1	11-MAR-2019 14:01	BRG		
2	ANALYZ	W1	SEM	15-MAR-2019 14:03	JWR	BRG	
3	STORE	SEM	A2	20-MAR-2019 15:15	BRG	CAS	

A1 - Sample Archive (COLD)
A2 - Sample Archive (AMBIENT)
F1 - Volatiles Freezer in Login
V1 - Volatiles Refrigerator in Login
W1 - Walkin Cooler in Login



Microbac Laboratories Inc.

Internal Chain of Custody Report

Login: L19030638

Account: 2886

Project: 2886.016

Samples: 8

Due Date: 22-MAR-2019

<u>Samplenum</u>	<u>Container ID</u>	<u>Products</u>
L19030638-07	307554	826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:18	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:18	EEA	BRG	

Bottle: 3

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:18	EEA	BRG	

<u>Samplenum</u>	<u>Container ID</u>	<u>Products</u>
L19030638-07	307555	6850

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	W1	11-MAR-2019 14:01	BRG		
2	ANALYZ	W1	SEM	15-MAR-2019 14:03	JWR	BRG	
3	STORE	SEM	A2	20-MAR-2019 15:15	BRG	CAS	

<u>Samplenum</u>	<u>Container ID</u>	<u>Products</u>
L19030638-08	307556	826-SPE

Bottle: 1

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:18	EEA	BRG	

Bottle: 2

Seq.	Purpose	From	To	Date/Time	Accept	Relinquish	pH
1	LOGIN	COOLER	V1	11-MAR-2019 14:01	BRG		
2	ANALYZ	V1	ORG1	11-MAR-2019 16:18	EEA	BRG	

A1 - Sample Archive (COLD)
A2 - Sample Archive (AMBIENT)
F1 - Volatiles Freezer in Login
V1 - Volatiles Refrigerator in Login
W1 - Walkin Cooler in Login



NELAP Addendum - January 3, 2019

Non-NELAP LIMS Product and Description

The following is a list of those tests that are not included in the Microbac – OVD NELAP Scope of Accreditation:

Heat of Combustion (BTU)
 Total Halide by Bomb Combustion (TX)
 Particle Sizing - 200 Mesh (PS200)
 Specific Gravity/Density (SPGRAV)
 Total Residual Chlorine (CL-TRL)
 Total Volatile Solids (all forms) (TVS)
 Total Coliform Bacteria (all methods)
 Fecal Coliform Bacteria (all methods)
 Sulfite (SO₃)
 Propionaldehyde (HPLC-UV)

SOLID AND HAZARDOUS CHEMICALS

Nitrogen, Ammonia by Method 350.1
 Chromium, Hexavalent, Leachable by SM3500 Cr-B 2009
 Phenolics, Total by Method 420.1
 ASTM D3987-06

NELAP Accreditation by Laboratory SOP

NONPOTABLE WATER

OVD HPLC02/HPLC-UV

Nitroglycerin
 Acetic acid
 Butyric acid
 Lactic acid
 Propionic acid
 Pyruvic acid

OVD MSS01/GC-MS

1,4-Phenylenediamine
 1-Methylnaphthalene
 1,4-Dioxane
 Atrazine
 Benzaldehyde
 Biphenyl
 Caprolactam
 Hexamethylphosphoramide (HMPA)
 Pentachlorobenzene
 Pentachloroethane

NELAP Accreditation by Laboratory SOP

NONPOTABLE WATEROVD MSV01/GC-MS

1, 1, 2-Trichloro-1,2,2-trifluoroethane
1,3-Butadiene
Cyclohexane
Cyclohexanone
Dimethyl disulfide
Dimethylsulfide
Ethyl-t-butylether (ETBE)
Isoprene
Methylacetate
Methylcyclohexane
T-amylmethylether (TAME)
Tetrahydrofuran (THF)

OVD HPLC07/HPLC-MS-MS

Hexamethylphosphoramide (XMPA-LCMS)

OVD HPLC12/HPLC/UV

Acetate
Formate

OVD RSK01/GC-FID

Acetylene
Propane

OVD K9305/ISE

Fluoroborate

NELAP Accreditation by Laboratory SOP**SOLID AND HAZARDOUS CHEMICALS**OVD MSS01/GC-MS

1-Methylnaphthalene
Benzaldehyde
Biphenyl
Caprolactam
Pentachloroethane

NELAP Accreditation by Laboratory SOP

SOLID AND HAZARDOUS CHEMICALSOVD MSV01/GC-MS

1.3-Butadiene
Cyclohexane
Cyclohexanone
Dimethyl disulfide
Dimethylsulfide
Ethyl-t-butylether (ETBE)
Isoprene
Methylacetate
Methylcyclohexane
n-Hexane
T-amylmethylether (TAME)



B

Data Validation Report



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DATA VALIDATION REPORT

Volatile Organic Compounds (VOCs) by USEPA SW-846 Method 8260B
Perchlorate by USEPA SW-846 Method 6850

Project: USACE Longhorn Army Ammunition Plant (LHAAP), Texas
Surface Waters – March 2019

Project/Task Number: 10097916 8.0

Sample Data Package: L19030638

Laboratory: Microbac Labs, Marietta, Ohio

Sample Matrix: Surface water

Sampling Dates: 07 March 2019

Validation Guidelines: Project QAPP (*Final Quality Assurance Project Plan for Longhorn Army Ammunition Plant, Karnack, Texas [July 2014]*); United States Environmental Protection Agency (USEPA) *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods, 3rd edition (SW-846)*; *National Functional Guidelines for Inorganic Superfund Data Review (USEPA, September 2016)*; *National Functional Guidelines for Organic Superfund Data Review (USEPA, September 2016)* and professional judgment

Validation Level: Stage 2BVM (100% of data), Stage 3VM (10% of data)

Data Reviewer: Jennifer Chandler, HDR

Sample ID	Lab ID	Matrix	VOCs	Perchlorate
SW1	L19030638-01	SW	X	X
SW1 MS	L19030638-02	SW	X	X
SW1 MSD	L19030638-03	SW	X	X
SW2	L19030638-04	SW	X	X
SW3	L19030638-05	SW	X	X
SW4	L19030638-06	SW	X	X
DUP01	L19030638-07	SW	X	X
TB01	L19030638-08	QC Water	X	

SUMMARY

One hundred percent of the data underwent Stage 2BVM level validation. Ten percent of the data underwent Stage 3VM level validation.

Laboratory data were acceptable with qualification as noted below. No data were rejected. Completeness was 100%, which met the goal of 90% for the project.

I. SAMPLE RECEIPT / CHAIN OF CUSTODY

The chains of custody (COC) were filled out and signed. Samples were received within the correct temperature range (at 3.0°C). No qualification was required.

II. SAMPLES ANALYZED

All planned samples were collected and all scheduled analyses were performed. The correct analyte lists were reported. No qualification was required.

VOCs: Samples were analyzed by USEPA SW-846 Method 8260B. Results are reported in µg/L.

Perchlorate: Samples were analyzed by USEPA Method SW-846 6850. Results are reported in µg/L.

III. HOLDING TIMES AND PRESERVATION

All holding time criteria were met and no qualification was required except as noted below.

VOCs: Samples were analyzed within 14 days of collection.

Perchlorate: Samples were analyzed within 28 days of collection. No qualification was required.

IV. BLANKS

METHOD BLANKS

Target analytes were not detected in method blanks except as noted below.

VOCs: VOCs were not detected in the MBs. No qualification was required.

Perchlorate: Perchlorate was not detected in the MBs. No qualification was required.

TRIP BLANKS

VOCs: One trip blank was collected (TB-01). Acetone was detected at 2.67 µg/L, below the limit of quantitation (LOQ) of 10 µg/L. The following samples were detected at less than five times the blank and has been qualified as estimated with a possible high bias (J+): SW-1, SW-2, SW-3, and DUP01.

Perchlorate: Not applicable.

EQUIPMENT RINSE BLANK

Equipment rinse blanks were not required.

INITIAL AND CONTINUING CALIBRATION BLANKS (ICBs and CCBs)

VOCs: Not applicable.

Perchlorate: Perchlorate was not detected in the CCBs. No qualification was required.

V. LABORATORY CONTROL SAMPLES / LABORATORY CONTROL SAMPLE DUPLICATE (LCS/LCSD)

Recoveries and relative percent differences (RPDs) were generally within control limits except as noted below.

VOCs: Recoveries were within control limits. No qualification was required.

Perchlorate: Recoveries were within control limits. No qualification was required.

VI. MATRIX SPIKE / MATRIX SPIKE DUPLICATE (MS/MSD)

VOCs: Sample SW-1 was designated on the COC for MS/MSD analysis. All recoveries and RPDs were within control limits. No qualification was required.

Perchlorate: Sample SW-1 was designated on the COC for MS/MSD analysis. The MSD recovery 80.5% was below the control limits of 84-119%. Sample results were qualified as estimated (J-).

VI. DILUTION TEST AND POST DIGESTION SPIKE (PDS)

A dilution test was performed for analytes detected at least 50 times the concentration of the MRL, and results agreed within ten percent. A post digestion spike (PDS) was performed on

samples and the recoveries were between 80 and 120 percent except as noted below. These tests are applicable only to metals by SW6010C and SW6020.

VOCs: Not applicable.

Perchlorate: Not applicable.

VIII. FIELD DUPLICATES

Field duplicate (FD) samples were collected as shown in the table below. Control limits listed in the QAPP were used.

Field Duplicate Sample	Parent Sample
DUP-01	SW1 (VOCs and perchlorate)

VOCs: Results were within the control limits. No qualification was required.

Perchlorate: Results were within the control limits. No qualification was required.

IX. LABORATORY DUPLICATES

Laboratory duplicate (LD) analyses were not performed on any samples from this project.

X. INSTRUMENT CALIBRATION

INITIAL CALIBRATION

VOCs: The calibration curve included a standard at or below the LOQ. Calibration requirements were met. No qualification was required.

Perchlorate: The calibration curve included a standard at or below the LOQ. Calibration requirements were met. No qualification was required.

INITIAL CALIBRATION VERIFICATION (ICV)

VOCs: ICV results were within the required limits. No qualification was required.

Perchlorate: ICV results were within the required limits. No qualification was required.

CONTINUING CALIBRATION VERIFICATION (CCV)

VOCs: CCV results were within the required limits. No qualification was required.

Perchlorate: CCV results were within the required limits. No qualification was required.

XI. SURROGATES, INTERNAL STANDARDS AND ION RATIOS

VOCs: Internal standard areas and retention times were within required limits. Surrogate recoveries were within control limits. No qualification was required.

Perchlorate: Internal standard areas and ion ratios were within required limits. No qualification was required.

XII. INTERFERENCE CHECK SAMPLES

Interference check sample solutions ICSA and ICSAB were analyzed (methods 6010C and 6020 only).

VOCs: Not applicable.

Perchlorate: Not applicable.

XIII. SAMPLE LIMITS OF QUANTITATION (LOQs)

Limits of quantitation (LOQs) should be less than the applicable cleanup levels as listed in the project QAPP.

VOCs: LOQs were below the groundwater MCLs/cleanup levels.

Perchlorate: LOQs were below the groundwater MCLs/cleanup levels.

XIV. SAMPLE RESULTS / TRANSCRIPTION VERIFICATION

Transcription between the data package and the EDDs was verified. Results reported below the LOQs but above the DLs were qualified as estimated (J). Non-detect results were reported down to the limits of detection (LODs).

VOCs: No issues.

Perchlorate: No issues.



DEPARTMENT OF THE ARMY
 LONGHORN ARMY AMMUNITION PLANT
 POST OFFICE BOX 220
 RATCLIFF, AR 72951

August 1, 2019

DAIM-ODB-LO

Ms. April Palmie
 Texas Commission on Environmental Quality,
 Superfund Section, MC-136
 12100 Park 35 Circle, Bldg D
 Austin, TX 78753

Re: August 2019 Underground Injection Control Substantive Requirements Notification for
 Remedy at LHAAP-04, Longhorn Army Ammunition Plant, Karnack, Texas

Dear Ms. Palmie,

The above-referenced document is being transmitted to you for your records. We are presumptively complying with the substantive requirements of 30 Texas Administrative Code (TAC) §331, Subchapters A, C, and H for Class V Injection Wells. Therefore, unless we hear in the negative within 30 calendar days (August 31, 2019), the injections will be implemented at Site LHAAP-04.

The document was prepared by Bhate Environmental Associates, Inc., on behalf of the Army as part of Bhate's Performance Based Remediation contract for the facility. I ask that Kim Nemmers, Bhate's Project Manager, be copied on any communications related to the project.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in cursive script that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
 Longhorn AAP Site Manager

Copies furnished:

R. Mayer, USEPA Region 6, Dallas, TX (1 hard copy and 1 CD)
 P. Bruckwicky, Caddo Lake NWR, TX (1 hard copy and 1 CD)
 A. Williams, USACE, Tulsa District, OK (1 CD)
 R. Smith, USACE, Tulsa District, OK (electronic only)
 A. Sherman, USAEC, San Antonio, TX (1 CD)
 K. Nemmers, Bhate, Lakewood, CO (1 CD)
 P. Srivastav, APTIM, Houston, TX (letter)



MEMORANDUM FOR RECORD

DATE: August 1, 2019

PROJECT NAME: Remedy for LHAAP-04, Longhorn Army Ammunition Plant, Karnack, Texas

TO: Rose Zeiler Site Manager
 Rick Smith Project Manager
 Aaron Williams Project Engineer
 Kimberly Nemmers Bhate Project Manager

FROM: Praveen Srivastav Bhate Team Deputy Project Manager, Aptim Federal Services, LLC (APTIM), Cell No. 281-639-8743

SUBJECT: **LHAAP-04 Underground Injection Control Substantive Requirements Notification, Longhorn Army Ammunition Plant, Karnack, TX (Contract No. W9128F-13-D-0012, Task Order No. W912BV17F0150)**

INTRODUCTION

Remedial activities are required under the Record of Decision (ROD) issued for the LHAAP-04 Former Pilot Wastewater Treatment Plant (AECOM 2016). The Draft Final (DF) Remedial Design/Remedial Action Work Plan (RD/RAWP) was submitted to Texas Commission on Environmental Quality (TCEQ) for review in April 2019 (U.S. Army 2019). No further comments were received from the regulators, and the RD/RAWP DF became the final document. The RAWP was developed using the basis and details of the RD for LHAAP-04.

As part of the selected remedy, in situ bioremediation (ISB) will be conducted in the area of perchlorate contaminated shallow groundwater with concentrations exceeding five times the TCEQ's Protective Concentration Level (PCL) of 17 µg/L and is planned to be conducted in September 2019. The ISB remedy will be implemented by injection of food-grade emulsified vegetable oil (EVO) to act as an electron donor to enhance populations of indigenous microorganisms that are capable of degrading perchlorate to harmless byproducts (chloride and oxygen).

The enclosed Class V Injection Well Inventory Form (**Enclosure 1**) and TCEQ Core Data Form (**Enclosure 2**) comply with the substantive requirements for construction, operation, and closure under 30 Texas Administrative Code (TAC) §331, Subchapters, A, C, and H (the Applicable or Relevant and Appropriate Requirements [ARARs] for underground injection control).

SITE HISTORY

LHAAP-04, known as Site 04 or the former pilot wastewater treatment plant, is approximately 0.5 acres and is located in the central portion of LHAAP (Class V Injection Well Inventory Form, **Enclosure 1**, Attachment A, Figure 1-2, LHAAP-04 Site Location). The demolition of the former pilot wastewater treatment facility structures, tanks, and piping, and the disposal of the associated wastes were completed in the summer of 1997 as part of the Resource Conservation and Recovery Act (RCRA) closure of the plant. Under the Comprehensive Environmental Response, Compensation, and Liability Act of

1980 (CERCLA) program, excavation of soil impacted with mercury and perchlorate at the LHAAP-04 site was completed in 2009 along the southern edge of the slab, which formerly housed storage tanks for the former pilot wastewater treatment facility.

The U.S. Department of the Army (U.S. Army) issued the Final ROD for LHAAP-04 (AECOM 2016) in October 2016 that was signed by the Army on December 15, 2016, and the U.S. Environmental Protection Agency (USEPA) on March 30, 2017. The Texas Commission on Environmental Quality (TCEQ) issued a letter concurring with the ROD on February 7, 2017. The ROD identified perchlorate as the only constituent of concern (COC) in groundwater for LHAAP-04. The remedy selected in the ROD included in-situ bioremediation (ISB) for perchlorate concentrations in groundwater, long-term monitoring (LTM) of groundwater, and LUCs to maintain the remedy and prohibit groundwater use until COC concentrations are reduced to levels supportive of unlimited use and unrestricted exposure.

PLANNED ACTION

The remedy will include ISB at LHAAP-04 to remediate the area of perchlorate contaminated groundwater with concentrations exceeding five times the PCL. Since the ROD, two shallow groundwater monitoring wells and one intermediate monitoring groundwater well were installed to better define the area of groundwater contamination prior to ISB. The Remedial Design was based on the most recent results and plume as shown on in the Class V Injection Well Inventory Form, **Enclosure 1**, Attachment E. Multiple injections of substrate may be needed based on effectiveness of the ISB.

The implementation of ISB will use a biogrid application for the injection of an electron donor to biodegrade the perchlorate. The Material Safety Data Sheet for the injectate (an emulsified vegetable oil mixture) is included in the Class V Injection Well Inventory Form, **Enclosure 1**, Attachment H.

Implementation of the remedy is planned for September 2019. Per Office of Solid Waste and Emergency Response (OSWER) Directive 9355.7, the Army is presumptively complying with substantive requirement of construction, operation and closure 30 TAC §331, Subchapters, A, C, and H (ARARs for underground injection control).

Enclosures:

1. Class V Injection Well Inventory Form
2. TCEQ Core Data Form

ENCLOSURE 1

CLASS V INJECTION WELL INVENTORY FORM

Texas Commission on Environmental Quality
Class V Injection Well
Inventory/Authorization Form

Submit To:**TCEQ**

UIC Permits Team

Radioactive Materials Division

MC233

PO Box 13087

Austin, Texas 78711-3087

512/239-6466

For TCEQ Use Only

Reg. No. _____

Date Received _____

Date Authorized _____

Section I General Information**Provide the information in items 1 through 8**

1. TCEQ Program Area (PST, VCP, IHW, etc.):
2. Agent/Consultant:
3. Facility Name: Longhorn Army Ammunition Plant, LHAAP-04
 Address (Street, City, County, State, and Zip Code) or location description (if no address is available): Northeast corner of Harrison County, located between State Highway 43 in Karnack, Texas, and the western shore of Caddo Lake
4. Latitude and Longitude (degrees-minutes-seconds) and method of determination (GPS, TOPO, etc.) (Attach a Topographic Quadrangle map which identifies the facility location relative to major streets or roadways as Attachment A.)
 Latitude: 32° 40' 58.67" N; Longitude: 94° 9' 5.80" W. LHAAP-04 Site Location Map (Figure 1-2 from the RD/RAWP) is also included in **Attachment A**.
5. Type of Well Construction (Vertical Injection, Subsurface Fluid Distribution System, Infiltration Gallery, Temporary Injection Points, etc.) and Number of Injection Wells:
 25 temporary direct-push injection points will be used to distribute the injectate mixture into the shallow zone.
6. Description regarding purpose of Injection System. Attach a Site Map as Attachment B (Attach the Approved Remediation Plan [if appropriate]):
 LHAAP-04 is a former pilot wastewater treatment plant located in the central portion of LHAAP near the former fire station. Wastewater treatment operations began at LHAAP-04 in 1984. The demolition of the former pilot wastewater treatment facility structures, tanks, and piping, and the disposal of the associated wastes were completed in the summer of 1997. Perchlorate is the only constituent of concern (COC) in groundwater

for LHAAP-04. The remedy selected in the ROD included in-situ bioremediation (ISB) for perchlorate concentrations in groundwater, long-term monitoring (LTM) of groundwater, and LUCs to maintain the remedy and prohibit groundwater use until COC concentrations are reduced to levels supportive of unlimited use and unrestricted exposure.

To implement ISB, emulsified vegetable oil (EVO), diammonium phosphate (microbial nutrient), and water will be injected into the subsurface at 25 locations shown on **Attachment B** (Figure 5-1 from the RD/RAWP) The injectate mixture will consist of approximately 1% (by volume) solution of EVO.

The Remedial Design/Remedial Action Work Plan (RD/RAWP) has been approved by TCEQ and USEPA and the approved injection plan from the RD/RAWP is included as **Attachment B**.

7. Water Well Driller/Installer:

Section II Proposed Down Hole Design

Attach a diagram signed and sealed by a licensed engineer as Attachment C

Name of String	Size	Setting Depth	Sacks Cement/Grout - Slurry Volume - Top of Cement	Hole Size	Weight (lbs/ft) PVC/Steel
Casing	N/A (See Note 1 below)	N/A	N/A	N/A	N/A
Tubing	N/A	N/A	N/A	N/A	N/A
Screen	N/A	N/A	N/A	N/A	N/A

Note 1: The injection of the amendments will be performed via direct-push injection points. The description of the direct-push injection points is provided in Section III below. There will be no permanent casing, tubing, or screens associated with the direct-push injection points.

Section III Proposed Trench System, Subsurface Fluid Distribution System, or Infiltration Gallery

Attach a diagram signed and sealed by a licensed engineer as Attachment D

12. System(s) Dimensions: Approximately 11,025 square feet based on the 85 µg/L perchlorate contour in shallow groundwater.
13. System(s) Construction: The fluid distribution system will consist of a temporary/mobile tanks/containers, hoses, a bulk storage tank, mixing equipment, injection pump, and volume and metering control equipment. Injection points will be advanced using via a tractor mounted rig with a direct-push technology (DPT) hammer. The injectate mixture will be injected using at each proposed direct-push point at 4-foot intervals in the target treatment area. Under this approach, drilling rods (injection probes) are advanced to the proposed injection interval. Injectate mixture will be pumped down through the DPT drilling rods (acting as a temporary well casing) to the injection interval and injectate will be forced through the ports in the rods to the surrounding formation. The tools are then advanced to the next injection depth and the material is again pumped through the rods. This cycle is repeated to provide coverage across the entire vertical treatment interval. Injection flow rates are expected to range from 2 to 6 gallons per minute and an attempt will be made to keep injection pressures below 25 pounds per square inch (psi).

The injection target intervals are shown in **Attachment C**, and a sample temporary DPT injection point diagram is included as **Attachment D**.

Section IV Site Hydrogeological and Injection Zone Data

Provide the information in items 14 through 31

14. Name of Contaminated Aquifer: Shallow Aquifer
15. Receiving Formation Name of Injection Zone: Unconsolidated Material
16. Well/Trench Total Depth: Injection point depths are vary from 6 to 20 feet (ft) below ground surface (bgs).
17. Surface Elevation: Approximately 205-215 ft
18. Depth to Ground Water: Depth to groundwater measured in monitoring wells ranges from approximately 2 to 10 ft bgs, with the saturated zone generally encountered between 6 and 15 feet bgs
19. Injection Zone Depth: 6 to 20 feet bgs
20. Injection Zone vertically isolated geologically? Y/N Impervious Strata between Injection Zone and nearest Underground Source of Drinking Water:
Name: Intermediate, Upper Deep and Deep Aquifer Zones (clay layers)
Thickness:
Intermediate Zone: 35 to 62 ft bgs
Upper Deep Groundwater: 80 to 151 ft bgs
Deep Groundwater: Extends below 220 ft bgs
21. Provide a list of contaminants and the levels (parts per million [ppm]) in contaminated aquifer.
Attach as Attachment E: Perchlorate is the only COC in groundwater. See **Attachment E** showing recent levels of contamination in parts per billion.
22. Horizontal and Vertical extent of contamination and injection plume
Attach as Attachment F: The horizontal extent of contamination shown in **Attachment E**. The contamination is limited to the shallow zone. There is no impact to the intermediate zone of the aquifer. A separate Attachment F is not provided.
23. Formation (Injection Zone) Water Chemistry (Background levels) TDS, etc.
Attach as Attachment G: Water chemistry data (dissolved oxygen, oxidation reduction potential, pH, turbidity, etc.) will be collected in the baseline sampling event and will be reported in the Remedial Action Completion Report. A separate Attachment G is not provided.
24. Injection Fluid Chemistry in ppm at point of injection
Attach as Attachment H: The injection fluid chemistry consists of approximately 1% (by volume) solution of EVO (carbon substrate). The EVO product, EDS-ER™, will be supplied by Tersus Environmental. See **Attachment H** for Safety Data Sheet.
Water for the solution will be procured from the Leigh Water Company in Karnack, Texas from their potable water supply.
25. Lowest Known Depth of Ground Water with < 10,000 ppm TDS: Unknown

26. Maximum injection Rate/Volume/Pressure: Rate: 2 to 10 gallons per minute; Volume: 1,478 gallons per point; Pressure: 10 to 40 psi (An attempt will be made to keep injection pressures below 25 psi). However, in some cases, depending on the lithology the injection pressure may be higher.
27. Water wells within 1/4 mile radius (attach map as Attachment I): None
28. Injection wells within 1/4 mile radius (attach map as Attachment I): None
29. Monitor wells within 1/4 mile radius (attach driller's logs and map as Attachment I): See **Attachment I**, quarter mile map which indicates approximately 21 monitoring wells within a quarter mile radius from the planned injections.
30. Sampling frequency: Baseline sampling will be conducted prior to the injections followed by quarterly sampling events for eight quarters. The sample results will be included in monitoring reports that will be submitted to TCEQ. Samples will be monitored for perchlorate, field parameters, and biogeochemical parameters to monitor the progress of ISB injections.
31. Known hazardous components in injection fluid: There are no hazardous components in the injection fluid.

Section V Site History

Provide the information in items 32 through 35

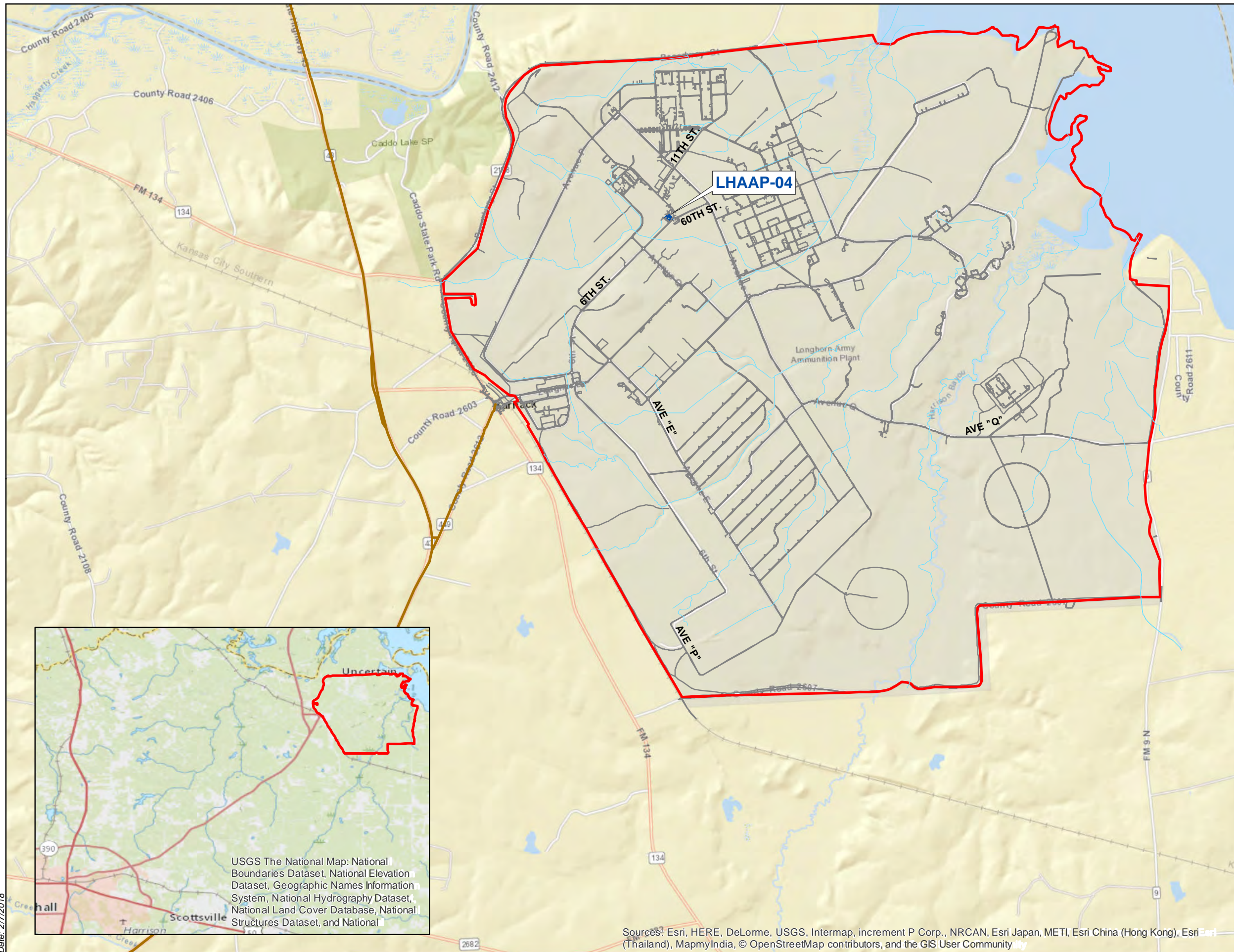
32. Type of Facility: Former Longhorn Army Ammunition Plant
33. Contamination Dates: 1940s to 1980s
34. Original Contamination (VOCs, TPH, BTEX, etc.) and Concentrations (Attach as Attachment J): The COC at LHAAP-04 is perchlorate from the wastewater treatment plant formerly located at the site. No separate Attachment J is provided.
35. Previous Remediation: Attach results of any previous remediation as Attachment K. Soil Remedial Action involved excavation of soil contaminated with perchlorate and was documented in the *Final Completion Report Non-Time-Critical Removal Action* dated August 2011. A separate Attachment K is not provided.





<<NOTE>> Authorization Form should be completed in detail and authorization given by TCEQ before construction, operation, and/or conversion can begin. Attach additional pages as necessary.

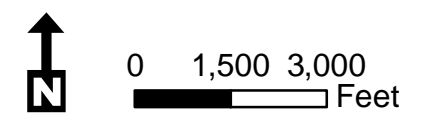
Attachment A

Site Topographic Map & Site Vicinity Map

- **Figure 1-2, LHAAP-04 Site Location Map**



-  Stream
-  Road
-  LHAAP Boundary
-  LHAAP-04 Site Boundary



U.S. ARMY CORP OF ENGINEERS
TULSA DISTRICT
TULSA, OKLAHOMA



Figure 1-2
LHAAP Site Location Map
LHAAP-04
Remedial Design/Remedial Action Work Plan
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

USGS The National Map: National Boundaries Dataset, National Elevation Dataset, Geographic Names Information System, National Hydrography Dataset, National Land Cover Database, National Structures Dataset, and National

Sources: Esri, HERE, DeLorme, USGS, Intermap, increment P Corp., NRCAN, Esri Japan, METI, Esri China (Hong Kong), Esri (Thailand), MapmyIndia, © OpenStreetMap contributors, and the GIS User Community

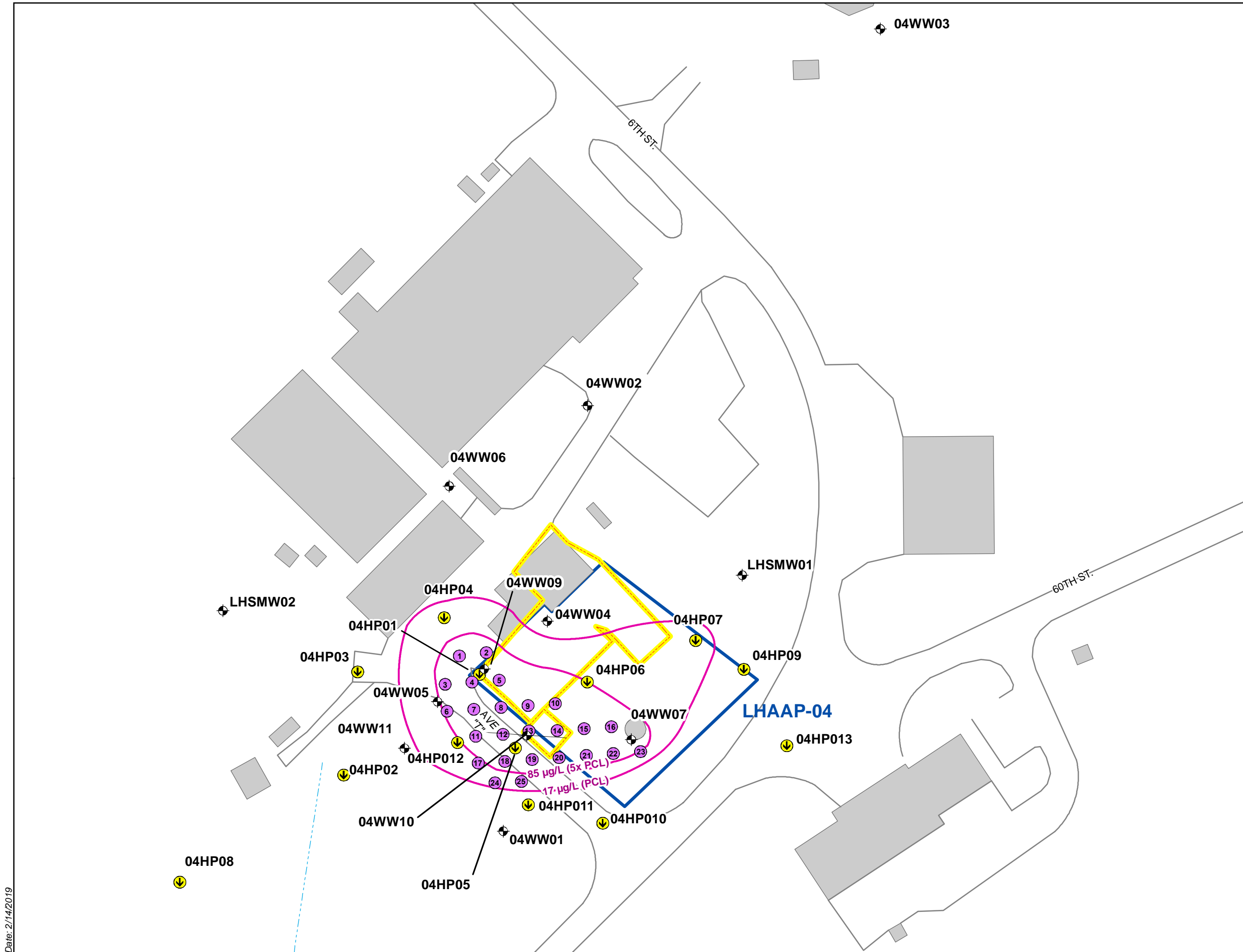
Date: 2/7/2018

Document Path: G:\Longhorn\LHAAP\Documents\Mxd\LHAAP04\RD_RAW\PI\Fig1-2_LHAAP_SiteLocationMap.mxd

Attachment B

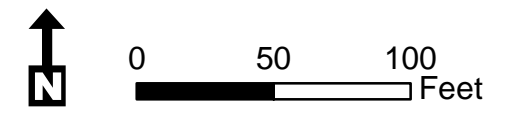
Proposed Injection Locations

- **Figure 5-1, LHAAP-04 Injection Plan, RD/RAWP**



- Shallow Monitoring Well
- Proposed Injection Location
- Hydropunch Location
- Perchlorate Plume
- Stream
- Limits of Excavation at Surface
(Source: LHAAP-04 Final Completion Report, Shaw, 2011)
- Road
- Building
- Site Boundary

Note:
 1. Plume boundaries based on most recent results available at each well (2018).
 2. PCL - Texas Risk Reduction Program Protective Concentration Level for Residential Groundwater
 3. The injection pattern generally targets the area within the 85 µg/L contour, but has been shifted to the southwest to account for potential future migration.



U.S. ARMY CORP OF ENGINEERS
 TULSA DISTRICT
 TULSA, OKLAHOMA



Figure 5-1

Injection Plan
 LHAAP-04

LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS

Date: 2/14/2019

Attachment C

Target Injection Intervals

- **Table 5-1, Injection Locations and Amendment Volumes, RD/RAWP**

Table 5-1
Injection Locations and Amendment Volumes

DPT Location	Amendment Volume per Location			Nearest Monitoring Well	DPT Injection Depths (ft bgs)
	Gallons of EVO (EDS-ER or Equivalent)	Gallons of Nutrients (DAP)	Gallons of Water		
04DPT03, -06, and -07	15	6	1,463	04WW05	12 - 20
04DPT01, -02, -04, and -05	15	6	1,463	04WW09	6 - 14
04DPT08, -09, -10, -11, -12, -13, -14, -17, -18, -19, -20, -24, and -25	15	6	1,463	04WW10	7 - 15
04DPT15, -16, -21, -22, and -23	15	6	1,463	04WW07	7 - 15

Notes:

ft bgs - feet below ground surface

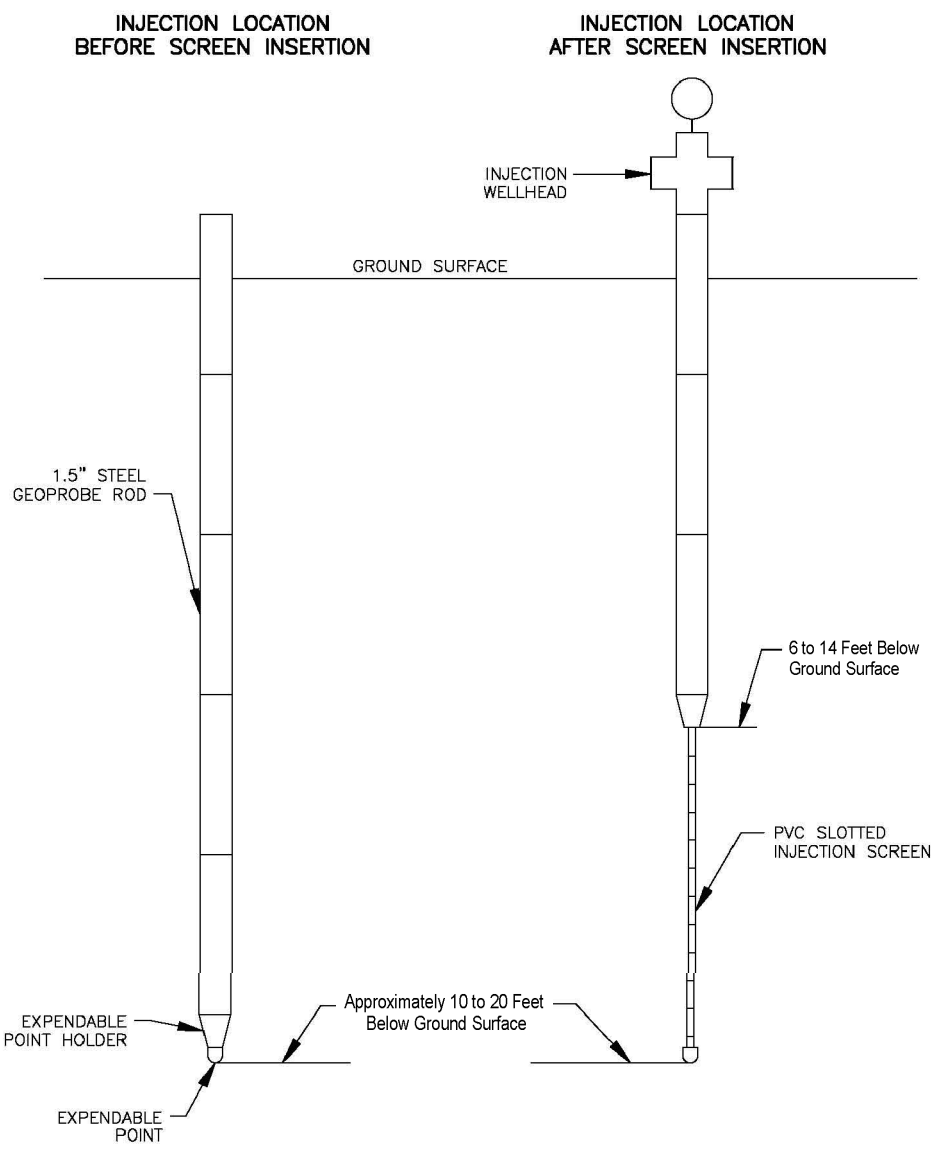
DAP - Diammonium phosphate

DPT - direct-push technology

evo - emulsified vegetable oil

Attachment D

Typical DPT Temporary Injection Point Diagram



Not to Scale

Project Number: 501032



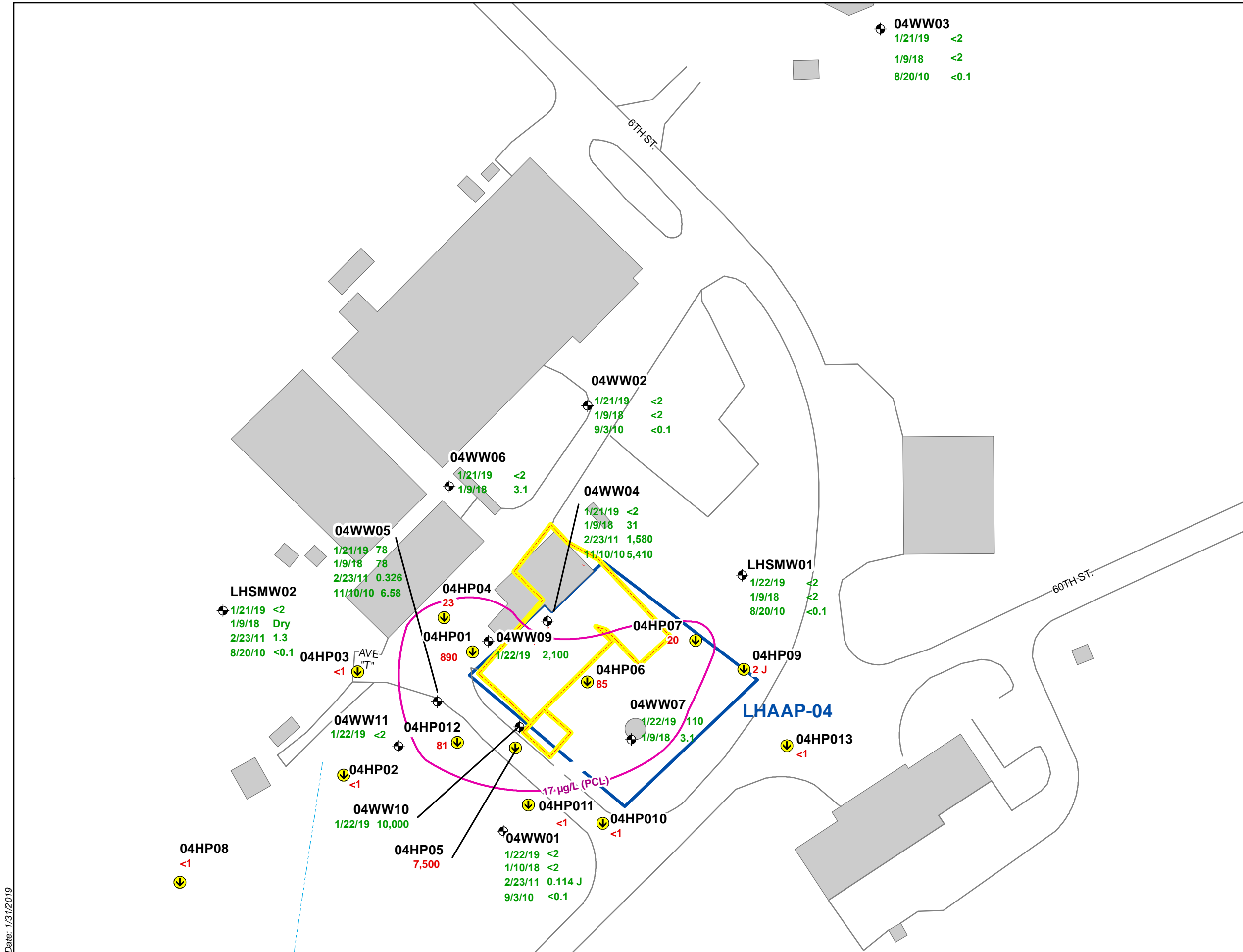
Attachment A
Typical DPT Temporary Injection Point
Diagram

Longhorn Army Ammunition Plant
Karnack, Texas

Attachment E

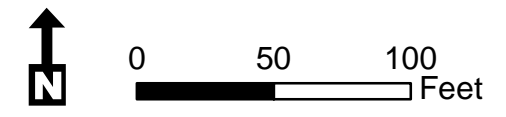
COC concentrations in Shallow Groundwater

- **Figure 2-3, Perchlorate Concentrations in Shallow Groundwater at LHAAP-04, RD/RAWP**



- ⊕ Shallow Monitoring Well
- ⬇️ Hydropunch Location
- Perchlorate Plume
- Stream
- Limits of Excavation at Surface
(Source: LHAAP-04 Final Completion Report, Shaw, 2011)
- Road
- ▭ Building
- ▭ Site Boundary

Note:
 1. Perchlorate concentrations reported in micrograms per liter (µg/L).
 2. Plume boundaries based on most recent results available at each well (2018-2019).
 3. PCL - Texas Risk Reduction Program Protective Concentration Level for Residential Groundwater
 4. 85 = Hydropunch November-December 2018 data
 110 = 2010, 2011, 2018, and 2019 monitoring well data



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 TULSA, OKLAHOMA

bhate
 ENVIRONMENT & INFRASTRUCTURE

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Figure 2-3
 Perchlorate Concentrations in Shallow Groundwater at LHAAP-04
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS

Attachment F
Not Applicable

Attachment G
Not Applicable

Attachment H

Safety Data Sheets

- **Electron Donor Solution (EDS-ER™)**



Material Safety Data Sheet

Electron Donor Solution

Section 1: Chemical Product and Company Identification

Product Name: Electron Donor Solution
Extended Release

Catalog Codes: EDS-ER

CAS#: 8001-22-7

TSCA: TSCA 8(b) inventory: Soybean oil

HMIS Code: H F R P: 10 0 A

Trade Name and Synonyms: EDS-ER

Chemical Family: Glyceride Oils

Contact Information:

Tersus Environmental, LLC

109 E. 17th Street, Suite #3880

Cheyenne, WY 82001

Ph: 307.638.2822 • info@tersusenv.com

www.tersusenv.com

For emergency assistance, call: 919.638.7892

Section 2: Composition and Information on Ingredients

COMPONANT	CAS #	OSHA TWA	OSHA STEL	ACGIH TWA	ACGIH STEL
Soybean Oil	8001-22-7	---	10 mg/m ³	---	---
Vegetable Oil Derived Fatty Acid Esters	Confidential	---	---	---	---

HAZARDOUS INGREDIENTS: NONE AS DEFINED UNDER THE U.S. OSHA HAZARD COMMUNICATION STANDARD (29 CFR 1910.1200) OR THE CANADIAN HAZARDOUS PRODUCTS ACT S.C. 1987, C.30 (PART 1).

THE PRECISE COMPOSITION OF THIS PRODUCT IS PROPRIETARY INFORMATION. A MORE COMPLETE DISCLOSURE WILL BE PROVIDED TO A PHYSICIAN IN THE EVENT OF A MEDICAL EMERGENCY.

SARA HAZARD: NONE NOTED (SECTION 311/312) TITLE III SECTION 313 - NOT LISTED
All components of this product are listed on the TSCA registry.

Section 3: Physical/Chemical Characteristics

BOILING RANGE: Not applicable VAPOR DENSITY: Exceeds 1.0

SPECIFIC GRAVITY (H₂O=1.0): 0.92 - 0.925 VAPOR PRESSURE: Not applicable

PERCENT VOLATILE BY VOLUME: 0% SOLUBILITY IN WATER: Miscible

EVAPORATION RATE: Not applicable

APPEARANCE AND ODOR: A pale yellow, oily liquid - only a faint odor.

WEIGHT PER GALLON: 7.7 lbs. at 60F.



Material Safety Data Sheet

Section 4: Fire and Explosion Data

FLAMMABILITY CLASSIFICATION: Combustible Liquid - Class IIIB.

FLASHPOINT: Greater than 550 F (288 C).

METHOD USED: Tag Closed Cup.

EXTINGUISHING MEDIA: CO₂, dry chemical, foam, sand.

SPECIAL FIREFIGHTING PROCEDURES: Avoid use of water as it may spread fire by dispersing oil. Use water to keep fire-exposed containers cool. Water spray may be used to flush spills away from fire.

UNUSUAL FIRE AND EXPLOSION HAZARDS: Rags soaked with any oil or solvent can present a fire hazard and should always be stored in UL Listed or Factory Mutual approved, covered containers. Improperly stored rags can create conditions that lead to oxidation. Oxidation, under certain conditions can lead to spontaneous combustion.

Section 5: Reactivity Data

STABILITY: Generally stable. Spontaneous combustion can occur. See Unusual Fire and Explosion Procedures, Section IV.

CONDITIONS TO AVOID: High surface area exposure to oxygen can result in polymerization and release of heat.

INCOMPATIBILITY (MATERIALS TO AVOID): Avoid contact with strong oxidizing agents.

HAZARDOUS DECOMPOSITIONS OR BY-PRODUCTS: Decomposition may produce carbon dioxide and carbon monoxide.

HAZARDOUS POLYMERIZATION: Will not occur.

Section 6: Health Hazard Data

THRESHOLD LIMIT VALUE: As a liquid - none. As oil mist - 10 mg/m³ total particulate.

INHALATION HEALTH RISKS AND SYMPTOMS OF EXPOSURE: Excessive inhalation of oil mist may affect the respiratory system. Oil mist is classified as a nuisance particulate by ACGIH.

SKIN ABSORPTION HEALTH RISKS AND SYMPTOMS OF EXPOSURE: Not classified as a primary skin irritant or corrosive material. Sensitive individuals may experience dermatitis after long exposure of oil on skin.

HEALTH HAZARDS (ACUTE AND CHRONIC): Acute: none observed by inhalation. Chronic: none reported.

EMERGENCY AND FIRST AID PROCEDURES FOR:

SKIN CONTACT: May be removed from skin by washing with soap and warm water.

EYE CONTACT: Immediately flush eyes with plenty of cool water for at least 15 minutes. Do NOT let victim rub eyes.

INHALATION: Immediately remove exposed individual to fresh air source. If victim has stopped breathing give artificial respiration, get medical attention immediately.



Material Safety Data Sheet

Section 7: Precautions for Safe Handling and Use

ENVIRONMENTAL PRECAUTIONS: Where large spills are possible, a comprehensive spill response plan should be developed and implemented.

STEPS TO BE TAKEN IN CASE MATERIAL IS RELEASED OR SPILLED: Wear appropriate respiratory protection and protective clothing as described in section VIII. Depending on quantity of spill: (a) Small spill - add solid adsorbent, shovel into disposable container and wash the area. Clean area with detergent. (b) Large spill - Squeegee or pump into holding container. Clean area with detergent. In the event of an uncontrolled release of this material, the user should determine if this release is reportable under applicable laws and regulations.

WASTE DISPOSAL METHOD: All recovered material should be packaged, labeled, transported, and disposed or reclaimed in accordance with local, state, and federal regulations and good engineering practices.

Section 8: Control Measures

RESPIRATORY PROTECTION: Not normally needed. A qualified health specialist should evaluate whether there is a need for respiratory protection under specific conditions.

VENTILATION: Handle in the presence of adequate ventilation. Intermittent clean air exchanges recommended, but not required.

PROTECTIVE GLOVES: Not normally needed. However, protective clothing is always recommended when handling chemicals.

EYE PROTECTION: Eye protection is always recommended when handling chemicals. Wear safety glasses meeting the specifications established in ANSI Standard Z87.1.

Section 9: Special Precautions

PRECAUTIONS TO BE TAKEN IN HANDLING AND STORAGE: Store away from flame, fire, and excessive heat.

Section 10: Disposal Considerations

General Information: Do not discharge into drains, watercourses or onto the ground. Discharge, treatment, or disposal may be subject to national, state, or local laws. Empty containers may contain product residues.

Disposal Methods: No specific disposal method required.

Container: Since emptied containers retain product residue, follow label warnings even after container is emptied.



Material Safety Data Sheet

Section 11: Transportation Information

DOT Not regulated.
 TDG Not regulated.
 IATA Not regulated.
 IMDG Not regulated.

Section 12: Other Information

Hazard Ratings

	Health Hazard	Fire Hazard	Instability	Special Hazard
NFPA	1	1	0	NONE

Hazard rating: 0 - Minimal; 1 - Slight; 2 - Moderate; 3 - Serious; 4 - Severe

NFPA Label colored diamond code: Blue - Health; Red - Flammability; Yellow - Instability; White - Special Hazards

	Health Hazard	Flammability	Physical Hazard	Personal Protection
HMIS	1	1	0	--

Hazard rating: 0 - Minimal; 1 - Slight; 2 - Moderate; 3 - Serious; 4 - Severe

HMIS Label colored bar code: Blue - Health; Red - Flammability; Orange - Physical Hazards; White - Special

Section 13: Disclaimer and/or Comments

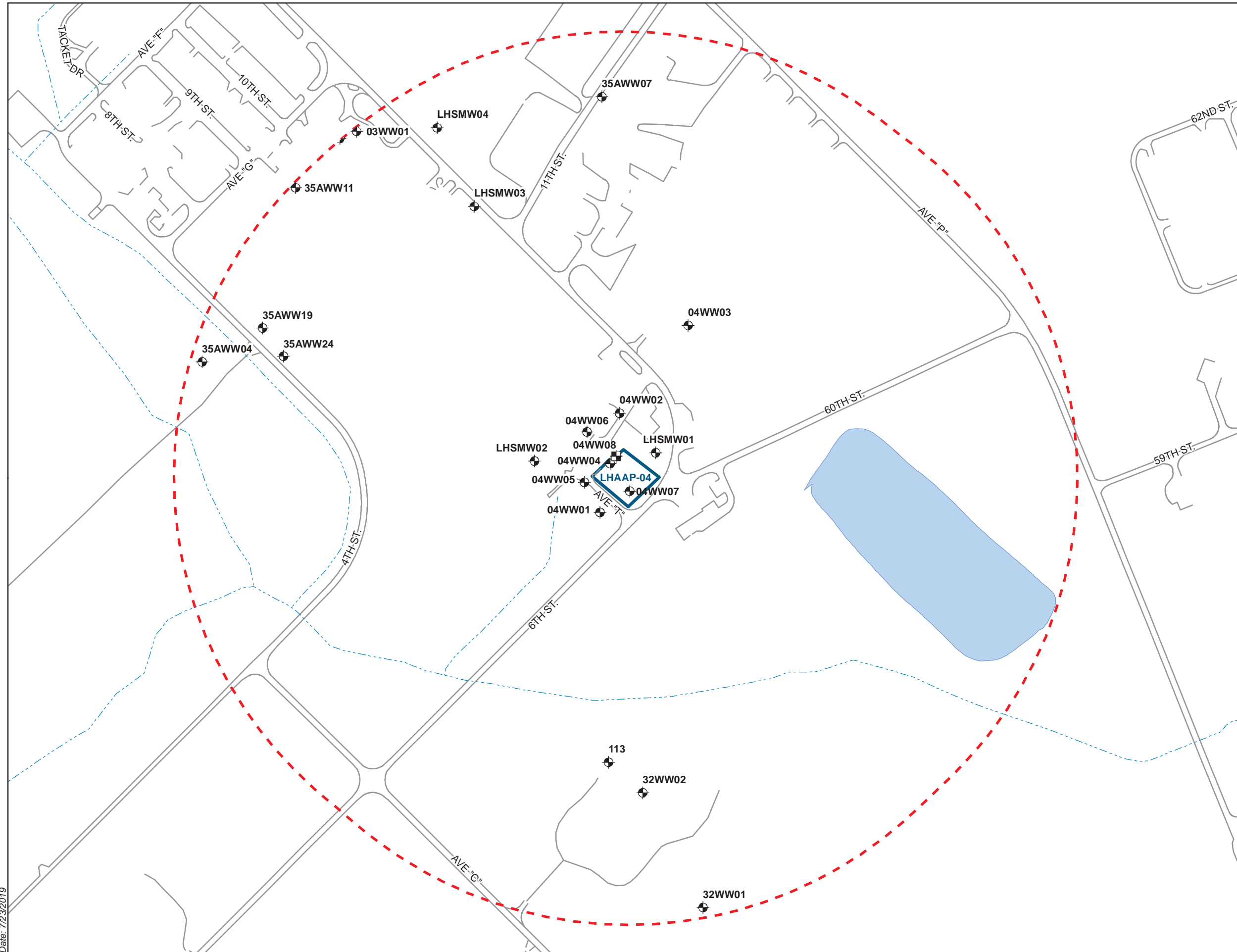
We suggest that containers be either professionally reconditioned for re-use by certified firms or properly disposed of by certified firms to help reduce the possibility of an accident. Disposal of containers should be in accordance with applicable federal, state and local laws and regulations. "Empty" drums should not be given to individuals.

The conditions of handling, storage, use and disposal of the product are beyond our control and may be beyond our knowledge. For this and other reasons, we do not assume responsibility and expressly disclaim liability for loss, damage or expense arising out of or in any way connected with the handling, storage, use or disposal of the product.

The information above is believed to be accurate and represents the best information currently available to us. However, we make no warranty of merchantability or any other warranty, express or implied, with respect to such information, and we assume no liability resulting from its use. Users should make their own investigations to determine the suitability of the information for their particular purposes. In no event shall Tersus Environmental be liable for any claims, losses, or damages of any third party or for lost profits or any special, indirect, incidental, consequential or exemplary damages, howsoever arising, even if Tersus Environmental has been advised of the possibility of such damages.

Attachment I

**Quarter Mile Radius Map
and Boring Logs**



- Shallow Monitor Well
- Intermediate Monitor Well
- Deep Monitor Well
- 1/4 Mile Radius
- Road
- Stream
- Site Boundary
- Lakes



0 150 300 Feet



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TULSA DISTRICT
TULSA, OKLAHOMA

Figure X

Wells within 1/4 Mile Radius
LHAAP-04

LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

Date: 7/23/2019

STATE OF TEXAS WELL REPORT for Tracking #162475

Owner: Longhorn Army Ammunition Plant	Owner Well #: 03WW01
Address: Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Grid #: 35-23-6
Well Location: Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Latitude: 32° 41' 10" N
Well County: Harrison	Longitude: 094° 09' 15" W
Elevation: No Data	GPS Brand Used: Garmin e-trex
<hr/>	
Type of Work: New Well	Proposed Use: Monitor

Drilling Date: Started: **11/18/2008**
 Completed: **11/18/2008**

Diameter of Hole: Diameter: **8.25 in From Surface To 30 ft**

Drilling Method: **Hollow Stem Auger**

Borehole
Completion: Gravel Packed From: **18 ft to 30 ft**
 Gravel Pack Size: **20/40**

Annular Seal Data: 1st Interval: **From 16 ft to 18 ft with 1 Bentonite (#sacks and material)**
 2nd Interval: **From 0 ft to 16 ft with 5 Cement (#sacks and material)**
 3rd Interval: **No Data**
 Method Used: **Tremmie Pipe**
 Cemented By: **Driller**
 Distance to Septic Field or other Concentrated Contamination: **No Data**
 Distance to Property Line: **No Data**
 Method of Verification: **No Data**
 Approved by Variance: **No Data**

Surface
Completion: **Surface Sleeve Installed**

Water Level: Static level: **No Data**
 Artesian flow: **No Data**

Packers: **No Data**

Plugging Info: Casing or Cement/Bentonite left in well: **No Data**

Type Of Pump: **No Data**

Well Tests: **No Data**

Water Quality: Type of Water: **No Data**
 Depth of Strata: **No Data**
 Chemical Analysis Made: **No Data**
 Did the driller knowingly penetrate any strata which contained undesirable constituents: **No**

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the log(s) being returned for completion and resubmittal.

Company
Information: **ETTL Engineers & Consultants Inc.**
 1717 E. Erwin

Tyler , TX 75702

Driller License Number: **2126**

Licensed Well Driller Signature: **H. Douglas Hinds**

Registered Driller Apprentice Signature: **No Data**

Apprentice Registration Number: **No Data**

Comments: **No Data**

IMPORTANT NOTICE FOR PERSONS HAVING WELLS DRILLED CONCERNING CONFIDENTIALITY

TEX. OCC. CODE Title 12, Chapter 1901.251, authorizes the owner (owner or the person for whom the well was drilled) to keep information in Well Reports confidential. The Department shall hold the contents of the well log confidential and not a matter of public record if it receives, by certified mail, a written request to do so from the owner.

Please include the report's Tracking number (Tracking #162475) on your written request.

Texas Department of Licensing & Regulation
P.O. Box 12157
Austin, TX 78711
(512) 463-7880

DESC. & COLOR OF FORMATION MATERIAL

From (ft) To (ft) Description
0-9.5 Sandy lean clay - brown
9.5-15.5 Silty sand - brown-gray
15.5-23.5 Clayey sand - brown
23.5-30 Sandy clay - gray

CASING, BLANK PIPE & WELL SCREEN DATA

Dia.	New/Used	Type	Setting From/To
2	New	PVC Sch. 40	0 - 20
2	New	PVC Sch. 40 - slotted	20 - 30 0.010"

DRILLING LOG		DIVISION USACE - Tulsa	INSTALLATION LHAAP-Karnack, TX	SHEET 1 OF 2 SHEETS
1. PROJECT LHAAP-Supplemental RI site 04		10. SIZE AND TYPE OF BIT Auger Bits		
2. LOCATION (Coordinates or Station)		11. DATUM FOR ELEVATION SHOWN (BSM or MSL)		
3. DRILLING AGENCY ETTL-Tyler, TX		12. MANUFACTURER'S DESIGNATION OF DRILL 35 = Soil Sample Mobil B-51 / 5' Cont. Sampler		
4. HOLE NO. (As shown on drawing title and file number) 04 WW01		13. TOTAL NO. OF OVER-BURDEN SAMPLES TAKEN DISTURBED 5 UNDISTURBED 0		
5. NAME OF DRILLER Tom Cook		14. TOTAL NUMBER CORE BOXES N/A		
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.		15. ELEVATION GROUND WATER		
7. THICKNESS OF OVERBURDEN Penetrated 24.0'		16. DATE HOLE STARTED 11-29-00 COMPLETED 11-29-00		
8. DEPTH DRILLED INTO ROCK N/A		17. ELEVATION TOP OF HOLE		
9. TOTAL DEPTH OF HOLE 23.0' below grade		18. TOTAL CORE RECOVERY FOR BORING N/A %		
		19. SIGNATURE OF INSPECTOR <i>[Signature]</i> PG C:Ka		

ELEVATION	DEPTH (ft)	LEGEND (USC)	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
13.00	0.0	CL	6" of Med. to dark brown topsoil with roots over		SS #1	1300 Commenced HSA Pilot hole 2 1/4" I.D. Augers for 04 WW01
12.00	3.8		Silty clay - Med. to dark brown and yellowish-brown - mottled, some Med. gray layering (1/2") trace to some fine sand - Damp	3.8 / 4.0	0.0 to 4.0	H2O col'd to isobutylerc (10.20v) Lamp
11.00	4.0					Ambient air = oppn
10.00	4.2		Silty clay Lt. to med. brown, gray, and yellowish-orange mottled, streaked, and stained, some occ DK brown streaks and inclusions, some fine sand throughout - Damp to sl moist (v. st.H)	4.2 / 5.0	4.0 to 9.0	H2O Reading in Sampler = 0ppn H2O Reading @ Augers = 0ppn Ambient Air = 0ppn
9.00	5.0					
8.00	5.0		Silty clay Lt. to med. brown and orange - Lower - mottled, some occ med. gray streaking and staining, trace fine sand throughout - Moist (v. st.H)	4.2 / 5.0	9.0 to 14.0	1325 H2O Reading in Sampler = 0 PPM H2O Reading @ Augers = 0 PPM
7.00	5.0					
6.00	5.0		Becoming Lt. to dark gray and layered @ 12.8' (Damp)			Ambient air = 0ppn
5.00	5.0	(CL)	Vis. moisture content inc. with depth		SS AA	H2O Reading in Sampler = 0ppn
4.00	5.0		Silty clay Lt. to med. gray v. thin layering and laminated texture/structure, trace lignite, horiz. bedding in thin to v. thin layers - Damp to Dry - v. stiff to Hard (clay shale properties) water in sample @ 18.5' along "fractured" texture of clays (shale) Note some v. thin fine sand layers - light gray - interbedded	5.0 / 5.0	14.0 to 19.0	H2O Reading @ Augers = 0ppn
3.00	5.0					
2.00	5.0				SS #5	1355

DRILLING LOG (Cont Sheet)			ELEVATION TOP OF HOLE		Hole No. 04WWD1		
PROJECT		SITE		INSTALLATION		SHEET	
Supplemental RI		Site 04		HAAP-Karnecker TX		2 of 2 SHEETS	
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)	
a	b	c	d	e	f	g	
	21	(Lk)	Silty clay med. to dark gray with abundant v. thin Lt. gray horiz. lenses, "breaks" along horiz. bedding, sample fractures easily along bedding - clay intervals	S.O. / S.O.	19.0 to 24.0	HNu Reading in Samplers = Oppm	
	22					HNu Reading @ Augers = Oppm	
	23						
	24		Damp and v. stiff			- Picked up augers off bottom @ 1400 for w.l. check from 18.5' zone	
	25		T.D. @ 24.0' below grade in Dry to			- Water in "hole" @ 1445	
	26		Damp layered silty clay			1455 Commenced overdrilling pilot hole with 11" O.D. "plugged" HSA for 4" SS monitoring well installation	
	27						
	28						
	29						
	30					- Bent./cement 8' to grade GROUT - Bent. Seal 11' - 8' (20-40) - Filterpack 23' - 11' - 4" SS #10 slot screen 23' - 13' T.D. @ 23.0' below grade	

DRILLING LOG		DIVISION USACE-Tulsa	INSTALLATION 04WU02	SHEET 1 OF 2 SHEETS
1. PROJECT LHAAP		10. SIZE AND TYPE OF BIT 8 1/2" OD HS / 10" IN OD HS		
2. LOCATION (Coordinates or Station) 04WU02		11. DATUM FOR ELEVATION SHOWN (TBM or MSL)		
3. DRILLING AGENCY FRTL		12. MANUFACTURER'S DESIGNATION OF DRILL FOREMOST B-5500		
4. HOLE NO. (As shown on drawing title and file number)		13. TOTAL NO. OF OVER-BURDEN SAMPLES TAKEN	DISTURBED 4	UNDISTURBED 0
5. NAME OF DRILLER DOUG HEINZ		14. TOTAL NUMBER CORE BOXES r/A		
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.		15. ELEVATION GROUND WATER		
7. THICKNESS OF OVERBURDEN		16. DATE HOLE STARTED 11-30-00 COMPLETED 11-30-00		
8. DEPTH DRILLED INTO ROCK		17. ELEVATION TOP OF HOLE		
9. TOTAL DEPTH OF HOLE		18. TOTAL CORE RECOVERY FOR BORING %		
		19. SIGNATURE OF INSPECTOR [Signature]		

ELEVATION a	DEPTH b	LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOVERY e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
	1	GC	0-3 BLACK ORGANIC TOPSOIL BROWN AND GRAY MOTTLED SILT CLAY, SOFT, MOIST, SOME ROUNDED GRAVEL SMALL TO MED. RIL	22" 24"	ST1	0738 Driller on location 0738 HNU OPPM
	2					
	3	SL	BROWN AND GRAY MOTTLED CLAYEY FINE SAND, MOIST, LOOSE, SOME ROOTLES, RUST STAINY	13" 12"	ST2	HNU OPPM
	4					
	5	CL	GRAY AND RED MOTTLED CLAY WITH FINE SAND STIFF, MOIST, PLASTIC	60" 60"	CS1	HNU OPPM
	6					
	7	CL	BROWN AND GRAY CLAY WITH FINE SAND, STIFF MOIST BROWN NODULES			
	8					0877
	9					
	10	OL	BROWN CLAY WITH FINE SAND INCREASE OF FINE SAND CONTENT, MOIST, LOOSE, NOT PLASTIC, TRACE BROWN NODULES, TRACE ROUND SMALL GRAVEL	60" 60"	CS2	HNU OPPM
	11					
	12					
	13					
	14	ML	BROWN CLAYEY SILT, FIRM DRY, NOT PLASTIC			HNU OPPM
	15					
	16					
	17	MH	BROWN FINE SAND, LOOSE, MOIST	60" 60"	CS3	
	18					
			BROWN CLAY, FIRM, MOIST, PLASTIC			

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 04ww02		
PROJECT LHAAP		INSTALLATION LHAAP		SHEET 2 OF 2 SHEETS		
ELEVATION a	DEPTH b	LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOVERY e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) g
	19	HL	GRAY CLAYEY FINE SAND, FIRM WET, HORIZONTAL DELINEATIONS			4200 02PM
	20			60"	654	0843 DECIDE TO
	21	MH	GRAY SILT, TRACE FINE SAND, STIFF, RTY NOT PLASTIC	60"		INSTALL WELL
	22					ALL 8 IN OD HS
	23					REAM HOLE W/
			EOB 23'			10 IN OD HS
						INSTALL SS
						23-13 SCREEN
						B - 11' SAND PACK
						ABOVE SCREEN
						11 - 8 BENTONITE
						8 - 0 GROUT
						0914 PIPPED OUT
						CASING PLUG AND
						INSTALLED WELL

028434

DRILLING LOG		DIVISION USACE - Tulsa	INSTALLATION LHAAP	SHEET 1 OF 2 SHEETS
1. PROJECT LHAAP		10. SIZE AND TYPE OF BIT 8 IN OD HS		
2. LOCATION (Coordinates or Station) SITE 4 04W03		11. DATUM FOR ELEVATION SHOWN (TBM or MSL)		
3. DRILLING AGENCY ETTL		12. MANUFACTURER'S DESIGNATION OF DRILL FOREMOST B-5500		
4. HOLE NO. (As shown on drawing title and file number)		13. TOTAL NO. OF OVER-BURDEN SAMPLES TAKEN		UNDISTURBED
5. NAME OF DRILLER DOUG HEINTZ		DISTURBED 7		UNDISTURBED 0
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.		14. TOTAL NUMBER CORE BOXES		
7. THICKNESS OF OVERBURDEN		15. ELEVATION GROUND WATER		
8. DEPTH DRILLED INTO ROCK N/A		16. DATE HOLE		STARTED 11-29-00 COMPLETED 11-29-00
9. TOTAL DEPTH OF HOLE		17. ELEVATION TOP OF HOLE		
		18. TOTAL CORE RECOVERY FOR BORING N/A		
		19. SIGNATURE OF INSPECTOR Soil Hecker		

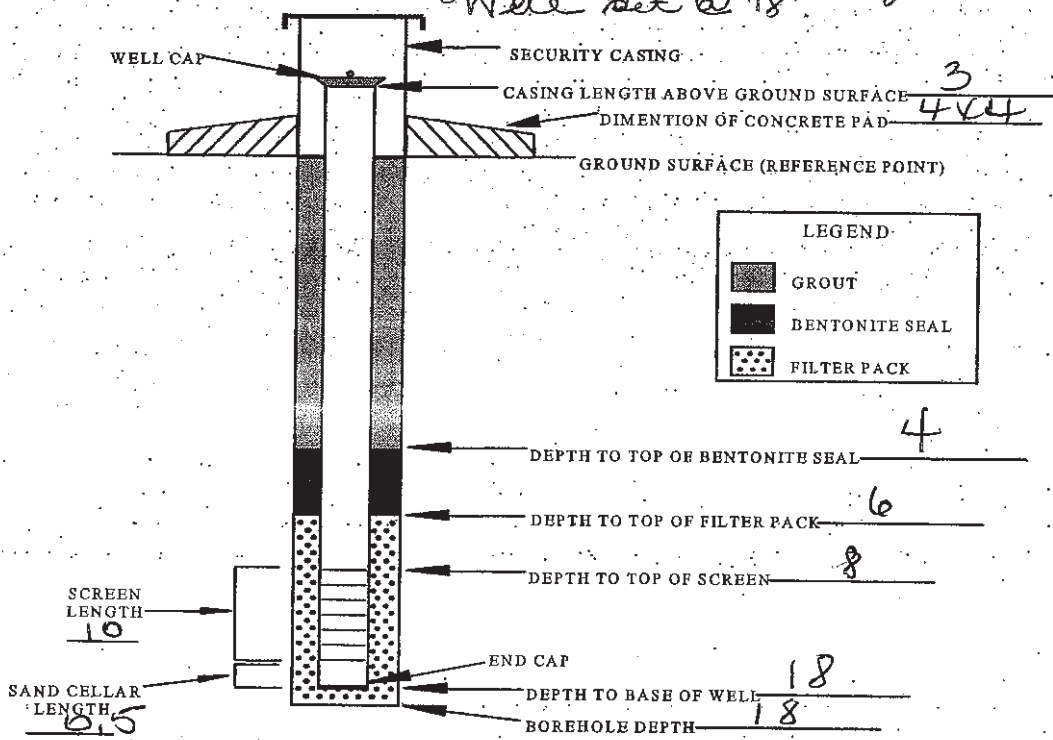
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
	1	SM	FILL BROWN & GRAY MOTTLED FINE SAND, LOOSE, MOIST, VEGETATION	22" / 24"	ST1	H2O 0 PPM 12/23 ST = Shelby TUBE
	2	SM	BROWN AND GRAY MOTTLED CLAYEY FINE SAND, LOOSE, MOIST, ROOTLETS, RUST STAINING, BLACK NODULES	17" / 12"	ST2	H2O 0 PPM
	3					
	4		BROWN AND GRAY MOTTLED FINE SANDY CLAY, STIFF, MOIST	54" / 60"	CS1	H2O 0 PPM CS = SFT CONTINUOUS SAMPLER
	5	SM	PLASTIC, RUST STAIN			
	6					
	7					
	8					
	9		BROWN AND GRAY MOTTLED SILTY CLAY WITH FINE SAND STIFF, DRY LOW PLASTICITY	60" / 60"	CS2	H2O 0 PPM
	10	SM-SC				
	11					
	12					
	13					
	14	SM-SC	SAME	55" / 60"	CS3	H2O 0 PPM
	15					
	16					
	17	OL	GRAY CLAYEY SILT, TRACE FINE SAND, VERY STIFF, DRY, NOT PLASTIC INCREASE IN FINE SAND IN DEPTH			
	18					

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 04 WU 03		
PROJECT LHAAP		INSTALLATION LHAAP		SHEET 2 OF 2 SHEETS		
ELEVATION a	DEPTH b	LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOVERY e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) g
	19		ALTERNATE LENSES OF BROWN AND GRAY CLAYEY FINE SAND	42"		THIS DRY HOLE. PULL UP AUGERS 2 FT TO SEE IF WATER PRESENT DRY HOLE BUT HEAR. 1456 ADVANCE
	20	SM	TRACE LIGNITE PIECES MOIST, LOOSE	30"		
	21	PT	BLACK LIGNITE, SOLID DRY, GRAY FINE SAND HORIZ. LAYERS		CSY	AUGERS TO 23 FT, AWAIT TO SEE IF WATER COMES IN CAN HEAR WATER.
	22					SS = SPIIT SPOON
	23	SH	GRAY CLAYEY SILT, STIFF DRY, HORIZ. LAYERS 2 IN	24"		1538 W/TAKE WATER level @ well on site
	24	SP	LIGHT GRAY FINE SAND	24"	SS1	5, 20 TO 17.8 FT
	25		FOB 25'			Done O.K.s Well
	26					23-13 screen
	27					13-11 over screen w/sand
	28					11-8 Bentonite
	29					5-0 grad. start to remove 8 in OD HS.
	30					Return w/10 in OD HS
						700 COMPLETE
						RETURN TO SITE 3.1.1
						Time 1 hour fast

WELL COMPLETION FORM (Stickup or Above Grade Completion Well)

FIELD REPRESENTATIVE: Olsen, S. TYPE OF FILTER PACK: Sand
 DRILLING CONTRACTOR: ETTL GRADIATION: 20/40
 AMOUNT OF FILTER PACK USED: 9
 DRILLING TECHNIQUE: Rotary TYPE OF BENTONITE: Pellets
 AUGER SIZE AND TYPE: 4" AMOUNT BENTONITE USED: 5 (1) for seal
 BOREHOLE IDENTIFICATION: 04WW04 TYPE OF CEMENT: Portland
 BOREHOLE DIAMETER: 4" AMOUNT CEMENT USED: 8
 WELL IDENTIFICATION: 04WW04 GROUT MATERIALS USED: -
 WELL CONSTRUCTION START DATE: 08/18/10
 WELL CONSTRUCTION COMPLETE DATE: _____ DIMENSIONS OF SECURITY CASING: 3x3
 SCREEN MATERIAL: Sch 40 PVC TYPE OF WELL CAP: Sch 40 PVC
 SCREEN DIAMETER: 2" TYPE OF END CAP: Sch 40 PVC
 STRATUM-SCREENED INTERVAL (FT): 8-18
 CASING MATERIAL: Sch 40 PVC COMMENTS:
 CASING DIAMETER: 2" 4 buckets of pellets used to fill from 18-28 ft. Well det @ 18'

SPECIAL CONDITIONS
(describe and draw)



INSTALLED BY: ETTL - B Reagin INSTALLATION OBSERVED BY: S. Olsen NOT TO SCALE
 DISCREPANCIES: NONE

HOLE NO. 04WW04

BORING LOG		DIVISION	INSTALLATION <u>LHAAP-04</u>	SHEET <u>1</u> OF <u>2</u> SHEETS
1. PROJECT <u>LHAAP</u>			9. DATUM FOR ELEVATION SHOWN (TBM or MSL) <u>MSL</u>	
2. LOCATION <u>LHAAP-04</u>			10. MANUFACTURER'S DESIGNATION OF DRILL <u>B5500</u>	
3. DRILLING AGENCY <u>ETTL</u>			11. OVERBURDEN SAMPLES	DISTURBED <u>—</u> UNDISTURBED <u>—</u>
4. HOLE NO. (As shown on drawing title and file number) <u>04WW04</u>			12. TOTAL NUMBER CORE BOXES <u>—</u>	
5. NAME OF DRILLER <u>B Reagan</u>			13. ELEVATION GROUND WATER <u>15.5 (DTW)</u>	
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED <u>0</u> DEG. FROM VERT.			14. DATE HOLE	STARTED <u>08/17/10</u> COMPLETED <u>08/18/10</u>
7. TOTAL DEPTH OF HOLE <u>27</u>			15. ELEVATION TOP OF HOLE <u>—</u>	
8. SIZE AND TYPE OF BIT <u>4" auger</u>			16. TOTAL CORE RECOVERY FOR BORING <u>100</u> %	
17. LOGGED BY <u>Olden, S</u>			QC <u>KE</u>	

PID (ppm) a	DEPTH b	USCS c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOVERY e	SAMPLE f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) g
	1		Backfill put in place during Shaw excavation activities.			
	2					
	3					
	4					
	5					
	6					
	7					
	8					
	9					
	10					
	11					
0.0	12	CL		Clay, sandy, low plasticity, dense, dry, friable, black	100%	

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 04WWD4		
PROJECT		INSTALLATION		SHEET 2 OF 2 SHEETS		
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
	13	SC	Sand, clayey, med grain, med to well sorted, dry, grey			
	14	CL	Clay, sandy, low to no plasticity, dense, dry, black to grey			
0.0	15	CL/CH	- becomes more moist, high plasticity, soft			
	16		- saturated			
	17	CL	- becomes dry, low to no plasticity, dense			Bentonite up to 18'. Set screen from 8 to 18.
	18					
	19					
0.0	20					
	21					
	22					
	23					
	24					
0.0	25					
	26					
	27		End of Boring			

STATE OF TEXAS WELL REPORT for Tracking #233508

Owner: USACE - LHAAP	Owner Well #: 04WW04
Address: Spur 449 @ Hwy 134 Karnack , TX 75661	Grid #: 35-23-6
Well Location: Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Latitude: 32° 40' 59" N
Well County: Harrison	Longitude: 094° 09' 06" W
Elevation: No Data	GPS Brand Used: Garmin 12 GPS
<hr/>	
Type of Work: New Well	Proposed Use: Monitor

Drilling Date: Started: **8/17/2010**
 Completed: **8/18/2010**

Diameter of Hole: Diameter: **7 7/8 in From Surface To 28 ft**

Drilling Method: **Mud Rotary Hollow Stem Auger**

Borehole
Completion: Gravel Packed From: **6 ft to 18 ft**
 Gravel Pack Size: **20/40**

Annular Seal Data: 1st Interval: **From 0 ft to 4 ft with 1 cement (#sacks and material)**
 2nd Interval: **From 4 ft to 6 ft with 1 bentonite (#sacks and material)**
 3rd Interval: **From 18 ft to 28 ft with 4 bentonite (#sacks and material)**
 Method Used: **Tremie pipe**
 Cemented By: **Driller**
 Distance to Septic Field or other Concentrated Contamination: **No Data**
 Distance to Property Line: **No Data**
 Method of Verification: **No Data**
 Approved by Variance: **No Data**

Surface
Completion: **Surface Sleeve Installed**

Water Level: Static level: **No Data**
 Artesian flow: **No Data**

Packers: **No Data**

Plugging Info: Casing or Cement/Bentonite left in well: **No Data**

Type Of Pump: **No Data**

Well Tests: **No Data**

Water Quality: Type of Water: **No Data**
 Depth of Strata: **No Data**
 Chemical Analysis Made: **No Data**
 Did the driller knowingly penetrate any strata which contained undesirable constituents: **No**

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the log(s) being returned for completion and resubmittal.

Company
Information: **ETTL Engineers & Consultants Inc.**
 1717 E. Erwin

Tyler , TX 75702
 Driller License Number: **54683**
 Licensed Well Driller Signature: **Wilburn Ragon, Jr.**
 Registered Driller Apprentice Signature: **No Data**
 Apprentice Registration Number: **No Data**
 Comments: **No Data**

IMPORTANT NOTICE FOR PERSONS HAVING WELLS DRILLED CONCERNING CONFIDENTIALITY

TEX. OCC. CODE Title 12, Chapter 1901.251, authorizes the owner (owner or the person for whom the well was drilled) to keep information in Well Reports confidential. The Department shall hold the contents of the well log confidential and not a matter of public record if it receives, by certified mail, a written request to do so from the owner.

Please include the report's Tracking number (Tracking #233508) on your written request.

Texas Department of Licensing & Regulation
P.O. Box 12157
Austin, TX 78711
(512) 463-7880

DESC. & COLOR OF FORMATION MATERIAL

From (ft) To (ft) Description
0-12 Fill
12-12.5 Sandy lean clay - black
12.5-13.5 Clayey sand - gray
13.5-28 Sandy lean clay - black & gray

CASING, BLANK PIPE & WELL SCREEN DATA

Dia.	New/Used	Type	Setting From/To
2	New	PVC Sch. 40	0 - 8
2	New	PVC Sch. 40 - slotted	8 - 18 0.010"

WELL COMPLETION FORM (Stickup or Above Grade Completion Well)

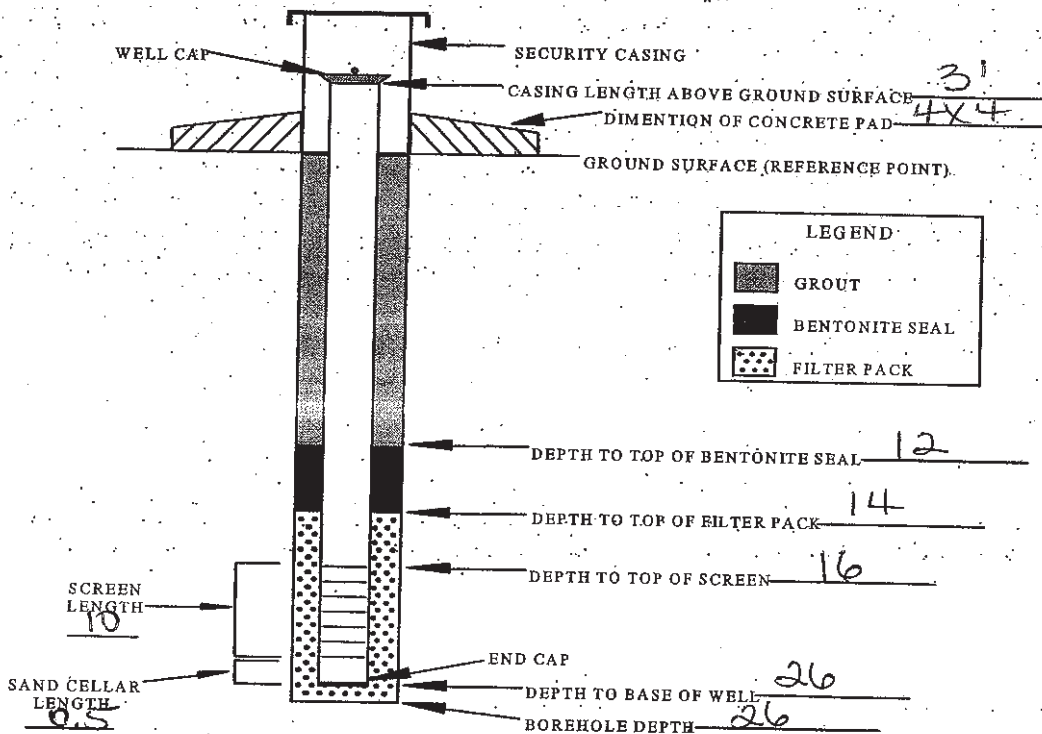
FIELD REPRESENTATIVE: J. Allen
 DRILLING CONTRACTOR: ETTL
 DRILLING TECHNIQUE: Rotary
 AUGER SIZE AND TYPE: 4"
 BOREHOLE IDENTIFICATION: 04W W05
 BOREHOLE DIAMETER: 4"
 WELL IDENTIFICATION: 04W W05
 WELL CONSTRUCTION START DATE: 08/18/10
 WELL CONSTRUCTION COMPLETE DATE: _____
 SCREEN MATERIAL: Sch 40 PVC
 SCREEN DIAMETER: 4" 2"
 STRATUM-SCREENED INTERVAL (FT): 16-26'
 CASING MATERIAL: Sch 40 PVC
 CASING DIAMETER: 4" 2"

TYPE OF FILTER PACK: Sand
 GRADATION: 20/40
 AMOUNT OF FILTER PACK USED: 7 bags
 TYPE OF BENTONITE: Pellet
 AMOUNT BENTONITE USED: 1 Bucket
 TYPE OF CEMENT: Portland
 AMOUNT CEMENT USED: 8
 GROUT MATERIALS USED: —

DIMENSIONS OF SECURITY CASING: 3" X 3"
 TYPE OF WELL CAP: Sch 40 PVC
 TYPE OF END CAP: Sch 40 PVC

COMMENTS: _____

SPECIAL CONDITIONS
(describe and draw)



INSTALLED BY: ETTL - B Reagen INSTALLATION OBSERVED BY: J. Allen NOT TO SCALE
 DISCREPANCIES: NONE

HOLE NO. 04 WW05

BORING LOG		DIVISION	INSTALLATION <u>LHAAP-04</u>	SHEET <u>1</u> OF <u>2</u> SHEETS
1. PROJECT <u>LHAAP</u>			9. DATUM FOR ELEVATION SHOWN (TBM or MSL) <u>MSL</u>	
2. LOCATION <u>LHAAP-04</u>			10. MANUFACTURER'S DESIGNATION OF DRILL <u>B5500</u>	
3. DRILLING AGENCY <u>ETTL</u>			11. OVERBURDEN SAMPLES	DISTURBED <u>-</u> UNDISTURBED <u>-</u>
4. HOLE NO. (As shown on drawing title and file number) <u>04 WW05</u>			12. TOTAL NUMBER CORE BOXES <u>-</u>	
5. NAME OF DRILLER <u>B. Reagan</u>			13. ELEVATION GROUND WATER <u>22'</u> (DTW)	
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED <u>0</u> DEG. FROM VERT.			14. DATE HOLE	STARTED <u>08/18/10</u> COMPLETED <u>08/18/10</u>
7. TOTAL DEPTH OF HOLE <u>26'</u>			15. ELEVATION TOP OF HOLE <u>-</u>	
8. SIZE AND TYPE OF BIT <u>4" auger</u>			16. TOTAL CORE RECOVERY FOR BORING <u>99.9%</u>	
			17. LOGGED BY <u>Olley S</u>	QC <u>KL</u>

PID (ppm)	DEPTH	USCS	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	SAMPLE	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
a	b	c	d	e	f	g
		OH	Topsail			
	1	CL	Clay, sandy, no plasticity, hard, dry, rocky, brown			
	2	SC	Sand, clayey, med grain med sorting, dry, rocky, brown			
0.0	3	CL	Clay, sandy, low to no plasticity, hard, dry, brown to reddish brown	100%		
	4					
	5					
	6					
	7					
0.0	8			100%		
	9					
	10					
	11					
	12					
		SC	Sand, clayey, rocky, med grain, poorly sorted,			

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 04WW05		
PROJECT			INSTALLATION		SHEET	
LHAAP			LHAAP-04		2	
			OF 2 SHEETS			
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
	13		friable w/ dark red-brown staining, dry, yellow to reddish brown			
0.0	14	CL	Clay, sandy, no plasticity, dense-hard, dry, black to grey	100%		
	15					
	16					
	17					
0.0	18			100%		
	19					
	20					
	21			0%		
0.0	22		saturated	▼		Expansive clay has caused core to become lodged in auger. Will go to shot pushed
	23					
	24					
0.0	25	CL	Clay, sandy, no plasticity, hard, dry, black-grey mottled			
	26		END of Boring			Set well @ 26' Screen 16-26'
	27					

STATE OF TEXAS WELL REPORT for Tracking #233509

Owner:	USACE - LHAAP	Owner Well #:	04WW05
Address:	Spur 449 @ Hwy 134 Karnack , TX 75661	Grid #:	35-23-6
Well Location:	Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Latitude:	32° 40' 58" N
Well County:	Harrison	Longitude:	094° 09' 07" W
Elevation:	No Data	GPS Brand Used:	Garmin 12 GPS
<hr/>			
Type of Work:	New Well	Proposed Use:	Monitor

Drilling Date: Started: **8/18/2010**
 Completed: **8/18/2010**

Diameter of Hole: Diameter: **8.25 in From Surface To 26 ft**

Drilling Method: **Hollow Stem Auger**

Borehole
Completion: Gravel Packed From: **14 ft to 26 ft**
 Gravel Pack Size: **20/40**

Annular Seal Data: 1st Interval: **From 0 ft to 12 ft with 3 cement (#sacks and material)**
 2nd Interval: **From 12 ft to 14 ft with 1 bentonite (#sacks and material)**
 3rd Interval: **No Data**
 Method Used: **Tremie pipe**
 Cemented By: **Driller**
 Distance to Septic Field or other Concentrated Contamination: **No Data**
 Distance to Property Line: **No Data**
 Method of Verification: **No Data**
 Approved by Variance: **No Data**

Surface
Completion: **Surface Sleeve Installed**

Water Level: Static level: **No Data**
 Artesian flow: **No Data**

Packers: **No Data**

Plugging Info: Casing or Cement/Bentonite left in well: **No Data**

Type Of Pump: **No Data**

Well Tests: **No Data**

Water Quality: Type of Water: **No Data**
 Depth of Strata: **No Data**
 Chemical Analysis Made: **No Data**
 Did the driller knowingly penetrate any strata which contained undesirable constituents: **No**

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the log(s) being returned for completion and resubmittal.

Company
Information: **ETTL Engineers & Consultants Inc.**
 1717 E. Erwin

Tyler, TX 75702

Driller License Number: **54683**

Licensed Well Driller Signature: **Wilburn Ragon, Jr.**

Registered Driller Apprentice Signature: **No Data**

Apprentice Registration Number: **No Data**

Comments: **No Data**

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Please include the report's Tracking number (Tracking #233509) on your written request.

Texas Department of Licensing & Regulation
P.O. Box 12157
Austin, TX 78711
(512) 463-7880

DESC. & COLOR OF FORMATION MATERIAL

CASING, BLANK PIPE & WELL SCREEN DATA

From (ft)	To (ft)	Description
0-0.5		Top soil
0.5-1.5		Sandy clay - brown
1.5-3		Clayey sand - brown
3-12.5		Sandy lean clay - brown & reddish brown
12.5-13.5		Clayey sand - red-brown, yellow, & reddish brown
13.5-26		Sandy lean clay - black & gray

Dia.	New/Used	Type	Setting From/To
2	New	PVC Sch. 40	0 - 16
2	New	PVC Sch. 40 - slotted	16 - 26 0.010"



Well / Boring Log

BORING NO.: _____
 MONITORING WELL NO.: 04WW06

SHEET 1 OF 1

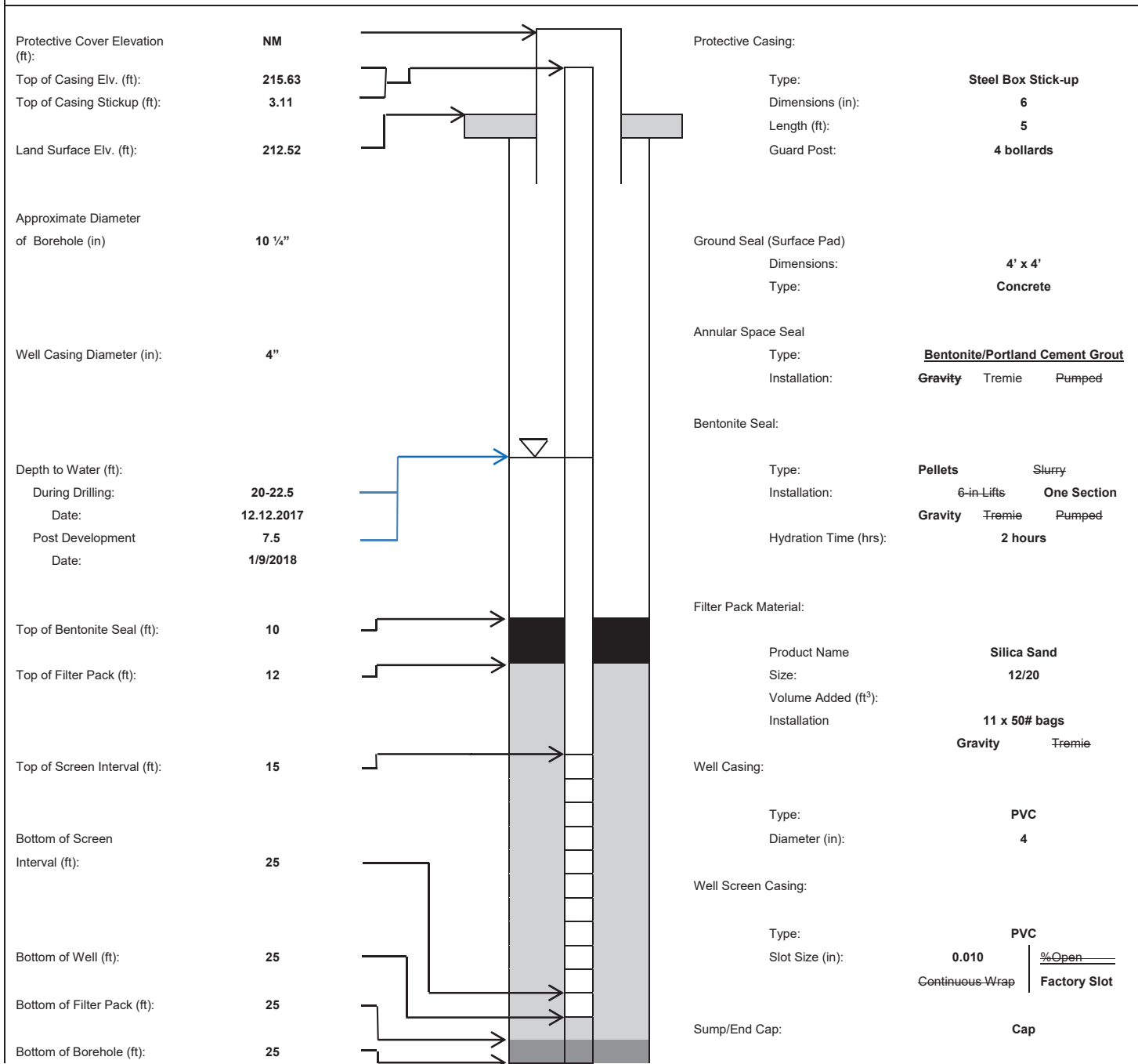
PROJECT NAME Longhorn Army Ammunition Plant	PROJECT NUMBER APTIM Project 501032	ELEVATION AND DATUM Northing: 6959225.38, Easting: 3305871.99	LOCATION LHAAP-04
DRILLING COMPANY Best Drilling	DRILLERS <u>Sonny Tobola</u>	DATE AND TIME STARTED <u>1030 : 12.12.2017</u>	DATE AND TIME COMPLETED <u>1150 : 12.12.2017</u>
DRILLING EQUIPMENT <u>CME 75 Mobile Rig</u>	DRILLING METHOD Hand Auger (0-5 feet) Direct Push (_____) HSA (<u>25</u>)	HOLE DIAMETER <u>10 1/4</u>	DP TOTAL DEPTH _____ HSA TOTAL DEPTH _____ TOTAL NO. OF SAMPLES / THICKNESS HA: _____ DP: _____ HSA: <u>13 / 2'6"</u>
DRILLING FLUID: NA	DRILLING ANGLE: 90 degrees (vertical)	WATER LEVEL DURING DRILLING: <u>~10'6"</u> FOLLOWING DRILLING: _____ ____ HRS AFTER DRILLING: <u>See Sample Form</u>	LOGGED BY: _____ DATE: <u>12.12.2017</u> D. Rowan CHECKED BY: _____ DATE: <u>12.12.2017</u> B. Foss

LITHOLOGY	DEPTH (FEET)	SAMPLE NO.	RECOVERY (INCHES)	STRENGTH (TCF)	PID READING (PPM)	DESCRIPTION Grading Term, Soil Group Name, Color (Code), Moisture, Consistency, Plasticity, Max Particle Sizes, Angularity, Shape, Other	USCS SYMBOL	ESTIMATED PERCENT OF			COMMENTS
								GRAVEL	SAND	FINES	
						Note: First 48" using post hole digger for utility clearance then probed to 60", boring drilled using ~10 1/4" diameter hollow stem augers, logged from split spoon @ 2.5' intervals					4" PVC Casing 10' 0.010 slot screen from 15 to 25'
	0	1	30	0		0 to 2'6" - Sandy clay, topsoil, some red silty clay	SC				
	2	2	18	0		2'6" to 4' - Gray silty clay, firm to soft, moist, mottled orange Fe staining, slight plasticity, platy layers	CL				
	4	3	12	0		4' to 5' - Tan sandy clay, moist, soft, mottled orange Fe staining, few calcite nodules, few intermittent pebbles	SC				
	6	4	18	0		5' to 6'6" - Dark gray to black silty clay, v. moist, lessening of sand concentration, orange Fe staining	CL				
	8	5	12	0		6'6" to 7'6" - Light gray silty clay, firm slightly plastic, moist, orange Fe staining, cont. lessening of sand concentration	CL				
	10	6	30	0		7'6" to 9' - Gray silty clay moist to v. moist, inter-bedded sand lenses, few calcite nodules	CL				
	12	7	12	0		9' to 10' - Tan sandy clay w/ inter-bedded gray clay lenses, v. moist, slightly mottled orange Fe staining	SC				
	14	8	18	0		10' to 12'6" - Light gray silty clay w/ inter-bedded sand lenses, moist to damp, firm to soft, plastic when wet	CL				
	16	9	30	0		12'6" to 15' - Same as above with the addition of int. orange Fe staining	CL				
	18	10	30	0		15' to 17'6" - Same as above	CL				
	20	11	30	0		17'6" to 20' - Same as above with the addition of int. black Fe Mn stained or black organics lenses	CL				
	22	12	30	0		20' to 22'6" - Dark gray to black silty clay w/ int. tan sand, firm, damp	CL				
	24	13	30	0		22'6" to 25' - Same as above w/o sand	CL				
	25					25' - End Boring					



MONITORING WELL CONSTRUCTION

Project: Longhorn Army Ammunition Plant Location: Karnack, Texas Client: US Army Corp of Engineers Subcontractor: Best Drilling Driller: Sonny Tobola Field Representative: B. Foss / D. Rowan	Well Number: 04WW06 Site Location: LHAAP-04 Installation Date: 12.12.2017 Northing: 6959225.38 Easting: 3305871.99
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Well / Boring Log

BORING NO.: _____
 MONITORING WELL NO.: 04WW07

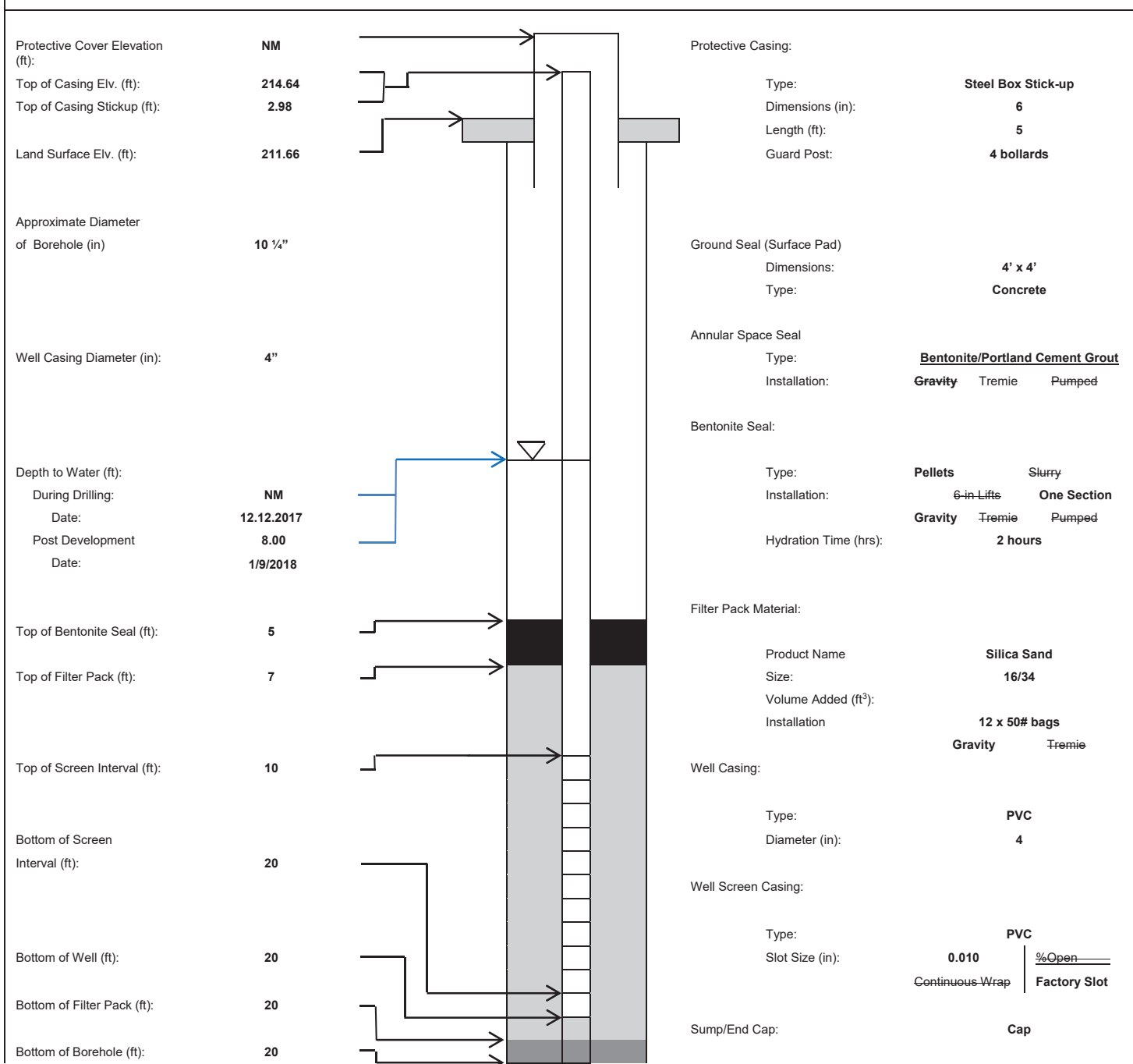
PROJECT NAME Longhorn Army Ammunition Plant	PROJECT NUMBER APTIM Project 501032	ELEVATION AND DATUM Orthing: 5 038. 3 asting 330 00 .11	LOCATION P-04
DRILLING COMPANY est rilling	DRILLERS _onn To ola	DATE AND TIME STARTED 1430 : 12.12.2017	DATE AND TIME COMPLETED 1530 : 12.12.2017
DRILLING EQUIPMENT 75 o ile Rig	DRILLING METHOD Hand Auger (0-5 feet) Direct Push (____) HSA (20)	HOLE DIAMETER 10 1/4	DP TOTAL DEPTH _____ HSA TOTAL DEPTH _____ TOTAL NO. OF SAMPLES / THICKNESS HA: _____ DP: _____ HSA: 11 / 2
DRILLING FLUID: NA	DRILLING ANGLE: 90 degrees (vertical)	WATER LEVEL DURING DRILLING: _____ FOLLOWING DRILLING: _____ ____ HRS AFTER DRILLING: _ee_a_ple or	LOGGED BY: DATE: _ Rowan 12.12.2017 CHECKED BY: DATE: _oss 12.12.2017

LITHOLOGY	DEPTH (FEET)	SAMPLE NO.	RECOVERY (INCHES)	STRENGTH (TCF)	PID READING (PPM)	DESCRIPTION Grading Term, Soil Group Name, Color (Code), Moisture, Consistency, Plasticity, Max Particle Sizes, Angularity, Shape, Other	USCS SYMBOL	ESTIMATED PERCENT OF			COMMENTS
								GRAVEL	SAND	FINES	
	0-10	1	18		0	ote: first 48 sing post hole digger for tilt clearance then pro ed to 0 oring drilled sing 10 1/4 dia eter hollow ste a gers logged fro split spoon 2.5 inter als					4 P asing 10 screen fro 10 to 20
	10-15	2	30		0	1 to 2 - Tan to gra silt cla w/ int. sand lenses orange e staining w/ lack e n or organic nod les oist w/ areas of . oist					
	15-20	3	30		0	2 to 5 - a e as a o e w/ increase in e staining higher concentration of sand in otto 1					
	20-25	4	30		0	5 to 7 - ilt la w/ loss of sand fir oist plastic ottled orange e staining lessening silt concentration with depth					
	25-30	5	30		0	7 to 10 - a e as a o e w/ so e int. sand lenses					
	30-35	6	30		0	10 to 12 - a e as a o e w/ lessening e staining					
	35-40	7	24		0	12 to 15 - ight gra to lack silt cla w/ ottled orange e staining nod les tr. sand/silica thro gho t fir to soft oist to da p little to no plasticit e en when wet					
	40-45	8	12		0	15 to 17 - a e as a o e w/ lack e n or organic nod les at 1 w/ darkening of color at 1					
	45-50	9	12		0	17 to 18 - sa e as a o e					
	50-55	10	12		0	18 to 18 - lack dr cla w/ few nod les of orange e staining					
	55-60	11	30		0	18 to 1 - lack dr silt cla fir .dense					
	60-65					1 to 22 - a e as a o e					
	65-70					20 - nd oring e tra 2 sa ple description fro split spoon sa ple onl did not ore down to 22 .					



MONITORING WELL CONSTRUCTION

Project: Longhorn Army Ammunition Plant Location: Karnack, Texas Client: US Army Corp of Engineers Subcontractor: Best Drilling Driller: Sonny Tobola Field Representative: B. Foss / D. Rowan	Well Number: 04WW07 Site Location: LHAAP-04 Installation Date: 12.12.2017 Northing: 6959038.63 Easting: 3306006.11
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Well / Boring Log

BORING NO.: _____
 MONITORING WELL NO.: **04WW0**

SHEET 1 OF 1

PROJECT NAME Longhorn Army Ammunition Plant	PROJECT NUMBER APTIM Project 501032	ELEVATION AND DATUM orthing: 95914 .15 asting: 33059 3.41	LOCATION P-04
DRILLING COMPANY est rilling	DRILLERS onn To ola	DATE AND TIME STARTED 0725 : 12.12.2017	DATE AND TIME COMPLETED 1100 : 12.14.2017
DRILLING EQUIPMENT 75 o ile Rig	DRILLING METHOD Hand Auger (0-5 feet) Direct Push (_____) HSA (45)	HOLE DIAMETER 14 10.25	DP TOTAL DEPTH _____ HSA TOTAL DEPTH _____ TOTAL NO. OF SAMPLES / THICKNESS HA: _____ DP: _____ HSA: 1 / 2 to 5
DRILLING FLUID: NA	DRILLING ANGLE: 90 degrees (vertical)	WATER LEVEL DURING DRILLING: 32 FOLLOWING DRILLING: _____ ____ HRS AFTER DRILLING: ee a ple or	LOGGED BY: DATE: . Rowan 12.12-14.2017 CHECKED BY: DATE: . oss 12.14.2017

LITHOLOGY	DEPTH (FEET)	SAMPLE NO.	RECOVERY (INCHES)	STRENGTH (TCF)	PID READING (PPM)	DESCRIPTION Grading Term, Soil Group Name, Color (Code), Moisture, Consistency, Plasticity, Max Particle Sizes, Angularity, Shape, Other	USCS SYMBOL	ESTIMATED PERCENT OF			COMMENTS
								GRAVEL	SAND	FINES	
	0					ote: first 4 sing post hole digger for tilt clearance then pro ed to 0 . first 23 drilled with 14 dia eter and logged 2.5 inter als sing split spoon. ro 23 to 41 was drilled with 10.25 and logged 5 inter als sing split spoon				12 P s rface casing set in the shallow one from 0 to 23 on 12.12.2017.	
	5	1			0.0	0 to 5 - ra cla e sand silt fir increasing cla concentration w/ decreasing sand with depth				4 P casing set 41 w/ 10 screen from 31 to 41 . on 12.14.17	
		2			0.0	5 to 7 - Red sand cla fir oist with depth there is increasing oist re content as well as decreasing sand concentration				Top 12 ost likel ackfill from e ca ation as denoted 04WW04 oring log.	
	10	3	30		0.0	7 to 10 - Red silt cla soft oist to . oist tr. orange e staining w/ so e nod les of lack e n or organics				ro 23 to T sa ples were collected for logging 5 inter als.	
		4			0.0	10 to 11 - a e as a o e				arrel eca e st ck in a ger 0930 d ring 23 to 25 sa pling r n. P lled a gers drilling res ed 1010	
		5	1		0.0	11 to 12 - Transition to nati e gra silt cla da p tr. sand ottled orange e staining w/ int. lack nod les of e n or organics				Water 32	
		6	12		0.0	12 to 15 - oss of red cla gra silt cla . fir da p sheeted plastic when wet tr. int. sand/silica particles					
	15	7	30		0.0	15 to 17 - a e as a o e w/ increasing tr. of sand starting at 1 so e orange e staining					
		8	30		0.0	17 to 20 - ight gra to dark gra cla w/silt and tr. sand fir oist plastic when wet					
	20	9	30		0.0	20 to 22 - a e as a o e w/ so e orange e staining starting at 21					
		10	30		0.0	22 to 23 - a e as a o e					
		11			0.0	23 - et shallow one casing					
		12	24		5.0	23 to 25 - ight to dark gra silt cla fir dr w/ int. lack organic nod les					
	25	13	0		3.0	25 to 30 - edi to dark gra silt cla . fir dense dr int. lack organic nod les. ncrease in oist re to a oist consistenc					
		14	12		.0	30 to 31 - ark gra cla oist so e orange e staining fir to soft plastic when wet					



Well / Boring Log

BORING NO.: _____
 MONITORING WELL NO.: **04WW0**

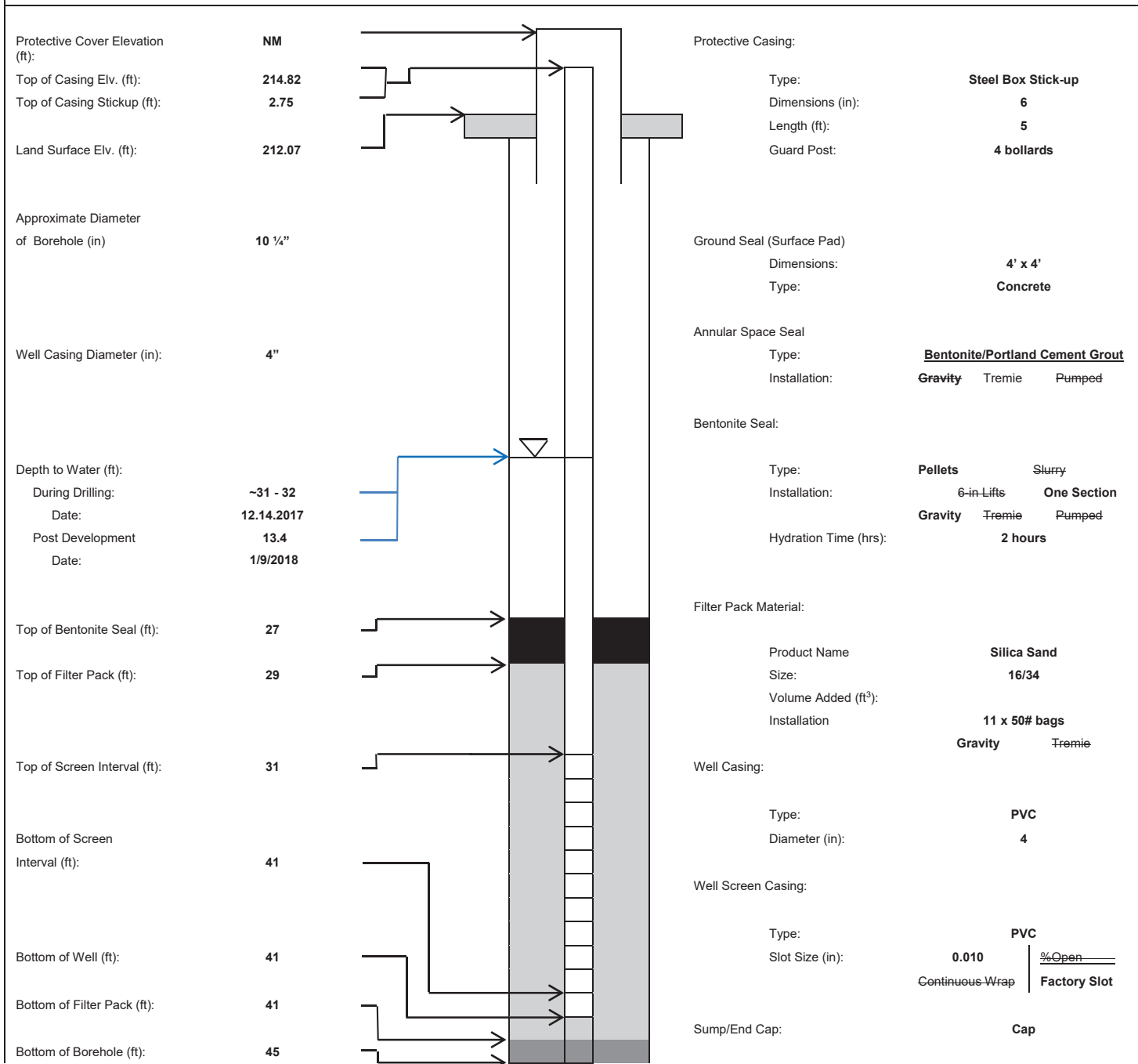
PROJECT NAME Longhorn Army Ammunition Plant	PROJECT NUMBER APTIM Project 501032	ELEVATION AND DATUM orthing: 95914 .15 asting: 33059 3.41		LOCATION P-04
DRILLING COMPANY est rilling	DRILLERS onn To ola	DATE AND TIME STARTED 0725 : 12.12.2017		DATE AND TIME COMPLETED 1100 : 12.14.2017
DRILLING EQUIPMENT 75 o ile Rig	DRILLING METHOD Hand Auger (0-5 feet) Direct Push () HSA (45)	HOLE DIAMETER 14 10.25	DP TOTAL DEPTH _____ HSA TOTAL DEPTH _____	TOTAL NO. OF SAMPLES / THICKNESS HA: _____ DP: _____ HSA: 1 / 2 to 5
DRILLING FLUID: NA	DRILLING ANGLE: 90 degrees (vertical)	WATER LEVEL DURING DRILLING: 32 FOLLOWING DRILLING: _____ HRS AFTER DRILLING: ee a ple or		LOGGED BY: . Rowan DATE: 12.12-14.2017 CHECKED BY: . oss DATE: 12.14.2017

LITHOLOGY	DEPTH (FEET)	SAMPLE NO.	RECOVERY (INCHES)	STRENGTH (TCF)	PID READING (PPM)	DESCRIPTION Grading Term, Soil Group Name, Color (Code), Moisture, Consistency, Plasticity, Max Particle Sizes, Angularity, Shape, Other	USCS SYMBOL	ESTIMATED PERCENT OF			COMMENTS
								GRAVEL	SAND	FINES	
	35	15	1		.0	31 to 32 - lack cla w/silt dr so e che ical odor . fir and dense.					32 to 35 no reco er hard cla pl g for ed in arrel. Partial sand reco er at otto inferred sat rated silt sand fro 32 to 35 .
		1	4		7.5	35 to 39 - ark gra to lack silt cla fro top to otto sat rated to oist tr. of che ical odor fir to stiff slight plasticit					
	40	17	12		9.2	39 to 40 - ight gra cla e silt da p no odor sheeted fir to soft no plasticit					
		1	0		.	40 to 45 - a e as a o e 45 - nd oring. Well et at 41					



MONITORING WELL CONSTRUCTION

<p>Project: Longhorn Army Ammunition Plant</p> <p>Location: Karnack, Texas</p> <p>Client: US Army Corp of Engineers</p> <p>Subcontractor: Best Drilling</p> <p>Driller: Sonny Tobola</p> <p>Field Representative: B. Foss / D. Rowan</p>	<p>Well Number: 04WW08</p> <p>Site Location: LHAAP-04</p> <p>Installation Date: 12.12-14.2017</p> <p>Northing: 6959148.15</p> <p>Easting: 3305963.41</p>
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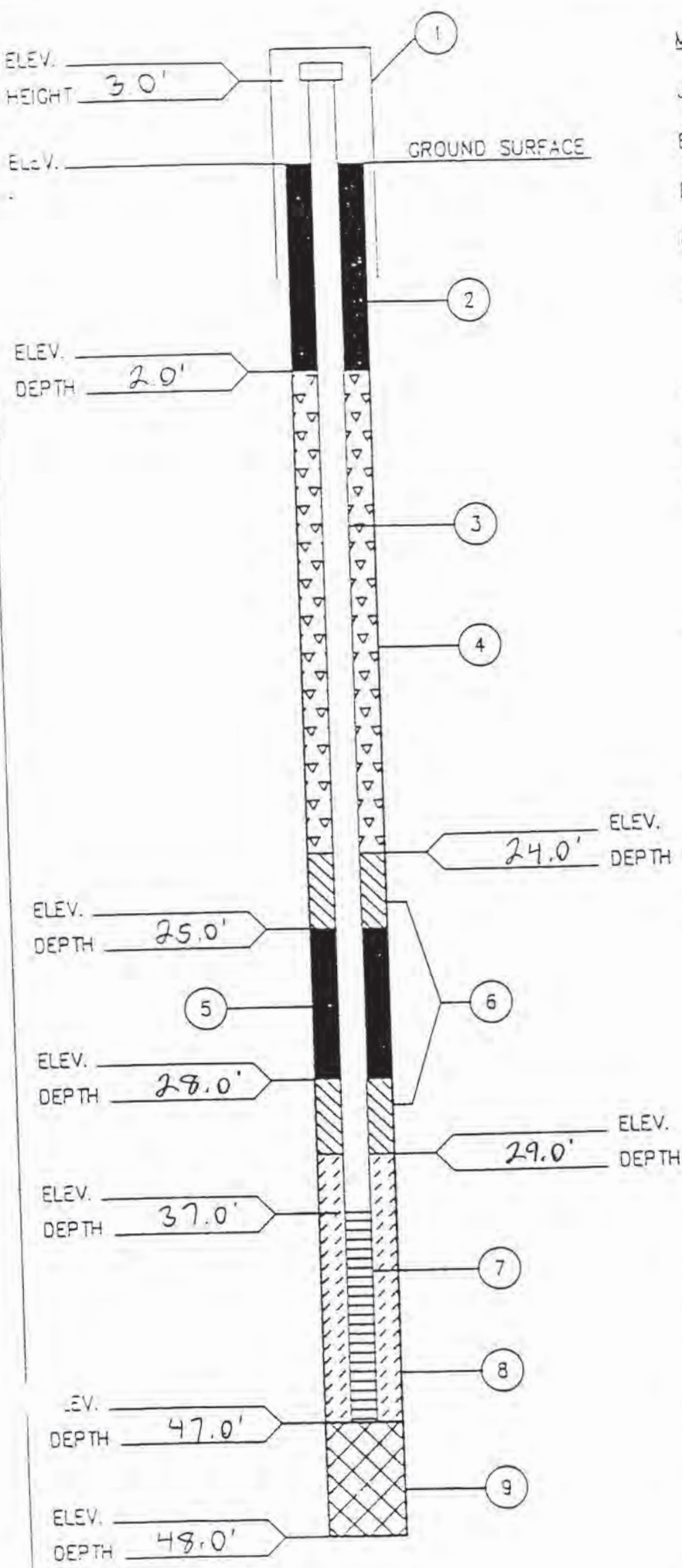


DRILLING LOG		DIVISION USACE, Tulsa	INSTALLATION LHAAP-Kornack TX	SHEET 1 OF 3 SHEETS
1. PROJECT LHAAP Phase III Site 32		10. SIZE AND TYPE OF BIT Anvil Bit		
2. LOCATION (Coordinates or Station) Southern Side of Site 32		11. DATUM FOR ELEVATION SHOWN (TBM or MSL)		
3. DRILLING AGENCY G.P.I. Inc. - Austin, TX		12. MANUFACTURER'S DESIGNATION OF DRILL CME-75 S.C.M.F. (with sample) 3 1/4" x 0.0 / 11" O.D.		
4. HOLE NO. (As shown on drawing title and file number) 32WW "01"		13. TOTAL NO. OF OVER-BURDEN SAMPLES TAKEN DISTURBED: 10 UNDISTURBED: -		
5. NAME OF DRILLER Jose Landeros		14. TOTAL NUMBER CORE BOXES N/A		
6. DIRECTION OF HOLE <input type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.		15. ELEVATION GROUND WATER -		
7. THICKNESS OF OVERBURDEN Penetrated		16. DATE HOLE STARTED: 7-22-97 COMPLETED: 7-24-98		
8. DEPTH DRILLED INTO ROCK N/A		17. ELEVATION TOP OF HOLE -		
9. TOTAL DEPTH OF HOLE Fl. Below Gravel		18. TOTAL CORE RECOVERY FOR BORING N/A		
		19. SIGNATURE OF INSPECTOR 1/ [Signature] 1/ [Signature]		

ELEVATION a	DEPTH (ft) b	USC LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOV- ERY e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) g
	1	OL CL	3" of Light blown DRP topsoil, some roots, over-		SS #1	Commenced HSA @ 1:16 3 1/4" x 0.0 / 11" O.D.
	2		- silty clay/clayey silt. Lt. to med. brown and gray, some fine sand throughout, trace roots - DRP (stiff)	10'	0.0 7.0 4.0'	HAW Reading = 9ppm in sampler
	3					
	4		- silty clay, Med brown, tan, Lt. gray, and orange-brown, mottled, streaked, and stained, some fine sand. Damp (stiff)		SS #2	HAW Reading = 9ppm in sampler
	5			5.0'	4.0 1.0 9.0'	
	6					
	7					
	8					
	9					
	10		- silty clay, Lt. to med. gray, tan, orange-brown and yellow-brown mottled streaked and stained some apparent layering, trace some fine sand Damp to Moist (stiff)	5.0'	SS #3 9.0 to 14.0'	HAW Reading = 9ppm in sampler
	11					
	12					
	13					
	14		- silty clay/clayey silt, Lt. gray, brown, tan, and yellow- brown mottled, streaked, and stained, some very silty sections from 14 to 16.0. Becoming moist to sl. wet @ 17' to 18' trace some fine sand throughout - Moist (stiff)	5.0'	SS #4 14.0 to 17.0'	HAW Reading = 9ppm in sampler
	15					
	16					
	17					
	18					
	19					
	20				SS #5	Water @ 17' B.E.G. w/1

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 32WW '01'		
PROJECT LHAAP Phase III Group IV		INSTALLATION LHAAP		SHEET 2 OF 3 SHEETS		
ELEVATION a	DEPTH ft. b	USC LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOV. e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) g
		CL	Med. gray color inc		19.0 to 24.0	Htu Readings open in sample
21			- silty clay / clayey silt, med. gray, tan, and yellow-brown, mottled, streaked, and stained, some abundant with horz layering of Lt. gray and tan silt + sand / sandy silt layers throughout sample, trace to some fine sand throughout, moist to wet (stiff)	S.O.		
22						
23						
24						
25					SS #6 24.0 to 29.0	Htu Readings = open in sample
26			- silty clay / clayey silt med. to dark gray with abundant U: thin horz. Lighter gray sandy silt / silty sand layers, some are tan and rust yellow-brown layers, trace to some fine sand throughout - Dry to Damp - (U: stiff)	S.O.		
27						
28						
29						
30			- Silty clay, some as a clay shale. Dark gray with abundant med. gray v. thin irregular fine sandy silt / silty sand laminations, some fine sand throughout, crumbles easily along laminations - Dry to Damp (Hard)	S.O.	SS #7 29.0 to 34.0	Stopped drilling @ HSA @ 1455 on 7/22/98 @ 29 ft. Hole Dry \$1000 7/24/98 - Resumed HSA @ 29.0'
31						
32						
33						
34			- Clayey silt, some as a sandy silt along laminations, med. gray with some to abundant U: thin silty sand irregular lenses / laminations, some clayey zones as clay shales, fine sand throughout sample - Dry to Damp (stiff to hard)		SS #8 34.0 to 39.0	1038 - 7/24/98 (@ 34 ft.) Htu Readings in samples open
35						
36						
37						
38		SM				
39			- Sandy silt, very clayey, in some zones, med. gray to med. blue-gray, fine grained sand throughout, some apparent layering, crumbles easily by hand, silt and sand contrast inc. with depth - Damp to sl. moist, (can shape by hand) (dense to loose)	S.O.	SS #9 39.0 to 44.0	Htu Readings in samples open
40						
41						
42						
43						
44						

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 32W401		
PROJECT			INSTALLATION		SHEET	
			LHAAP - Karnack, TX		3	
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
a	b	c	d	e	f	g
	45	SM	- Sandy silt, some as silty sand, med. gray to med. blue-gray, fine grained. Some apparent layering some med. gray fine v. sand layers - <u>Damp to moist</u> - (Dense to loose)	5.0'	SS	HAM Reading: Super off
	46	#10				
	47	44.0 to 47.0'				
	48					
	49					
	50					1304 Commenced Overdrilling of Pilot hole 2 1/2" O.D. HSA 4" Mon. well install. Bent Grout 21' to grade Sugar Seal 25'-24' Bent. seal 28' x 25' Sugar Seal 29'-28' (50-70) Primary Seal 48'-29' (6-30) 4" SS Screen 47'-31' J.O. - 48'



MONITORING WELL CONSTRUCTION INFORMATION

JOB. NO. 000187
 BORING/WELL NO. 32WW01(S)
 DATE 7-24-98
 CHIEF/UNIT DWC/CME-75

1. PROTECTIVE CASING YES NO
 LOCKING YES NO

2. TYPE OF SURFACE SEAL (IF INSTALLED)
Cement

3. SOLID PIPE TYPE Stainless Steel
 SOLID PIPE LENGTH 39.0 ft.
 JOINT TYPE SLIP/GLUED/THREADED

4. TYPE OF BACKFILL Bent./cement Grout
 HOW INSTALLED TREMIE Pipe
 FROM SURFACE

5. TYPE OF LOWER SEAL (IF INSTALLED)
Bentonite (hydrated)

6. TYPE OF SECONDARY FILTER PACK
Sand (30-70)

7. SCREEN TYPE Stainless steel
 SCREEN LENGTH 10.0 ft.
 SLOT SIZE #10 machine in.
 SCREEN DIAMETER 4 in. (I.D.)

8. TYPE OF PRIMARY FILTER PACK
Primary Sand (16-30)

9. TYPE OF BACKFILL Primary Filter pack (16-30)

10. DRILLING METHOD HSA

WATER LEVEL _____ DATE _____

•ALL DEPTHS MEASURED FROM GROUND SURFACE

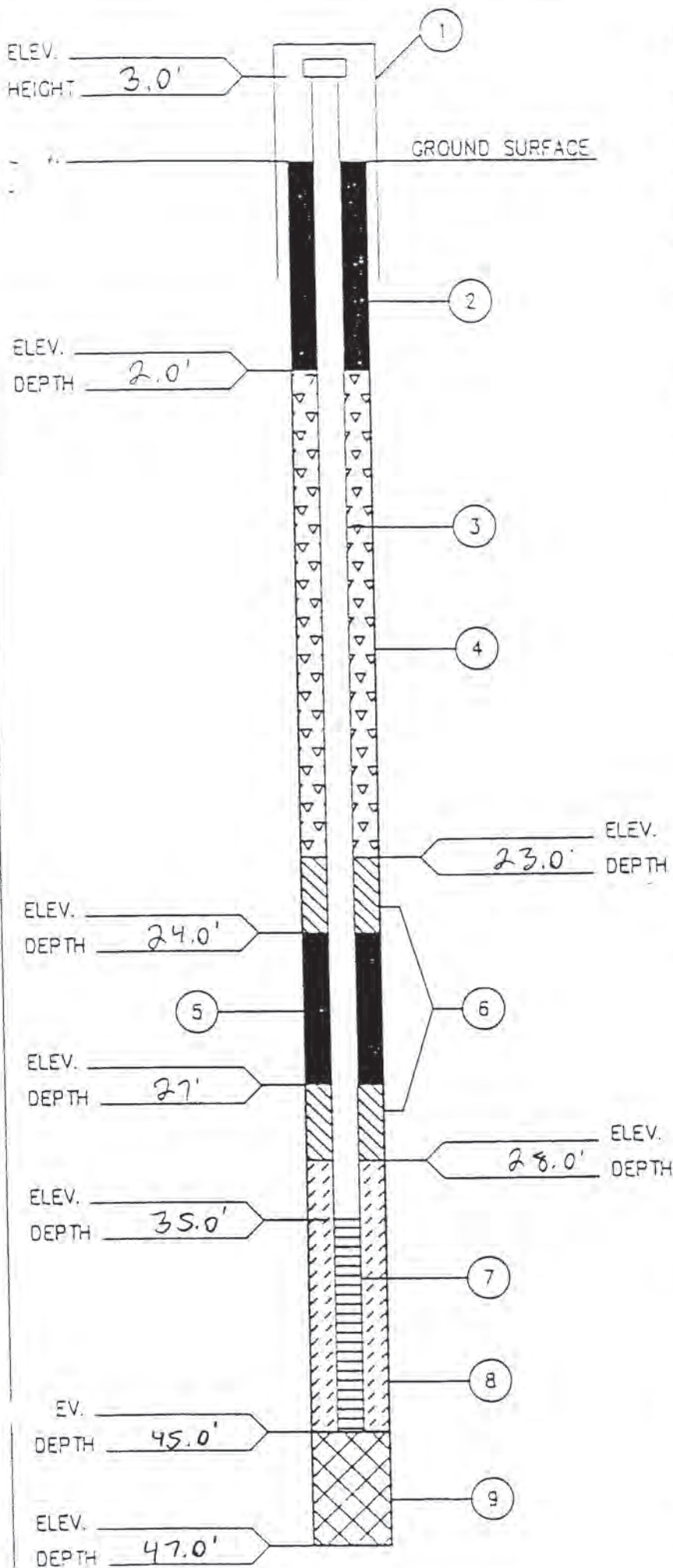
DRILLING LOG		DIVISION 16AcE-Tulsa	INSTALLATION CHAAP-Karnack TX	SHEET 1 OF 3 SHEETS
1. PROJECT CHAAP - Site 32			10. SIZE AND TYPE OF BIT Auger Bit	
2. LOCATION (Coordinates or Station) South of Well 113 - Northern side of site			11. DATUM FOR ELEVATION SHOWN (TBM or MSL)	
3. DRILLING AGENCY GPI, Inc. - Austin, TX			12. MANUFACTURER'S DESIGNATION OF DRILL CME-75/5 continuous CME sampler 3/4" ID OD Auger	
4. HOLE NO. (As shown on drawing title and file number) 32 LW "02"			13. TOTAL NO. OF OVER-BURDEN SAMPLES TAKEN DISTURBED: 10 UNDISTURBED: -	
5. NAME OF DRILLER Jose Lenceros			14. TOTAL NUMBER CORE BOXES N/A	
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.			15. ELEVATION GROUND WATER	
7. THICKNESS OF OVERBURDEN Penetrated 49 Ft			16. DATE HOLE STARTED: 7-23-98 COMPLETED: 7-24-98	
8. DEPTH DRILLED INTO ROCK N/A			17. ELEVATION TOP OF HOLE	
9. TOTAL DEPTH OF HOLE 49 Ft. Pilot hole B.E.G.			18. TOTAL CORE RECOVERY FOR BORING N/A	
			19. SIGNATURE OF INSPECTOR J. Lenceros	

11" OD Augers

ELEVATION a	DEPTH (Ft) b	U/C LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOV- ERY e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) g
	1	DL	6" of sandy med. brown and gray topsoil, some roots: leaves, damp overlying		SS #1	Commenced HSA @ 3 1/4" ID HSA @ 0840
	2	CL	- silt-clay/clayey silt, Lt. to med. tan and gray, some dk redd. sh-mottling from 3.0 to 4.0'. some fine sand throughout - <u>Dry to Damp</u> (v. stiff)	4.0'	4.0'	Hu Reading @ sampler = (102 ed PID) opp
	3					
	4					
	5		- silt-clay and clayey silt, Lt. to med. brown, gray, tan, yellow-brown, or reddish brown mottled, streaked, and stained, some apparent layering, some less plastic (clayey silts), some fine sand throughout - <u>Damp to Sl. moist</u> (stiff)		SS #2	
	6			5.0'	4.0'	Hu Reading @ sampler = opp
	7				9.0'	
	8					
	9					
	10		(trace roots/veg. @ 9.0')		SS #3	
	11		(Competent red-brown clay-shale @ 10.0')			
	12		- clayey silt, Lt. to med. gray and tan, some darker gray streaking staining, no mottling, some apparent layering with less sl. plastic thin layers. some fine sand throughout - <u>Damp (med. stiff)</u>	5.0'	9.0'	Hu Reading @ sampler = opp
	13				14.0'	
	14					
	15		- clayey silt, Lt. to dk gray, tan and orange-brown v. thin layering with laminated v. thin Lt. gray sandy silt/silty sand layers, clays: sl. plastic		SS #4	
	16		<u>Damp to Sl. moist</u> (med. stiff) No vis. water while Drilling and sampling	5.0'	14.0'	Hu Reading @ sampler = opp
	17				19.0'	
	18	SM-CL				
	19					
	20				SS #5	

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 326W "02"		
PROJECT LHAAP			INSTALLATION LHAAP - Karnack, TX			
			SHEET 2 OF 3 SHEETS			
ELEVATION a	DEPTH (ft) b	USL LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOV- ERY e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) g
	21	SM- CL	- Sandy silt, within interlayered with clayey silt, some irregular layering, med. blue gray to dark gray.	5.0'	19.0' to 24.0'	HNU Reading @ sampler oppn
	22		sl. plastic to non-plastic sandy silts, fine sandy silts, some to abundant fine sand throughout			
	23		clayey silts - Damp (No vis. H ₂ O w/d. sampling)			
	24	CL	- Silty clay/clayey silt, med. to dark gray with abundant within lighter gray silty sand/sandy silt layers.	5.0'	SS #6 24.0' to 29.0'	HNU Reading @ sampler oppn
	25		horiz. and irregular, some fine sand throughout - Dry to Damp (stiff) No vis. H ₂ O w/d.			
	26		- black shaley "lignite" coal seam - Damp			
	27	CL	- clayey silt, blue gray to med. gray, within horiz. and irregular fine sandy silt and silty sand layers - Dry to Damp (stiff)			Bottom of hole dry @ 29 ft. @ 0900 - pulled augers up 3 ft. & waiting on water till 1000 - still dry inside augers (called cliff murray - Tulsa Corp saw advised of drilling to 50 ft. to find a shallow water zone)
	28	(cont)	- silty clay, dark gray with some med. gray within sandy silt lenses - horiz. and irreg., some apparent layering, trace "lignite" fragments embedded in more stiffer clays, some fine sand throughout - sl. Damp (V. stiff) No vis. water w/d. and sampling.	5.0'	SS #7 29.0' to 34.0'	
	29		- silty clay, (Approaching) clay-shale in texture, dark gray, some within med. gray sandy silt lenses, breaks into layers, some fine sand throughout - Dry to sl. Damp (HARD) (No vis. water while drilling sampling)			HNU Reading in sampler oppn
	30		silt content increasing			
	31		- Sandy silt (sl. clayey), med. blue gray, fine grained sand, some sl. compacted clayey layering - Damp (Med. Dense to loose) No vis. water w/d. sampling	5.0'	SS #8 34.0' to 39.0'	HNU Reading in sampler oppn
	32					
	33					
	34					
	35					
	36					
	37					
	38					
	39					
	40					
	41	SM				
	42					
	43					
	44					

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 32LW '02"		
PROJECT LHAAP		INSTALLATION LHAAP - Karnack, TX		SHEET 3 OF 3 SHEETS		
ELEVATION a	DEPTH b	LEGEND c	CLASSIFICATION OF MATERIALS (Description) d	% CORE RECOVERY e	BOX OR SAMPLE NO. f	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant) Time g
	45	SM	- Sandy silt, Med. to darker blue-gray and gray, some more competent interbedded U. thin clay-shale layers, fine grained sand, sil. moist @ 47' to 48' - Damp overall (Dense to loose) (crumbles easily in hand)	S.O	55	HNU feeding in sample = oppm
	46				10	
	47				44.0	
	47				49.0	
	49					
	50		T.O. = 49.0 ft. BEG in Damp sandy silt			1035 - T.O. @ 50.0 ft. below grade - Hole dry inside augers and at base of hole. Pulled up augers 4 ft. Waiting on water yield
						1300 - water @ 47 ft. BEG in side of augers and dripping from 45' (augers)
						1405 overdrilled hole with 11" HSA augers (plugged) to 47.0 ft. To install 4" well
						22' Grout to surface
						23' Top of sugar sand
						24' Top of Bentonite seal
						27' Top of sugar sand
						28' Top of Primary Filter pack
						35' Top of 4" 55 # 10 slot screen
						45' Bottom of 4" 55 # 10 slot screen
						47' Bottom of 11" hole
						47' Bottom of Primary Filter pack
						7-24-98
						*0715 - Water Level @ 33.9 ft. below grade



MONITORING WELL CONSTRUCTION INFORMATION

JOB. NO. 000187

BORING/WELL NO. 32WW02 (S)

DATE 7-24-98

CHIEF/UNIT DWC / CME-75

1. PROTECTIVE CASING YES NO
 LOCKING YES NO

2. TYPE OF SURFACE SEAL (IF INSTALLED)
Cement

3. SOLID PIPE TYPE Stainless steel
 SOLID PIPE LENGTH 38.0' ft.
 JOINT TYPE SLIP/GLUED/THREADED

4. TYPE OF BACKFILL Bent./cement Grout
 HOW INSTALLED - TREMIE Pipe
 FROM SURFACE

5. TYPE OF LOWER SEAL (IF INSTALLED)
Bentonite (hydrated)

6. TYPE OF SECONDARY FILTER PACK
Sand (30-70)

7. SCREEN TYPE stai
 SCREEN LENGTH 10.0 ft.
 SLOT SIZE #10 machine in.
 SCREEN DIAMETER 4 in. (C.I.D.)

8. TYPE OF PRIMARY FILTER PACK
Primary Sand (16-30)

9. TYPE OF BACKFILL Primary Sand

10. DRILLING METHOD HSA

WATER LEVEL _____ DATE _____

•ALL DEPTHS MEASURED FROM GROUND SURFACE

Hole No. 35AW04

DRILLING LOG	DIVISION LISACE-Tulsa	INSTALLATION LHAAP Karmack, TX	SHEET / OF 2 SHEETS
PROJECT LHAAP		10. SIZE AND TYPE OF BIT Auger B-7	
LOCATION (Coordinates or Station)		11. DATUM FOR ELEVATION SHOWN (TBM = MSL)	
DRILLING AGENCY GPI, Inc. Muskogee, TX		12. MANUFACTURER'S DESIGNATION OF DRILL AME 75 / 3 1/2" ID HSA / 11" G DMSA	
HOLE NO. (As shown on drawing title and title number) 35AW04		13. TOTAL NO. OF OVER-BURDEN SAMPLES TAKEN 7	
NAME OF DRILLER Jose Landros		14. TOTAL NUMBER CORE BOXES N/A	
DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.		15. ELEVATION GROUND WATER	
THICKNESS OF OVERBURDEN Penetrated 24 ft.		16. DATE HOLE STARTED: 8-23-98 COMPLETED: 8-23-98	
DEPTH DRILLED INTO ROCK N/A		17. ELEVATION TOP OF HOLE	
TOTAL DEPTH OF HOLE 24.0 ft. Below grade		18. TOTAL CORE RECOVERY FOR BORING N/A	
		19. SIGNATURE OF INSPECTOR Steve S. ... / Dave C. ...	

ELEVATION	DEPTH (ft)	LOG LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
1000	0.0	SM	Sandy silt - Lt. brown, tan, and Lt. gray. Fine grain and sand, poorly sorted, some roots - Dry to St. Damp (loose)	4/4	SS #1	1000 Commenced HSA 35AW04 Z 7' 0.0. HSA
	4.0					HNA Reading in sampler = 9ppm
	3.0	CL	Silty clay - med. gray and dark reddish-brown (mottled) trace roots and fine sand. v. plastic - Damp (v. stiff)		SS #2	
	5.0			5/5	4.0 to 9.0	HNA Reading in sampler = 6ppm
	6.0		Silty clay - Lt. gray, Lt. greenish gray, tan, yellowish-orange and tan (mottled) and med. brown mottled, streaked, and stained, trace to some fine sand, some apparent v. thin layering. trace roots - Damp (stiff to v. stiff)		SS #3	
	9.0			5/5	9.0 to 14.0	HNA Reading in sampler = 9ppm
	10.0		Silty clay - Lt. gray, Lt. greenish-gray, yellowish-orange and tan (mottled) abundant black v. thin root-like structures throughout, some fine sand. plastic - Damp to Moist @ 13' (stiff)			
	11.0		v. Moisture content inc. (sticking to sampler)			
	14.0	CC CL	Sandy clay - Lt. gray and greenish gray (mottled) layers in silty clay, fine grained sand - wet/saturated (soft)		SS #4	Water @ 14' W.A.
	17.0			5/5	14.0 to 17.0	HNA Reading in sampler = 8ppm
	17.0	SC	Clayey sand - Lt. brown, gray, tan (mottled). some silty, fine grained sand, trace med. gravel - wet/saturated (med. dense to loose)			
	18.0					
	19.0	CL	Silt/silty sand - med. gray, tan, and brown, fine to med. grained, some coarse, subangular sub rounded, poorly sorted, trace clay zones - wet/saturated (loose)		SS #5	
	20.0	SP		1/1	17' to 20'	HNA Reading in sampler = 10ppm

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE		Hole No. 3SAWWS04		
PROJECT			INSTALLATION		SHEET 2 OF 2 SHEETS	
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
a	b	c	d	e	f	g
	21	SP	- Sand-Mud brown and reddish-brown, fine to coarse grains - trace gravel, poorly sorted - Wet/Saturated	1 1/1	SS#6 20' to 21'	Hnu Racking in sample = 11 Apr
	22	SP-GP SD	- Sand and gravel - Dark "rust" to brown fine to coarse sanding gravel; some Cobble - Wet-Saturated	3 1/3	SS#7 21' to 24'	
	23	CL	I Silty clay - Lt grey, greenish grey, and tan, M-A-L, trace to some fine sand - Damp (U.S.I.H)			
	24					
			T.D. @ 24.0 ft below grade in Silty clay - Drop to moist			Overdrilled pilot hole with 1 1/2" O.D. HSA Plugger augers to T.D. of 27 ft. below grade. Complete details for man. installation of 3SAWWS04 Continent groot 7' to 9' (Bent Seal) 10' to 7' Sugar sand (30-20) 11' to 10' Primary filterpack 15' to 11' (16-30) 4" 10' #10 slot screen 23' to 13' T.D. -23.0' below grade

HOLE NO. 35ANW07

DRILLING LOG		DIVISION FEDERAL	INSTALLATION Longhorn AAP	SHEET 1 OF 3 SHEETS
1. PROJECT Longhorn MARC		10. SIZE AND TYPE OF BIT HSA 10"		
2. LOCATION (Coordinates or Station) BARNACK, TX		11. DATUM FOR ELEVATION SHOWN (TBM or MSL) MSL		
3. DRILLING AGENCY ETEL		12. MANUFACTURER'S DESIGNATION OF DRILL FOREMOST 5500		
4. HOLE NO. (As shown on drawing title and file number) 35ANW07		13. OVERBURDEN SAMPLES	DISTURBED <input checked="" type="checkbox"/>	UNDISTURBED
5. NAME OF DRILLER Billy Ragon		14. TOTAL NUMBER CORE BOXES NR		
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED 0° DEG. FROM VERT.		15. ELEVATION GROUND WATER NR		
7. THICKNESS OF OVERBURDEN —		16. DATE HOLE STARTED 11/13/08 COMPLETED 11/14/08		
8. DEPTH DRILLED INTO ROCK NA		17. ELEVATION TOP OF HOLE NR		
9. TOTAL DEPTH OF HOLE 30'		18. TOTAL CORE RECOVERY FOR BORING NR %		
Allen Wilmore				INSPECTOR

ELEVATION PID	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVER- ERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
a	b	c	d	e	f	g
0.0			TOP SOIL			
0.0			CLAY, SILTY, LOW TO NO PLASTICITY Brown, SOFT, NO ODOR	65%		
0.0			-BECOMES SANDY, STIFF, MOIST			
0.0	5			100%		
0.0			SAND, SILTY, DENSE, BROWN, IRON STAINING MOIST, NO ODOR TO GRAY	60%		
0.0	10					
0.0						
0.0						

DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE NA		Hole No. 35AWW07		
PROJECT Loughorn MARC			INSTALLATION Loughorn AAP		SHEET 2 OF 3 SHEETS	
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
0.0			SAME AS ABOVE			
0.0	15	CL	CLAY, SILTY, BROWN, SLIGHT PLASTICITY, MOIST, SOFT, NO ODOR	70%		
0.0			SAND, SILTY, MOIST, GRAY, IRON STAINING, MOTTLED, NO ODOR, MODERATELY DENSE			
0.0	20		- MORE IRON, MORE ORGANICS	100%		
0.0		CL	CLAY, SILTY, STIFF, MOIST, HIGH PLASTICITY, NO ODOR			
0.0	25					
0.0		MLS	SILT, SAND, SILTY, MOIST, IRON STAINING, GRAY, DENSE	100%		
0.0				100%		

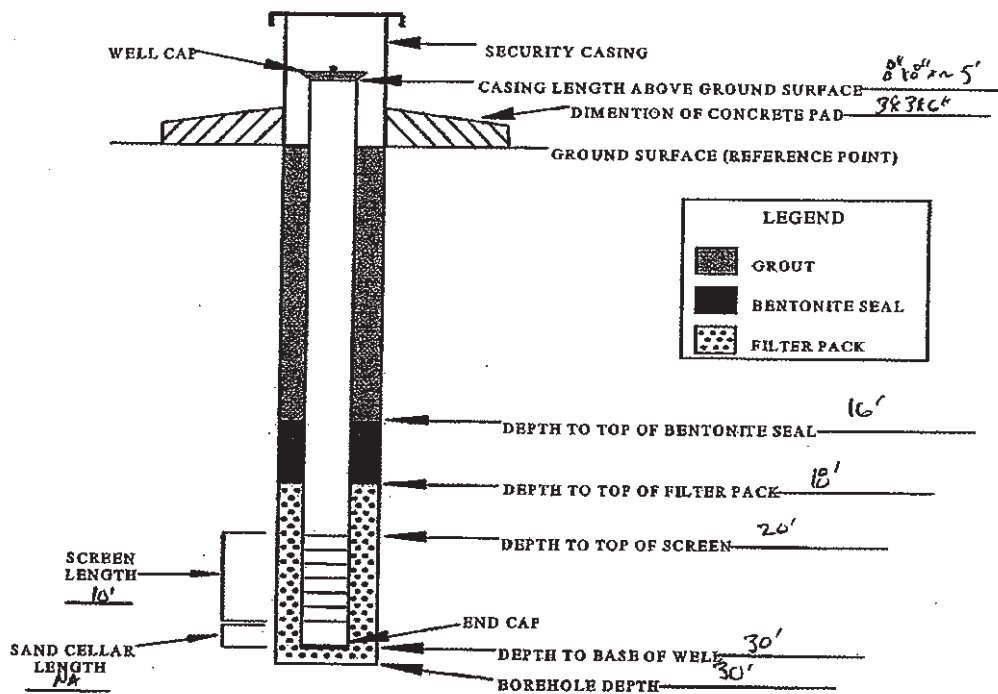
DRILLING LOG (Cont Sheet)		ELEVATION TOP OF HOLE NA		Hole No. 35AWW07		
PROJECT Longhorn MARC			INSTALLATION Longhorn AAP		SHEET 3 OF 3 SHEETS	
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
b-2	30		- BECOMES MOAC CLAY END OF BORING @ 30' bgs	100%		

35AWW07

WELL COMPLETION FORM (Stickup or Above Grade Completion Well)

FIELD REPRESENTATIVE: Allen Willmore TYPE OF FILTER PACK: silica sand
 DRILLING CONTRACTOR: ETC GRADIATION: 26/40
 AMOUNT OF FILTER PACK USED: 12-bgs (50 lbs)
 DRILLING TECHNIQUE: HSA TYPE OF BENTONITE: pellets
 AUGER SIZE AND TYPE: 10" AMOUNT BENTONITE USED: 1-50lb bucket
 BOREHOLE IDENTIFICATION: 35AWW07 TYPE OF CEMENT: portland-cement
 BOREHOLE DIAMETER: 10" AMOUNT CEMENT USED: N/A
 WELL IDENTIFICATION: 35AWW07 GROUT MATERIALS USED: bentonite slurry
 WELL CONSTRUCTION START DATE: 11/13/08 DIMENSIONS OF SECURITY CASING: 8" x 8" x 3'
 WELL CONSTRUCTION COMPLETE DATE: 11/14/08
 SCREEN MATERIAL: sch. 40 PVC TYPE OF WELL CAP: plugging cap
 SCREEN DIAMETER: 4" TYPE OF END CAP: sch. 40 PVC
 STRATUM-SCREENED INTERVAL (FT): 10'
 CASING MATERIAL: sch. 40 PVC COMMENTS:
 CASING DIAMETER: 4"

SPECIAL CONDITIONS
(describe and draw)



LEGEND	
	GROUT
	BENTONITE SEAL
	FILTER PACK

NOT TO SCALE
 INSTALLED BY: Billy Ragon INSTALLATION OBSERVED BY: Allen Willmore
 DISCREPANCIES: none

STATE OF TEXAS WELL REPORT for Tracking #162467

Owner: Longhorn Army Ammunition Plant	Owner Well #: 35AWW07
Address: Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Grid #: 35-23-6
Well Location: Hwy 143 @ Spur 449, LHAAP Karnack , TX 75661	Latitude: 32° 41' 10" N
Well County: Harrison	Longitude: 094° 09' 06" W
Elevation: No Data	GPS Brand Used: Garmin e-trex
<hr/>	
Type of Work: New Well	Proposed Use: Monitor

Drilling Date: Started: **11/13/2008**
 Completed: **11/14/2008**

Diameter of Hole: Diameter: **10.25 in From Surface To 30 ft**

Drilling Method: **Hollow Stem Auger**

Borehole
Completion: Gravel Packed From: **18 ft to 30 ft**
 Gravel Pack Size: **20/40**

Annular Seal Data: 1st Interval: **From 16 ft to 18 ft with 1 Bentonite (#sacks and material)**
 2nd Interval: **From 0 ft to 16 ft with 7 Cement (#sacks and material)**
 3rd Interval: **No Data**
 Method Used: **Tremmi pipe**
 Cemented By: **Driller**
 Distance to Septic Field or other Concentrated Contamination: **No Data**
 Distance to Property Line: **No Data**
 Method of Verification: **No Data**
 Approved by Variance: **No Data**

Surface
Completion: **Surface Sleeve Installed**

Water Level: Static level: **No Data**
 Artesian flow: **No Data**

Packers:

Plugging Info: Casing or Cement/Bentonite left in well: **No Data**

Type Of Pump: **No Data**

Well Tests: **No Data**

Water Quality: Type of Water: **No Data**
 Depth of Strata: **No Data**
 Chemical Analysis Made: **No Data**
 Did the driller knowingly penetrate any strata which contained undesirable constituents: **No**

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the log(s) being returned for completion and resubmittal.

Company
Information: **ETTL Engineers & Consultants Inc.**
 1717 E. Erwin

Tyler , TX 75702

Driller License Number: **54683**

Licensed Well Driller Signature: **Wilburn Ragon, Jr.**

Registered Driller Apprentice Signature: **No Data**

Apprentice Registration Number: **No Data**

Comments: **No Data**

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Please include the report's Tracking number (Tracking #162467) on your written request.

Texas Department of Licensing & Regulation
P.O. Box 12157
Austin, TX 78711
(512) 463-7880

DESC. & COLOR OF FORMATION MATERIAL

From (ft) To (ft) Description

0-8.5 Silty clay - brown

8.5-14.5 Silty sand - brown to gray

14.5-17 Silty clay - brown

17-22 Silty sand - gray

22-25.5 Silty clay

25.5-30 Silty sand - gray

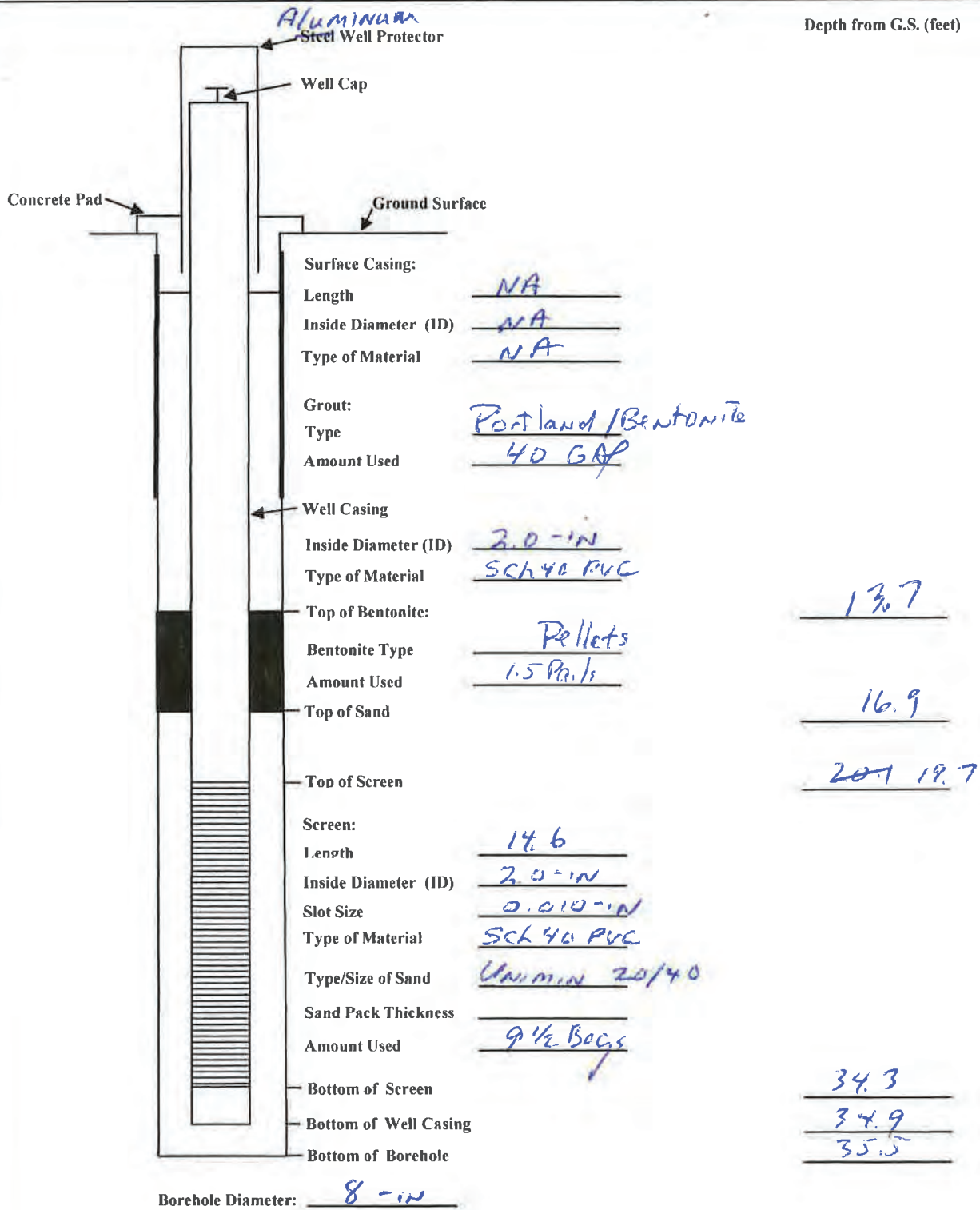
CASING, BLANK PIPE & WELL SCREEN DATA

Dia.	New/Used	Type	Setting From/To
4	New	PVC Sch. 40	0 - 20
4	New	PVC Sch. 40 - slotted	20 - 30 0.010"

AZCOM

Client: <u>USAGE</u>	WELL ID: <u>35AWW11</u>
Project Number: <u>60256135</u>	Date Installed: <u>8-23-13</u>
Site Location: <u>Site 58</u>	Borehole Diameter: <u>8-in</u>
Well Location: _____ Coords: _____	Contractor: <u>FUGRO</u>
Method: <u>HSA</u>	

MONITORING WELL CONSTRUCTION DETAIL



Comments: _____

Installation Observed By: M. Hartford

STATE OF TEXAS WELL REPORT for Tracking #352610

Owner: Longhorn AAP- USACE	Owner Well #: 35AWW11
Address: Highway 43 Karnack , TX 75661	Grid #: 35-23-6
Well Location: Highway 43 Karnack , TX 75661	Latitude: 32° 41' 08" N
Well County: Harrison	Longitude: 094° 09' 17" W
Elevation: 218 ft.	GPS Brand Used: No Data
<hr/>	
Type of Work: New Well	Proposed Use: Monitor

Drilling Date: Started: **8/23/2013**
Completed: **8/23/2013**

Diameter of Hole: Diameter: **8 in From Surface To 35.5 ft**

Drilling Method: **Hollow Stem Auger**

Borehole Completion: Gravel Packed From: **35.5 ft to 16.9 ft**
Gravel Pack Size: **20 / 40**

Annular Seal Data: 1st Interval: **From 35.5 ft to 16.9 ft with 9.5 (#sacks and material)**
2nd Interval: **From 16.9 ft to 13.7 ft with 1.5 (#sacks and material)**
3rd Interval: **From 13.7 ft to 0 ft with 2 Portland (#sacks and material)**
Method Used: **Tremie**
Cemented By: **Donald Edwards**
Distance to Septic Field or other Concentrated Contamination: **No Data**
Distance to Property Line: **No Data**
Method of Verification: **No Data**
Approved by Variance: **No Data**

Surface Completion: **Surface Slab Installed**

Water Level: Static level: **No Data**
Artesian flow: **No Data**

Packers: **No Data**

Plugging Info: Casing or Cement/Bentonite left in well: **No Data**

Type Of Pump: **No Data**

Well Tests: **No Data**

Water Quality: Type of Water: **No Data**
Depth of Strata: **No Data**
Chemical Analysis Made: **No Data**
Did the driller knowingly penetrate any strata which contained undesirable constituents: **No Data**

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the log(s) being returned for completion and resubmittal.

Company Information: **Fugro
6105 Rookin St
Houston , TX 77074**

Driller License Number: **56013**

Licensed Well Driller Signature: **Donald Edwards**

Registered Driller Apprentice Signature: **No Data**


Apprentice Registration Number: **No Data**

Comments: **No Data**

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Texas Department of Licensing & Regulation
P.O. Box 12157
Austin, TX 78711
(512) 463-7880 

DESC. & COLOR OF FORMATION MATERIAL

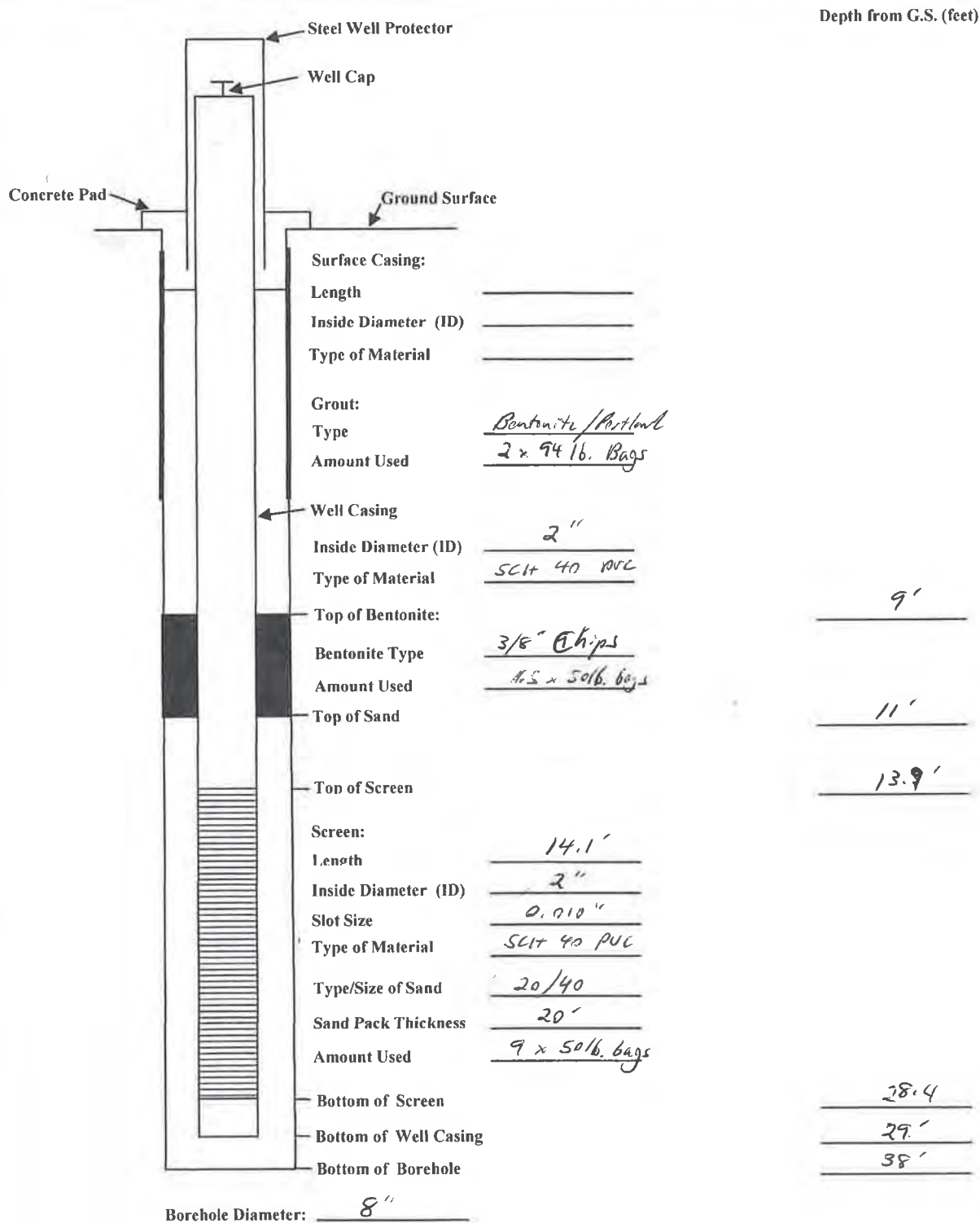
From (ft) To (ft) Description
Gray / Brown clay with layers of sand

CASING, BLANK PIPE & WELL SCREEN DATA

Dia.	New/Used	Type	Setting From/To
2in	new	pvc screen	34.9ft to 19.7ft .010 slot
2in	new	pvc casing	19.7ft to 0

AZCOM	Client: <i>USACE</i>	WELL ID:
	Project Number: <i>60258135</i>	<i>90 3SAWW19</i>
	Site Location: <i>Site 58</i>	Date Installed: <i>9/19/13</i>
	Well Location: _____ Coords: _____	Borehole Diameter: <i>8"</i>
	Method: <i>HSA</i>	Contractor: <i>Fygro</i>

MONITORING WELL CONSTRUCTION DETAIL



Comments:

*Back-filled w/ Bentonite chips from 28' to 31' Bgs
Filter sand from 31' to 11' Bgs*

Installation Observed By:

Mr. Law

STATE OF TEXAS WELL REPORT for Tracking #352655

Owner: Longhorn AAP-USACE	Owner Well #: 35AWW19
Address: Highway 43 Houston , TX 75661	Grid #: 35-23-6
Well Location: Highway 43 Karnack , TX 75661	Latitude: 32° 41' 04" N
Well County: Harrison	Longitude: 094° 09' 18" W
Elevation: 217 ft.	GPS Brand Used: No Data
<hr/>	
Type of Work: New Well	Proposed Use: Monitor

Drilling Date: Started: **9/19/2013**
Completed: **9/19/2013**

Diameter of Hole: Diameter: **8 in From Surface To 38 ft**

Drilling Method: **Hollow Stem Auger**

Borehole Completion: Gravel Packed From: **38 ft to 11 ft**
Gravel Pack Size: **20/40**

Annular Seal Data: 1st Interval: **From 38 ft to 11 ft with 9 sand (#sacks and material)**
2nd Interval: **From 11 ft to 9 ft with 1 bentonite (#sacks and material)**
3rd Interval: **From 9 ft to 0 ft with 2 Portland (#sacks and material)**
Method Used: **Tremie**
Cemented By: **Donald Edwards**
Distance to Septic Field or other Concentrated Contamination: **No Data**
Distance to Property Line: **No Data**
Method of Verification: **No Data**
Approved by Variance: **No Data**

Surface Completion: **Surface Slab Installed**

Water Level: Static level: **No Data**
Artesian flow: **No Data**

Packers: **No Data**

Plugging Info: Casing or Cement/Bentonite left in well: **No Data**

Type Of Pump: **No Data**

Well Tests: **No Data**

Water Quality: Type of Water: **No Data**
Depth of Strata: **No Data**
Chemical Analysis Made: **No Data**
Did the driller knowingly penetrate any strata which contained undesirable constituents: **No Data**

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the log(s) being returned for completion and resubmittal.

Company Information: **Fugro
6105 Rookin St
Houston , TX 77074**

Driller License Number: **56013**

Licensed Well Driller Signature: **Donald Edwards**

Registered Driller Apprentice Signature: **No Data**


Apprentice Registration Number: **No Data**

Comments: **No Data**

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Texas Department of Licensing & Regulation
P.O. Box 12157
Austin, TX 78711
(512) 463-7880 

DESC. & COLOR OF FORMATION MATERIAL

From (ft) To (ft) Description
0 to 38ft Gray / Brown clay with layers of sand

CASING, BLANK PIPE & WELL SCREEN DATA

Dia.	New/Used	Type	Setting From/To
2in	new	pvc screen	28.4ft to 13.9ft .010 slot
2in	new	pvc casing	13.9 to 0



BORING NUMBER 35AWW24

CLIENT AFCEC **PROJECT NAME** Longhorn Army Ammunition Plant
PROJECT NUMBER NWO1312.0150.012.0001.02 **PROJECT LOCATION** 15600 FM 134 Karnack, Texas 75661
DATE STARTED 2/20/18 **COMPLETED** 2/20/18 **GROUND ELEVATION** _____ **HOLE SIZE** 6
DRILLING CONTRACTOR Walker Hill **GROUND WATER LEVELS:**
DRILLING METHOD Geoprobe - Hollow Stem Auger **AT TIME OF DRILLING** ---
LOGGED BY J. Cook **AT END OF DRILLING** ---
NOTES _____ **AFTER DRILLING** ---

BHATE ENVIRONMENTAL ASSOCIATES LONGHORN AAP.GPJ GINT US LAB.GDT 4/17/18

DEPTH (ft)	SAMPLE TYPE NUMBER	RECOVERY %	U.S.C.S.	GRAPHIC LOG	MATERIAL DESCRIPTION	PID (ppm)
0						
		100	CL-ML		Dark brown silty clay/ organics. High plasticity. Wet. 7.5 YR 2.5/2	0
					2.0	
		100	CL		Light brown high plasticity clay. High plasticity. Moist. 7.5 YR 4/4	0
					4.0	
5		100	CL		Grey/red brown clay. Medium plasticity. Dry. 7.5 YR 4/1	0
					6.0	
		100	CL-ML		Light grey silty clay. Low plasticity. Dry. 7.5 YR 5/1	0
		100	CL-ML			0
10					10.0	
		100	CL		Light grey/brown silty clay. Dry. 7.5 YR 6/1	0
		100	CL			0
					14.0	
15		100	CL-ML		Light brown silty clay. Medium plasticity. Dry. 7.5 YR 5/6	0
		100	CL-ML			0
		100	CL-ML			0
20					20.0	
		100			Light brown sandy clay. Dry. 7.5 YR 5/6	0
					22.0	
		100	CL-ML		Brown high plasticity silty clay. High plasticity. Moist. 7.5 YR 4/3	0
					24.0	
25		100	CL-ML		Light brown silty clay. High plasticity. Wet at 26'. 7.5 YR 4/4	0
					26.0	
		100			Light grey/brown silty clay. High plasticity. Wet. 7.5 YR 5/1	0
		100	CL-ML			0
30						
		100				0
					32.0	
					Bottom of hole at 32.1 feet.	

STATE OF TEXAS WELL REPORT for Tracking #472959

Owner: USACE	Owner Well #: 35AWW24
Address: 1645 S. 101st. East Ave. Tulsa, OK 74128	Grid #: 35-23-6
Well Location: 15600 FM 134 Karnack, TX 75661	Latitude: 32° 41' 02.9" N
Well County: Harrison	Longitude: 094° 09' 18.16" W
	Elevation: No Data
Type of Work: New Well Proposed Use: Monitor	

Drilling Start Date: **2/19/2018** Drilling End Date: **2/23/2018**

	Diameter (in.)	Top Depth (ft.)	Bottom Depth (ft.)
Borehole:	8	0	32

Drilling Method: **Hollow Stem Auger**

Borehole Completion: **Filter Packed**

	Top Depth (ft.)	Bottom Depth (ft.)	Filter Material	Size
Filter Pack Intervals:	20	32	Sand	20/40

	Top Depth (ft.)	Bottom Depth (ft.)	Description (number of sacks & material)
Annular Seal Data:	0	15	Cement 5 Bags/Sacks
	15	20	Bentonite 0 Bags/Sacks

Seal Method: **Tremie**

Sealed By: **Driller**

Distance to Property Line (ft.): **No Data**

Distance to Septic Field or other concentrated contamination (ft.): **No Data**

Distance to Septic Tank (ft.): **No Data**

Method of Verification: **No Data**

Surface Completion: **Surface Slab Installed**

Surface Completion by Driller

Water Level: 26.98 ft. below land surface on 2018-02-23	Measurement Method: Electric Line
Packers: No Data	
Type of Pump: No Data	
Well Tests: No Test Data Specified	

Water Quality:	<i>Strata Depth (ft.)</i>	<i>Water Type</i>
	No Data	No Data

Chemical Analysis Made: **No**Did the driller knowingly penetrate any strata which contained injurious constituents?: **No**

The driller did certify that while drilling, deepening or otherwise altering the above described well, injurious water or constituents was encountered and the landowner or person having the well drilled was informed that such well must be completed or plugged in such a manner as to avoid injury or pollution.

Certification Data: The driller certified that the driller drilled this well (or the well was drilled under the driller's direct supervision) and that each and all of the statements herein are true and correct. The driller understood that failure to complete the required items will result in the report(s) being returned for completion and resubmittal.

Company Information: **Walker-Hill Environmental, Inc.**

**PO 1147
Foxworth, MS 39483**

Driller Name: **Gary P Hill** License Number: **58141**Apprentice Name: **Eric Meitzler**Comments: **No Data**

Lithology:
DESCRIPTION & COLOR OF FORMATION MATERIAL

Casing:
BLANK PIPE & WELL SCREEN DATA

<i>Top (ft.)</i>	<i>Bottom (ft.)</i>	<i>Description</i>
0	2	CL-ML, Dark Brown, Silty Clay
2	4	CL, Light brown, high plasticity clay
4	8	CL grey,reddish brown, Clay
8	12	CL-ML, Light grey, silty clay
12	18	CL-ML, Light grey/brown, silty clay
18	20	CL-ML, Light Brown, Silty Clay
20	22	CLS, Light Brown, Sandy Clay
22	24	CL-ML, Brown, High plasticity, Silty clay
24	28	CL-ML, Light Brown, silty clay
28	32	CL-ML, Light Grey/Brown, Silty Clay

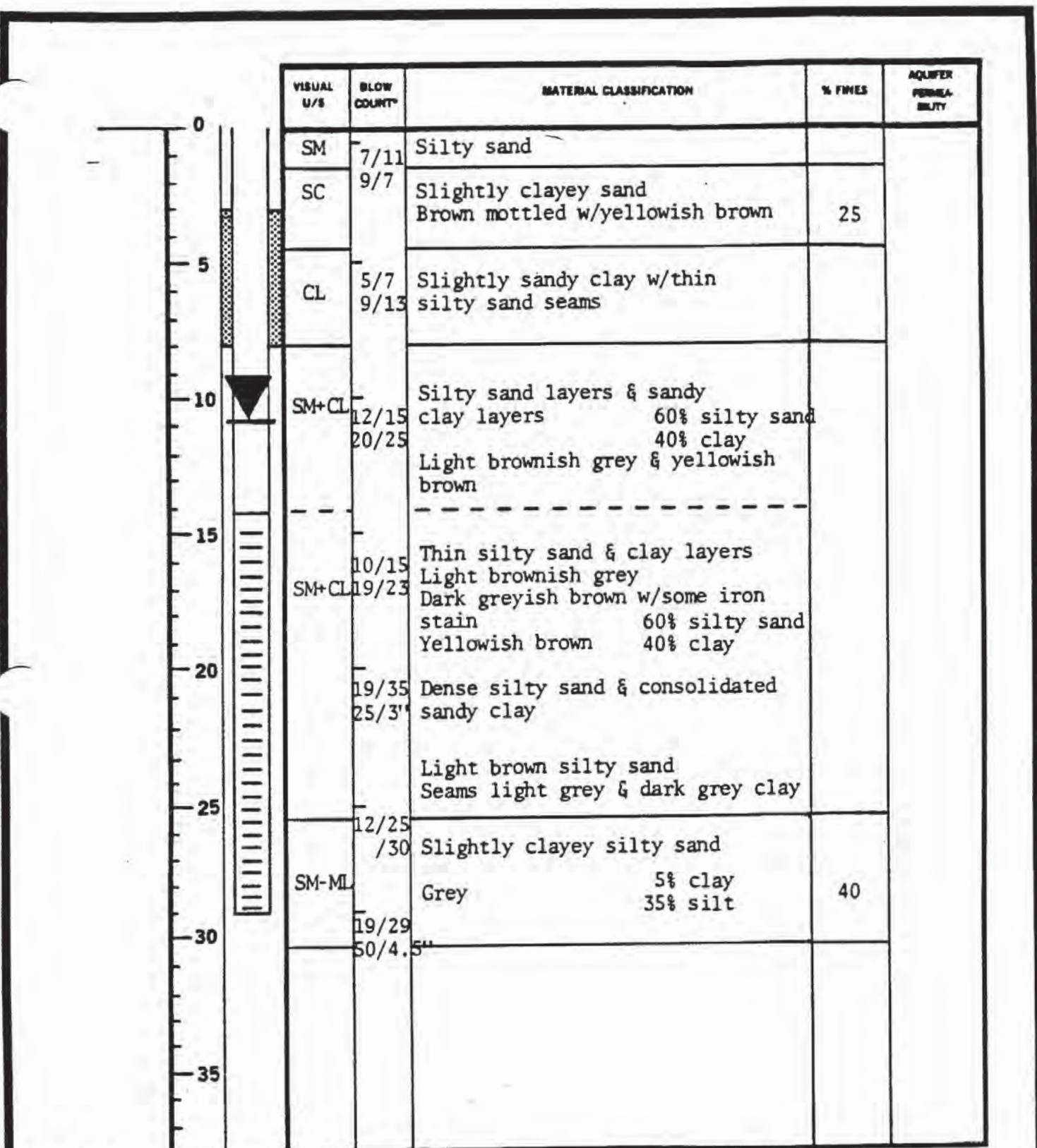
<i>Dia (in.)</i>	<i>Type</i>	<i>Material</i>	<i>Sch./Gage</i>	<i>Top (ft.)</i>	<i>Bottom (ft.)</i>
2	Riser	New Plastic (PVC)	40	0	22
2	Screen	New Plastic (PVC)	40 0.010	22	32

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Texas Department of Licensing and Regulation
P.O. Box 12157
Austin, TX 78711
(512) 334-5540



*BLOWS PER 6 INCH USING
18, 24, or 36 SPLIT SPOON

VERTICAL SCALE 1" = 5.0'

WELL 113 DATE DRILLED 6/29/82
STUDY AREA Former TNT Waste Disp. Plant

ELEVATION TOP OF STEEL CASING = 214.87 ft
ELEVATION TOP OF PVC RISER = 214.53 ft
ELEVATION OF GROUND SURFACE = 211.97 ft

SOURCE: ENVIRONMENTAL PROTECTION SYSTEMS, INC. - 1983

CONTAMINATION SURVEY
LONGHORN ARMY AMMUNITION PLANT
MARSHALL, TEXAS

U. S. ARMY TOXIC AND HAZARDOUS
MATERIALS AGENCY
ABERDEEN PROVING GROUND, MARYLAND

HOLE NO. LHS-MW1

DRILLING LOG		DIVISION	INSTALLATION	SHEET		
PROJECT: LHAAP-WASTE SUMPS		SOUTHWEST	LHAAP	1 of 1 SHEETS		
2. LOCATION (Coordinates or Station) 6959157.90 3306087.10		10. SIZE AND TYPE OF BIT 8" AUGER				
3. DRILLING AGENCY TULSA DISTRICT COE		11. DATUM FOR ELEVATION SHOWN (TBM or MSL) MSL				
4. HOLE NO. (As shown on drawing file and file number) LHS-MW1		12. MANUFACTURER'S DESIGNATION OF DRILL CME 75				
5. NAME OF DRILLER TOM BEAVERS		13. OVERBURDEN SAMPLES DISTURBED 7 UNDISTURBED 0				
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.		14. TOTAL NUMBER CORE BOXES 0				
7. THICKNESS OF OVERBURDEN 16.0		15. ELEVATION GROUND WATER SEE REMARKS				
8. DEPTH DRILLED INTO ROCK 0.0		16. DATE HOLE STARTED 08/19/1994 COMPLETED 08/19/1994				
9. TOTAL DEPTH OF HOLE 16.0		17. ELEVATION TOP OF HOLE 211.2				
		18. TOTAL CORE RECOVERY FOR BORING 0.0 %				
MERLIN DEAN						
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
	2		LEAN CLAY (CL) (0.0 - 4.5) WITH SAND, LIGHT YELLOWISH BROWN, MOIST.		J-1	WATER ENCOUNTERED @ 11.7'
	4					SAMPLE TYPE ZONE AUGER 0.0- 10.0 SPLITSPOON 10.0- 15.0
206.7			FAT CLAY (CH) (4.5 - 11.7) WITH SAND, LIGHT BROWNISH GRAY MOIST.		J-2	SAMPLE DEPTH J-1 0.0- 4.5 J-2 4.5- 7.2 J-3 7.2- 11.7 J-4 11.7- 12.7 J-5 13.0- 14.0 J-6 14.0- 15.0 J-7 15.0- 15.0
	6				J-3	
	8				J-4	
	10				J-5	
199.5			FAT CLAY (CH) (11.7 - 16.0) DARK GRAY TO OLIVE BROWN, MOIST, 12.8-14.0 VERY MOIST.		J-6	
	12				J-7	
	14					
195.2						
	16					
	18					
	20					

PROJECT
LHAAP-WASTE SUMPS

HOLE NO.
LHS-MW1

WELL NO. LHS-MW1

MONITORING WELL SHEET

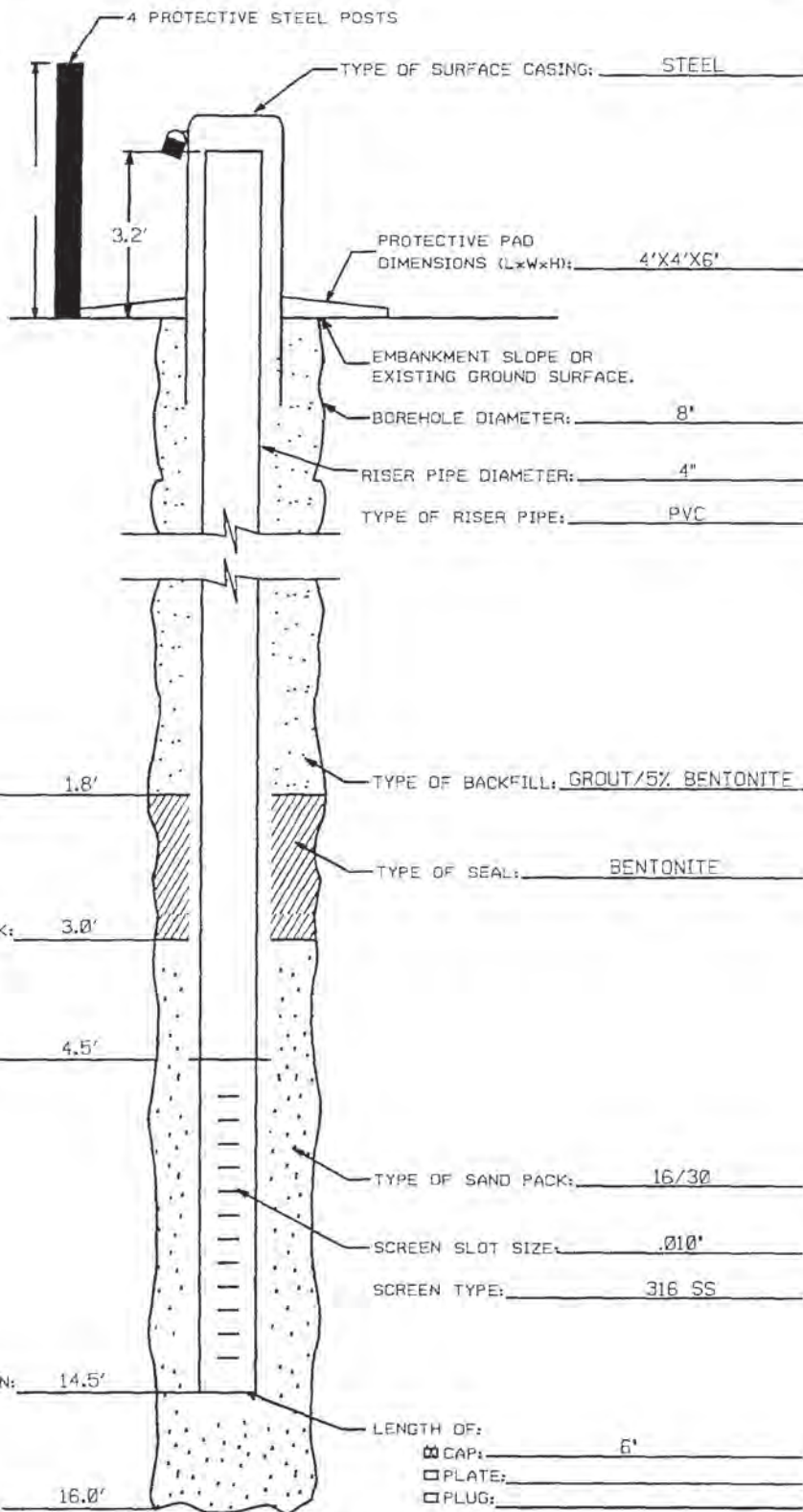
PROJECT & INSTALLATION:
LONGHORN ARMY AMMUNITION PLANT - SUMPS


DATE STARTED:

8-19-1994

DATE COMPLETED:

8-19-1994




 GROUNDWATER DEPTH: 11.7'
 GROUNDWATER DATE: 8-19-1994

INSPECTOR:

MERLIN DEAN

HOLE NO. LHS-MW2

DRILLING LOG		DIVISION	INSTALLATION	SHEET		
1. PROJECT		SOUTHWEST	LHAAP	1		
LHAAP-WASTE SUMPS				OF 1 SHEETS		
2. LOCATION (Coordinates or Station)		3305704.50	16. SIZE AND TYPE OF BIT	8" AUGER		
8959135.90			11. DATUM FOR ELEVATION SHOWN (TBM or MSL)	MSL		
3. DRILLING AGENCY		TULSA DISTRICT COE	12. MANUFACTURER'S DESIGNATION OF DRILL			
			FALLING 1500			
4. HOLE NO. (As shown on drawing title and file number)		LHS-MW2	13. OVERBURDEN SAMPLES	DISTURBED 0 UNDISTURBED 11		
5. NAME OF DRILLER		TOM BEAVERS	14. TOTAL NUMBER CORE BOXES	0		
6. DIRECTION OF HOLE		<input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.	15. ELEVATION GROUND WATER	NOT DETERMINED		
7. THICKNESS OF OVERBURDEN		16.0	16. DATE HOLE	STARTED 05/18/1994 COMPLETED 05/20/1994		
8. DEPTH DRILLED INTO ROCK		0.0	17. ELEVATION TOP OF HOLE	211.7		
9. TOTAL DEPTH OF HOLE		16.0	18. TOTAL CORE RECOVERY FOR BORING	0.0 %		
MERLIN DEAN						
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
211.2			LEAN CLAY (CL) (0.0 - 1.5) SANDY, BROWN, MOIST, VERY STIFF NUMEROUS ROOTS IN TOP, FEW SCATTERED GRAVELS TO 3/4", MORE SAND IN BOTTOM 0.3'.		ST-1	SAMPLE TYPE ZONE SHELBY 0.0- 16.0
208.7	2		LEAN CLAY (CL) (1.5 - 3.0) GRAYISH BROWN, MOIST, STIFF TO VERY STIFF, VERY FRIABLE, VERY SANDY IN TOP 0.4'.		ST-2	SAMPLE DEPTH ST-1 0.0- 1.5 ST-2 1.5- 3.0 ST-3 3.0- 4.5 ST-4 4.5- 6.0 ST-5 6.0- 7.5 ST-6 7.5- 9.0 ST-7 9.0- 10.5 ST-8 10.5- 12.0 ST-9 12.0- 13.5 ST-10 13.5- 15.0 ST-11 15.0- 16.0
205.7	4		LEAN CLAY (CL) (3.0 - 6.0) WITH SAND, BROWNISH GRAY AND YELLOW, MOIST, STIFF, VERY FRIABLE, VERY SANDY IN TOP 0.4'.		ST-3	
	6				ST-4	
	8		FAT CLAY (CH) (6.0 - 13.5) OLIVE AND GRAY WITH YELLOW, MOIST, STIFF TO VERY STIFF, FRIABLE, LAMINATIONS OF VERY FINE SAND AND SILT, CALCAREOUS TO SLIGHTLY CALCAREOUS, FRIABLE FISSILITY, 10.5-12.0 VERY WET.		ST-5	
	10				ST-6	
	12				ST-7	
	14		FAT CLAY (CH) (13.5 - 15.0) DARK OLIVE GRAY, MOIST, HARD, FISSILITY, SLIGHTLY SLICKEN- SIDED, LAMINATIONS OF SILT.		ST-8	
198.2	16		FAT CLAY (CH) (15.0 - 16.0) DARK OLIVE GRAY, MOIST, VERY STIFF, FRIABLE, FISSILITY, LAMINATIONS OF VERY FINE SAND AND SILT, TOP 0.2' VERY WET.		ST-9	
196.7					ST-10	
195.7					ST-11	
	18					
	20					

PROJECT
LHAAP-WASTE SUMPS

HOLE NO.
LHS-MW2

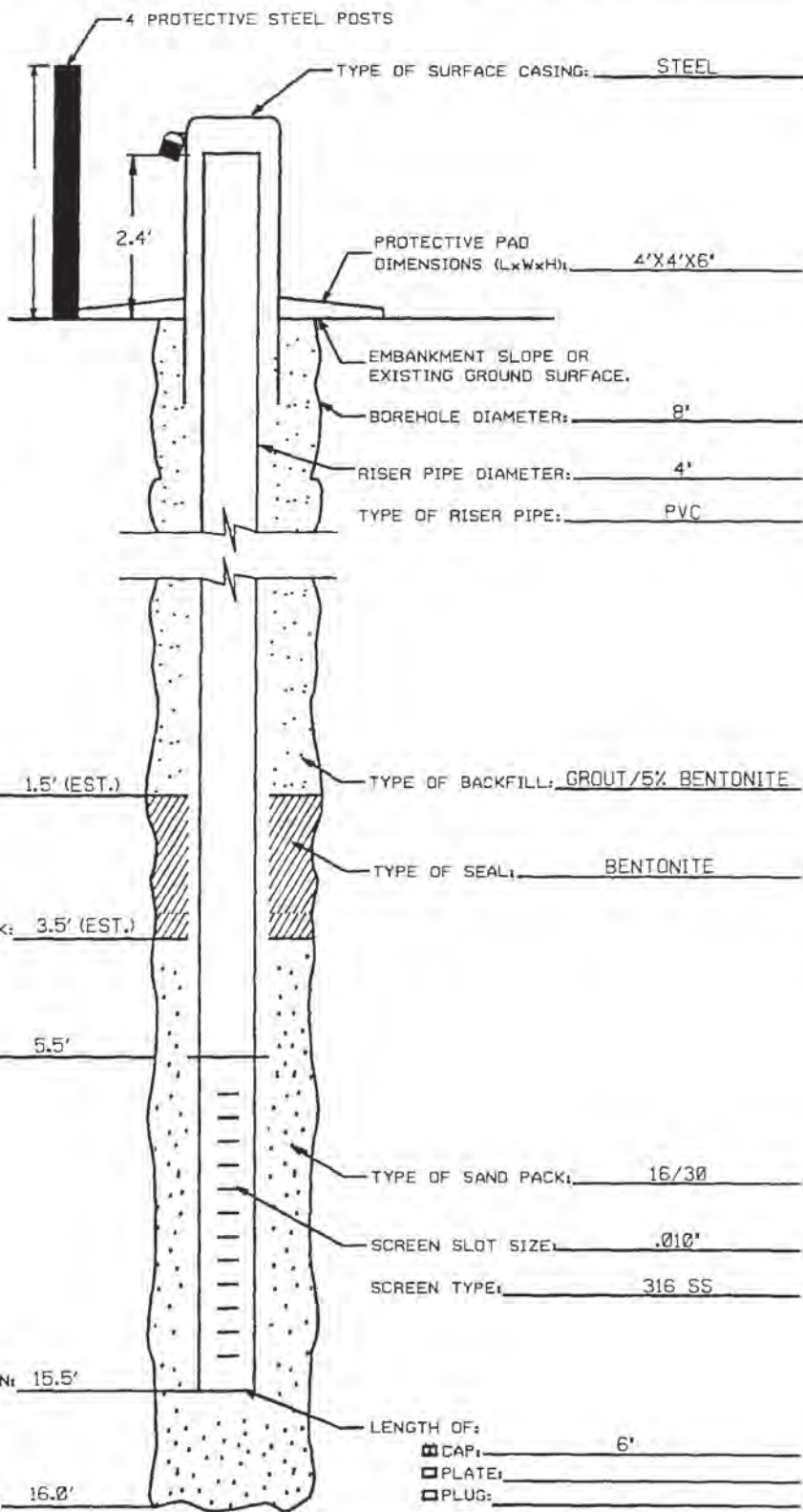
WELL NO. LHS-MW2

MONITORING WELL SHEET

PROJECT & INSTALLATION:
LONGHORN ARMY AMMUNITION PLANT - SUMPS

DATE STARTED:
8-19-1994

DATE COMPLETED:
8-20-1994



GROUNDWATER DEPTH: NOT OBSERVED

GROUNDWATER DATE: _____

INSPECTOR:
MERLIN DEAN

HOLE NO. LHS-MW3

DRILLING LOG		DIVISION	INSTALLATION		SHEET	
		SOUTHWEST	LHAAP		1	
1. PROJECT: LHAAP-WASTE SUMPS		10. SIZE AND TYPE OF BIT: 8" AUGER		OF 2 SHEETS		
2. LOCATION (Coordinates or Station): 6959935.80 3305514.20		11. DATUM FOR ELEVATION SHOWN (TBM or MSL): MSL				
3. DRILLING AGENCY: TULSA DISTRICT COE		12. MANUFACTURER'S DESIGNATION OF DRILL: CME 75				
4. HOLE NO. (As shown on drawing title and file number): LHS-MW3		13. OVERBURDEN SAMPLES: DISTURBED 4 UNDISTURBED 13				
5. NAME OF DRILLER: TOM BEAVERS		14. TOTAL NUMBER CORE BOXES: 0				
6. DIRECTION OF HOLE: <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.		15. ELEVATION GROUND WATER: NOT DETERMINED				
7. THICKNESS OF OVERBURDEN: 32.0		16. DATE HOLE: STARTED 08/20/1994 COMPLETED 08/20/1994				
8. DEPTH DRILLED INTO ROCK: 0.0		17. ELEVATION TOP OF HOLE: 214.1				
9. TOTAL DEPTH OF HOLE: 32.0		18. TOTAL CORE RECOVERY FOR BORING: 0.0		x		
MERLIN DEAN						
ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOV-ERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water loss, depth of weathering, etc., if significant)
212.0	2		LEAN CLAY (CL) (0.0 - 2.0) SANDY, BROWN, MOIST, VERY STIFF, FEW GRAVELS TO 3/8", TOP 0.5' SILTY SAND (SM) DARK BROWN, CALCAREOUS, ROOTS.		ST-1	SAMPLE TYPE ZONE SHELBY 0.0 - 22.8 SPLITSPOON 22.8 - 32.0
210.0	4		LEAN CLAY (CL) (2.0 - 4.0) SANDY, BROWNISH YELLOW WITH GRAY, VERY MOIST, VERY STIFF, FEW GRAVELS TO 3/8", FEW BLACK IRON-OXIDE STAINS.		ST-2	SAMPLE DEPTH ST-1 0.0 - 2.0 ST-2 2.0 - 4.0 ST-3 4.0 - 4.4 ST-4 6.0 - 8.0 ST-5 8.0 - 9.7 ST-6 9.7 - 10.2 ST-7 10.2 - 12.0 ST-8 12.0 - 13.8 ST-9 13.8 - 15.5 ST-10 15.5 - 17.4 ST-11 17.4 - 18.9 ST-12 18.9 - 21.0 ST-13 21.0 - 22.8
209.6	5		LEAN CLAY (CL) (4.0 - 4.4) SANDY, GRAY AND YELLOW WITH RED, MOIST, HARD.		ST-3	J-1 22.8 - 23.4 J-2 24.2 - 24.8 J-3 24.8 - 26.6 J-4 31.5 - 32.0
208.0	6		FAT CLAY (CH) (4.4 - 6.0) SANDY, RED WITH GRAY, MOIST, VERY STIFF.			
206.6	8		LEAN CLAY (CL) (6.0 - 7.4) WITH SAND, YELLOW WITH GRAY, MOIST, HARD.		ST-4	
206.0	8		FAT CLAY (CH) (7.4 - 8.0) GRAY WITH YELLOW, MOIST, HARD.			
205.4	10		CLAY SAND (SC) (8.0 - 8.6) WITH GRAVEL, BROWN, MOIST, GRAVELS TO 3/4".		ST-5	
204.6	10		LEAN CLAY (CL) (8.6 - 9.4) SANDY, GRAY WITH YELLOW AND BROWN, MOIST, HARD.		ST-6	
204.3	10		CLAY SAND (SC) (9.4 - 9.7) WITH GRAVEL, MOIST, GRAVEL TO 1/2".		ST-7	
	12		LEAN CLAY (CL) (9.7 - 13.8) WITH SAND, GRAY AND YELLOW, MOIST, HARD, POCKETS OF SAND THROUGHOUT, 12.0-13.8 VERY STIFF TO HARD.		ST-8	
200.2	14		LEAN CLAY (CL) (13.8 - 17.4) YELLOW WITH GRAY, MOIST, VERY STIFF.		ST-9	
	16				ST-10	
196.6	18		LEAN CLAY (CL) (17.4 - 21.0) YELLOW WITH GRAY, VERY MOIST TO WET, STIFF.		ST-11	
	20				ST-12	

PROJECT LHAAP-WASTE SUMPS

HOLE NO. LHS-MW3

HOLE NO. LHS-MW3

DRILLING LOG		DIVISION	SOUTHWEST		INSTALLATION	LHAAP		SHEET	2				
1. PROJECT		LHAAP-WASTE SUMPS			10. SIZE AND TYPE OF BIT		8" AUGER						
2. LOCATION		(Coordinates or Station)			11. DATUM FOR ELEVATION SHOWN		(TBM or MSL)						
3. DRILLING AGENCY		TULSA DISTRICT COE			12. MANUFACTURER'S DESIGNATION OF DRILL		CME 75						
4. HOLE NO.		(As shown on drawing title and file number)			13. OVERBURDEN SAMPLES		DISTURBED		UNDISTURBED				
5. NAME OF DRILLER		TOM BEAVERS			14. TOTAL NUMBER CORE BOXES		0						
6. DIRECTION OF HOLE		<input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.			15. ELEVATION GROUND WATER		NOT DETERMINED						
7. THICKNESS OF OVERBURDEN		32.0			16. DATE HOLE		STARTED		COMPLETED				
8. DEPTH DRILLED INTO ROCK		0.0			17. ELEVATION TOP OF HOLE		214.1						
9. TOTAL DEPTH OF HOLE		32.0			18. TOTAL CORE RECOVERY FOR BORING		0.0 %						
ELEVATION		DEPTH		LEGEND		CLASSIFICATION OF MATERIALS		% CORE RECOVERY		BOX OR SAMPLE NO.		REMARKS	
a		b		c		d		e		f		g	
191.0							LEAN CLAY (CL) (17.4 - 21.0) YELLOW WITH GRAY, VERY MOIST TO WET, STIFF.				ST-12		
			22				LEAN CLAY (CL) (21.0 - 22.8) WITH SAND, YELLOW WITH GRAY VERY MOIST, STIFF, SCATTERED BLACK IRON-OXIDE STAINS, BOTTON 0.4' SANDY LEAN CLAY.				ST-13		
							LEAN CLAY (CL) (22.8 - 24.2) WITH SAND, LIGHT OLIVE BROWN, MOIST.				J-1		
189.8			24				LEAN CLAY (CL) (24.2 - 24.6) OLIVE YELLOW, MOIST.				J-2		
							LEAN CLAY (CL) (24.8 - 31.5) WITH SAND, LIGHT OLIVE BROWN, MOIST.				J-3		
			26										
			28										
			30										
182.5							LEAN CLAY (CL) (31.5 - 32.0) LIGHT OLIVE BROWN, MOIST.				J-4		
182.1			32										
			34										
			36										
			38										
			40										

PROJECT LHAAP-WASTE SUMPS

HOLE NO. LHS-MW3

WELL NO. LHS-MW3

MONITORING WELL SHEET

PROJECT & INSTALLATION:
LONGHORN ARMY AMMUNITION PLANT - SUMPS

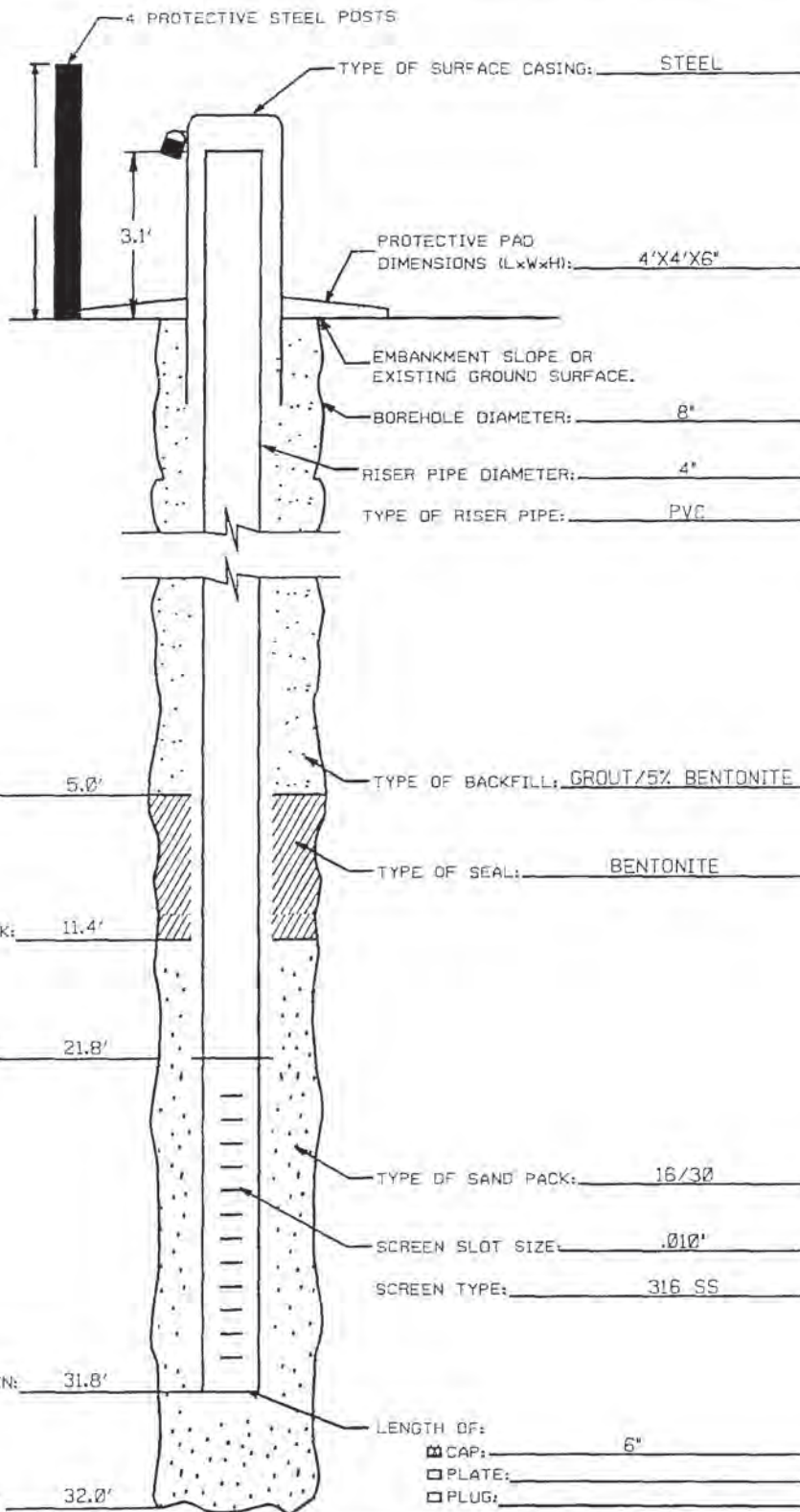
DATE STARTED:

8-20-1994

DATE COMPLETED:

8-20-1994

GROUNDWATER
DEPTH: NOT OBSERVED
GROUNDWATER
DATE: _____



INSPECTOR:

MERLIN DEAN

HOLE NO. LHS-MW4

DRILLING LOG		DIVISION SOUTHWEST	INSTALLATION LHAAP	SHEET OF 1 SHEETS
1. PROJECT LHAAP-WASTE SUMPS			10. SIZE AND TYPE OF BIT 8" AUGER	
2. LOCATION (Coordinates or Station) 6960183.80 3305398.40			11. DATUM FOR ELEVATION SHOWN (TBM or MSL) MSL	
3. DRILLING AGENCY TULSA DISTRICT COE			12. MANUFACTURER'S DESIGNATION OF DRILL CME 75	
4. HOLE NO. (As shown on drawing title and file number) LHS-MW4			13. OVERBURDEN SAMPLES DISTURBED 12 UNDISTURBED 0	
5. NAME OF DRILLER TOM BEAVERS			14. TOTAL NUMBER CORE BOXES 0	
6. DIRECTION OF HOLE <input checked="" type="checkbox"/> VERTICAL <input type="checkbox"/> INCLINED _____ DEG. FROM VERT.			15. ELEVATION GROUND WATER SEE REMARKS	
7. THICKNESS OF OVERBURDEN 30.0			16. DATE HOLE STARTED 08/22/1994 COMPLETED 08/23/1994	
8. DEPTH DRILLED INTO ROCK 0.0			17. ELEVATION TOP OF HOLE 214.0	
9. TOTAL DEPTH OF HOLE 30.0			18. TOTAL CORE RECOVERY FOR BORING 0.0 <input checked="" type="checkbox"/>	
STEVE BREWER				

ELEVATION	DEPTH	LEGEND	CLASSIFICATION OF MATERIALS (Description)	% CORE RECOVERY	BOX OR SAMPLE NO.	REMARKS (Drilling time, water level, depth of weathering, etc., if significant)
c	d	e	f	g	h	i
			LEAN CLAY (CL) (0.0 - 9.0) SANDY, OLIVE BROWN TO BROWNISH GRAY, MOIST, ROOTS TO 3.4'		J-1	WATER ENCOUNTERED @ 13.2'
	3				J-2	SAMPLE TYPE ZONE SPUTSPOON 0.0- 30.0
	6					SAMPLE DEPTH J-1 0.3- 1.2 J-2 1.2- 3.4 J-3 3.4- 9.0 J-4 9.0- 13.2 J-5 13.2- 16.0 J-6 16.0- 18.9 J-7 18.9- 23.8 J-8 23.8- 25.2 J-9 25.2- 27.2 J-10 27.2- 28.1 J-11 28.1- 29.0 J-12 29.0- 30.0
205.0	9		LEAN CLAY (CL) (9.0 - 16.0) OLIVE YELLOW, MOIST.		J-4	
	12					
	15				J-5	
198.0	18		LEAN CLAY (CL) (16.0 - 18.9) SANDY, LIGHT YELLOWISH BROWN, MOIST.		J-6	
195.1	21		LEAN CLAY (CL) (18.9 - 23.8) WITH SAND, YELLOWISH BROWN, MOIST, SILTY.		J-7	
190.2	24		CLAY SAND (SC) (23.8 - 25.2) WITH GRAVEL, REDDISH BROWN, MOIST, GRAVELS TO 1/2".		J-8	
188.8	27		CLAY SAND (SC) (25.2 - 27.2) PALE BROWN, MOIST, ROOTS.		J-9	
186.8			LEAN CLAY (CL) (27.2 - 28.1) SANDY, REDDISH BROWN, MOIST, GRAVELS TO 1/2".		J-10	
185.9			FAT CLAY (CH) (28.1 - 29.0) SANDY, REDDISH BROWN, MOIST, GRAVEL TO 1/2".		J-11	
185.0			FAT CLAY (CH) (29.0 - 30.0) WITH SAND, LIGHT BROWNISH GRAY MOIST.		J-12	
184.0	30					

PROJECT LHAAP-WASTE SUMPS HOLE NO. LHS-MW4

WELL NO. LHS-MW4

MONITORING WELL SHEET

PROJECT & INSTALLATION:
LONGHORN ARMY AMMUNITION PLANT - SUMPS

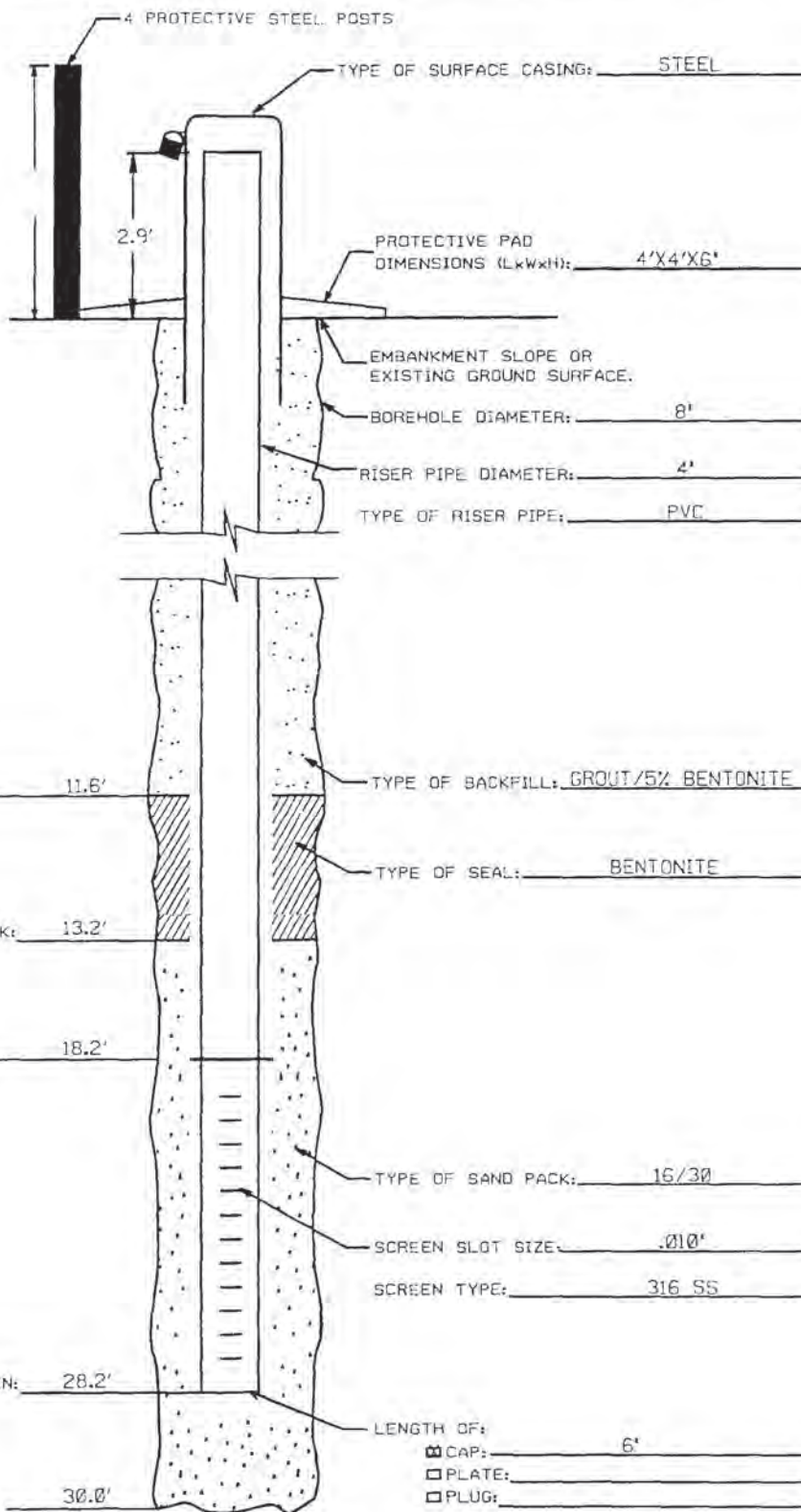
DATE STARTED:

8-22-1994

DATE COMPLETED:

8-23-1994

GROUNDWATER
DEPTH: 13.2'
GROUNDWATER
DATE: 8-23-1994



TOP OF SEAL: 11.6'

TOP OF SAND PACK: 13.2'

TOP OF SCREEN: 18.2'

BOTTOM OF SCREEN: 28.2'

BOTTOM OF HOLE: 30.0'

TYPE OF BACKFILL: GROUT/5% BENTONITE

TYPE OF SEAL: BENTONITE

TYPE OF SAND PACK: 16/30

SCREEN SLOT SIZE: .010"

SCREEN TYPE: 316 SS

LENGTH OF: 6'

CAP:

PLATE:

PLUG:

INSPECTOR:

STEVE BREWER

ENCLOSURE 2

TCEQ CORE DATA FORM



TCEQ Core Data Form

TCEQ Use Only

For detailed instructions regarding completion of this form, please read the Core Data Form Instructions or call 512-239-5175.

SECTION I: General Information

1. Reason for Submission <i>(If other is checked please describe in space provided.)</i>		
<input type="checkbox"/> New Permit, Registration or Authorization <i>(Core Data Form should be submitted with the program application.)</i>		
<input type="checkbox"/> Renewal <i>(Core Data Form should be submitted with the renewal form)</i>	<input type="checkbox"/> Other	
2. Customer Reference Number <i>(if issued)</i>	Follow this link to search for CN or RN numbers in Central Registry**	3. Regulated Entity Reference Number <i>(if issued)</i>
CN		RN

SECTION II: Customer Information

4. General Customer Information		5. Effective Date for Customer Information Updates (mm/dd/yyyy)		
<input type="checkbox"/> New Customer		<input type="checkbox"/> Update to Customer Information		<input type="checkbox"/> Change in Regulated Entity Ownership
<input type="checkbox"/> Change in Legal Name (Verifiable with the Texas Secretary of State or Texas Comptroller of Public Accounts)				
<i>The Customer Name submitted here may be updated automatically based on what is current and active with the Texas Secretary of State (SOS) or Texas Comptroller of Public Accounts (CPA).</i>				
6. Customer Legal Name <i>(If an individual, print last name first: eg: Doe, John)</i>			<i>If new Customer, enter previous Customer below:</i>	
7. TX SOS/CPA Filing Number	8. TX State Tax ID (11 digits)	9. Federal Tax ID (9 digits)	10. DUNS Number <i>(if applicable)</i>	
11. Type of Customer:		Partnership: <input type="checkbox"/> General <input type="checkbox"/> Limited		
<input type="checkbox"/> Corporation		<input type="checkbox"/> Individual		
Government: <input type="checkbox"/> City <input type="checkbox"/> County <input type="checkbox"/> Federal <input type="checkbox"/> State <input type="checkbox"/> Other		<input type="checkbox"/> Sole Proprietorship <input type="checkbox"/> Other:		
12. Number of Employees		13. Independently Owned and Operated?		
<input type="checkbox"/> 0-20 <input type="checkbox"/> 21-100 <input type="checkbox"/> 101-250 <input type="checkbox"/> 251-500 <input type="checkbox"/> 501 and higher		<input type="checkbox"/> Yes <input type="checkbox"/> No		
14. Customer Role (Proposed or Actual) – <i>as it relates to the Regulated Entity listed on this form. Please check one of the following:</i>				
<input type="checkbox"/> Owner		<input type="checkbox"/> Operator		<input type="checkbox"/> Owner & Operator
<input type="checkbox"/> Occupational Licensee		<input type="checkbox"/> Responsible Party		<input type="checkbox"/> Voluntary Cleanup Applicant <input type="checkbox"/> Other:
15. Mailing Address:				
City				
State				
ZIP				
ZIP + 4				
16. Country Mailing Information <i>(if outside USA)</i>			17. E-Mail Address <i>(if applicable)</i>	
18. Telephone Number		19. Extension or Code		20. Fax Number <i>(if applicable)</i>
() -				() -

SECTION III: Regulated Entity Information

21. General Regulated Entity Information <i>(If 'New Regulated Entity' is selected below this form should be accompanied by a permit application)</i>	
<input type="checkbox"/> New Regulated Entity <input type="checkbox"/> Update to Regulated Entity Name <input type="checkbox"/> Update to Regulated Entity Information	
<i>The Regulated Entity Name submitted may be updated in order to meet TCEQ Agency Data Standards (removal of organizational endings such as Inc, LP, or LLC.)</i>	
22. Regulated Entity Name <i>(Enter name of the site where the regulated action is taking place.)</i>	

23. Street Address of the Regulated Entity: <i>(No PO Boxes)</i>											
		City		State		ZIP		ZIP + 4			
24. County											
Enter Physical Location Description if no street address is provided.											
25. Description to Physical Location:		LHAAP is located in the northeast corner of Harrison County between SH43 and the western shore of Caddo Lake									
26. Nearest City						State		Nearest ZIP Code			
Karnack						TX		75661			
27. Latitude (N) In Decimal:			32.6665			28. Longitude (W) In Decimal:			94.1265		
Degrees		Minutes		Seconds		Degrees		Minutes		Seconds	
32		39		59.64		94		07		35.4	
29. Primary SIC Code (4 digits)			30. Secondary SIC Code (4 digits)			31. Primary NAICS Code (5 or 6 digits)			32. Secondary NAICS Code (5 or 6 digits)		
33. What is the Primary Business of this entity? <i>(Do not repeat the SIC or NAICS description.)</i>											
34. Mailing Address:											
		City		State		ZIP		ZIP + 4			
35. E-Mail Address:											
36. Telephone Number				37. Extension or Code			38. Fax Number <i>(if applicable)</i>				
() -							() -				

39. TCEQ Programs and ID Numbers Check all Programs and write in the permits/registration numbers that will be affected by the updates submitted on this form. See the Core Data Form instructions for additional guidance.

<input type="checkbox"/> Dam Safety	<input type="checkbox"/> Districts	<input type="checkbox"/> Edwards Aquifer	<input type="checkbox"/> Emissions Inventory Air	<input type="checkbox"/> Industrial Hazardous Waste
<input type="checkbox"/> Municipal Solid Waste	<input type="checkbox"/> New Source Review Air	<input type="checkbox"/> OSSF	<input type="checkbox"/> Petroleum Storage Tank	<input type="checkbox"/> PWS
<input type="checkbox"/> Sludge	<input type="checkbox"/> Storm Water	<input type="checkbox"/> Title V Air	<input type="checkbox"/> Tires	<input type="checkbox"/> Used Oil
<input type="checkbox"/> Voluntary Cleanup	<input type="checkbox"/> Waste Water	<input type="checkbox"/> Wastewater Agriculture	<input type="checkbox"/> Water Rights	<input checked="" type="checkbox"/> Other:

SECTION IV: Preparer Information

40. Name:	Praveen Srivastav			41. Title:	Project Manager		
42. Telephone Number	43. Ext./Code	44. Fax Number	45. E-Mail Address				
(281) 639-8743		() -	praveen.srivastav@aptim.com				

SECTION V: Authorized Signature

46. By my signature below, I certify, to the best of my knowledge, that the information provided in this form is true and complete, and that I have signature authority to submit this form on behalf of the entity specified in Section II, Field 6 and/or as required for the updates to the ID numbers identified in field 39.

Company:		Job Title:	
Name <i>(In Print)</i> :		Phone:	() -
Signature:		Date:	

**Subject: Final Minutes, Monthly Managers' Meeting (MMM),
Longhorn Army Ammunition Plant (LHAAP)**
Location of Meeting: Karnack Community Center, Karnack, Texas
Date of Meeting: 25 July 2019– 10:30 AM Central Daylight Time (CDT)

Attendees:

Army BRAC: Rose Zeiler (RMZ)
 EPA: Dorelle Harrison
 TCEQ: April Palmie (AP)
 USACE: Aaron Williams (AW)
 AEC: Amanda Sherman (AS)
 USGS: Kent Becher (KB)
 Bhate: Kim Nemmers (KN)
 USFWS: Paul Bruckwicki (PB)
 APTIM: Praveen Srivastav (PS) and Susan Watson (SW)

Action Items

Bhate/APTIM: PS stated that a proposed subset of wells for the baseline resampling at LHAAP-16 was provided to the Army for review. RMZ stated that the proposed wells could be discussed with the team later in the day. PS stated that the sites are drying out and waste characterization soil samples for LHAAP-17 were collected. KN stated that a field kick-off meeting was planned for later in the day.

KN stated that the water line to well 18WW17 needed to be repaired which had been held up by wet conditions. KN stated that Scott Beesinger was assessing the area and working with the contractor to schedule the repairs. RMZ asked how long the line has been down. KN stated that she would have to confirm the date, but that repairs in June 2019 led to the discovery that the line to well 18WW17 was leaking (the 2-inch line within the 4-inch line).

For the Oil and Gas Water Well Drilling Impact Update, PB stated that no additional information was available. RMZ stated that the action item is with the USFWS Regional office. AS requested more information on the subject. RMZ explained that this action item stems from a comment made during the Five Year Review (FYR). The question was whether there was a land use control for water resources during oil and gas water well drilling that would restrict drilling near a plume. RMZ stated that the main concern is where there is not a clay over the Wilcox Formation, which narrows down the sites to LHAAP-18/24. RMZ stated that the property at LHAAP-18/24 has not been transferred.

Defense Environmental Restoration Program (DERP) Performance Based Remediation (PBR) Update

KN stated the groundwater treatment plant (GWTP) is operating via the generator so LHAAP-16 remains down. AW stated that the transformer will be installed by August 20, 2019 followed by completing connections and testing for a few days. RMZ asked what takes so long. AW clarified that a whole new transformer is being installed and thus entire system needs to be tested. AP asked how long LHAAP-16 will be operated after the transformer is operational. KN stated that she recalled two weeks being the agreed upon time period. AP then confirmed that baseline groundwater samples would be collected after the system was operational for two weeks of time to

which KN concurred. KN explained that poles and lines damaged over the past few months are still requiring repairs, which will be completed ahead of the transformer installation to ensure that the system is operational as soon as possible. PS stated that the water elevations at LHAAP-16, collected in June 2019, show that water level elevations are up which may be a function of lack of pumping and the rainfall. KN stated that most months have received at least 10-inches of rain since September 2018. PS asked if the goal was to have the water table return to previous conditions. RMZ stated that the design was based upon having an operational extraction system and the goal is to have the aquifer behave like the system is operating again. PS stated that the groundwater flow directions based upon the June 2019 data have not changed significantly compared to 2018 groundwater flow, even though the system is not operating. SW clarified that the system has to operate per the remedy so ensuring that the extraction system is operational is important. AP concurred stating that the extraction system is part of the remedy for recirculation also. KN stated that the baseline sampling will allow for evaluation of possible changes from the October 2018 sampling event also. PS stated that approximately 19 of the original 39 monitoring wells (approximately 50-percent) were selected for baseline resampling at LHAAP-16 based upon detections and/or potential influence from extraction. RMZ stated that the wells selected could be discussed further during the kick-off meeting. KB asked how deep the injection was going to be along the bayou wall. PS stated injections along wall would be around 20 feet. KB expressed concern that the intermediate contamination in that area might be missed. RMZ stated that deeper injections were not part of the Remedial Design (RD) being implemented.

PS stated that monitoring wells were being installed, weather allowing, the following week. The intermediate well at LHAAP-16 will be completed because only the outer casing had been installed in October 2018. If conditions are right, the two LHAAP-16 wells on east side of Harrison Bayou will be installed. PS stated that wells are also planned to be installed at LHAAP-12, -50 and -67. PS stated that Scott Beesinger had scouted the location of the wells, and all of the locations at LHAAP-12, -50 and -67 looked to be accessible except a well at LHAAP-50, which is near the creek. PS also discussed the monitoring well with tree roots at LHAAP-46, which likely will need to be replaced if rehabilitation is not effective. RMZ then clarified that installation of wells at LHAAP-16 should be the highest priority, followed by LHAAP-50. SW confirmed that the priority was understood. PS stated that Scott Beesinger will attempt to make an alternate route for installation of the two monitoring wells across the creek at LHAAP-16. PS stated that Scott Beesinger planned to use the dozer to cut the alternative route for the monitoring wells at LHAAP-16 the following week, if the alternative route looks good. PS stated that a surveyor would be onsite to survey the new wells installed in the fall of 2018. RMZ mentioned the replacement well at LHAAP-12 needs to be installed at the same location and depth as the old abandoned monitoring well 12WW10. The new well will be labeled as 12WW10R.

KN asked everyone to refer to the Document and Issues Tracking Table dated July 25, 2019.

- **Task 1** (Project Management) -
 - KN stated that the next MMM is schedule for August 15, 2019 and requested the meeting be moved to August 22, 2019. RMZ concurred with the change to the date. KN stated that the June 2019 MMM Minutes were final.
 - KN stated that the Restoration Advisory Board (RAB) Meeting is being held at the Community Center tonight. The April 2019 meeting minutes were ready for approval at the RAB Meeting. The next RAB meeting is planned for October 17, 2019.

- **Task 3** (LHAAP-03) – PS stated that the Regulators (TCEQ and EPA) have approved the Draft Final RD/Remedial Action Work Plan (RAWP) for LHAAP-03.
- **Task 4** (LHAAP-04) – PS stated that there are no current documents.
- **Task 5** (LHAAP-12) – PS stated that the 2018 Annual Remedial Action Operation [RA (O)] Report was finalized. RMZ then asked if the new well at LHAAP-12 would be installed prior to the next sampling event.
- **Task 6** (LHAAP-16) – KN stated that the 1st Quarter 2019 GWTP Report that was just delivered to the Regulators contains the groundwater sample results from select wells.
- **Task 7** (LHAAP-17) – PS stated that no documents are currently in process for LHAAP-17.
- **Task 9** (LHAAP-37) – PS stated that the validated data from the May 2019 sampling event is provided with the data for this July 2019 MMM. AW stated that the data still shows a blip to the north. KN stated that the Year 2 Quarter 4 RA-O sampling event is planned for early August 2019. KN clarified that the priority, however, is the other work PS discussed, and so it is possible that the sampling events may be pushed until later in the month.
- **Task 10** (LHAAP-46) – PS stated that the Year 5 2nd Semi-Annual RA-O event is planned for August 2019. RMZ asked about the operating properly and successfully (OPS) determination. PS stated that a monitoring natural attenuation (MNA) evaluation will be completed in conjunction with the Year 5 RA-O Report, using all the prior data including the August 2019 data. PS stated that he thought an Army Draft could be provided in October 2019 and then the OPS report could be provided after that. SW clarified that the validated data was due in October 2019 and not the Army Draft report. RMZ stated to AP that the OPS determination was not a statutory requirement but is a transfer requirement for USFWS. RMZ continued that the EPA typically reviews the OPS determination. PS further discussed the monitoring well that has tree roots down into the screen interval, which is surprising since this is approximately 20 feet below the ground surface (bgs). PS explained that the driller has indicated that there is no easy way to remove the tree roots without damaging the well. PS stated that the best option is to install another well within 5 feet of this well. AS asked if the well could be “roto-rooted.” SW stated that the concern is damaging the screen. PS stated that the groundwater is not higher than the roots. KN asked if the well could be replaced because that seemed like the best alternative for the issue. AP stated that it seemed reasonable but that she wanted to look at the monitoring well logs.
- **Task 11** (LHAAP-50) – PS stated that Year 5 2nd semi-annual RA-O sampling event was completed in May and the validated data was presented for this July 2019 MMM meeting. PS stated that TCEQ approved the Draft Final Explanation of Significant Difference (ESD) for a contingency remedy. The EPA issued an email on July 15, 2019 indicated no comment on the Draft Final ESD. PS stated that the Army Draft RD/RAWP was under Army review. PS stated that the new monitoring well is being installed next week and the RD/RAWP might be modified based upon the data from the new well. PS stated that the monitoring well will be sampled after it is installed and then any necessary revisions to the RD/RAWP will be completed.
- **Task 12** (LHAAP-58) – KN stated that the LHAAP-58 sampling was completed in June 2019, and that validated data was included with the July 2019 validated data. KN stated that the Year 5 RA(O)Report is being prepared to evaluate the first four quarters of the western plume remedy implementation as well as the continued performance of the eastern plume remedy. KN stated that the Year 5 RA(O) Report was planned for submittal to the Regulators in October 2019. The next groundwater sampling event is the quarterly groundwater monitoring for the western plume in September 2019. KN clarified that the

western plume has another full year of quarterly groundwater monitoring prior to evaluation for OPS.

- **Task 13** (LHAAP-67) – PS stated that the Year 5 2nd semi-annual RA-O validated data was included in the package for this July 2019 MMM.
- **Task 14** (LHAAP-001-R and –003-R) - KN stated that no documents are in process but that the land use control (LUC) inspections were planned for July 2019.
- **Task 16** (GWTP) – KN stated that the Regulator comments were received on the 4th Quarter 2018 GWTP report, which are discussed in the 1st Quarter 2019 GWTP Report that was just submitted. KN indicated that the 2nd Quarter 2019 GWTP was being prepared and is anticipated for delivery to the Regulators in October 2019.
- **Task 17** (LHAAP-18/24) – KN stated that two of the monitoring wells from LHAAP-18/24 still need to be sampled and that the hope was that the samples would be collected the following week. KN explained that one monitoring well also had to be sampled in July 2019 due to the wet conditions. AS asked if there was any issues with the wells being sampled over such a long period of time, KN stated that sampling of wells over a month after a majority of the monitoring wells were sampled was not ideal but the only other option was to not sample the wells. KN stated that discussion of the effects of sampling at different times would be discussed in the report. KN also stated that data is compared to historic data and could identify if there was a significant variances. KN stated that the report will likely be prepared with or without this data, at least for the Army Draft. KN stated that the LHAAP-18/24 data will be included the 2nd Quarter 2019 GWTP Report, which is due to the Regulators in October 2019.
- **Task 18** (Surface Water) – KN stated that surface water was collected for analysis in July 2019.
- **Task 19** (LUC Management) - RMZ stated that there will be the three RDs (LHAAP-03, LHAAP-04, and LHAAP-17) that need to be included in the update to the LUC Management Plan. SW asked what the cutoff date was to which RMZ stated September 30, 2019. RMZ stated that the LUC portion of the RD is what goes into the LUC Management Plan. RMZ stated that nothing changed on LHAAP-50 for the LUC for the site. RMZ noted that the recordation for LHAAP-16 will need to eventually be included. RMZ stated that even if LHAAP-16 gets re-surveyed that the new LUC boundary would not go into the 2019 revision to the LUC Management Plan due to timing for completion likely after the cutoff date for the 2019 LUC Management Plan update. RMZ stated that LHAAP-03 did not have a recordation so it is just the RD (LUC portion).
- **Administrative Record (AR)** – SW said that the December 2018 AR hard copies were placed into the site trailer before the meeting. SW said that the list of documents for the AR is being reviewed for the 1st quarter. SW stated that the April through June document list is also being prepared. RMZ stated that the letters for the draft submittal has to be included in the AR for primary documents.

Update on other DERP Sites

- **LHAAP 18/24** – AW explained that the Final Proposed Plan (PP) for LHAAP-18/24 will be placed into the AR. AW stated that he could provide the letter for the draft PP, if needed, for the AR. SW stated that the PP is in the January through March 2019 list and that she would double check if she had the letter transmitting the draft. AW stated that the Record of Decision (ROD) is being prepared for submittal to the Regulators on September 6, 2019.
- **LHAAP-29** – AW stated that the LHAAP-29 PP is ready for AR. Regulator comments on the ROD have been addressed and the Draft Final ROD was issued to the Regulators. RMZ stated that the Army had noticed text was added that should not have been included on one

of the pages with the applicable or relevant and appropriate requirements (ARARs). AW stated that a change page would be issued. AP stated that an email with the change page would be sufficient.

- **LHAAP-47** – AW stated that the Post-Screening Investigation (PSI) Report is ready for the AR. AW stated that the EPA had three comments on the Addendum to the PSI Report, which was submitted on July 1, 2019. AP stated that the EPA comments cover the TCEQ comments. KB stated that Rich Mayer (not present) had mentioned an issue with LHAAP-47. RMZ stated that she felt the issue was the one she called him about and that it might be resolved. RMZ stated that there is one replacement monitoring well with 120,000 parts per billion (ppb) of trichloroethylene (TCE), which may be indicative of dense non-aqueous phase liquid (DNAPL). RMZ stated that only having one point with those levels does not tell us much about how contaminated the aquifer is with TCE. KB suggested understanding the distribution of the contamination along the screen length would be helpful and suggested using passive samplers. PS then explained that the bioremediation process is slow, and high TCE concentrations can be toxic to bacteria. Bioremediation is known to release chlorinated solvents from the soil matrix, allowing contamination to be treated by the bacteria over time, but the process is very slow. RMZ stated that the Army and Regulators don't want to sign a ROD knowing that the remedy will not work, to which AP concurred. AP proposed that the Army write a letter stating that the recently collected data changes the site information and may require that the remedy selected previously also be changed. RMZ asked if there was a quicker way to complete the remedy change other than completing a new PP. PS asked how deep the contamination was located, to which RMZ stated that the contamination was 20 to 30 feet bgs. AW stated that there are borings in the area that were clean that could be used to help bound the contamination. KN asked if the Army could dig the area to which RMZ said excavation would be her preference and that excavation was an element of the selected remedy, but for perchlorate, not TCE. PS cautioned that the contamination will likely rebound with just bioremediation based upon his experience. AS suggested an interim removal action and continue with the current ROD since the issue is limited to one area. Options under CERCLA were discussed for LHAAP-47, but the main issue identified by RMZ is whether the EPA will approve a change to the enforceable schedule. AP pointed out that this is a site held up by the dispute and that the data represents a change condition for LHAAP-47. AS asked if the site would need to be dewatered. RMZ stated that the groundwater was at the clay layer. AP stated that there is no measurable groundwater in many of the wells, which is why the investigation was completed.
- **FYR** – AW stated that the FYR was ready for the AR with the insertion of the errata.
- **Other items** - AP asked about the underground injection control (UIC) submittals. SW explained that the LHAAP-50 submittal is awaiting the additional groundwater sampled to determine if there will be any change in the remedial design. However, the LHAAP-04 UIC was under Army review and will be submitted to TCEQ soon. AP clarified that injection field work can begin only 30 days after UIC submittal. This should be factored into the field work planning.

Schedule Next Managers' Meeting

The next MMM will be held on August 22, 2019 at 10:00 am CDT via a conference call.

Meeting concluded at 11:35 am CDT.

ACRONYM LIST

AP April Palmie

APTIM	APTIM Federal Services, LLC
AR	Administrative Record
ARAR	Applicable or relevant and appropriate requirements
AS	Amanda Sherman
AW	Aaron Williams
bgs	below ground surface
Bhate	Bhate Environmental Associates, Inc.
BRAC	Base Realignment and Closure
CDT	Central Daylight Time
DERP	Defense Environmental Restoration Program
DNAPL	Dense Non-Aqueous Phase Liquid
EPA	United States Environmental Protection Agency
ESD	Explanation of Significant Differences
FYR	Five Year Review
GWTP	Ground Water Treatment Plant
KN	Kim Nemmers
LHAAP	Longhorn Army Ammunition Plant
MMM	Monthly Managers' Meeting
PBR	Performance-Based Remediation
PP	Proposed Plan
ppb	Parts per billion
PS	Praveen Srivastav
PSI	Post-Screening Investigation
RAB	Restoration Advisory Board
RA(O)	remedial action – operation
RAWP	Remedial Action Work Plan
RD	Remedial Design
RMZ	Rose M. Zeiler
ROD	Record of Decision
RTC	Response to Comment
SW	Susan Watson
TCE	trichloroethylene
TCEQ	Texas Commission on Environmental Quality
UIC	Underground Injection Control
USACE	United States Army Corps of Engineers
USFWS	United States Fish and Wildlife Service

**LHAAP Validated Data Packages for
July 2019 Monthly Manager's Meeting**

LHAAP Area	Analytic Method
LHAAP-35B(37)	<p><i>Year 2 Quarter 3 Remedial Action Operation Sampling - May 2019</i> VOCs (SW8260)</p>
LHAAP-50	<p><i>Year 5 Semiannual Remedial Action Operation Sampling Event #2 - May 2019</i> VOCs (SW8260) Perchlorate (SW6850)</p>
LHAAP-67	<p><i>Year 5 Semiannual Remedial Action Operation Sampling Event #2 - May 2019</i> VOCs (SW8260)</p>
GWTP Effluent	<p><i>Weekly Perchlorate Sampling – June 2019</i> Perchlorate (6850)</p>
GWTP Effluent	<p><i>Weekly, Bi-Weekly, and Monthly Sampling – June 2019</i> Ammonia (350.3) Ortho-Phosphate (365.3) Organic Carbon (415.1) VOC (8260C) Metals (6020A) Hexavalent Chromium (7196A) 1,4-Dioxane (8270D-SIM) Anions (9056)</p>
GWTP Influent	<p><i>Monthly Sampling – June 2019</i> Metals (6020A) Perchlorate (6850) Hexavalent Chromium (7196A)</p>
LHAAP-58	<p><i>Semi-Annual Sampling Event– June 2019</i> Anions (9056) VOC (8260C) Total Organic Carbon (415.1) Metabolic Acids (HPLC-METACIDS) Dechlorinating Bacteria (CENSUS) Dissolved Gases (RSK-175) Arsenic (6020A)</p>

LHAAP-37 RA-O Sampling - Year 2, Quarter 3, May 2019

Location Code			35BWW01				35BWW04		35BWW05		35BWW07		35BWW08	
Sample ID			35BWW01-190520		35BWW01-190520-FD		35BWW04-190520		35BWW05-190521		35BWW07-190522		35BWW08-190520	
Sample Date			5/20/2019		5/20/2019		5/20/2019		5/21/2019		5/22/2019		5/20/2019	
Location Description			Shallow zone, unimpacted, within site boundary		Shallow zone, unimpacted, within site boundary		Shallow zone, unimpacted, within site boundary		Shallow zone, impacted, within site boundary		Shallow zone, unimpacted downgradient		Shallow zone, unimpacted, within site boundary	
Parameter	Units	MCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
VOCs														
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Tetrachloroethene	µg/L	5	< 0.5	U	< 0.5	U	4.7		1.1		< 0.5	U	< 0.5	U
Trichloroethene	µg/L	5	< 0.5	U	< 0.5	U	0.58	J	5.5		< 0.5	U	< 0.5	U
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

Blue highlighting indicates concentrations above the MCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

Val Qual - validation qualifier

VOCs - volatile organic compounds

LHAAP-37 RA-O Sampling - Year 2, Quarter 3, May 2019

Location Code			35BWW09		35BWW10		35BWW11		35BWW12				35BWW13	
Sample ID			35BWW09-190521		35BWW10-190520		35BWW11-190521		35BWW12-190521		35BWW12-190521-FD		35BWW13-190522	
Sample Date			5/21/2019		5/20/2019		5/21/2019		5/21/2019		5/21/2019		5/22/2019	
Location Description			Shallow zone, impacted outside site boundary		Shallow, impacted, within site boundary		Shallow zone, unimpacted, within site boundary		Shallow zone, impacted, within site boundary		Shallow zone, impacted, within site boundary		Shallow zone, unimpacted, crossgradient	
Parameter	Units	MCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
VOCs														
1,1-Dichloroethene	µg/L	7	5.5		< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	1.8		< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Tetrachloroethene	µg/L	5	41		5.5		< 0.5	U	6.2		5.7		< 0.5	U
Trichloroethene	µg/L	5	20		9.4		< 0.5	U	0.55	J	0.53	J	< 0.5	U
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

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MCL - Maximum Contaminant Limit

Val Qual - validation qualifier

VOCs - volatile organic compounds

LHAAP-37 RA-O Sampling - Year 2, Quarter 3, May 2019

Location Code			35BWW14		35BWW15		35BWW16		35BWW17		35BWW18		35BWW19	
Sample ID			35BWW14-190521		35BWW15-190522		35BWW16-190523		35BWW17-190523		35BWW18-190520		35BWW19-190520	
Sample Date			5/21/2019		5/22/2019		5/23/2019		5/22/2019		5/20/2019		5/20/2019	
Location Description			Shallow zone, impacted, within site boundary		Shallow zone, impacted, within site boundary		Shallow zone, impacted, outside site boundary		Shallow zone, unimpacted, within site boundary		Shallow zone, unimpacted, outside site boundary		Shallow zone, unimpacted, outside site boundary	
Parameter	Units	MCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
VOCs														
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	0.66	J	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Tetrachloroethene	µg/L	5	< 0.5	U	7.8		9.5		< 0.5	U	< 0.5	U	< 0.5	U
Trichloroethene	µg/L	5	110		7.1		3.6		< 0.5	U	< 0.5	U	< 0.5	U
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

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MCL - Maximum Contaminant Limit

Val Qual - validation qualifier

VOCs - volatile organic compounds

LHAAP-37 RA-O Sampling - Year 2, Quarter 3, May 2019

Location Code			35BWW20		35BWW23		35BWW24		35BWW25		35BWW26		LHSMW58			
Sample ID			35BWW20-190522		35BWW20-190522-FD		35BWW23-190521		35BWW24-190522		35BWW25-190522		35BWW26-190520		LHSMW58-190522	
Sample Date			5/22/2019		5/22/2019		5/21/2019		5/22/2019		5/22/2019		5/20/2019		5/22/2019	
Location Description			Shallow zone, impacted, within site boundary		Shallow zone, impacted, within site boundary		Shallow zone, unimpacted, outside site boundary		Shallow zone, unimpacted, outside site boundary		Shallow zone, impacted, outside site boundary		Shallow zone, unimpacted, within site boundary		Shallow zone, unimpacted, within site boundary	
Parameter	Units	MCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
VOCs																
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Tetrachloroethene	µg/L	5	12		12		< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	3.6	
Trichloroethene	µg/L	5	2.8		2.9		< 0.5	U	1.1		6.3		< 0.5	U	< 0.5	U
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

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µg/L - micrograms per liter

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U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

Val Qual - validation qualifier

VOCs - volatile organic compounds

LHAAP-50 Year 5, Semiannual RA-O Sampling Event #2

Location Code Sample ID Sample Date Location Description			50WW05		50WW06		50WW08		50WW09			
			50WW05-190507		50WW06-190506		50WW08-190507		50WW09-190507		50WW09-190507-FD	
			5/7/2019		5/6/2019		5/7/2019		5/7/2019		5/7/2019	
			Site 50 - NE, lower shallow, outside site boundary		Site 50 - ENE, outside site boundary		Site 50 - E, upper shallow, inside site boundary		Site 50 - E, lower shallow, inside site boundary		Site 50 - E, lower shallow, inside site boundary	
Parameter	Units	MCL/ PCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
Perchlorate												
Perchlorate	µg/L	17	< 2	U	4.3		230		7.2		7.8	
Volatiles												
1,1-Dichloroethene	µg/L	7	1.2		< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
1,2-Dichloroethane	µg/L	5	0.94	J	< 0.5	U	2.1		< 0.5	U	< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	28		< 0.5	U	20		2.2		2.1	
Tetrachloroethene	µg/L	5	< 0.5	U	< 0.5	U	0.75	J	< 0.5	U	< 0.5	U
Trichloroethene	µg/L	5	110		1.4		210		16		16	
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

Blue highlighting indicates concentrations above the MCL/PCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

PCL - Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level.

RA-O - Remedial Action Operation

Val Qual - validation qualifier

LHAAP-50 Year 5, Semiannual RA-O Sampling Event #2

Location Code Sample ID Sample Date Location Description			50WW10		50WW11		50WW12				50WW13	
			50WW10-190507		50WW11-190506		50WW12-190516		50WW12-190516-FD		50WW13-190506	
			5/7/2019		5/6/2019		5/16/2019		5/16/2019		5/6/2019	
			Site 50 - E, intermediate, inside site boundary		Site 50 - ENE, upper shallow, outside site boundary		Site 50 - ENE, upper shallow, outside site boundary		Site 50 - ENE, upper shallow, outside site boundary		Site 50 - E, upper shallow, outside site boundary	
Parameter	Units	MCL/ PCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
Perchlorate												
Perchlorate	µg/L	17	< 2	U	450		65,000		76,000		160	
Volatiles												
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	2.2		2.2		0.57	J
1,2-Dichloroethane	µg/L	5	< 0.5	U	0.75	J	1.5		1.5		1.3	
cis-1,2-Dichloroethene	µg/L	70	< 0.5	U	3.1		< 0.5	U	< 0.5	U	12	
Tetrachloroethene	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Trichloroethene	µg/L	5	< 0.5	U	95		170		170		250	
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

Blue highlighting indicates concentrations above the MCL/PCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

PCL - Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level.

RA-O - Remedial Action Operation

Val Qual - validation qualifier

LHAAP-50 Year 5, Semiannual RA-O Sampling Event #2

Location Code			50WW14		50WW15		50WW16		50WW17		50WW18	
Sample ID			50WW14-190506		50WW15-190523		50WW16-190506		50WW17-190516		50WW18-190516	
Sample Date			5/6/2019		5/23/2019		5/6/2019		5/16/2019		5/16/2019	
Location Description			Site 50 - E, lower shallow, outside site boundary, along S. Crockett Ave		Site 50 - NNE, upper shallow, outside site boundary, along Goose Prairie Creek bridge		Site 50 - NE, upper shallow, outside site boundary, along Goose Prairie Creek		Site 50 - NE, fully penetrating shallow, outside site boundary, near Goose Prairie Creek		Site 50 - NE, upper shallow, outside site boundary, along Goose Prairie Creek	
Parameter	Units	MCL/ PCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
Perchlorate												
Perchlorate	µg/L	17	< 2	U	< 2	U	< 2	U	< 2	U	< 2	U
Volatiles												
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
1,2-Dichloroethane	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	1.9		2.9		< 0.5	U	< 0.5	U	< 0.5	U
Tetrachloroethene	µg/L	5	< 0.5	U	0.87	J	< 0.5	U	< 0.5	U	< 0.5	U
Trichloroethene	µg/L	5	27		2.3		0.61	J	< 0.5	U	< 0.5	U
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

Blue highlighting indicates concentrations above the MCL/PCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

PCL - Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level.

RA-O - Remedial Action Operation

Val Qual - validation qualifier

LHAAP-50 Year 5, Semiannual RA-O Sampling Event #2

Location Code			50WW21	50WW22	50WW23	50WW24	50WW27					
Sample ID			50WW21-190516	50WW22-190506	50WW23-190507	50WW24-190507	50WW27-190523					
Sample Date			5/16/2019	5/6/2019	5/7/2019	5/7/2019	5/23/2019					
Location Description			Site 50 - E, upper shallow, outside site boundary, east side of S. Crockett Ave	Site 50 - SE, upper shallow, outside site boundary	Site 50 - E, upper shallow, outside site boundary	Site 50 - ENE, upper shallow, outside site boundary	Site 50 - N, upper shallow, outside site boundary, east of S. Crockett Ave					
Parameter	Units	MCL/ PCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
Perchlorate												
Perchlorate	µg/L	17	< 2	U	< 2	U	< 2	U	< 2	U	< 2	U
Volatiles												
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
1,2-Dichloroethane	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	< 0.5	U	< 0.5	U	< 0.5	UJ	< 0.5	U	0.74	J
Tetrachloroethene	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Trichloroethene	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

Blue highlighting indicates concentrations above the MCL/PCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

PCL - Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level.

RA-O - Remedial Action Operation

Val Qual - validation qualifier

LHAAP-67 Year 5, Semiannual RA-O Sampling Event #2

Location Code			67WW01		67WW02		67WW05		67WW07		67WW08		67WW09	
			Sample ID		67WW01-190503		67WW02-190501		67WW05-190502		67WW07-190503		67WW08-190501	
Sample Date			5/3/2019		5/1/2019		5/2/2019		5/3/2019		5/1/2019		5/3/2019	
			Location Description			Site 67-Central, within site boundary		Site 67 - NW, within site boundary		Site 67 - WNW, outside site boundary		Site 67 - E, outside site boundary		Site 67-S, within site boundary
Parameter	Units	MCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual
Volatiles														
1,1,1-Trichloroethane	µg/L	200	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
1,1,2-Trichloroethane	µg/L	5	1.4		1.2		< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
1,1-Dichloroethene	µg/L	7	280		180		< 0.5	U	< 0.5	U	110		< 0.5	U
1,2-Dichloroethane	µg/L	5	21		4.7		< 0.5	U	< 0.5	U	5.1		< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	0.86	J	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Trichloroethene	µg/L	5	1.3		0.87	J	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U
Vinyl chloride	µg/L	2	0.92	J	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

Blue highlighting indicates concentrations above the MCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

RA-O - Remedial Action Operation

Val Qual - validation qualifier

LHAAP-67 Year 5, Semiannual RA-O Sampling Event #2

Location Code			67WW09A		67WW10		67WW11		67WW12		67WW13					
			Sample ID		67WW09A-190502		67WW10-190502		67WW11-190501		67WW12-190502		67WW13-190501		67WW13-190501-FD	
			Sample Date		5/2/2019		5/2/2019		5/1/2019		5/2/2019		5/1/2019		5/1/2019	
			Location Description			Site 67 - S outside site boundary		Site 67 - SE, outside site boundary		Site 67- SW, outside site boundary		Site 67 - NE, within site boundary		Site 67 - NE, within site boundary		Site 67 - NE, within site boundary
Parameter	Units	MCL	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual	Result	Val Qual		
Volatiles																
1,1,1-Trichloroethane	µg/L	200	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U		
1,1,2-Trichloroethane	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	3.9		3.8			
1,1-Dichloroethene	µg/L	7	< 0.5	U	< 0.5	U	2.8		2.1		200		160			
1,2-Dichloroethane	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	24		23			
cis-1,2-Dichloroethene	µg/L	70	< 0.5	U	< 0.5	U	< 0.5	UJ	< 0.5	U	0.94	J	0.81	J		
Trichloroethene	µg/L	5	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	1.1		1			
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U	< 0.5	U		

Notes:

Blue highlighting indicates concentrations above the MCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

RA-O - Remedial Action Operation

Val Qual - validation qualifier

LHAAP-67 Year 5, Semiannual RA-O Sampling Event #2

Location Code			67WW14	67WW15	67WW16I			
Sample ID			67WW14-190502	67WW15-190503	67WW16I-190501			
Sample Date			5/2/2019	5/3/2019	5/1/2019			
Location Description			Site 67 - SW, outside the site boundary beside Ignatius Ave	67-W, outside site boundary	67 - S, within site boundary			
Parameter	Units	MCL	Result	Val Qual	Result	Val Qual	Result	Val Qual
Volatiles								
1,1,1-Trichloroethane	µg/L	200	< 0.5	U	< 0.5	U	< 0.5	U
1,1,2-Trichloroethane	µg/L	5	< 0.5	U	5.4		< 0.5	U
1,1-Dichloroethene	µg/L	7	5.3		330		< 0.5	U
1,2-Dichloroethane	µg/L	5	1.8		22		< 0.5	U
cis-1,2-Dichloroethene	µg/L	70	< 0.5	U	1.3		< 0.5	U
Trichloroethene	µg/L	5	< 0.5	U	1.1		< 0.5	U
Vinyl chloride	µg/L	2	< 0.5	U	< 0.5	U	< 0.5	U

Notes:

Blue highlighting indicates concentrations above the MCL.

µg/L - micrograms per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

FD - field duplicate

MCL - Maximum Contaminant Limit

RA-O - Remedial Action Operation

Val Qual - validation qualifier

GWTP Weekly/Effluent Perchlorate Sampling - June 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_060419_BIX 6/4/19	LH18/24- SP650_060419_BIX 6/4/19	LH18/24- SP650_061219_BIX 6/12/19
Location Description		Collected from a spigot on the discharge of effluent TK-650.			
		Monthly EFF	Weekly	Weekly	
Perchlorate (6850)					
Perchlorate	µg/L	589	< 2.0 U	< 2.0 U	< 2.0 U

µg/L - micrograms per liter

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

BIX - before ion exchange

GWTP Weekly Sampling - June 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_060419 6/4/19	LH18/24-SP650_061219 6/12/19
Location Description			GWTP—Collected from a spigot on the discharge of effluent TK-650. Sampled Weekly.	
Ammonia as N (350.3)				
Ammonia as N	mg/L	NV	7.5	10
Ortho-Phosphate (365.3)				
Ortho-Phosphate	mg/L	NV	2.11	2.37
Organic Carbon (415.1)				
Total Organic Carbon (TOC)	mg/L	NV	2.21	2.04

mg/L - milligrams per liter

NV - No Value

GWTP Bi-Weekly Sampling - June 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_061219 6/12/19
Location Description			GWTP – Collected from a spigot on the discharge of effluent TK-650. Sampled Biweekly.
Volatile Organic Compounds (8260C)			
1,1,1-Trichloroethane	µg/L	7,230	< 0.5 U
1,1,2-Trichloroethane	µg/L	216.9	< 0.5 U
1,1-Dichloroethane	µg/L	14,032	< 0.5 U
1,1-Dichloroethene	µg/L	253	< 0.5 U
1,2-Dichloroethane	µg/L	181	0.46 J
1,2-Dichloropropane	µg/L	5	< 0.5 U
Acetone	µg/L	2,395	< 1.0 UJ
Benzene	µg/L	181	< 0.5 U
Carbon tetrachloride	µg/L	181	< 0.5 U
Chlorobenzene	µg/L	47,180	< 0.5 U
Chloroform	µg/L	3,615	< 0.5 U
Ethylbenzene	µg/L	57,025	< 0.5 U
m,p-Xylene	µg/L	83.6	< 1.0 U
Methylene chloride	µg/L	1,699	< 1.0 U
o-Xylene	µg/L	83.6	< 0.5 U
Styrene	µg/L	5,987	< 0.5 U
Tetrachloroethene	µg/L	180.7	< 0.5 U
Toluene	µg/L	4,189	< 0.5 U
Trichloroethene	µg/L	181	0.64 J
Vinyl chloride	µg/L	72	< 0.5 U
Anions (9056)			
Chloride	mg/L	NV	362
Sulfate	mg/L	NV	33.4

µg/L - micrograms per liter

mg/L - milligrams per liter

U - Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

NV - No Value

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

UJ - estimated non-detect due to quality control issues

GWTP Monthly Effluent Sampling - June 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_060419 6/4/19
Location Description			GWTP – Collected from a spigot on the discharge of effluent TK-650. Sampled monthly
Volatile Organic Compounds (8260C)			
1,1,1-Trichloroethane	µg/L	7,230	< 0.5 U
1,1,2-Trichloroethane	µg/L	216.9	< 0.5 U
1,1-Dichloroethane	µg/L	14,032	< 0.5 U
1,1-Dichloroethene	µg/L	253	< 0.5 U
1,2-Dichloroethane	µg/L	181	< 0.5 U
1,2-Dichloropropane	µg/L	5	< 0.5 U
Acetone	µg/L	2,395	< 1.0 U
Benzene	µg/L	181	< 0.5 U
Carbon tetrachloride	µg/L	181	< 0.5 U
Chlorobenzene	µg/L	47,180	< 0.5 U
Chloroform	µg/L	3,615	< 0.5 U
Ethylbenzene	µg/L	57,025	< 0.5 U
m,p-Xylene	µg/L	83.6	< 1.0 U
Methylene chloride	µg/L	1,699	< 1.0 U
o-Xylene	µg/L	83.6	< 0.5 U
Styrene	µg/L	5,987	< 0.5 U
Tetrachloroethene	µg/L	180.7	< 0.5 U
Toluene	µg/L	4,189	< 0.5 U
Trichloroethene	µg/L	181	0.76 J
Vinyl chloride	µg/L	72	< 0.5 U
Metals (6020A)			
Barium	mg/L	2	0.175
Lead	mg/L	0.0046	< 0.00100 U
Selenium	mg/L	0.012	< 0.00250 UJ
Silver	mg/L	0.003	< 0.000500 U
Hexavalent Chromium (7196A)			
Hexavalent Chromium	mg/L	0.1244	< 0.0100 U
Semi-Volatile Organic Compounds (8270D SIM)			
1,4-Dioxane	µg/L	134.2	6.2

µg/L - micrograms per liter

mg/L - milligrams per liter

U - Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

UJ - estimated non-detect due to quality control issues

GWTP Monthly Influent Sampling - June 2019

Location ID: Sample Date:	Units	LH18/24-SP140_060419 6/4/19
Location Description		GWTP – Collected from a spigot on the influent to TK-140. Sampled Monthly.
Metals (6020A)		
Selenium	mg/L	< 0.00250 U
Silver	mg/L	< 0.000500 U
Hexavalent Chromium (7196A)		
Hexavalent Chromium	mg/L	< 0.0100 U
Perchlorate (6850)		
Perchlorate	µg/L	6,800

mg/L - milligrams per liter

µg/L - micrograms per liter

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

LHAAP-58 Remedial Action Operation Validated Data - June 2019

Location ID: Sample Date:	Units	MCL/MSL	03WW01_06041 9 6/4/19	35AWW01_061019 6/10/19	35AWW05_061019 6/10/19	35AWW06_060419 6/4/19	35AWW08_060419 6/4/19	35AWW09_060319 6/3/19	35AWW10_060419 6/4/19	35AWW10_060419_a 6/4/19	35AWW11_060319 6/3/19	35AWW12_061019 6/10/19	35AWW12_061019a 6/10/19	35AWW13_060719 6/7/19	35AWW14_060719 6/7/19	35AWW15_060719 6/7/19	
Location Description				Site 58 - E, inside site boundary.	Site 58 - E, inside site boundary.	Site 58 - SW, outside site boundary.	Site 58 - SW, outside site boundary.	Site 58 - E, inside site boundary.	Site 58 - E, inside site boundary.	Site 58 - ESE, inside site boundary.	Site 58 - ESE, inside site boundary. Duplicate.	Site 58 - SE, inside site boundary.	Site 58 - E, outside site boundary.	Site 58 - E, outside site boundary. Field duplicate.	Site 58 - SE, outside site boundary.	Site 58 - SE, outside site boundary.	Site 58 - W, inside site boundary.
Location Depth				Shallow	Intermediate	Intermediate	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow
Dissolved Gases (RSK-175)																	
Carbon Dioxide	µg/L	NV	520,000	NA	NA	750,000	260,000	620,000	670,000	620,000	750,000	NA	NA	NA	NA	NA	NA
Ethane	µg/L	NV	< 0.47 U	NA	NA	< 0.47 U	< 0.47 U	< 0.47 U	< 0.47 U	< 0.47 U	< 0.47 U	NA	NA	NA	NA	NA	NA
Ethane	µg/L	NV	1.6	NA	NA	< 0.55 U	4.1	< 0.55 U	< 0.55 U	< 0.55 U	< 0.55 U	NA	NA	NA	NA	NA	NA
Methane	µg/L	NV	2,900	NA	NA	1,500	3,000	2	< 1.0 U	< 1.0 U	590	NA	NA	NA	NA	NA	NA
Dechlorinating Bacteria																	
BAV1 Vinyl Chloride Reductase	cells/mL	NV	< 0.50 U	NA	NA	0.80 J	< 3.0 U	< 0.7 U	< 0.5 U	NA	< 0.5 U	NA	NA	NA	NA	NA	NA
Dehalobacter spp.	cells/mL	NV	292	NA	NA	159	139000	73.3	261	NA	2570000	NA	NA	NA	NA	NA	NA
Dehalococoides	cells/mL	NV	6040	NA	NA	26.8	3790000	5.8	96.6	NA	2100	NA	NA	NA	NA	NA	NA
tceA Reductase	cells/mL	NV	2.1	NA	NA	< 1.6 U	480000	< 0.7 U	6.5	NA	< 0.5 U	NA	NA	NA	NA	NA	NA
Vinyl Chloride Reductase	cells/mL	NV	2020	NA	NA	0.90 J	265000	< 0.7 U	6.4	NA	174	NA	NA	NA	NA	NA	NA

Blue Highlighting Indicates concentrations above the MCL/MSL

MCL/MSL - Maximum Contaminant Limit/Medium-Specific Concentrations

NA - Not Analyzed

µg/L - micrograms per liter

mg/L - milligrams per liter

J - Estimated: Between the method detection limit and reporting limit and/or due to discrepancies in meeting certain analyte-specific quality control criteria.

U - The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

NV - No Value

UB - considered an artifact of blank contamination

LHAAP-58 Remedial Action Operation Validated Data - June 2019

Location ID: Sample Date:	Units	MCL/MSL	35AWW16_061019 6/10/19	35AWW17_061019 6/10/19	35AWW18_060419 6/4/19	35AWW19_060319 6/3/19	35AWW20_060419 6/4/19	34AWW21_061019 6/10/19	34AWW21_061019_a 6/10/19	35AWW22_061019 6/10/19	35AWW23_060319 6/3/19	35AWW24_060319 6/3/19	LHSMW06_060719 6/7/19	LHSMW07_060319 6/3/19	35ASW03_060719 6/7/19
Location Description			Site 58 - SW, outside site boundary, near Building 744-A.	Site 58 - SW, outside site boundary.	Site 58 - SSW, outside site boundary.	Site 58 - S, outside site boundary.	Site 58 - SW, inside site boundary.	Site 58 - ESE, outside site boundary, beside Building 725.	Site 58 - ESE, outside site boundary, beside Building 725. Field Duplicate.	Site 58 - ENE, outside site boundary.	Site 58 - SW, outside site boundary.	Downgradient Western Plume well	Site 58 - SW, inside site boundary, beside Building 715.	Site 58 - SW, outside site boundary.	Site 58 - SW, outside site boundary.
Location Depth			Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow	Shallow
Dissolved Gases (RSK-175)															
Carbon Dioxide	µg/L	NV	NA	NA	NA	560,000	730,000	NA	NA	NA	750,000	530,000	NA	660,000	NA
Ethane	µg/L	NV	NA	NA	NA	< 0.47 U	0.19 J	NA	NA	NA	< 0.47 U	< 0.47 U	NA	0.20	NA
Ethene	µg/L	NV	NA	NA	NA	< 0.55 U	1.3	NA	NA	NA	< 0.55 U	< 0.55 U	NA	3.9	NA
Methane	µg/L	NV	NA	NA	NA	1.4	920	NA	NA	NA	1.300	4.2	NA	78	NA
Dechlorinating Bacteria															
BAV1 Vinyl Chloride Reductase	cells/mL	NV	NA	NA	NA	< 0.50 U	2370	NA	NA	NA	2.5 J	< 0.5 U	NA	< 1.0 U	NA
Dehalobacter spp.	cells/mL	NV	NA	NA	NA	62700	< 21.7 U	NA	NA	NA	< 38.5 U	24.7	NA	< 9.5 U	NA
Dehalococoides	cells/mL	NV	NA	NA	NA	3.30	15500	NA	NA	NA	652	0.90	NA	7580	NA
tceA Reductase	cells/mL	NV	NA	NA	NA	< 0.50 U	< 2.20 U	NA	NA	NA	< 3.8 U	< 0.5 U	NA	< 1.0 U	NA
Vinyl Chloride Reductase	cells/mL	NV	NA	NA	NA	0.10 J	< 2.20 U	NA	NA	NA	72.7	< 0.5 U	NA	1620.0	NA

Blue Highlighting Indicates concentrations above the MCL/MSL

MCL/MSL - Maximum Contaminant Limit/Medium-Specific Concentrations

NA - Not Analyzed

µg/L - micrograms per liter

mg/L - milligrams per liter

J - Estimated: Between the method detection limit and reporting limit and/or due to disc

U - The analyte was not detected; however, the result is estimated due to discrepancies

U - Undetected: The analyte was analyzed for, but not detected.

NV - No Value

UB - considered an artifact of blank contamination

Subject: Final Minutes, Monthly Managers' Meeting (MMM),
Longhorn Army Ammunition Plant (LHAAP)
Location of Meeting: Karnack Community Center, Karnack, Texas
Date of Meeting: 21 August 2019– 2:00PM Central Daylight Time (CDT)

Attendees:

Army BRAC: Rose Zeiler (RMZ)
 EPA: Rich Mayer (RM) and Dorelle Harrison
 TCEQ: April Palmie (AP)
 USACE: Aaron Williams (AW)
 AEC: Amanda Sherman
 USGS: Kent Becher (KB)
 Bhate: Kim Nemmers (KN)
 USFWS: Paul Bruckwicki (PB)
 APTIM: Praveen Srivastav (PS) and Bill Foss (BF)

Action Items

Bhate/APTIM: KN stated that the proposed subset of wells for baseline sampling at LHAAP-16 had been discussed with the Army and that the proposed subset would be provided to the TCEQ and EPA by the end of the week for comment.

KN stated that the water line to extraction well 18WW17 repairs had not been scheduled. KN indicated that the Regulators would be notified once the repairs were scheduled.

Defense Environmental Restoration Program (DERP) Performance Based Remediation (PBR) Update

KN stated the groundwater treatment plant (GWTP) is operating properly. KN stated that the repairs to the conveyance lines at LHAAP-18/24 had resulted in improvements in the difference between the influent and effluent readings for the GWTP based upon the July 2019 evaluation of flow readings. KN also stated that the effluent started to be discharged to the INF pond as of August 20, 2019. KN explained that the effluent is going through the Fluidized Bed Reactor (FBR) and then the ion exchange vessels before discharging to the pond.

AW stated that the transformer was delivered and was to be tested on Wednesday, August 21, 2019. It is expected to be fully operational on Thursday, August 22, 2019. RM asked if the transformer had a warranty on it. AW confirmed that the transformer has a one year warranty. AP asked if there was additional surge protection. AW responded that the transformer should be able to withstand a power surge from lightning strike or storm.

PS stated that monitoring wells were installed at LHAAP-16, -12, -50 and -67. PS stated that the monitoring well at LHAAP-46 needs to be rehabilitated, which is planned for the next Tuesday, August 27, 2019. PS stated that the new wells have been surveyed. PS stated that the groundwater samples have been collected from new wells at LHAAP-16 and LHAAP-50, and data is expected by the end of next week. PS stated that LHAAP-17 is being excavated with mobilization on Monday, August 19, 2019. PS stated that the first loads of soil from LHAAP-17 went off on August 21, 2019. BF stated that 10 total loads were expected to go offsite based upon the turnaround time for each truck of 3.25 hours. PS stated that the plan was to increase to six trucks running the next day (Thursday, August 22, 2019). RMZ asked about the capacity of each truck. BF stated that the trucks are about 18 cubic yards each. KN stated that work was planned for Saturdays also. PS

stated that the landfill is open for part of a day on Saturdays. KN stated that the excavation is starting on the outside of the site. BF stated that the area being excavated is along the western edge of Area H and the excavation is working inwards. AP stated that the daily reports have been helpful.

KN asked everyone to refer to the Document and Issues Tracking Table dated August 21, 2019.

- **Task 1** (Project Management) -KN stated that concurrence/comments were received by the TCEQ on the MMM minutes and the Restoration Advisory Board (RAB) minutes. RM stated the DH and KB would review and that he would provide concurrence/comments.
- **Task 3** (LHAAP-03) – PS stated that there are no current documents. BF stated that the plan is to mark the areas to excavate at LHAAP-03, following removal at LHAAP-17.
- **Task 4** (LHAAP-04) – PS stated that there are no current documents.
- **Task 5** (LHAAP-12) – PS stated that the monitoring well was installed and developed in August 2019, and that the new monitoring well would be sampled in December 2019 during the annual RA-O event.
- **Task 6** (LHAAP-16) – PS stated that injections will begin mid-September and is likely to occur the week of September 23, 2019. PS stated that pre-injection, baseline sampling will be completed prior to the injections and following two weeks of extraction. KN stated that once power is restored, the system at LHAAP-16 will be assessed to determine if or what repairs are needed. Repairs to the extraction system at LHAAP-16 could push the schedule out. KN stated that the requirements for repairs should be known by Friday.
- **Task 7** (LHAAP-17) – PS stated that no documents are currently in progress for LHAAP-17. PS explained that even the area on the northern eastern end of the site is dried up.
- **Task 9** (LHAAP-37) – PS stated that the Year 2, Quarter 4 sampling was completed in August 2019.
- **Task 10** (LHAAP-46) – PS stated that the Year 5 2nd Semi-Annual RA-O event was completed in August 2019. PS stated that the monitoring well rehabilitation/ replacement is planned for Tuesday, 8/27/19. If this well can be sampled, PS explained that the data will be included in the Year 5 2nd Semi-Annual sampling event and in the Year 5 Annual RA-O Report.
- **Task 11** (LHAAP-50) – PS stated that the annual report will be prepared with the last sampling event that occurred in May 2019. PS stated that the new monitoring well had been installed, developed and sampled on 8/15/19. PS explained that the results were due by Friday based upon a 5-day turnaround time. The results will be evaluated to determine if changes to the Contingency RD/RAWP were needed. PS asked RM if there was an update on the EPA's signature of the ESD. RM stated that he would find out the status of the ESD signatures.
- **Task 12** (LHAAP-58) – KN stated that the Year 5 RA(O)Report was in Bhate's internal quality control (QC) review and was planned for delivery to the Regulators in October 2019.
- **Task 13** (LHAAP-67) – PS stated that the Year 5 Annual RA-O Report will be prepared with the last sampling event that occurred in May 2019. PS stated that the two new monitoring wells had been installed and developed in August 2019. PS stated that the monitoring well would be sampled in November 2019 during the Year 6 Annual sampling event.
- **Task 14** (LHAAP-001-R and -003-R) - KN stated that the Annual LTM Report was in Bhate's internal QC review and was planned for delivery to the Regulators in October 2019.
- **Task 16** (GWTP) –KN indicated that the 2nd Quarter 2019 GWTP was under Bhate's internal QC and is anticipated for delivery to the Regulators in October 2019.

- **Task 17** (LHAAP-18/24) – KN stated that the validated data available for LHAAP-18/24 had been provided with the validated data provided for the MMM. KN explained that the data from two monitoring wells that could not be sampled due to physically not being able to access the area. KN stated that those wells were sampled in August 2019.
- **Task 18** (Surface Water) – KN stated that surface water had been sampled in July 2019. KN asked if that data had been shared with the TCEQ and EPA. AP and RM stated that they knew of the surface water sample that had exceeded and understood that the resample did not exceed. KN confirmed this to be correct.
- **Task 19** (LUC Management Plan) – KN stated that the LUC Management Plan Update was being completed, but that the DI tracker did not show it being delivered to the Regulators until December 2019. KN stated that the goal was to beat this date.
- **Administrative Record (AR)** – BF stated that additional documents were received for the April 2019 list but that he didn't have an update on the production of January through March 2019 AR. PS stated that he thought the AR for that time period was being compiled still.

Update on other DERP Sites

- **LHAAP 18/24** – AW explained that the Final Proposed Plan (PP) for LHAAP-18/24 will be placed into the AR. AW stated that Record of Decision (ROD) is with Army's Environmental Law Division (ELD) and Assistant Chief of Staff for Installation Management (ACSIM) and that the Army is working to meet the enforceable schedule date of September 6, 2019.
- **LHAAP-29** – AW stated that the LHAAP-29 PP was placed into the AR. AW stated that the ROD was submitted by the Army for EPA signature and TCEQ concurrence in August 2019.
- **LHAAP-47** – AW stated that the Post-Screening Investigation (PSI) Report will be placed into the latest March 2019 AR. AW stated that the Addendum to the PSI Report, will be included in the follow on update to the AR. AW stated that based upon the results at monitoring well WW25R [replacement monitoring well with 120,000 parts per billion of trichloroethylene] that the ROD is being delayed for additional investigation in the area, which will be included in Addendum Number 2 to the PSI Report. AW pointed out that the ROD submittal data was pushed out in the DI tracker to September 30, 2020.
- **Other items** –
RMZ asked PB if he knew the transfer status of the Environmental Condition of Property 7 parcels and PB said he had not.
KB stated that two of the samplers had been removed from the Bayou and Goose Prairie Creek. KB stated that the pipe was left in these locations however.

Schedule Next Managers' Meeting

The next MMM will be held on September 18, 2019 at 1:00 CDT via a conference call. In general, calls will be the third Wednesday of each month.

Meeting concluded at 2:38 pm CDT.

ACRONYM LIST

AP	April Palmie
APTIM	APTIM Federal Services, LLC
AR	Administrative Record
AW	Aaron Williams
BF	Bill Foss

Bhate	Bhate Environmental Associates, Inc.
BRAC	Base Realignment and Closure
CDT	Central Daylight Time
DERP	Defense Environmental Restoration Program
DNAPL	Dense Non-Aqueous Phase Liquid
ELD	Environmental Law Division
EPA	United States Environmental Protection Agency
ESD	Explanation of Significant Differences
FBR	Fluidized Bed Reactor
FYR	Five Year Review
GWTP	Ground Water Treatment Plant
KN	Kim Nemmers
LHAAP	Longhorn Army Ammunition Plant
MMM	Monthly Managers' Meeting
PB	Paul Bruckwicki
PBR	Performance-Based Remediation
PP	Proposed Plan
PS	Praveen Srivastav
PSI	Post-Screening Investigation
RAB	Restoration Advisory Board
RA(O)	remedial action – operation
RAWP	Remedial Action Work Plan
RD	Remedial Design
RMZ	Rose M. Zeiler
ROD	Record of Decision
TCEQ	Texas Commission on Environmental Quality
USACE	United States Army Corps of Engineers
USFWS	United States Fish and Wildlife Service

**LHAAP Data Validated
August MMM Validated Data**

LHAAP Surface Water	<i>Quarterly Perchlorate Sampling - July 2019</i> Perchlorate (6850)
GWTP Effluent	<i>Weekly Perchlorate Sampling – July 2019</i> Perchlorate (6850)
GWTP Effluent	<i>Weekly, Bi-Weekly, and Monthly Sampling – July 2019</i> Ammonia (350.3) Ortho-Phosphate (365.3) Organic Carbon (415.1) VOC (8260C) Metals (6020A) Hexavalent Chromium (7196A) 1,4-Dioxane (8270D-SIM) Anions (9056)
GWTP Influent	<i>Monthly Sampling – July 2019</i> Metals (6020A) Perchlorate (6850) Hexavalent Chromium (7196A)
LHAAP-18/24	<i>Sampling – June 2019</i> Perchlorate (6850) Metals (6020A) VOCs (8260C) 1,4- Dioxane (8270D SIM)

LHAAP-Quarterly Surface Water Sampling - July 2019

Location ID: Sample Date:	Units	PCL	HBW7-071119 7/11/19	HBW7-073019 7/30/19*	HBW10_071119 7/11/19	HBW1_071119 7/11/19	GPW1_071119 7/11/19	GPW1_071119_a 7/11/19	GPW3_071119 7/11/19	
Perchlorate (6850)				Harrison Bayou				Goose Prairie Creek		
Perchlorate	µg/L	17	27	1.2 J	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U	< 2.0 U	

PCL – Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level

µg/L - micrograms per liter

U - Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

Exceeded PCL screening criteria

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

*resampled

GWTP Weekly Sampling - July 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_070919 7/9/19	LH18/24-SP650_071619 7/16/19	LH18/24-SP650_072319 7/23/19	LH18/24-SP650_073019 7/30/19
Location Description			GWTP—Collected from a spigot on the discharge of effluent TK-650. Sampled Weekly.			
Ammonia as N (350.3)						
Ammonia as N	mg/L	NV	6.9	7.5	9.2	7.2
Ortho-Phosphate (365.3)						
Ortho-Phosphate	mg/L	NV	2.19	0.276	2.68	0.19
Organic Carbon (415.1)						
Total Organic Carbon (TOC)	mg/L	NV	5.09 J	2.52	1.96	2.33

mg/L - milligrams per liter

NV - No Value

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

GWTP Bi-Weekly Sampling - July 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_070919 7/9/19	LH18/24-SP650_072319 7/23/19
Location Description		GWTP – Collected from a spigot on the discharge of effluent TK-650. Sampled Biweekly.		
Volatile Organic Compounds (8260C)				
1,1,1-Trichloroethane	µg/L	7,230	< 0.5 U	< 0.5 U
1,1,2-Trichloroethane	µg/L	216.9	< 0.5 U	< 0.5 U
1,1-Dichloroethane	µg/L	14,032	< 0.5 U	< 0.5 U
1,1-Dichloroethene	µg/L	253	< 0.5 U	< 0.5 U
1,2-Dichloroethane	µg/L	181	< 0.5 U	< 0.5 U
1,2-Dichloropropane	µg/L	5	< 0.5 U	< 0.5 U
Acetone	µg/L	2,395	< 1.0 U	4.6
Benzene	µg/L	181	< 0.5 U	< 0.5 U
Carbon tetrachloride	µg/L	181	< 0.5 U	< 0.5 U
Chlorobenzene	µg/L	47,180	< 0.5 U	< 0.5 U
Chloroform	µg/L	3,615	< 0.5 U	< 0.5 U
Ethylbenzene	µg/L	57,025	< 0.5 U	< 0.5 U
m,p-Xylene	µg/L	83.6	< 1.0 U	< 1.0 U
Methylene chloride	µg/L	1,699	< 1.0 U	< 1.0 U
o-Xylene	µg/L	83.6	< 0.5 U	< 0.5 U
Styrene	µg/L	5,987	< 0.5 U	< 0.5 U
Tetrachloroethene	µg/L	180.7	< 0.5 U	< 0.5 U
Toluene	µg/L	4,189	< 0.5 U	< 0.5 U
Trichloroethene	µg/L	181	1.0	0.88 J
Vinyl chloride	µg/L	72	< 0.5 U	< 0.5 U
Anions (9056)				
Chloride	mg/L	NV	292	494
Sulfate	mg/L	NV	29.9	32.3

µg/L - micrograms per liter

mg/L - milligrams per liter

U - Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

NV - No Value

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

GWTP Monthly Effluent Sampling - July 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24-SP650_071619 7/16/19
Location Description		GWTP – Collected from a spigot on the discharge of effluent TK-650. Sampled monthly	
Volatile Organic Compounds (8260C)			
1,1,1-Trichloroethane	µg/L	7,230	< 0.5 U
1,1,2-Trichloroethane	µg/L	216.9	< 0.5 U
1,1-Dichloroethane	µg/L	14,032	< 0.5 U
1,1-Dichloroethene	µg/L	253	< 0.5 U
1,2-Dichloroethane	µg/L	181	0.52 J
1,2-Dichloropropane	µg/L	5	< 0.5 U
Acetone	µg/L	2,395	< 1.0 U
Benzene	µg/L	181	< 0.5 U
Carbon tetrachloride	µg/L	181	< 0.5 U
Chlorobenzene	µg/L	47,180	< 0.5 U
Chloroform	µg/L	3,615	< 0.5 U
Ethylbenzene	µg/L	57,025	< 0.5 U
m,p-Xylene	µg/L	83.6	< 1.0 U
Methylene chloride	µg/L	1,699	< 1.0 U
o-Xylene	µg/L	83.6	< 0.5 U
Styrene	µg/L	5,987	< 0.5 U
Tetrachloroethene	µg/L	180.7	< 0.5 U
Toluene	µg/L	4,189	< 0.5 U
Trichloroethene	µg/L	181	0.99 J
Vinyl chloride	µg/L	72	< 0.5 U
Metals (6020A)			
Barium	mg/L	2	0.0903
Lead	mg/L	0.0046	< 0.00100 U
Selenium	mg/L	0.012	< 0.00250 U
Silver	mg/L	0.003	< 0.000500 U
Hexavalent Chromium (7196A)			
Hexavalent Chromium	mg/L	0.1244	0.0100
Semi-Volatile Organic Compounds (8270D SIM)			
1,4-Dioxane	µg/L	134.2	12

µg/L - micrograms per liter

mg/L - milligrams per liter

U - Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

J - estimated value between the detection limit and limit of quantitation and/or due to quality control issues

GWTP Weekly/Effluent Perchlorate Sampling - July 2019

Location ID: Sample Date:	Units	Daily Maximum Conc	LH18/24- SP650_070919_BIX 7/9/19	LH18/24-SP650_071619- BIX 7/16/19	LH18/24- SP650_071619_BIX 7/16/19	LH18/24- SP650_072319_BIX 7/23/19	LH18/24- SP650_073019_BIX 7/30/19
Location Description		Collected from a spigot on the discharge of effluent TK-650.					
			Weekly	Monthly EFF	Weekly	Weekly	Weekly
Perchlorate (6850)							
Perchlorate	µg/L	589	5.8	< 2.0 U	< 2.0 U	42	< 2.0 U

µg/L - micrograms per liter

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

BIX - before ion exchange

GWTP Monthly Influent Sampling - July 2019

Location ID:		LH18/24-SP140_071619
Sample Date:	Units	7/16/19
Location Description		GWTP – Collected from a spigot on the influent to TK-140. Sampled Monthly.
Metals (6020A)		
Selenium	mg/L	< 0.00250 U
Silver	mg/L	< 0.00250 U
Hexavalent Chromium (7196A)		
Hexavalent Chromium	mg/L	< 0.0100 U
Perchlorate (6850)		
Perchlorate	µg/L	6,900

mg/L - milligrams per liter

µg/L - micrograms per liter

U- Undetected: The analyte was analyzed for, but not detected and reported to the limit of detection.

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/PC CL	AWD1_062019 6/20/19	AWD3_062119 6/21/19	AWD4_062619 6/26/19	AWD4_062619_a 6/26/19	18CPTMW01DW _061919 6/19/19	18CPTMW01SW _061919 6/19/19	18CPTMW03SW_ _062119 6/21/19	18CPTMW04 _061919 6/19/19	18CPTMW04SW _061919 6/19/19	18CPTMW06 _062119 6/21/19	18CPTMW06 _062119_a 6/21/19	18CPTMW07 _062519 6/25/19	18CPTMW08SW _061819 6/18/19	18CPTMW08DW _061819 6/18/19	18CPTMW10SW _062019 6/20/19	18CPTMW10DW _062019 6/20/19	18CPTMW12SW _061719 6/17/19	18CPTMW12SW_a _061719_a 6/17/19	18CPTMW12DW _061719 6/17/19	18CPTMW14 _062519 6/25/19
Lab Package	Well ID		HS19061164	HS19061210	HS19061482	HS19061482	HS19061083	HS19061083	HS19061210	HS19061083	HS19061083	HS19061210	HS19061210	HS19061383	HS19060986	HS19060986	HS19061157	HS19061157	HS19060929	HS19060929	HS19060929	HS19061383
Perchlorate (6850)			AWD-1	AWD-3	AWD-4	AWD-4	18CPTMW01DW	18CPTMW01SW	18CPTMW03SW	18CPTMW04	18CPTMW04SW	18CPTMW06	18CPTMW06	18CPTMW07	18CPTMW08SW	18CPTMW08DW	18CPTMW10SW	18CPTMW10DW	18CPTMW12SW	18CPTMW12SW	18CPTMW12DW	18CPTMW14
Perchlorate	µg/L	17*	NA	1.8 J	23	27	<2.0 U	<2.0 U	9.9	620	<2.0 U	1.7 J	<2.0 U	1.4 J	23,000	4,300	<2.0 U	9.1	<2.0 U	<2.0 U	1.2 J	1,300
Volatile Organic Compounds (8260C)																						
1,1,1,2-Tetrachloroethane	µg/l	110	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,1,1-Trichloroethane	µg/l	200	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,1,2,2-Tetrachloroethane	µg/l	14	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,1,2-Trichloroethane	µg/l	5	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,1-Dichloroethane	µg/l	10,000	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	2.1	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,1-Dichloroethane	µg/l	7	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,1-Dichloropropene	µg/l	2.9	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2,3-Trichlorobenzene	µg/l	310	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2,3-Trichloropropane	µg/l	0.041	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2,4-Trichlorobenzene	µg/l	70	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2,4-Trimethylbenzene	µg/l	5,100	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2-Dibromo-3-chloropropane	µg/l	0.2	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2-Dibromoethane	µg/l	0.05	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2-Dichlorobenzene	µg/l	600	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,2-Dichloroethane	µg/l	5	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	1.3	0.75 J	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	2.1
1,2-Dichloropropane	µg/l	5	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,3,5-Trimethylbenzene	µg/l	5,100	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,3-Dichlorobenzene	µg/l	3,100	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,3-Dichloropropane	µg/l	29	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
1,4-Dichlorobenzene	µg/l	75	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
2,2-Dichloropropane	µg/l	42	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
2-Butanone	µg/l	61,000	NA	<10 U	<1.0 U	<1.0 U	<1.0 U	<25 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	NA	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U
2-Chlorotoluene	µg/l	2,000	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
2-Hexanone	µg/l	6,100	NA	<10 U	<1.0 U	<1.0 U	<1.0 U	<25 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	NA	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U
4-Chlorotoluene	µg/l	2,000	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
4-Isopropyltoluene	µg/l	10,000	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
4-Methyl-2-pentanone	µg/l	8,200	NA	<10 U	<1.0 U	<1.0 U	<1.0 U	<25 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	NA	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U
Acetone	µg/l	92,000	NA	<10 U	<1.0 U	6.8	5.1 UB	<25 U	5.2	4.6 UB	5 UB	3.1	<1.0 U	<1.0 U	2.7 UB	NA	5.4 UB	3.3 UB	2.9 UB	3 UB	3.2 UB	<1.0 U
Benzene	µg/l	5	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	1.3
Bromobenzene	µg/l	2,000	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Bromochloromethane	µg/l	4,100	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Bromodichloromethane	µg/l	4.6	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Bromoform	µg/l	36	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Bromomethane	µg/l	140	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Carbon disulfide	µg/l	10,000	NA	<10 U	<1.0 U	<1.0 U	<1.0 U	<25 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	NA	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U
Carbon tetrachloride	µg/l	5	NA	35	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Chlorobenzene	µg/l	100	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Chloroethane	µg/l	41,000	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
Chloroform	µg/l	1,000	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	6.7 J
Chloromethane	µg/l	220	NA	<5.0 U	<0.5 U	<0.5 U	<0.5 U	<12 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	NA	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U
cis-1,2-Dichloroethane	µg/l	70</																				

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Location ID: Sample Date:	Units	MCL/MSC/P CL	AWD1_062019 6/20/19	AWD3_062119 6/21/19	AWD4_062619 6/26/19	AWD4_062619_a 6/26/19	18CPTMW01DW _061919 6/19/19	18CPTMW01SW _061919 6/19/19	18CPTMW03SW_ 062119 6/21/19	18CPTMW04 _061919 6/19/19	18CPTMW04SW _061919 6/19/19	18CPTMW06 _062119 6/21/19	18CPTMW06 _062119_a 6/21/19	18CPTMW07 _062519 6/25/19	18CPTMW08SW -061819 6/18/19	18CPTMW08DW -061819 6/18/19	18CPTMW10SW _062019 6/20/19	18CPTMW10DW _062019 6/20/19	18CPTMW12SW _061719 6/17/19	18CPTMW12SW _061719_a 6/17/19	18CPTMW12DW _061719 6/17/19	18CPTMW14 _062519 6/25/19
n-Propylbenzene	µg/l	4,100	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
o-Xylene	µg/l	10,000**	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
sec-Butylbenzene	µg/l	4,100	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
Styrene	µg/l	100	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
tert-Butylbenzene	µg/l	4,100	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
Tetrachloroethene	µg/l	5	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	1.1
Toluene	µg/l	1,000	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
trans-1,2-Dichloroethene	µg/l	100	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	1.4	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
trans-1,3-Dichloropropene	µg/l	29	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
Trichloroethene	µg/l	5	NA	580	< 0.5 U	< 0.5 U	< 0.5 U	32	19	510	< 0.5 U	0.98 J	0.88 J	0.66 J	40	NA	< 0.5 U	1.2	0.65 J	0.60 J	< 0.5 U	700
Trichlorofluoromethane	µg/l	31,000	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
Vinyl chloride	µg/l	2	NA	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 12 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	5.0	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U
Metals (6020A)																						
Aluminum	mg/L	100	NA	0.157	20.1 J	9.41 J	0.110	0.0504	0.0446	NA	0.0645	NA	NA	NA	NA	NA	NA	0.611	0.0204	0.0248	0.0201	0.427
Antimony	mg/L	0.006	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000408 J	< 0.000500 U	< 0.000500 U	NA	< 0.000500 U	NA	NA	NA	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000406 J	0.000513 UB
Arsenic	mg/L	0.01	NA	< 0.000500 U	0.00257 J	0.00158 J	0.00152 J	0.00871	0.00272 J	NA	0.00326 J	NA	NA	NA	NA	NA	NA	0.00262 J	0.000866 J	0.00106 J	0.00401 J	0.00240 J
Barium	mg/L	2	NA	0.0352	0.267	0.239	0.412	0.172	0.172	NA	0.911	NA	NA	NA	NA	NA	NA	0.108	0.803	0.825	0.133	6.03
Beryllium	mg/L	0.004	NA	< 0.000500 U	0.000982 J	0.000691 J	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	< 0.000500 U	NA	NA	NA	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Cadmium	mg/L	0.005	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000345 J	< 0.000500 U	0.000367 J	NA	< 0.000500 U	NA	NA	NA	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000243 J
Calcium	mg/L	NV	NA	0.817	11.3	10.7	40.3	32.9	21.0	NA	30.3	NA	NA	NA	NA	NA	NA	6.74	58.5	58.7	9.65	466
Chromium	mg/L	0.1	NA	0.0808	0.387 J	0.193 J	0.0949	0.00638	0.0163	NA	0.0219	NA	NA	NA	NA	NA	NA	0.00736	0.0159	0.0160	0.0136	0.00598
Cobalt	mg/L	6.1	NA	0.000846 J	0.0121 J	0.00828 J	0.00121 J	0.000511 J	0.00263 J	NA	0.00473 J	NA	NA	NA	NA	NA	NA	0.000393 J	0.00248 J	0.00304 J	0.000115 J	0.0142
Copper	mg/L	1.3	NA	< 0.00100 U	0.0215 J	0.0140 J	0.00332 UB	< 0.00100 U	< 0.00100 U	NA	< 0.00100 U	NA	NA	NA	NA	NA	NA	0.00188 UB	< 0.00100 U	< 0.00100 U	< 0.00100 U	0.00130 J
Iron	mg/L	NV	NA	0.430	22.3 J	11.0 J	1.65	51.3	1.75	NA	10.4	NA	NA	NA	NA	NA	NA	2.80	0.540	0.669	1.10	2.89
Lead	mg/L	0.015	NA	< 0.00100 U	0.00842 J	0.00448 J	0.00104 J	< 0.00100 U	< 0.00100 U	NA	< 0.00100 U	NA	NA	NA	NA	NA	NA	0.000967 J	< 0.00100 U	< 0.00100 U	< 0.00100 U	< 0.00100 U
Magnesium	mg/L	NV	NA	0.586	5.94	5.31	11.7	22.6	7.22	NA	18.3	NA	NA	NA	NA	NA	NA	5.34	38.8	39.2	4.67	157
Manganese	mg/L	1.1*	NA	0.00771	0.0703	0.0536	0.115	0.635	0.0478	NA	0.455	NA	NA	NA	NA	NA	NA	0.0448	0.181	0.232	0.0329	0.793
Nickel	mg/L	0.49*	NA	0.0253	0.240	0.227	0.00669	0.00112 J	0.00956	NA	0.00641	NA	NA	NA	NA	NA	NA	0.00202 J	0.00382 J	0.00361 J	0.000995 J	0.00836
Potassium	mg/L	NV	NA	0.746	1.29 J	0.827 J	147	18.4	233	NA	108	NA	NA	NA	NA	NA	NA	111	75.5	75.0	108	17.5
Selenium	mg/L	0.05	NA	0.00434 J	0.00245 J	0.00328 J	< 0.00250 U	< 0.00250 U	< 0.00250 U	NA	< 0.00250 U	NA	NA	NA	NA	NA	NA	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U
Silver	mg/L	0.51	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	< 0.000500 U	NA	NA	NA	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Sodium	mg/L	NV	NA	30.1	62.6	62.3	370	115	265	NA	130	NA	NA	NA	NA	NA	NA	184	234	229	184	605
Thallium	mg/L	0.002	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	< 0.000500 U	NA	NA	NA	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Vanadium	mg/L	0.72	NA	0.00191 UB	0.0366 J	0.0207 J	0.00102 UB	< 0.00100 U	0.00183 UB	NA	< 0.00100 U	NA	NA	NA	NA	NA	NA	0.00163 UB	0.00102 UB	0.00150 UB	0.00169 UB	0.00212 J
Zinc	mg/L	31	NA	0.0177	0.0883 J	0.0588 J	0.0266	0.0209	0.0987	NA	0.0400	NA	NA	NA	NA	NA	NA	0.0244	0.0181	0.0159	0.0198	0.0184
Mercury	mg/L	0.002	NA	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	NA	< 0.000100 U	NA	NA	NA	NA	NA	NA	< 0.000100 U	< 0.000100 U	< 0.000100 U	0.0000310 J	< 0.000100 U
1,4-Dioxane (8270D SIM)																						
1,4-Dioxane	µg/l	9.1	0.99	NA	< 0.010 U	< 0.010 U	0.27	< 0.010 U	0.67	1.3	< 0.010 U	0.46 J	0.052 J	0.077	1.7	0.38	< 0.010 U	NA	0.073 J	0.18 J	0.11	0.13

Notes:
 Blue highlighting indicates concentrations above the MCL/MSC/PCL
 MCL/MSC - Maximum Contaminant Limit/Medium-Specific Concentrations/Protective Concentration Level
 NA - Not Analyzed
 µg/L - micrograms per liter a - duplicate sample
 mg/L - milligrams per liter
 J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.
 UJ - The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.
 U - Undetected: The analyte was analyzed for, but not detected.
 NV - No Value
 UB - considered a non-detect due to blank contamination
 *Perchlorate, manganese, and nickel compared to the PCL
 ** Value is for total xylenes
 PCL – Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential Protective Concentration Level

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	18CPTMW15 _062119 6/21/19	18CPTMW16 _062719 6/27/19	18CPTMW18 _062619 6/26/19	18CPTMW19 _062019 6/20/19	18CPTMW19SW _062019 6/20/19	18CPTMW22R _061319 6/13/19	18CPTMW22SW_0 61319 6/13/19	18CPTMW22DW_ 061319 6/13/19	18CPTMW23 -061719 6/17/19	18CPTMW23SW -061719 6/17/19	18CPTMW24 _062519 6/25/19	18CPTMW26SW_ 061319 6/13/19	17WW08 _062519 6/25/19	18WW02 _062019 6/20/19	18WW03 _062419 6/24/19	18WW03 _062419_a 6/24/19	18WW06 _062019 6/20/19	18WW08 _062719 6/27/19	18WW09 _062719 6/27/19	18WW10 _062519 6/25/19	18WW10 _062519-a 6/25/19	18WW14 _061719 6/17/19		
n-Propylbenzene	µg/l	4,100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
o-Xylene	µg/l	10,000**	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
sec-Butylbenzene	µg/l	4,100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
Styrene	µg/l	100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
tert-Butylbenzene	µg/l	4,100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
Tetrachloroethene	µg/l	5	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
Toluene	µg/l	1,000	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
trans-1,2-Dichloroethene	µg/l	100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
trans-1,3-Dichloropropene	µg/l	29	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
Trichloroethene	µg/l	5	2.3	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	2,900	< 0.5 U	9.1	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
Trichlorofluoromethane	µg/l	31,000	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
Vinyl chloride	µg/l	2	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	NA	
Metals (6020A)																										
Aluminum	mg/L	100	NA	NA	0.0122 UB	NA	0.0745	1.54	0.284	0.133	NA	NA	NA	NA	0.0212 UB	1.56	0.0156 UB	0.0175 UB	NA	NA	0.0863	NA	NA	NA	0.138	
Antimony	mg/L	0.006	NA	NA	< 0.000500 U	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	NA	NA	< 0.005000 U	< 0.005000 U	< 0.005000 U	< 0.005000 U	NA	NA	< 0.005000 U	NA	NA	< 0.005000 U	NA	
Arsenic	mg/L	0.01	NA	NA	0.00127 J	NA	0.00543	0.00317 J	0.00358 J	0.00316 J	NA	NA	NA	NA	0.00576	0.000675 J	0.000834 J	0.000750 J	NA	NA	0.00754	NA	NA	0.000663 J	NA	
Barium	mg/L	2	NA	NA	0.608	NA	0.198	0.0493	0.0799	0.0930	NA	NA	NA	NA	0.539	0.0506	0.194	0.185	NA	NA	0.239	NA	NA	NA	1.19	
Beryllium	mg/L	0.004	NA	NA	< 0.000500 U	NA	< 0.000500 U	0.000337 J	< 0.000500 U	< 0.000500 U	NA	NA	NA	NA	< 0.005000 U	< 0.000500 U	< 0.005000 U	< 0.005000 U	NA	NA	< 0.005000 U	NA	NA	< 0.005000 U	NA	
Cadmium	mg/L	0.005	NA	NA	< 0.000500 U	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	NA	NA	< 0.005000 U	< 0.000500 U	< 0.005000 U	< 0.005000 U	NA	NA	0.000214 J	NA	NA	NA	0.000783 J	
Calcium	mg/L	NV	NA	NA	281	NA	11.2	0.683	24.5	12.4	NA	NA	NA	NA	129	7.84	8.98	8.70	NA	NA	17.9	NA	NA	NA	50.8	
Chromium	mg/L	0.1	NA	NA	0.00200 J	NA	0.00171 J	0.00400 J	0.0486	0.000531 J	NA	NA	NA	NA	0.438	0.00851	0.0540 J	0.0356 J	NA	NA	0.0175	NA	NA	0.409	NA	
Cobalt	mg/L	6.1	NA	NA	0.0139	NA	0.0104	0.00659	< 0.000500 U	0.000402 J	NA	NA	NA	NA	0.0156	0.000341 J	0.000142 J	< 0.005000 U	NA	NA	0.00123 J	NA	NA	0.0132	NA	
Copper	mg/L	1.3	NA	NA	< 0.00100 U	NA	< 0.00100 U	0.00172 UB	< 0.00100 U	< 0.00100 U	NA	NA	NA	NA	0.00484 J	0.00663 UB	< 0.00100 U	< 0.00100 U	NA	NA	< 0.00100 U	NA	NA	0.0119	NA	
Iron	mg/L	NV	NA	NA	1.06	NA	16.1	2.90	0.0723 J	0.832	NA	NA	NA	NA	29.0	2.46	5.60	5.40	NA	NA	39.4	NA	NA	NA	1.90	
Lead	mg/L	0.015	NA	NA	< 0.00100 U	NA	< 0.00100 U	0.000894 J	< 0.00100 U	< 0.00100 U	NA	NA	NA	NA	< 0.00100 U	0.00123 J	< 0.00100 U	< 0.00100 U	NA	NA	< 0.00100 U	NA	NA	< 0.00100 U	NA	
Magnesium	mg/L	NV	NA	NA	194	NA	5.98	0.800	2.20	6.18	NA	NA	NA	NA	85.6	1.44	6.34	6.12	NA	NA	9.91	NA	NA	NA	40.4	
Manganese	mg/L	1.1*	NA	NA	2.21	NA	0.477	0.0807	0.0223	0.0603	NA	NA	NA	NA	1.62	0.0988	0.117	0.112	NA	NA	0.792	NA	NA	NA	0.876	
Nickel	mg/L	0.49*	NA	NA	0.0131	NA	0.00246 J	0.00370 J	0.00100 J	0.000918 J	NA	NA	NA	NA	0.370	0.00574	0.00325 J	0.00335 J	NA	NA	0.0111	NA	NA	NA	0.306	
Potassium	mg/L	NV	NA	NA	2.98	NA	1.68	0.542	379	2.85	NA	NA	NA	NA	2.12	2.12	1.97	1.85	NA	NA	2.58	NA	NA	NA	4.38	
Selenium	mg/L	0.05	NA	NA	< 0.00250 U	NA	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	NA	NA	NA	NA	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	NA	NA	< 0.00250 U	NA	NA	< 0.00250 U	NA	
Silver	mg/L	0.51	NA	NA	< 0.000500 U	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	< 0.000500 U	NA	NA	< 0.000500 U	NA	
Sodium	mg/L	NV	NA	NA	746	NA	28.1	20.8	313	234	NA	NA	NA	NA	527	25.4	111	106	NA	NA	42.5	NA	NA	NA	137	
Thallium	mg/L	0.002	NA	NA	< 0.000500 U	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	< 0.000500 U	NA	NA	< 0.000500 U	NA	
Vanadium	mg/L	0.72	NA	NA	< 0.00100 U	NA	0.00184 UB	0.00954	0.00200 UB	0.00216 UB	NA	NA	NA	NA	0.00169 J	0.00394 UB	0.00237 UB	0.00240 UB	NA	NA	0.00236 UB	NA	NA	0.00422 UB	NA	
Zinc	mg/L	31	NA	NA	0.0225	NA	0.0247	0.0183	0.0294	0.0220	NA	NA	NA	NA	0.0291	0.0147	0.0171	0.0163	NA	NA	0.0131	NA	NA	NA	0.0390	
Mercury	mg/L	0.002	NA	NA	< 0.000100 U	NA	< 0.000100 U	< 0.000100 U	0.0000360 J	< 0.000100 U	NA	NA	NA	NA	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	NA	NA	< 0.000100 U	NA	NA	< 0.000100 U	NA	
1,4-Dioxane (8270D SIM)																										
1,4-Dioxane	µg/l	9.1	0.43	0.048	0.021	0.068	< 0.010 U	< 0.010 U	0.95	0.15	8.1	< 0.010 U	< 0.010 U	0.024	NA	< 0.010 U	< 0.010 U	< 0.010 U	< 0.010 U	< 0.010 U	< 0.010 U	< 0.010 U	< 0.010 U	< 0.010 U	NA	

Notes:
 Blue highlighting indicates concentrations above the MCL/MSC/PCL
 MCL/MSC - Maximum Contaminant Limit/Medium-Specific Concentrations/Protective Concentration Level
 NA - Not Analyzed
 µg/L - micrograms per liter a - duplicate sample
 mg/L - milligrams per liter
 J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.
 UJ - The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.
 U - Undetected: The analyte was analyzed for, but not detected.
 NV - No Value
 UB - considered a non-detect due to blank contamination
 *Perchlorate, manganese, and nickel compared to the PCL
 ** Value is for total xylenes
 PCL - Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	18WW16 _062619 6/26/19	18WW17 _062419 6/24/19	18WW18 _062419 6/24/19	18WW19	18WW20	18WW22 _062019 6/20/19	18WW24 _062619 6/26/19	18WW25 _062719 6/27/19	C01_062119 6/21/19	C02_062519 6/25/19	C03_062619 6/26/19	C04_062019 6/20/19	C06_062619 6/26/19	C08_062119 6/21/19	C09_062619 6/26/19	MW1_061919 6/19/19	MW2_061919 6/19/19	MW2_061919_ a 6/19/19	MW3_062519 6/25/19	MW5_061819 6/18/19	MW6_061819 6/18/19	MW7_061719 6/17/19		
n-Propylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
o-Xylene	µg/l	10,000**	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
sec-Butylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
Styrene	µg/l	100	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
tert-Butylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
Tetrachloroethene	µg/l	5	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	43 J	44 J	0.7 J	< 0.5 U	< 0.5 U	< 2.5 U		
Toluene	µg/l	1,000	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	30 J	31 J	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
trans-1,2-Dichloroethene	µg/l	100	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	150	150	0.99 J	< 0.5 U	< 0.5 U	< 2.5 U		
trans-1,3-Dichloropropene	µg/l	29	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
Trichloroethene	µg/l	5	< 0.5 U	34	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	1.1	< 0.5 U	< 0.5 U	15,000	1,400	1,400	240	30	14	1,300		
Trichlorofluoromethane	µg/l	31,000	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	< 0.5 U	< 0.5 U	< 0.5 U	< 2.5 U		
Vinyl chloride	µg/l	2	< 0.5 U	< 0.5 U	< 0.5 U			< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 25 U	< 25 U	3.0	1.6	< 0.5 U	< 2.5 U		
Metals (6020A)																										
Aluminum	mg/L	100	0.649	0.0219 UB	0.0389 UB			0.213	0.482	0.255	NA	NA	0.00694 UB	NA	0.0222	NA	0.370	0.0775	NA	NA	0.00644 J	NA	0.0152	NA		
Antimony	mg/L	0.006	0.00244 J	< 0.005000 U	< 0.005000 U			< 0.000500 U	< 0.00500 U	< 0.00500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA	0.000800 J	< 0.000500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA		
Arsenic	mg/L	0.01	0.00640	0.00328 J	0.00567			0.00175 J	0.000998 J	0.00770	NA	NA	0.00318 J	NA	0.00101 J	NA	< 0.000500 U	< 0.000500 U	NA	NA	0.000625 J	NA	0.00204 J	NA		
Barium	mg/L	2	0.141	4.13	1.34			0.145	0.0582	0.316	NA	NA	1.39	NA	1.03	NA	0.238	0.406	NA	NA	0.370	NA	0.700	NA		
Beryllium	mg/L	0.004	0.00310	< 0.005000 U	0.000322 J			< 0.000500 U	< 0.00500 U	< 0.00500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA	< 0.000500 U	< 0.000500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA		
Cadmium	mg/L	0.005	0.00858	0.000710 J	< 0.005000 U			< 0.000500 U	0.000207 J	< 0.00500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA	< 0.000500 U	< 0.000500 U	NA	NA	< 0.000500 U	NA	0.000205 J	NA		
Calcium	mg/L	NV	182	371	58.2			15.2	8.52	20.6	NA	NA	46.7	NA	19.7	NA	76.3	17.9	NA	NA	26.5	NA	26.9	NA		
Chromium	mg/L	0.1	9.14	1.80	0.00520			0.0245	0.000791 J	0.000744 J	NA	NA	< 0.000500 U	NA	0.000674 J	NA	0.00659	0.358	NA	NA	0.00222 J	NA	0.228	NA		
Cobalt	mg/L	6.1	0.418	0.0122	0.000122 J			0.000243 J	0.00641	0.00410 J	NA	NA	0.000279 J	NA	0.00214 J	NA	0.000529 J	0.0170	NA	NA	0.00313 J	NA	0.00561	NA		
Copper	mg/L	1.3	0.755	0.109	< 0.00100 U			< 0.00100 U	0.00151 J	< 0.00100 U	NA	NA	< 0.00100 U	NA	< 0.00100 U	NA	0.00119 J	0.00256 UB	NA	NA	0.00658	NA	0.00520 UB	NA		
Iron	mg/L	NV	29.7	25.2	113			0.0577 J	0.483	27.0	NA	NA	87.5	NA	63.2	NA	0.354	5.03	NA	NA	0.218	NA	2.15	NA		
Lead	mg/L	0.015	0.00364 J	0.00130 J	0.000647 J			< 0.00100 U	< 0.00100 U	< 0.00100 U	NA	NA	< 0.00100 U	NA	< 0.00100 U	NA	< 0.00100 U	< 0.00100 U	NA	NA	< 0.00100 U	NA	< 0.00100 U	NA		
Magnesium	mg/L	NV	166	249	36.3			3.50	6.41	12.5	NA	NA	30.1	NA	12.4	NA	3.79	14.4	NA	NA	16.4	NA	22.9	NA		
Manganese	mg/L	1.1*	3.12	0.308	2.33			0.00169 J	2.89	2.92	NA	NA	1.52	NA	0.994	NA	0.0798	0.703	NA	NA	0.295	NA	0.137 J	NA		
Nickel	mg/L	0.49*	14.3	0.886	0.00103 J			< 0.00100 U	0.206	0.00273 J	NA	NA	< 0.00100 U	NA	0.00170 J	NA	0.00408 J	1.07	NA	NA	0.00306 J	NA	0.168	NA		
Potassium	mg/L	NV	4.36	1.55	3.49			4.81	1.08	1.51	NA	NA	3.38	NA	4.20	NA	0.511	2.47	NA	NA	1.56	NA	2.16	NA		
Selenium	mg/L	0.05	0.0130	0.00181 J	< 0.00250 U			< 0.00250 U	< 0.00250 U	< 0.00250 U	NA	NA	< 0.00250 U	NA	< 0.00250 U	NA	< 0.00250 U	< 0.00250 U	NA	NA	< 0.00250 U	NA	< 0.00250 U	NA		
Silver	mg/L	0.51	0.000338 J	0.000215 J	< 0.000500 U			< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA	< 0.000500 U	< 0.000500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA		
Sodium	mg/L	NV	532	1,270	148			39.1	208	35.7	NA	NA	168	NA	233	NA	14.3	170	NA	NA	254	NA	203	NA		
Thallium	mg/L	0.002	< 0.000500 U	< 0.000500 U	< 0.000500 U			< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA	0.000881 J	< 0.000500 U	NA	NA	< 0.000500 U	NA	< 0.000500 U	NA		
Vanadium	mg/L	0.72	0.0771	0.0203	< 0.00100 U			0.0148	0.00210 J	0.000983 UB	NA	NA	< 0.00100 U	NA	< 0.00100 U	NA	0.00199 J	0.00117 UB	NA	NA	0.00152 J	NA	0.00378 UB	NA		
Zinc	mg/L	31	0.175	0.185	0.0145			0.0168	0.138	0.0183	NA	NA	0.0216	NA	0.0188	NA	0.0252	0.00816	NA	NA	0.0217	NA	0.0358	NA		
Mercury	mg/L	0.002	0.0000520 J	< 0.000100 U	< 0.000100 U			< 0.000100 U	< 0.000100 U	< 0.000100 U	NA	NA	< 0.000100 U	NA	< 0.000100 U	NA	< 0.000100 U	< 0.000100 U	NA	NA	< 0.000100 U	NA	< 0.000100 U	NA		
1,4-Dioxane (8270D SIM)																										
1,4-Dioxane	µg/l	9.1	NA	NA	< 0.010 U			0.017	NA	< 0.010 U	NA	NA	0.012	NA	NA	< 0.010 U	NA	NA	3.5 J	6.2 J	NA	< 0.010 U	NA	9.7		

Notes:
 Blue highlighting indicates concentrations above the MCL/MSC/PCL
 MCL/MSC - Maximum Contaminant Limit/Medium-Specific Concentrations/Protective Concentration Level
 NA - Not Analyzed
 µg/L - micrograms per liter a - duplicate sample
 mg/L - milligrams per liter
 J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.
 UJ - The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.
 U - Undetected: The analyte was analyzed for, but not detected.
 NV - No Value
 UB - considered a non-detect due to blank contamination
 *Perchlorate, manganese, and nickel compared to the PCL
 ** Value is for total xylenes
 PCL - Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	MW8_062119 6/21/19	MW9_062519 6/25/19	MW10_062519 6/25/19	MW12_062019 6/20/19	MW13_062519 6/25/19	MW14_062019 6/20/19	MW14_062019- a 6/20/19
Lab Package			HS19061210	HS19061386	HS19061386	HS19061164	HS19061386	HS19061164	HS19061164
Well ID			MW-8	MW-9	MW-10	MW-12	MW-13	MW-14	MW-14
Perchlorate (6850)									
Perchlorate	µg/L	17*	7,100	520	<2.0 U	<2.0 U	<2.0 U	240,000	230,000
Volatile Organic Compounds (8260C)									
1,1,1,2-Tetrachloroethane	µg/l	110	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,1,1-Trichloroethane	µg/l	200	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,1,2,2-Tetrachloroethane	µg/l	14	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,1,2-Trichloroethane	µg/l	5	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,1-Dichloroethane	µg/l	10,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,1-Dichloroethene	µg/l	7	<0.5 U	<0.5 U	<0.5 U	1.0	<0.5 U	130	140 J
1,1-Dichloropropene	µg/l	2.9	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2,3-Trichlorobenzene	µg/l	310	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2,3-Trichloropropane	µg/l	0.041	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2,4-Trichlorobenzene	µg/l	70	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2,4-Trimethylbenzene	µg/l	5,100	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2-Dibromo-3-chloropropane	µg/l	0.2	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2-Dibromoethane	µg/l	0.05	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2-Dichlorobenzene	µg/l	600	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,2-Dichloroethane	µg/l	5	0.75 J	<0.5 U	<0.5 U	0.57 J	<0.5 U	100	110 J
1,2-Dichloropropane	µg/l	5	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,3,5-Trimethylbenzene	µg/l	5,100	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,3-Dichlorobenzene	µg/l	3,100	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,3-Dichloropropane	µg/l	29	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
1,4-Dichlorobenzene	µg/l	75	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
2,2-Dichloropropane	µg/l	42	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
2-Butanone	µg/l	61,000	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<20 U	<20 U
2-Chlorotoluene	µg/l	2,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
2-Hexanone	µg/l	6,100	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<20 U	<20 U
4-Chlorotoluene	µg/l	2,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
4-Isopropyltoluene	µg/l	10,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
4-Methyl-2-pentanone	µg/l	8,200	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<20 U	<20 U
Acetone	µg/l	92,000	<1.0 U	2.1	<1.0 U	3.5	<1.0 U	<20 U	<20 U
Benzene	µg/l	5	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Bromobenzene	µg/l	2,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Bromochloromethane	µg/l	4,100	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Bromodichloromethane	µg/l	4.6	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Bromoform	µg/l	36	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Bromomethane	µg/l	140	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Carbon disulfide	µg/l	10,000	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<20 U	<20 U
Carbon tetrachloride	µg/l	5	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Chlorobenzene	µg/l	100	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Chloroethane	µg/l	41,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Chloroform	µg/l	1,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	25 J
Chloromethane	µg/l	220	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
cis-1,2-Dichloroethene	µg/l	70	<0.5 U	25	<0.5 U	15	<0.5 U	1,800	2,000 J
cis-1,3-Dichloropropene	µg/l	5.3	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Dibromochloromethane	µg/l	34	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Dibromomethane	µg/l	380	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Dichlorodifluoromethane	µg/l	20,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Ethylbenzene	µg/l	700	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
Hexachlorobutadiene	µg/l	20	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<20 U	<20 U
Isopropylbenzene	µg/l	10,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
m,p-Xylene	µg/l	10,000**	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<20 U	<20 U
Methylene chloride	µg/l	5	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<1.0 U	<20 U	<20 U
Naphthalene	µg/l	2,000	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U
n-Butylbenzene	µg/l	4,100	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<0.5 U	<10 U	<10 U

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	MW8_062119 6/21/19	MW9_062519 6/25/19	MW10_062519 6/25/19	MW12_062019 6/20/19	MW13_062519 6/25/19	MW14_062019 6/20/19	MW14_062019- a 6/20/19
n-Propylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
o-Xylene	µg/l	10,000**	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
sec-Butylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
Styrene	µg/l	100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
tert-Butylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
Tetrachloroethene	µg/l	5	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
Toluene	µg/l	1,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
trans-1,2-Dichloroethene	µg/l	100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
trans-1,3-Dichloropropene	µg/l	29	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
Trichloroethene	µg/l	5	120	1,100	22	160	< 0.5 U	11,000	11,000
Trichlorofluoromethane	µg/l	31,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
Vinyl chloride	µg/l	2	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 10 U
Metals (6020A)									
Aluminum	mg/L	100	NA	0.0255	NA	NA	0.336	0.0408	0.0353
Antimony	mg/L	0.006	NA	< 0.000500 U	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U
Arsenic	mg/L	0.01	NA	< 0.000500 U	NA	NA	0.000513 J	0.00702	0.00636
Barium	mg/L	2	NA	0.119	NA	NA	0.234	0.311	0.297
Beryllium	mg/L	0.004	NA	< 0.000500 U	NA	NA	< 0.000500 U	0.000305 J	0.000276 J
Cadmium	mg/L	0.005	NA	< 0.000500 U	NA	NA	0.000407 J	0.000877 J	0.000834 J
Calcium	mg/L	NV	NA	15.0	NA	NA	14.8	122	116
Chromium	mg/L	0.1	NA	0.156	NA	NA	0.738	0.0974	0.0866
Cobalt	mg/L	6.1	NA	0.00255 J	NA	NA	0.00531	0.0252	0.0240
Copper	mg/L	1.3	NA	< 0.00100 U	NA	NA	0.0274	0.00394 UB	0.00362 UB
Iron	mg/L	NV	NA	0.830	NA	NA	37.1	128	120
Lead	mg/L	0.015	NA	< 0.00100 U	NA	NA	< 0.00100 U	< 0.00100 U	< 0.00100 U
Magnesium	mg/L	NV	NA	3.78	NA	NA	7.96	72.8	68.8
Manganese	mg/L	1.1*	NA	0.0495	NA	NA	0.844	4.58	4.81
Nickel	mg/L	0.49*	NA	0.127	NA	NA	0.247	0.158	0.150
Potassium	mg/L	NV	NA	0.438	NA	NA	3.58	14.0	13.5
Selenium	mg/L	0.05	NA	< 0.00250 U	NA	NA	< 0.00250 U	< 0.00250 U	< 0.00250 U
Silver	mg/L	0.51	NA	< 0.000500 U	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U
Sodium	mg/L	NV	NA	17.2	NA	NA	56.0	449	482
Thallium	mg/L	0.002	NA	< 0.000500 U	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U
Vanadium	mg/L	0.72	NA	0.000919 J	NA	NA	0.00339 J	< 0.00100 U	< 0.00100 U
Zinc	mg/L	31	NA	0.0279	NA	NA	0.0257	0.467	0.439
Mercury	mg/L	0.002	NA	< 0.000100 U	NA	NA	< 0.000100 U	< 0.000100 U	< 0.000100 U
1,4-Dioxane (8270D SIM)									
1,4-Dioxane	µg/l	9.1	0.5	2	0.12	NA	NA	280 J	390 J

Notes:

Blue highlighting indicates concentrations above the MCL/MSC/PCL

MCL/MSC - Maximum Contaminant Limit/Medium-Specific Concentrations/Protective Concentration Level

NA - Not Analyzed

µg/L - micrograms per liter a - duplicate sample

mg/L - milligrams per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

UJ - The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

NV - No Value

UB - considered a non-detect due to blank contamination

*Perchlorate, manganese, and nickel compared to the PCL

** Value is for total xylenes

PCL – Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	MW16_062619 6/26/19	MW17_062519 6/25/19	MW18_062619 6/26/19	MW19_062619 6/26/19	MW20_061319 6/13/19	MW21-061819 6/18/19	MW22_062119 6/21/19	MW23_062419 6/24/19	102_062419 6/24/19	109_061819 6/18/19	120_062019 6/20/19	125_062419 6/24/19	126_062419 6/24/19	129_062419 6/24/19	130_062519 6/25/19	130_062519- a 6/25/19	ICT2_062819 6/28/19	ICT4_062819 6/28/19	ICT7_062819 6/28/19	ICT8_062819 6/28/19	ICT11_062819 6/28/19	ICT11_062819- a 6/28/19	ICT12B_062819 6/28/19		
n-Propylbenzene	µg/l	4,100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
o-Xylene	µg/l	10,000**	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
sec-Butylbenzene	µg/l	4,100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
Styrene	µg/l	100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
tert-Butylbenzene	µg/l	4,100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
Tetrachloroethene	µg/l	5	1.4	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	0.90 J	1.8	< 0.5 U	240	< 0.5 U	< 0.5 U	6.6 J	< 0.5 U		
Toluene	µg/l	1,000	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
trans-1,2-Dichloroethene	µg/l	100	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	1.0	5.2	< 0.5 U	9.8 J	< 0.5 U	< 0.5 U	5.3 J	< 0.5 U		
trans-1,3-Dichloropropene	µg/l	29	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
Trichloroethene	µg/l	5	1,200	NA	25	0.63 J	< 0.5 U	6,300	230	2,600	< 0.5 U	130	7,900	< 0.5 U	< 0.5 U	820	< 0.5 U	670	6,000	1.0	7,500	270	210	14,000	< 0.5 U		
Trichlorofluoromethane	µg/l	31,000	< 0.5 U	NA	< 0.5 U	< 0.5 U	< 0.5 U	< 10 U	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U		
Vinyl chloride	µg/l	2	2.1	NA	< 0.5 U	< 0.5 U	< 0.5 U	18 J	< 0.5 U	< 2.5 U	< 0.5 U	< 0.5 U	< 10 U	< 5.0 U	< 0.5 U	< 0.5 U	< 0.5 U	0.95 J	10	< 0.5 U	13	< 0.5 U	< 0.5 U	28	< 0.5 U		
Metals (6020A)																											
Aluminum	mg/L	100	NA	NA	NA	1.54	1.12	0.0299	0.0158	NA	10.3	NA	NA	0.324	0.0917	2.44	5.22	5.34	0.139	0.0112	0.211	0.0188	0.00846 J	0.0149 J	0.0271	< 0.000500 U	
Antimony	mg/L	0.006	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	< 0.000500 U	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000493 UB	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	
Arsenic	mg/L	0.01	NA	NA	NA	0.0137	0.0104 J	0.00241 J	0.00145 J	NA	0.00262 J	NA	NA	0.000450 J	0.00462 J	0.00120 J	0.00360 J	0.00349 J	0.00229 J	0.000680 J	0.000594 J	0.00649	< 0.000500 U	0.000576 J	< 0.000500 U	< 0.000500 U	
Barium	mg/L	2	NA	NA	NA	0.467	0.479	9.21	0.683	NA	0.276	NA	NA	0.0647	10.4	0.119	0.0930	0.0899	0.167	0.134	0.217	0.303	0.443	0.446	0.286	< 0.000500 U	
Beryllium	mg/L	0.004	NA	NA	NA	0.000294 J	< 0.000500 U	0.000301 J	< 0.000500 U	NA	0.00113 J	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000454 J	0.000469 J	< 0.000500 U	< 0.000500 U	0.000549 J	< 0.000500 U	0.000285 J	0.000280 J	< 0.000500 U	< 0.000500 U	
Cadmium	mg/L	0.005	NA	NA	NA	0.00227	0.000359 J	0.000837 J	< 0.000500 U	NA	0.000225 J	NA	NA	0.000218 J	0.000370 J	0.000738 J	0.000499 J	0.000533 J	< 0.000500 U	0.000394 J	< 0.000500 U	0.000354 J	0.000314 J	< 0.000500 U	< 0.000500 U		
Calcium	mg/L	NV	NA	NA	NA	44.1	44.7	223	76.1	NA	3.53	NA	NA	1.25	344	4.49	8.03	8.07	19.5	35.8	2.41	12.8	10.3	10.4	26.6	< 0.000500 U	
Chromium	mg/L	0.1	NA	NA	NA	0.791	0.0188	1.09	0.0930	NA	0.0141	NA	NA	0.00412 J	0.000825 J	0.00567	0.00929	0.00918	0.00284 J	0.0321	0.00142 J	0.00144 J	0.142 J	0.219 J	0.0412	< 0.000500 U	
Cobalt	mg/L	6.1	NA	NA	NA	0.0171	0.0154	0.117	0.00142 J	NA	0.00925	NA	NA	0.000344 J	0.00777	0.00936	0.00947	0.00954	0.0116	0.00928	0.00298 J	0.00314 J	0.00481 J	0.00468 J	0.00545	< 0.000500 U	
Copper	mg/L	1.3	NA	NA	NA	0.0244	0.00576 UB	0.219	0.00249 J	NA	0.0139	NA	NA	< 0.00100 U	< 0.00100 U	0.00326 J	0.00755	0.00743	0.0501	< 0.00100 U	0.00281 J	< 0.00100 U	0.00472 J	0.00424 J	0.0824	< 0.000500 U	
Iron	mg/L	NV	NA	NA	NA	67.1	3.14	15.2	1.13	NA	15.7	NA	NA	0.466	4.05	3.29	7.93	7.87	4.17	0.404	0.836	16.0	1.00	1.58	1.68	< 0.000500 U	
Lead	mg/L	0.015	NA	NA	NA	0.00323 J	0.00155 J	0.000663 J	< 0.00100 U	NA	0.0113	NA	NA	0.00128 J	< 0.00100 U	0.00297 J	0.00546	0.00565	0.00446 J	< 0.00100 U	< 0.00100 U	< 0.00100 U	< 0.00100 U	< 0.00100 U	< 0.00100 U	< 0.00100 U	
Magnesium	mg/L	NV	NA	NA	NA	26.1	12.5	178	34.5	NA	3.81	NA	NA	0.937	269	3.31	5.19	5.20	20.0	31.8	2.46	10.3	7.04	7.20	18.6	< 0.000500 U	
Manganese	mg/L	1.1*	NA	NA	NA	1.50	0.274	2.59	0.0164	NA	0.0964	NA	NA	0.00625	0.150	0.176	0.285	0.287	0.336	0.311	0.0961	0.228	0.205	0.200	0.165	< 0.000500 U	
Nickel	mg/L	0.49*	NA	NA	NA	0.214	0.540	1.49	0.179	NA	0.0255	NA	NA	0.0022 J	0.0137	0.00812	0.00933	0.00981	0.0139	0.265	0.0181	0.0213	0.564	0.547	0.345	< 0.000500 U	
Potassium	mg/L	NV	NA	NA	NA	4.36	0.862	3.20	1.93	NA	1.73	NA	NA	0.321	3.57	0.647	1.88	1.89	1.24	1.54	1.45	1.48	0.934	0.968	0.743	< 0.000500 U	
Selenium	mg/L	0.05	NA	NA	NA	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	NA	0.00243 J	NA	NA	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	
Silver	mg/L	0.51	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	< 0.000500 U	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	
Sodium	mg/L	NV	NA	NA	NA	453	48.6	552	383	NA	17.0	NA	NA	28.8	949	69.6	123	126	155	259	13.3	69.6	64.8	67.2	195	< 0.000500 U	
Thallium	mg/L	0.002	NA	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	NA	0.000275 J	NA	NA	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000618 J	< 0.000500 U	< 0.000500 U	< 0.000500 U	
Vanadium	mg/L	0.72	NA	NA	NA	0.0102	0.00507 UB	0.00433 UB	0.00257 UB	NA	0.0237	NA	NA	0.00142 UB	0.00257 UB	0.00659	0.0165	0.0168	0.00189 UB	0.00174 UB	0.00136 UB	0.00195 UB	0.00109 UB	0.00243 UB	0.000667 J	< 0.000500 U	
Zinc	mg/L	31	NA	NA	NA	0.0441	0.102	0.0462	0.0241	NA	0.0748	NA	NA	0.0207	0.0423	0.0356	0.0478	0.0456	0.196	0.0164	0.0553	0.0101	0.0902	0.0793	0.105	< 0.000500 U	
Mercury	mg/L	0.002	NA	NA	NA	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	NA	< 0.000100 U	NA	NA	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	
1,4-Dioxane (8270D SIM)																											
1,4-Dioxane	µg/l	9.1	12	0.31	NA	NA	NA	1.2 J	NA	NA	NA	0.500	9.9	< 0.010 U	< 0.010 U	1.2	NA	NA	NA	NA	NA	NA	NA	NA	NA	NA	

Notes:
 Blue highlighting indicates concentrations above the MCL/MSC/PCL
 MCL/MSC - Maximum Contaminant Limit/Medium-Specific Concentrations/Protective Concentration Level
 NA - Not Analyzed
 µg/L - micrograms per liter

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	ICT12B_062819- a 6/28/19	ICT12C_062819 6/28/19	ICT12D_062819 6/28/19	ICT12E_062819 6/28/19	ICT13A_071118 7/11/18
Lab Package			HS19070016	HS19070016	HS19070016	HS19070016	HS19070622
Well ID			ICT-12B	ICT-12C	ICT-12D	ICT-12E	ICT-13A
Perchlorate (6850)							
Perchlorate	µg/L	17*	150,000	6,500	13,000	28,000	540
Volatile Organic Compounds (8260C)							
1,1,1,2-Tetrachloroethane	µg/l	110	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,1,1-Trichloroethane	µg/l	200	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,1,2,2-Tetrachloroethane	µg/l	14	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,1,2-Trichloroethane	µg/l	5	< 5.0 U	< 5.0 U	9.8 J	14	< 0.5 U
1,1-Dichloroethane	µg/l	10,000	12	< 5.0 U	75	100	1.3 J
1,1-Dichloroethene	µg/l	7	< 5.0 U	< 5.0 U	370	550	5.5 J
1,1-Dichloropropene	µg/l	2.9	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,2,3-Trichlorobenzene	µg/l	310	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,2,3-Trichloropropane	µg/l	0.041	< 5.0 U	< 5.0 U	350	< 5.0 U	< 0.5 U
1,2,4-Trichlorobenzene	µg/l	70	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,2,4-Trimethylbenzene	µg/l	5,100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,2-Dibromo-3-chloropropane	µg/l	0.2	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,2-Dibromoethane	µg/l	0.05	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,2-Dichlorobenzene	µg/l	600	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,2-Dichloroethane	µg/l	5	68	27	160	63	18 J
1,2-Dichloropropane	µg/l	5	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,3,5-Trimethylbenzene	µg/l	5,100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,3-Dichlorobenzene	µg/l	3,100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,3-Dichloropropane	µg/l	29	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
1,4-Dichlorobenzene	µg/l	75	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
2,2-Dichloropropane	µg/l	42	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
2-Butanone	µg/l	61,000	< 10 U	< 10 U	< 10 U	< 10 U	< 1.0 U
2-Chlorotoluene	µg/l	2,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
2-Hexanone	µg/l	6,100	< 10 U	< 10 U	< 10 U	< 10 U	< 1.0 U
4-Chlorotoluene	µg/l	2,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
4-Isopropyltoluene	µg/l	10,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
4-Methyl-2-pentanone	µg/l	8,200	< 10 U	< 10 U	< 10 U	< 10 U	< 1.0 U
Acetone	µg/l	92,000	< 10 U	< 10 U	< 10 U	180	< 1.0 U
Benzene	µg/l	5	11 J	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Bromobenzene	µg/l	2,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Bromochloromethane	µg/l	4,100	< 5.0 U	< 5.0 U	55	40	< 0.5 U
Bromodichloromethane	µg/l	4.6	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Bromoform	µg/l	36	< 5.0 U	< 5.0 U	7.8 J	< 5.0 U	< 0.5 U
Bromomethane	µg/l	140	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Carbon disulfide	µg/l	10,000	< 10 U	< 10 U	< 10 U	< 10 U	< 1.0 U
Carbon tetrachloride	µg/l	5	49 J	11	< 5.0 U	< 5.0 U	< 0.5 U
Chlorobenzene	µg/l	100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Chloroethane	µg/l	41,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Chloroform	µg/l	1,000	70	16	61	84	< 0.5 U
Chloromethane	µg/l	220	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
cis-1,2-Dichloroethene	µg/l	70	1,100	140	10,000	17,000	86 J
cis-1,3-Dichloropropene	µg/l	5.3	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Dibromochloromethane	µg/l	34	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Dibromomethane	µg/l	380	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Dichlorodifluoromethane	µg/l	20,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Ethylbenzene	µg/l	700	< 5.0 U	< 5.0 U	6.0 J	7.7 J	< 0.5 U
Hexachlorobutadiene	µg/l	20	< 10 U	< 10 U	< 10 U	< 10 U	< 1.0 U
Isopropylbenzene	µg/l	10,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
m,p-Xylene	µg/l	10,000**	< 10 U	< 10 U	7.1 J	12 J	< 1.0 U
Methylene chloride	µg/l	5	11 J	< 10 U	11,000	120,000	< 1.0 U
Naphthalene	µg/l	2,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
n-Butylbenzene	µg/l	4,100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	ICT12B_062819- a 6/28/19	ICT12C_062819 6/28/19	ICT12D_062819 6/28/19	ICT12E_062819 6/28/19	ICT13A_071118 7/11/18
n-Propylbenzene	µg/l	4,100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
o-Xylene	µg/l	10,000**	< 5.0 U	< 5.0 U	5.3 J	8.6 J	< 0.5 U
sec-Butylbenzene	µg/l	4,100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Styrene	µg/l	100	< 5.0 U	< 5.0 U	< 5.0 U	41	< 0.5 U
tert-Butylbenzene	µg/l	4,100	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Tetrachloroethene	µg/l	5	< 5.0 U	< 5.0 U	10	14	2.2 J
Toluene	µg/l	1,000	< 5.0 U	< 5.0 U	17	28	< 0.5 U
trans-1,2-Dichloroethene	µg/l	100	5.2 J	< 5.0 U	29	43	1.1 J
trans-1,3-Dichloropropene	µg/l	29	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Trichloroethene	µg/l	5	15,000	3,900	44,000	84,000	920
Trichlorofluoromethane	µg/l	31,000	< 5.0 U	< 5.0 U	< 5.0 U	< 5.0 U	< 0.5 U
Vinyl chloride	µg/l	2	32	< 5.0 U	320	440	6.4 J
Metals (6020A)							
Aluminum	mg/L	100	0.0285	0.0415	0.595	0.0433	0.677
Antimony	mg/L	0.006	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Arsenic	mg/L	0.01	< 0.000500 U	< 0.000500 U	0.00524	0.00106 J	0.00149 J
Barium	mg/L	2	0.285	0.123	0.444	0.197	0.0656 J
Beryllium	mg/L	0.004	< 0.000500 U	< 0.000500 U	0.000829 J	0.000205 J	0.000221 J
Cadmium	mg/L	0.005	< 0.000500 U	< 0.000500 U	0.000738 J	0.000355 J	< 0.000500 U
Calcium	mg/L	NV	27.2	6.85	20.5	37.7	3.93
Chromium	mg/L	0.1	0.0449	0.0972	0.338	0.142	0.0155
Cobalt	mg/L	6.1	0.00515	0.000998 J	0.0404	0.0315	0.00696
Copper	mg/L	1.3	0.0802	< 0.00250 U	0.721	0.0229	0.00803
Iron	mg/L	NV	1.58	0.969	20.4	3.22	5.49
Lead	mg/L	0.015	< 0.00100 U	< 0.00100 U	0.0259	< 0.00100 U	0.000925 J
Magnesium	mg/L	NV	18.6	5.56	17.2	29.8	3.51
Manganese	mg/L	1.1*	0.155	0.0176	1.13	1.28	0.132
Nickel	mg/L	0.49*	0.328	0.0408	0.850	0.178	0.0145
Potassium	mg/L	NV	0.749	0.301	1.05	2.43	0.708
Selenium	mg/L	0.05	< 0.00250 U	< 0.00250 U	0.00224 J	0.00349 J	0.00127 J
Silver	mg/L	0.51	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Sodium	mg/L	NV	192	96.2	175	339	70.0
Thallium	mg/L	0.002	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000243 J
Vanadium	mg/L	0.72	0.000792 J	0.00125 J	0.00303 J	< 0.000500 U	0.00425 J
Zinc	mg/L	31	0.120	0.0188	0.0862	0.0387	0.288
Mercury	mg/L	0.002	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U
1,4-Dioxane (8270D SIM)							
1,4-Dioxane	µg/l	9.1	NA	NA	NA	NA	NA

Notes:

Blue highlighting indicates concentrations above the MCL/MSC/PCL

MCL/MSC - Maximum Contaminant Limit/Medium-Specific Concentrations/Protective Concentration Level

NA - Not Analyzed

µg/L - micrograms per liter a - duplicate sample

mg/L - milligrams per liter

J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.

UJ - The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.

U - Undetected: The analyte was analyzed for, but not detected.

NV - No Value

UB - considered a non-detect due to blank contamination

*Perchlorate, manganese, and nickel compared to the PCL

** Value is for total xylenes

PCL - Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	ICT13B_062819 6/28/19	ICT13D_062819 6/28/19	ICT13E_062819 6/28/19	ICT13F_062819 6/28/19	ICT14B_062819 6/28/19	ICT14C_062819 6/28/19	ICT14D_062819 6/28/19
Lab Package			HS19070016	HS19070016	HS19070016	HS19070016	HS19070016	HS19070016	HS19070016
Well ID			ICT-13B	ICT-13D	ICT-13E	ICT-13F	ICT-14B	ICT-14C	ICT-14D
Perchlorate (6850)									
Perchlorate	µg/L	17*	< 2.0 U	43	3.8 J	< 2.0 U	12,000	< 2.0 U	210
Volatile Organic Compounds (8260C)									
1,1,1,2-Tetrachloroethane	µg/l	110	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,1,1-Trichloroethane	µg/l	200	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,1,2,2-Tetrachloroethane	µg/l	14	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,1,2-Trichloroethane	µg/l	5	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	5.1 J	< 5.0 U
1,1-Dichloroethane	µg/l	10,000	1.0	< 0.5 U	< 0.5 U	< 0.5 U	1.4	10	25
1,1-Dichloroethene	µg/l	7	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	140	< 5.0 U
1,1-Dichloropropene	µg/l	2.9	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2,3-Trichlorobenzene	µg/l	310	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2,3-Trichloropropane	µg/l	0.041	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2,4-Trichlorobenzene	µg/l	70	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2,4-Trimethylbenzene	µg/l	5,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2-Dibromo-3-chloropropane	µg/l	0.2	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2-Dibromoethane	µg/l	0.05	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2-Dichlorobenzene	µg/l	600	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,2-Dichloroethane	µg/l	5	42	10	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	13
1,2-Dichloropropane	µg/l	5	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,3,5-Trimethylbenzene	µg/l	5,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,3-Dichlorobenzene	µg/l	3,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,3-Dichloropropane	µg/l	29	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
1,4-Dichlorobenzene	µg/l	75	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
2,2-Dichloropropane	µg/l	42	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
2-Butanone	µg/l	61,000	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 10 U	< 10 U
2-Chlorotoluene	µg/l	2,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
2-Hexanone	µg/l	6,100	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 10 U	< 10 U
4-Chlorotoluene	µg/l	2,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
4-Isopropyltoluene	µg/l	10,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
4-Methyl-2-pentanone	µg/l	8,200	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 10 U	< 10 U
Acetone	µg/l	92,000	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	6.2	< 10 U	< 10 U
Benzene	µg/l	5	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Bromobenzene	µg/l	2,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Bromochloromethane	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	22
Bromodichloromethane	µg/l	4.6	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Bromoform	µg/l	36	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Bromomethane	µg/l	140	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Carbon disulfide	µg/l	10,000	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 10 U	< 10 U
Carbon tetrachloride	µg/l	5	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Chlorobenzene	µg/l	100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Chloroethane	µg/l	41,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Chloroform	µg/l	1,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	39	29
Chloromethane	µg/l	220	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
cis-1,2-Dichloroethene	µg/l	70	140	37	< 0.5 U	< 0.5 U	31	11,000	21,000
cis-1,3-Dichloropropene	µg/l	5.3	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Dibromochloromethane	µg/l	34	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Dibromomethane	µg/l	380	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Dichlorodifluoromethane	µg/l	20,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Ethylbenzene	µg/l	700	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Hexachlorobutadiene	µg/l	20	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 0.5 U	< 10 U	< 10 U
Isopropylbenzene	µg/l	10,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
m,p-Xylene	µg/l	10,000**	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 10 U	< 10 U
Methylene chloride	µg/l	5	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	< 1.0 U	1,000	11,000 J
Naphthalene	µg/l	2,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
n-Butylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U

LHAAP-18/24 Sampling Event - June 2019

Location ID: Sample Date:	Units	MCL/MSC/P CL	ICT13B_062819 6/28/19	ICT13D_062819 6/28/19	ICT13E_062819 6/28/19	ICT13F_062819 6/28/19	ICT14B_062819 6/28/19	ICT14C_062819 6/28/19	ICT14D_062819 6/28/19
n-Propylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
o-Xylene	µg/l	10,000**	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
sec-Butylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Styrene	µg/l	100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
tert-Butylbenzene	µg/l	4,100	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Tetrachloroethene	µg/l	5	1.2	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	13
Toluene	µg/l	1,000	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
trans-1,2-Dichloroethene	µg/l	100	1.2	< 0.5 U	< 0.5 U	< 0.5 U	0.6 J	51	44
trans-1,3-Dichloropropene	µg/l	29	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 0.5 U	< 5.0 U	< 5.0 U
Trichloroethene	µg/l	5	1,200	160	< 0.5 U	< 0.5 U	190	120	1,400
Trichlorofluoromethane	µg/l	31,000	< 0.5 U	< 0.5 U	0.5 U	0.5 U	0.5 U	< 5.0 U	< 5.0 U
Vinyl chloride	µg/l	2	3.5	1.5	0.5 U	0.5 U	0.68 J	1,600	1,100
Metals (6020A)									
Aluminum	mg/L	100	0.0158	1.41	6.70	3.09	0.0254	0.00742 J	0.0223 J
Antimony	mg/L	0.006	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Arsenic	mg/L	0.01	0.00630	0.00193 J	0.00294 J	0.00162 J	0.00114 J	0.0108	0.00810
Barium	mg/L	2	0.203	0.0962	0.136	0.115	0.447	1.07	1.18
Beryllium	mg/L	0.004	0.000547 J	< 0.000500 U	0.00102 J	0.000645 J	0.000259 J	< 0.000500 U	0.000518 J
Cadmium	mg/L	0.005	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	0.000225 J	0.000219 J	0.000514 J
Calcium	mg/L	NV	13.3	3.50	1.16	1.05	12.4	78.9	80.0
Chromium	mg/L	0.1	0.0230	0.121	0.0248	0.0124	0.0489	0.0341	0.00422 J
Cobalt	mg/L	6.1	0.105	0.00316 J	0.00491 J	0.00302 J	0.00274 J	0.0331	0.0141
Copper	mg/L	1.3	0.0276	0.00533	0.0168	0.0127	0.00307 J	0.00281 J	0.00411 J
Iron	mg/L	NV	20.3	9.93	12.1	6.24	0.842	274	19.5
Lead	mg/L	0.015	< 0.00100 U	0.00505	0.00889	0.00438 J	< 0.00100 U	< 0.00100 U	< 0.00100 U
Magnesium	mg/L	NV	17.7	4.38	1.63	1.14	11.6	80.4	57.2
Manganese	mg/L	1.1*	4.05	0.0744	0.0546	0.0357	0.239	2.33	1.31
Nickel	mg/L	0.49*	0.0245	0.0578	0.0192	0.00929	0.0822	0.120	0.0474
Potassium	mg/L	NV	2.37	0.918	2.09	1.90	0.565	1.41	1.46
Selenium	mg/L	0.05	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	< 0.00250 U	0.0109	< 0.000500 U
Silver	mg/L	0.51	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Sodium	mg/L	NV	103	34.2	5.75	4.44	85.8	265	295
Thallium	mg/L	0.002	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U	< 0.000500 U
Vanadium	mg/L	0.72	0.00112 J	0.00524	0.0267	0.0146	0.00184 UB	< 0.000500 U	0.000634 J
Zinc	mg/L	31	0.0910	0.491	0.0557	0.0596	0.0305	0.0439	0.141
Mercury	mg/L	0.002	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U	< 0.000100 U
1,4-Dioxane (8270D SIM)									
1,4-Dioxane	µg/l	9.1	NA	NA	NA	NA	NA	NA	NA

Notes:

- Blue highlighting indicates concentrations above the MCL/MSC/PCL
- MCL/MSC - Maximum Contaminant Limit/Medium-Specific Concentrations/Protective Concentration Level
- NA - Not Analyzed
- µg/L - micrograms per liter
- a - duplicate sample
- mg/L - milligrams per liter
- J - Estimated: The analyte was positively identified, the quantitation is an estimation due to discrepancies in meeting certain analyte-specific quality control criteria.
- UJ - The analyte was not detected; however, the result is estimated due to discrepancies in meeting certain analyte-specific quality control criteria.
- U - Undetected: The analyte was analyzed for, but not detected.
- NV - No Value
- UB - considered a non-detect due to blank contamination
- *Perchlorate, manganese, and nickel compared to the PCL
- ** Value is for total xylenes
- PCL – Texas Risk Reduction Program (TRRP) Tier 1 Groundwater Residential



DEPARTMENT OF THE ARMY
LONGHORN ARMY AMMUNITION PLANT
POST OFFICE BOX 220
RATCLIFF, AR 72951

September 13, 2019

DAIM-ODB-LO

Mr. Rich Mayer
U.S. Environmental Protection Agency, Region 6
1201 Elm Street, Suite 500
Dallas, TX 75270-2102

Re: Final Signature Page for the Explanation of Significant Differences, Record of Decision
for Contingency Remedy at LHAAP-50, Former Sump Water Tank
Longhorn Army Ammunition Plant, July 2019

Dear Mr. Mayer,

Please find attached two replacement signature pages signed by the Army and EPA, and one electronic copy (compact disc) of the Final Explanation of Significant Differences for LHAAP-50, Former Sump Water Tank for your records.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in cursive script that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
Longhorn AAP Site Manager

Copies furnished:

- A. Palmie, TCEQ, Austin (letter)
- P. Bruckwicki, Caddo Lake NWR, TX (1 hard copy and 1 CD)
- A. Williams, USACE, Tulsa District, OK (1 CD)
- A. Sherman, USAEC, San Antonio, TX (1 CD)
- K. Nemmers, Bhate, Lakewood, CO (1 hard copy and 1 CD for project files)
- P. Srivastav, APTIM, Houston, TX



DEPARTMENT OF THE ARMY
LONGHORN ARMY AMMUNITION PLANT
POST OFFICE BOX 220
RATCLIFF, AR 72951

September 13, 2019

DAIM-ODB-LO

Ms. April Palmie
Texas Commission on Environmental Quality (TCEQ)
Superfund Section, MC-136
12100 Park 35 Circle, Bldg D
Austin, TX 78753

Re: Final Signature Page for the Explanation of Significant Differences, Record of Decision
for Contingency Remedy at LHAAP-50, Former Sump Water Tank
Longhorn Army Ammunition Plant, July 2019

Dear Ms. Palmie,

Please find attached one replacement signature page signed by the Army and EPA, and one electronic copy (compact disc) of the Final Explanation of Significant Differences for LHAAP-50, Former Sump Water Tank for your records.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

A handwritten signature in cursive script that reads "Rose M. Zeiler".

Rose M. Zeiler, Ph.D.
Longhorn AAP Site Manager

Copies furnished (letter only):
R. Mayer, USEPA, Region 6, Dallas, TX
P. Bruckwicki, Caddo Lake NWR, TX
A. Williams, USACE, Tulsa District, OK
A. Sherman, USAEC, San Antonio, TX
K. Nemmers, Bhate, Lakewood, CO (for project files)
P. Srivastav, APTIM, Houston, TX

**FINAL
EXPLANATION OF SIGNIFICANT DIFFERENCES
RECORD OF DECISION FOR
CONTINGENCY REMEDY AT LHAAP-50
FORMER SUMP WATER TANK
LONGHORN ARMY AMMUNITION PLANT**

July 2019

**Contract Number: W9128F-13-D-0012
Task Order Number: W912BV17F0150**

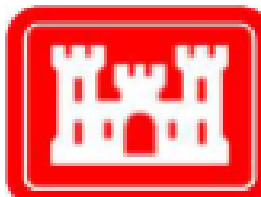
**Performance-Based Remediation
Longhorn Army Ammunition Plant
Karnack, Texas**

Prepared For:



**Longhorn Army Ammunition Plant
Karnack, Texas**

Under Contract To:



**U.S. Army Corps of Engineers
Tulsa District
Tulsa, Oklahoma**

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ACRONYMS AND ABBREVIATIONS

µg/kg	micrograms per kilogram
µg/L	micrograms per liter
AECOM	AECOM Technical Services, Inc.
APTIM	Aptim Federal Services, LLC
ARAR	applicable or relevant and appropriate requirement
AST	aboveground storage tank
BRAC	Base Realignment and Closure
CERCLA	Comprehensive Environmental Response, Compensation, and Liability Act
C.F.R.	Code of Federal Regulations
COC	chemical of concern
DCA	dichloroethane
DCE	dichloroethene
EPA	see USEPA
ESD	Explanation of Significant Differences
GWP-Ind	soil MSC for industrial use based on groundwater protection
GW-Res	groundwater MSC for residential use
ISB	in situ bioremediation
Jacobs	Jacobs Engineering Group, Inc.
LHAAP	Longhorn Army Ammunition Plant
LTM	long-term monitoring
LUC	land use control
MCL	maximum contaminant level
MNA	monitored natural attenuation
MSC	medium-specific concentration
NCP	National Oil and Hazardous Substances Pollution Contingency Plan
O&M	operations and maintenance
PCE	tetrachloroethene
PCL	protective concentration level
RAO	remedial action objective
RCRA	Resource Conservation and Recovery Act
RD	remedial design
ROD	Record of Decision
SDC-9™	APTIM's dechlorinating culture
Shaw	Shaw Environmental, Inc.
TCE	trichloroethene
TCEQ	Texas Commission on Environmental Quality
TNRCC	Texas Natural Resources Conservation Commission
TRRP	Texas Risk Reduction Program
U.S. Army	U.S. Department of the Army
U.S.C.	United States Code
USEPA	U.S. Environmental Protection Agency
VOC	volatile organic compound

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1 INTRODUCTION AND STATEMENT OF PURPOSE

Site and Location: LHAAP-50 is in the north-central section of Longhorn Army Ammunition Plant (LHAAP) and covers approximately 1 acre.

Lead Agency and Supporting Agency:

Lead Agency – U.S. Department of the Army (U.S. Army)

Lead Oversight Agency – U.S. Environmental Protection Agency (USEPA) Region 6

Support Agency – Texas Commission on Environmental Quality (TCEQ)

This Explanation of Significant Differences (ESD) is in compliance with the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA) §117 (c), 42 United States Code (U.S.C.) Section (§) 9617 (c) and National Oil and Hazardous Substances Pollution Contingency Plan (NCP) 40 Code of Federal Regulations (C.F.R.) §300.435(c)(2)(i).

Date of Record of Decision (ROD) Signature: September 2010, Administrative Record, Bate Stamp 00098892-00098976.

Need for ESD: The September 2010 ROD (Shaw, 2010), Section 1.4, specified the implementation of monitored natural attenuation (MNA) for the volatile organic compounds (VOCs) and perchlorate plume to verify that the plumes are stable, and that natural attenuation is occurring. The ROD also specified that performance objectives will be evaluated after two years of monitoring MNA and if MNA is found to be ineffective, a contingency remedy to enhance MNA will be implemented and documented in an ESD.

The 3rd Annual Remedial Action Operation Report for LHAAP-50 (APTIM, 2018) evaluated the performance of MNA for the VOCs and perchlorate plume. The report presented evidence of plume migration, increasing concentrations of perchlorate and trichloroethene (TCE), and geochemical conditions that are not optimal for MNA and recommended that an in situ bioremediation (ISB) contingency remedy be implemented at this time to enhance MNA.

The purpose of this ESD is to document the significant change from the ROD selected remedy of MNA and the proposed implementation of the ISB contingency remedy to enhance MNA. The ISB contingency remedy is consistent with the ROD requirement to enhance MNA, and is capable of degrading perchlorate and TCE to address the increased concentrations detected.

This ESD will become part of the Administrative Record file in accordance with NCP 40 CFR §300.825(a)(2). The file will be located at the Marshall Public Library:

Marshall Public Library
300 South Alamo Blvd.
Marshall, Texas 75670
Phone: 903 935 4465

Hours:
Monday, Tuesday, and Thursday 9:30 am to 7:30 pm
Wednesday and Friday 9:30 am to 5:30 pm
Saturday 9:30 am to 3:30 pm

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2 SITE HISTORY, CONTAMINATION, AND SELECTED REMEDY

2.1 SITE HISTORY AND CONTAMINATION

LHAAP-50, Former Sump Water Tank, contained a 47,000-gallon capacity aboveground storage tank (AST) which received industrial wastewater from various industrial waste production sumps throughout LHAAP between 1955 and 1988. After the solids were filtered, the storage tank contents were discharged into Goose Prairie Creek upstream of the Goose Prairie Creek bridge on South Crockett Avenue, south of 51st Street. The flow in the creek was sufficient to dilute the water to safe levels (Jacobs, 2002). If natural flow in the creek was considered insufficient, clean water was reportedly pumped into the creek to dilute the contents. The AST is no longer present.

The chemicals of concern (COCs) identified in the LHAAP-50 ROD include dissolved phase perchlorate and VOCs including tetrachloroethene (PCE), TCE, 1,1-dichloroethene (1,1-DCE), 1,2-dichloroethane (1,2-DCA), cis-1,2-dichloroethene (cis-1,2-DCE), and vinyl chloride in groundwater, and perchlorate in soil. There are no COCs in other environmental media at the site. The presence of these COCs at concentrations exceeding the human health cleanup levels in the soil and groundwater represented the primary driver for remedial action, as there were no ecological risks at the site.

Approximately 183 cubic yards of perchlorate-contaminated soil was removed and disposed offsite in September 2013 as described in the Final Remedial Action Completion Report (AECOM, 2016). An area of groundwater contamination is present in the shallow groundwater (upper and lower zones) that poses an unacceptable carcinogenic risk and non-carcinogenic hazard to a future maintenance worker under an industrial exposure scenario. There is no groundwater contamination in the intermediate groundwater zone.

The cleanup levels are the maximum contaminant level (MCL) for the VOCs and the Texas Risk Reduction Program (TRRP) protective concentration level (PCL) for residential use for perchlorate (USEPA, 2014). Concentrations of perchlorate, TCE, 1,1-DCE, and 1,2-DCA have been detected above the cleanup levels. The maximum concentration of perchlorate was detected in the eastern portion of the plume at 91,000 micrograms per liter ($\mu\text{g/L}$) in May 2018 above the Texas Residential Groundwater PCL. The maximum concentration of TCE as of May 2018 was detected at 620 $\mu\text{g/L}$ in the central part of the plume; 1,1-DCE concentrations were below the MCL as of May 2018; an isolated detection of 1,2-DCA was observed at 83 $\mu\text{g/L}$ as of May 2018.

A Federal Facility Agreement became effective December 30, 1991, among USEPA, the U.S. Army, and the Texas Natural Resources Conservation Commission (TNRCC), now the TCEQ. LHAAP-50 has been added to the list of National Priorities List sites at LHAAP with concurrence from the U.S. Army and USEPA Headquarters.

2.2 SELECTED REMEDY

The selected remedy identified as Alternative 2 in Section 2.12.2 of the ROD for LHAAP-50 includes soil removal, MNA and land use controls (LUCs). This alternative was selected because it was consistent with the intended future use of the site as a wildlife refuge. The alternative also satisfied the remedial action objectives (RAOs) for the site through LUC groundwater restriction, which would ensure protection of human health by preventing human exposure to contaminated groundwater. MNA, and a contingency remedy to enhance MNA, if MNA is found to be ineffective, would return the contaminated water to its potential beneficial use, wherever practicable. Furthermore, long-term monitoring (LTM) would assure that human health and the environment are being protected by verifying that contaminated groundwater does not migrate into nearby surface water bodies at levels that exceed MCLs. This alternative offered a high degree of long-term effectiveness that could be easily implemented at a lower cost than other alternatives.

The following language is taken from Section 2.12.2, Description of the Selected Remedy, of the ROD (Shaw, 2010):

- Excavation of the contaminated soil and disposal in a Resource Conservation and Recovery Act (RCRA)-permitted landfill will remove soil that is considered to be a contaminant source to groundwater, thereby, protecting groundwater. The estimated volume of soil to be removed is 150 cubic yards and is based on the conservative TCEQ soil medium-specific concentration (MSC) for industrial use based on groundwater protection (GWP-Ind) of 7,200 micrograms per kilogram ($\mu\text{g}/\text{kg}$) for perchlorate in soil. The removal of soil contamination will be verified by collecting confirmation samples from the walls and floors of the excavation area and submitting them for laboratory analysis for perchlorate.
- Semi-annual performance monitoring of Goose Prairie Creek adjacent to the LHAAP-50 will be conducted at two locations after excavation of the contaminated perchlorate pathway. Evaluation of this data will be included in the annual reports. The frequency and locations of sampling may be modified after evaluation of data. If perchlorate levels in the creek are consistently above the groundwater MSC for residential use (GW-Res) after two years of monitoring, then additional evaluation will be conducted, and any proposed actions will be included in the annual evaluation report to be submitted after Year 2. The need to continue creek sampling will be evaluated during the five-year reviews.
- *MNA to return groundwater to its potential beneficial use, wherever practicable.* Historic data suggest that natural attenuation of COCs is occurring at the site; however, additional data collection is necessary to fully evaluate natural attenuation. Monitoring wells will be sampled for eight consecutive quarters to evaluate and confirm the occurrence of natural attenuation in conjunction with historical data. Data from the eight quarterly events will be combined with historic data to evaluate the effectiveness of various natural physical, chemical, and biological processes in reducing contaminant concentrations.

- *Performance objectives to evaluate the MNA remedy performance after two years.* Each of the general performance objectives must be met as indicated below. If the criteria are not met to illustrate that MNA is an effective remedy, a contingency action would be initiated. If MNA is effective, a baseline will be established from the data to this point in time. Specific evaluation criteria will be developed in the Remedial Design (RD). The MNA evaluation will be based on the USEPA lines of evidence (USEPA, 1999) and the anaerobic screening (USEPA, 1998) as follows:
 - MNA potential based on evaluating biodegradation screening scores using USEPA guidance.
 - Plume stability (i.e., the plume concentrations are decreasing in the majority of performance wells, and the plume is not expanding in area as demonstrated with compliance wells).
 - MNA Process Evaluation demonstrated based on an attenuation rate calculated with empirical performance monitoring data, and MNA Process Demonstration based on the presence of daughter products and bacterial culture counts.
- *A contingency remedy to enhance MNA to reach the RAO if MNA is found to be ineffective.* The contingency remedy will use elements from other active remedial alternatives included in this ROD to address the ineffective aspects of MNA. The area and the elements of the contingency remedy would be selected based on the entire data set available. If a contingency remedy is implemented, it will be documented in an ESD.
- *Initiate LTM.* If MNA is determined to be effective, monitoring will be conducted to evaluate the remedy performance and determine if the plume conditions remain constant, improve or worsen after the baseline is established. Monitoring will continue after the initial eight quarters at a frequency of semiannual for three years, then annually until the next five-year review. The performance monitoring plan will be developed in the RD and will be based on USEPA guidance (USEPA, 2004).
- *Continue LTM every five years* to evaluate remedy performance and determine if plume conditions remain constant, improve, or worsen. The baseline of the plume for future five-year reviews will be established as part of the MNA evaluation program. The initial LTM plan will be developed during RD.
- *LUC to restrict access to the contaminated groundwater to environmental monitoring and testing only until cleanup levels are reached.* LUC implementation details will be included in the RD. The recordation notification for the site to be filed with Harrison County will include a description of the LUC. The boundary of the LUC will encompass the site boundaries and the plume boundaries.

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3 BASIS FOR THE DOCUMENT

The September 2010 ROD (Shaw, 2010), Section 2.12.2, contingency remedy component states that if a contingency remedy is implemented, it will be documented in an ESD. The 3rd Annual Remedial Action Operation Report for LHAAP-50 (APTIM, 2018) presented data that indicated geochemical conditions were not optimal for MNA. The 3rd Annual Remedial Action Operation Report for LHAAP-50 documented that MNA was found to be ineffective through several lines of evidence (APTIM, 2018). The TCE groundwater plume has expanded beyond its baseline footprint and now extends to 50WW12, the current downgradient monitoring well within the plume. At 50WW12, TCE concentrations increased from 0.5 µg/L in October 2013 to 79 µg/L in May 2018 while perchlorate concentrations increased from 23,600 µg/L in October 2013 to 91,000 µg/L in May 2018 suggesting continued migration within the current plume boundary. Though the perchlorate plume was relatively stable, a decreasing trend was observed only at one well (APTIM, 2018). The U.S. Army, USEPA, and TCEQ agree that MNA is currently not effective and that the contingency remedy should be implemented at this time (APTIM, 2018).

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4 DESCRIPTION OF SIGNIFICANT DIFFERENCES

As stated in Section 2.9.1 of the ROD, the groundwater remedy includes Excavation, MNA and LUC. The ROD states that a contingency remedy to enhance MNA will be implemented to reach the RAOs, if MNA is found to be ineffective, and will be documented in an ESD. The contingency remedy selected in this ESD to enhance MNA is ISB.

Change to Remedy Scope Presented in the ROD:

The only change to the remedy proposed in the ROD is the implementation of the contingency remedy (ISB) to enhance MNA. After three years of MNA performance monitoring, MNA was found to be ineffective (APTIM, 2018). The contingency remedy will consist of injection of emulsified vegetable oil, a microbial culture (SDC-9™), and nutrients to enhance the MNA remedy in an approximately 6,000 square foot area near monitoring wells 50WW12 and 50WW13. Upon implementation of the contingency remedy (ISB), two years of quarterly performance monitoring will be conducted. ISB will be the contingency remedy implemented for the impacted groundwater.

ROD Performance Objectives for the Groundwater Remedy, Section 2.12.2 Description of the Selected Remedy, paragraph 2:

The MNA evaluation will be based on the USEPA lines of evidence (USEPA, 1999) and the anaerobic screening (USEPA, 1998) as follows:

- MNA potential based on evaluating biodegradation screening scores using USEPA guidance.
- Plume stability (i.e., the plume concentrations are decreasing in the majority of performance wells, and the plume is not expanding in area as demonstrated with compliance wells).
- MNA Process Evaluation demonstrated based on an attenuation rate calculated with empirical performance monitoring data and MNA Process Demonstration based on the presence of daughter products and bacterial culture counts.

Change to Performance Objectives:

No change to the performance objectives in the ROD is proposed. MNA is not currently meeting the performance objectives. The 3rd Annual Remedial Action Operation Report for LHAAP-50 (APTIM, 2018) presented evidence of plume migration, increasing COC trends and geochemical conditions that are not optimal for MNA. The contingency remedy (ISB) will enhance MNA and performance objectives will be re-evaluated after two years of quarterly monitoring.

ROD Implementability Determination, Section 2.10.6, Implementability, paragraph 2:

Alternative 2 (MNA) is easily implemented from a technical standpoint with minimal construction activities followed by long-term sampling, maintenance and enforcement of the LUC.

Change to Implementability:

The contingency remedy (ISB) would be somewhat more difficult to implement due to the specialized expertise required for design and construction. However, ISB is effective in creating conditions conducive to reductive dechlorination and reducing VOC and perchlorate concentrations.

ROD Protection of Human Health and the Environment, Section 2.13.1, Protection of Human Health and the Environment, paragraph 1:

The selected remedy, Alternative 2, will protect human health and the environment, and achieve the RAOs for LHAAP-50. Although this alternative does not provide for human intervention to remediate groundwater, the alternative is a passive subsurface remedial action conducted by natural processes and mechanisms. The contaminated groundwater will be reduced to protective applicable or relevant and appropriate requirement (ARAR) levels, and the soil above protective ARAR levels will be removed. LUC would prevent human exposure to the contaminated groundwater by prohibiting the construction of potable wells within the LUC boundaries. Surface water monitoring of the creek will verify that the soil removal effectively mitigated the soil-to-groundwater pathway.

Change to Protection of Human Health and the Environment:

Currently, based on the 3rd Annual Remedial Action Operation Report for LHAAP-50 (APTIM, 2018), there is evidence of plume migration, increasing COC trends and geochemical conditions that are not optimal for MNA. The implementation of the contingency remedy (ISB) will enhance MNA and reduce groundwater contaminant concentrations which will prevent contaminated groundwater from migrating into nearby surface water at levels that may present an unacceptable risk to human health and the environment. Monitoring activities associated with the enhanced MNA would assure the protection of human health and the environment by documenting the return of the groundwater to its potential beneficial use as a drinking water supply, and by documenting reduction of the contaminant mass and protection of surface water through containment of the plume.

ROD Cost Estimate for the Selected Remedy, Section 2.12.3, paragraphs 1 and 2:

Table 2-8 in the ROD (Shaw, 2010) presents the present worth analysis of the cost for the selected remedy, Alternative 2. The information in this table is based on the best available information regarding the anticipated scope of the remedial alternative. The quantities used in the estimate are for estimating purposes only. Changes in the cost estimates are likely to occur as a result of new information and data collected during the engineering design of the remedial alternative. Major changes may be documented in the form of a memorandum in the Administrative Record, an ESD, or a ROD amendment. This is an order of magnitude engineering cost estimate that is expected to be within -30 to +50 percent of the actual project cost.

The total project present worth cost of this alternative is approximately \$639,000, using a discount rate of 2.8%. The capital cost is estimated at \$215,000. The total operations and maintenance (O&M) present value cost is estimated at approximately \$424,000. The O&M cost includes evaluation of MNA, maintenance of LUC, and LTM through Year 30. The LTM would support the required CERCLA five-year reviews.

Change to Cost Estimate for the Selected Remedy:

The implementation of the contingency remedy (ISB) for Alternative 2 will increase the overall costs associated with this remedial alternative. This increase in cost is due to the capital cost associated with the use of ISB technology to enhance MNA.

It is estimated that implementation of the contingency remedy (ISB) associated with Alternative 3 will increase the original estimate for this alternative by approximately \$280,000.

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5 SUPPORT AGENCY COMMENTS

The USEPA and TCEQ have reviewed this ESD and support the changes to the selected remedy.

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6 STATUTORY DETERMINATION

The modification presented herein satisfies CERCLA §121, 42 U.S.C. §9621. The contingency remedy (ISB) will enhance MNA and reduce groundwater contaminant concentrations.

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7 PUBLIC PARTICIPATION

A notice summarizing the ESD shall be published in the Marshall News Messenger upon finalization of the ESD. This ESD and all supporting ESD documentation will be made a part of the Administrative Record file in accordance with the NCP at 40 C.F.R. §300.825(a)(2). The Administrative Record will be located at the repository identified in Section 1.0 of this document. All public participation requirements set out in NCP at 40 C.F.R. §300.435(c)(2)(i) have been met.

The Proposed Plan for Remedial Action at the site was released for public comments on 25 January 2010. The Proposed Plan identified the preferred alternative to be Alternative 2:

- Excavation and off-site disposal of perchlorate contaminated soil
- MNA of groundwater and LUCs
- A contingency remedy to enhance MNA, if MNA is found to be ineffective

The U.S. Army reviewed all written and oral comments submitted during the public comment period. There were no significant comments captured related to the groundwater remedy.

Authorizing Signature:

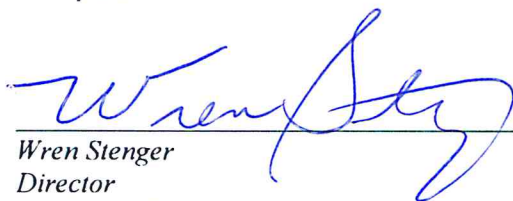


Thomas E. Lederle
Chief, ACSIM BRAC Division
U.S. Department of the Army

Date: 6 Aug 2019

Authorizing Signature:

I have reviewed this document, and any comments I had have been addressed and/or incorporated:



Wren Stenger
Director
Superfund Division
U.S. Environmental Protection Agency, Region 6

Date: 8/29/19

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8 REFERENCES

AECOM Technical Services, Inc. (AECOM), 2016, Final Remedial Action Report, LHAAP-50, Former Sump Water Tank, Longhorn Army Ammunition Plant, Karnack, Texas. June.

Aptim Federal Services, LLC (APTIM), 2018, *Draft Final 3rd Annual Remedial Action Operation Report, LHAAP-50 Shops Former Sump Water Tank, Longhorn Army Ammunition Plant, Karnack, Texas.* July.

Jacobs Engineering Group, Inc. (Jacobs), 2002, Final Remedial Investigation Report for the Group 4 Sites, Sites 35A, 35B, 35C, 46, 47, 48, 50, 60, 67, and Goose Prairie Creek, Longhorn Army Ammunition Plant, Karnack, Texas, Oak Ridge, TN, January (RI).

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U.S. Environmental Protection Agency (USEPA), 1998, *Technical Protocol for Evaluating Natural Attenuation of Chlorinated Solvents in Ground Water, EPA/600/R-98/128,* September.

USEPA, 1999, *Use of Monitored Natural Attenuation at Superfund, RCRA Corrective Action, and Underground Storage Tank Sites, Directive 9200.4-17P,* April.

USEPA, 2004, *Performance Monitoring of MNA Remedies for VOCs in Ground Water, EPA/600/R-04/027,* April.

USEPA, 2014, Transmittal of Final Dispute Decision, Letter dated October 31, 2014, from Gina McCarthy, USEPA Administrator, to John McHugh, Secretary of the Army, and Bryan Shaw, Chairman of TCEQ. Longhorn Army Ammunition Plant, Karnack, Texas, Administrative Record, Volume 4, 2014, 00192717-00192750.

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DEPARTMENT OF THE ARMY
 LONGHORN ARMY AMMUNITION PLANT
 POST OFFICE BOX 220
 RATCLIFF, AR 72951

September 26, 2019

DAIM-ODB-LO

Ms. April Palmie
 Texas Commission on Environmental Quality
 Remediation Division
 12100 Park 35 Circle, Bldg D
 Austin, TX 78753

**Re: LHAAP-29 Record of Decision, August 2019
 Longhorn Army Ammunition Plant, Karnack, Texas**

Dear Ms. Palmie,

Enclosed please find the replacement pages for the August 2019 LHAAP-29 Record of Decision (ROD) for your records. This is the final ROD and includes the completed signature page with Army and EPA signatures, and the TCEQ concurrence.

The Final document transmitted for signature and concurrence on 16 August 2019 was revised in response to EPA's 4 September 2019 request to remove "Major" from the fourth sentence in Section 2.12.3 Cost Estimate of the Selected Remedy. The sentence "Major changes may be documented in the form of a memorandum in the Administrative Record, an Explanation of Significant Difference (ESD), or a ROD amendment" was revised to "Changes will be documented in accordance with 40 CFR 300.435(c)(2) in the form of a memorandum in the Administrative Record, an Explanation of Significant Difference (ESD), or a ROD amendment, as necessary." The revision also required changes to the notes for Tables 2-12 and 2-13. Other than these changes, the document remains the same as the 16 August 2019 Final ROD.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

Rose M. Zeiler, Ph.D.
 Longhorn AAP Site Manager

One Enclosure
 Copies furnished:

R. Mayer, USEPA, Dallas, TX
 R. Smith, USACE, Tulsa District, OK

A. Williams, USACE, Tulsa District, OK
 A. Sherman, USAEC, San Antonio, TX

K.Nemmers, Bhat
 P. Werner, HDR



DEPARTMENT OF THE ARMY
 LONGHORN ARMY AMMUNITION PLANT
 POST OFFICE BOX 220
 RATCLIFF, AR 72951

September 26, 2019

DAIM-ODB-LO

Mr. Rich Mayer
 U.S. Environmental Protection Agency
 Federal Facilities Section R6
 1201 Elm Street, Suite 500
 Dallas, TX 75202-2102

**Re: LHAAP-29 Record of Decision, August 2019
 Longhorn Army Ammunition Plant, Karnack, Texas**

Dear Mr. Mayer,

Enclosed please find the August 2019 LHAAP-29 Record of Decision (ROD) for your records. This is the final ROD and includes the completed signature page with Army and EPA signatures, and the TCEQ concurrence.

The Final document transmitted for signature and concurrence on 16 August 2019 was revised in response to EPA's 4 September 2019 request to remove "Major" from the fourth sentence in Section 2.12.3 Cost Estimate of the Selected Remedy. The sentence "Major changes may be documented in the form of a memorandum in the Administrative Record, an Explanation of Significant Difference (ESD), or a ROD amendment" was revised to "Changes will be documented in accordance with 40 CFR 300.435(c)(2) in the form of a memorandum in the Administrative Record, an Explanation of Significant Difference (ESD), or a ROD amendment, as necessary." The revision also required changes to the notes for Tables 2-12 and 2-13. Other than these changes, the document remains the same as the 16 August 2019 Final ROD.

The point of contact for this action is the undersigned. I may be contacted at 479-635-0110, or by email at rose.m.zeiler.civ@mail.mil.

Sincerely,

Rose M. Zeiler, Ph.D.
 Longhorn AAP Site Manager

One Enclosure
 Copies furnished:

A. Palmie, TCEQ, Austin, TX
 R. Smith, USACE, Tulsa District, OK

A. Williams, USACE, Tulsa District, OK
 A. Sherman, USAEC, San Antonio, TX

K.Nemmers, Bhat
 P. Werner, HDR



Jon Niermann, *Chairman*
Emily Lindley, *Commissioner*
Toby Baker, *Executive Director*

TEXAS COMMISSION ON ENVIRONMENTAL QUALITY

Protecting Texas by Reducing and Preventing Pollution

September 13, 2019

Mr. Thomas E. Lederle
Chief, ACSIM BRAC Division
2530 Crystal Drive, Room 5000
Taylor Bldg / NC3
Arlington, VA 22202

Subject: Record of Decision for LHAAP-29, Former TNT Production Area,
Longhorn Army Ammunition Plant Federal Superfund Site TX6213820529
Karnack, Harrison County, Texas

Dear Mr. Lederle:

The Texas Commission on Environmental Quality (TCEQ) received the final Record of Decision (ROD) for LHAAP-29, Former TNT Production Area, Longhorn Army Ammunition Plant Federal Superfund Site in Karnack, Texas on August 16, 2019. The TCEQ has completed the review of the above referenced document and concurs that the described action is appropriate.

Sincerely,

A handwritten signature in black ink, appearing to read "Toby Baker".

Toby Baker
Executive Director

Cc: Ms. Wren Stenger, Director, Superfund Division, US Environmental Protection Agency,
Region 6



Final

Record of Decision

for LHAAP-29,
Former TNT Production Area, Group 2
Longhorn Army Ammunition Plant
Karnack, Texas

August 2019

Prepared For:



U.S. Army Corps of Engineers – Tulsa District

Prepared By:

HDR

**9871 S. Meridian Blvd, Suite 400
Englewood, CO 80112**

**Contract No. W912BV-15-D-0014
Task Order No. W912BV18F0023**

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Final
RECORD OF DECISION
FOR
LHAAP-29, FORMER TNT PRODUCTION AREA, GROUP 2
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

Prepared For:
U.S. Army Corp of Engineers Tulsa District

Prepared By:
HDR, Inc.
9871 S. Meridian Blvd, Suite 400
Englewood, CO 80112

Contract No. W912BV-15-D-0014
Task Order No. W912BV18F0023

August 2019

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Appendix A: Public Notice Affidavits



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Acronyms and Abbreviations

ARAR	applicable or relevant and appropriate requirement
BERA	baseline ecological risk assessment
bgs	below ground surface
BHHRA	baseline human health risk assessment
°C	degrees Celsius
CDI	chronic daily intake
CERCLA	Comprehensive Environmental Response, Compensation, and Liability Act
C. F. R.	Code of Federal Regulations
COC	chemical of concern
COPEC	chemical of potential ecological concern
COPC	chemical of potential concern
CSM	conceptual site model
CWA	Clean Water Act of 1972
DCA	dichloroethane
DCE	dichloroethene
DNAPL	dense non-aqueous phase liquid
DNT	dinitrotoluene
ECP	environmental condition of property
EcoPRG	ecological preliminary remediation goal
EEQ	ecological effects quotient
EISB	Enhanced in-situ bioremediation
EPC	exposure point concentration
ERH	Electrical resistance heating
ESD	Explanation of Significant Differences
FFA	Federal Facility Agreement
FR	Federal Register
FS	feasibility study
g/L	grams per liter
GWP-Ind	medium-specific concentration for industrial use based on groundwater protection
GWTP	Groundwater treatment plant
HEAST	Health Effects Assessment Summary Tables
HI	hazard index
HQ	hazard quotient
IRIS	Integrated Risk Information System
ISTD	In-situ thermal desorption
Jacobs	Jacobs Engineering Group
LHAAP	Longhorn Army Ammunition Plant
LTM	long-term monitoring
LUC	land use control
MC	methylene chloride
MCL	maximum contaminant level
µg/L	Micrograms per liter
mg/kg	milligrams per kilogram
mg/L	Milligrams per liter
mg/kg-day	milligrams per kilogram per day
MNA	monitored natural attenuation
MOA	memorandum of agreement
MSC	medium-specific concentration
NCP	National Oil and Hazardous Substances Pollution Contingency Plan



NOAEL	no-observed adverse effect level
NPDES	National Pollutant Discharge Elimination System
NPL	National Priorities List
NT	nitrotoluene
O&M	operation and maintenance
PCB	polychlorinated biphenyl
PCL	Protective concentration level
PP	Proposed Plan
RA	remedial action
RAB	Restoration Advisory Board
RAO	remedial action objective
RAWP	Remedial Action Work Plan
RCRA	Resource Conservation and Recovery Act
RD	remedial design
RFA	RCRA Facility Assessment
RfD	reference dose
RI	remedial investigation
ROD	record of decision
SAI-Ind	soil MSC for industrial use based on inhalation, ingestion, and dermal contact
SARA	Superfund Amendments and Reauthorization Act
scfm	standard cubic feet per minute
SDWA	Safe Drinking Water Act
SF	slope factor
Shaw	Shaw Environmental, Inc.
STEP	Solutions to Environmental Problems, Inc.
SVE	Soil vapor extraction
SVOC	semivolatile organic compound
TAC	Texas Administrative Code
TCDD	tetrachlorodibenzo-p-dioxin
TCE	trichloroethene
TCEQ	Texas Commission on Environmental Quality
TCH	Thermal conductance heating
TDS	Total dissolved solids
TEC	toxicity equivalence concentration
TNT	trinitrotoluene
TRRP	Texas Risk Reduction Program
TRRP Residential Groundwater PCL	Texas Risk Reduction Program Tier 1 Residential Groundwater Protective Concentration Level
TRV	toxicity reference level
UCL	upper confidence limit
U.S. Army	U.S. Department of the Army
USACE	U.S. Army Corps of Engineers
USAEHA	U.S. Army Environmental Hygiene Agency
USATHAMA	U.S. Army Toxic and Hazardous Materials Agency
USC	U.S. Code
USEPA	U.S. Environmental Protection Agency
USFWS	U.S. Fish and Wildlife Service
VC	vinyl chloride
VOC	volatile organic compound



1. The Declaration

1.1 Site Name and Location

Longhorn Army Ammunition Plant (LHAAP)-29, Former TNT Production Area, Group 2
Longhorn Army Ammunition Plant
Karnack, Texas

Comprehensive Environmental Response, Compensation, and Liability Information System, U.S. Environmental Protection Agency (USEPA) Identification Number: TX6213820529.

1.2 Statement of Basis and Purpose

This Record of Decision (ROD) presents the selected remedy for LHAAP-29, Former Trinitrotoluene (TNT) Production Area, Group 2 located at the Longhorn Army Ammunition Plant (LHAAP) in Karnack, Texas. The remedy was chosen in accordance with the Comprehensive Environmental Response, Compensation, and Liability Act (CERCLA) of 1980, as amended by the Superfund Amendments and Reauthorization Act (SARA) of 1986, and, to the extent practicable, the National Oil and Hazardous Substances Pollution Contingency Plan (NCP), Code of Federal Regulations (C. F. R.) Title 40 §300.

The remedy selection was based on documentation available in the Administrative Record for the site, including the remedial investigation (RI) (Jacobs Engineering Group, Inc. [Jacobs], 2001), baseline human health risk assessment (BHHRA) report (Jacobs, 2002), installation-wide baseline ecological risk assessment (BERA) report (Shaw Environmental, Inc. [Shaw], 2007a), feasibility study (FS) (Shaw, 2010), RI Addendum (AECOM Technical Services [AECOM], 2016), Final Baseline Ecological Risk Assessment Addendum (AGEISS, 2014), FS Addendum (AECOM, 2017) and Revised Proposed Plan (PP) (U.S. Department of the Army [U.S. Army], 2018).

This document is issued by the U.S. Army, the lead agency for this installation. The U.S. Army, USEPA, and the Texas Water Commission (currently known as the TCEQ) entered into the FFA for remedial activities at LHAAP which became effective on December 30, 1991. The USEPA (Region 6) and the Texas Commission on Environmental Quality (TCEQ) are the regulatory agencies providing technical support, project review and comment, and oversight of the LHAAP cleanup program. The USEPA and the U.S Army jointly select the remedy and TCEQ concurs with the selected remedy in this Record of Decision (ROD).

1.3 Assessment of the Site

The response action selected in this ROD is necessary to protect the public health or welfare or the environment from actual or threatened releases of hazardous substances, pollutants, or contaminants into the environment.

1.4 Description of the Selected Remedy

The final selected remedy for LHAAP-29 includes excavation and off-site disposal of contaminated soil, flushing, inspection, and plugging of the trinitrotoluene (TNT) cooling water and wastewater lines, in situ thermal desorption (ISTD) treatment of the intermediate groundwater zone dense non-



aqueous phase liquid (DNAPL) plume, monitored natural attenuation (MNA) for the shallow zone groundwater plumes and for the intermediate groundwater plume following ISTD, and land use controls (LUCs) for soil and groundwater.

The final selected remedy for LHAAP-29 protects human health and the environment by preventing human and ecological receptor exposure to contaminated soil and contaminated groundwater. The human health risk assessment scenarios evaluated were based on the hypothetical future maintenance worker. In the soil, chemicals of concern (COCs) and chemicals of potential ecological concern (COPECs) are explosives (2,4,6-TNT, 2,4-dinitrotoluene [DNT], and 2,6-DNT). Perchlorate is considered a potential soil COC based on groundwater concentrations. In the shallow groundwater zone, the COCs are Volatile Organic Compounds (VOCs) trichloroethene [TCE], 1,2-dichloroethane (DCA), TCE's daughter products cis-1,2-dichloroethene [DCE]), trans 1,2 DCE, and vinyl chloride [VC]), metals (arsenic, mercury, nickel, selenium), explosives (2,4-DNT, 2,6-DNT, 2-nitrotoluene [NT], 3-NT, 4-NT), and perchlorate. In the intermediate groundwater zone, the COCs are VOCs (methylene chloride (MC), TCE, 1,2-DCA, TCE's daughter products 1,1-DCE, cis-1,2-DCE, trans-1,2 DCE, and VC) and arsenic. COCs in the transite TNT wastewater and vitrified clay cooling water lines include explosives (2,4,6-TNT, 2,4-DNT, 2,6-DNT, 2-amino-4,6-DNT, and 4-amino-2,6-DNT, 1,3-dinitrobenzene). Residual MC DNAPL acting as a source material in the intermediate zone may be considered a principal threat waste at LHAAP-29. There are no COCs associated with the deep groundwater zone.

The components of the selected remedy are summarized below:

- Contaminated soil removal with off-site disposal to protect the hypothetical future maintenance worker and ecological receptors and eliminate the soil-to-groundwater pathway. Additional confirmation soil sampling during the remedial design (RD) will be needed to define the final excavation extent and volume of soil contaminated with explosives near former Building 812-F and in the cooling water outfall/ditch and may identify additional areas for soil removal adjacent to the North and South Cooling water lines as well as the wooden and transite TNT wastewater lines, see **Section 2.12.2**.
- Flushing, inspection, and plugging of the transite TNT wastewater line and the vitrified clay cooling water lines to eliminate potential exposure from residual contamination. Confirmation sampling of the wooden TNT wastewater line during the RD may result in excavation, see **Section 2.12.2**.
- MNA in the shallow groundwater zone to confirm protection of human health and the environment by documenting that contaminated groundwater remains localized with minimal migration and that COCs are being reduced to cleanup levels.
 - Performance objectives will be evaluated after two years of MNA. During those two years, monitoring will be quarterly. If MNA is found to be effective, it will be continued, and long-term monitoring (LTM) will be semiannual for three years. In subsequent years, LTM will be annual until the next five-year review and annually thereafter until recommended otherwise by the five-year review. The monitoring and reporting associated with this remedy will be used to track the effectiveness of MNA and will continue until recommended otherwise at the five-year review.
 - If MNA is found to be ineffective, a contingency remedy to enhance MNA would be developed. The contingency remedy would consist of injection of bioremediation



amendments to enhance degradation of the groundwater contaminants at selected locations based on data available at the time it is determined MNA is not successful. Development and specific description of the contingency remedy would be presented in a Remedial Design/Remedial Action Work Plan (RD/RAWP).

- ISTD treatment will be performed for the DNAPL MC plume in the intermediate groundwater zone to reduce concentrations to levels amenable to MNA. One of two ISTD treatment process options, Electrical Resistance Heating (ERH) or Thermal Conduction Heating (TCH), will be selected during the RD.
- MNA will be implemented in the intermediate groundwater zone following successful implementation of ISTD to confirm protection of human health and the environment by documenting that the contaminated groundwater remains localized with minimal migration and that contaminant concentrations are being reduced to cleanup levels. Trigger level or target value for successful ISTD treatment of MC in the intermediate zone is 8,000 micrograms per liter ($\mu\text{g/L}$). Performance monitoring will be conducted at a frequency necessary to evaluate the effectiveness of the in-situ treatment, with sampling and LTM conducted as described for the shallow groundwater MNA.
- Groundwater monitoring will be conducted to evaluate inorganic COCs. The need to continue groundwater monitoring for this purpose will be evaluated at five year reviews.
- The LUC objectives include maintaining the integrity of any current or future remedial or monitoring systems, and preventing the use of groundwater contaminated above cleanup levels as a potable water source. The groundwater treatment and MNA remedial components include a groundwater monitoring system that will be used to characterize the condition of the groundwater during the period the groundwater remedy is in place until the groundwater remediation goals are achieved, and to demonstrate achievement of the groundwater remediation goals when the groundwater remedy is complete. As a part of this groundwater remedy, the Army will maintain the remedial and monitoring systems associated with the groundwater remedies until these components of the remedy are no longer needed to achieve cleanup levels, and cleanup levels have been achieved. During the period of operation of the groundwater remedy, if any of the elements of the remedial and groundwater monitoring systems are damaged, destroyed, or become ineffective, they will be repaired or replaced with suitable components to assure that the remedial and groundwater monitoring systems are able to provide data of the quality necessary to determine the progress of and eventual completion of this component of the remedy. The actions to be taken to implement these LUC objectives and requirements will be provided through modifying the “Comprehensive Land Use Control (LUC) Management Plan, Former Longhorn Army Ammunition Plant, Karnack, Texas” and detailed in the LUC RD.¹
- The LUC for prohibition of groundwater use (except for monitoring and testing) shall be implemented and shall remain in place at the Site until the COCs (i.e. including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soil and groundwater remaining at the site, are reduced below levels that would support unlimited use and unrestricted exposure. A LUC RD will be finalized as the land use component of the RD. Within 21 days of the issuance of the ROD, the Army will

¹ This paragraph is October 31, 2014 Dispute Decision language that is included despite the ROD not being subject to the dispute.



propose deadlines for completion of the RD Work Plan, RD and Remedial Action Work Plan. The documents will be prepared and submitted to the EPA and the TCEQ pursuant to the FFA. The LUC RD will contain implementation and maintenance actions, including periodic inspections. The long-term groundwater and surface water monitoring and MNA performance monitoring plan will also be presented in the RD. The recordation notification for the Site, which will be filed with Harrison County, will include a description of the LUCs.¹ The preliminary boundary for the groundwater LUC is shown on **Figure 2-16**.

- The LUC restricting land use to nonresidential shall be implemented until it is demonstrated that surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure.¹
- The LUC to maintain the integrity of any current or future remedial or monitoring systems will remain in place until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met. The LUC to prohibit groundwater use (except for environmental monitoring and testing) as a potable source will remain in place until the levels of COCs (i.e., all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soil and groundwater allow for unlimited use and unrestricted exposure.¹

CERCLA five-year reviews are required until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soil and groundwater allow for unlimited use and unrestricted exposure.

Based on a preliminary natural attenuation evaluation and groundwater modeling, cleanup levels are expected to be met through natural attenuation in the shallow zone groundwater in approximately 70 years (Shaw, 2010), and 5-10 years following ISTD treatment in the intermediate zone groundwater (AECOM, 2017). Specifically, based on the attenuation of 1,2-DCA, MNA is estimated to take approximately 70 years in the shallow groundwater zone. The ISTD treatment in the intermediate zone is estimated to take 65-87 days if ERH is used, and 180 days if TCH is used. MNA will follow the in-situ treatment, and is estimated to take 5-10 years based on the attenuation of TCE. Other COCs are expected to require less time to attenuate, based on the natural attenuation study presented in the Final FS (Shaw, 2010). Considering the lithologic variability, particularly the lateral and vertical change from sand to clay, the time to achieve cleanup levels may vary. In the course of the remedy, the additional monitoring results will allow more accurate time estimates.

No adverse impact is expected to the surface water during the time it would take natural attenuation to reduce contaminant concentrations to cleanup levels.

A LUC RD will be finalized as the land use component of the Remedial Design. Within 21 days of the issuance of the ROD, the Army will propose deadlines for completion of the RD Work Plan, RD, and Remedial Action Work Plan. The documents will be prepared and submitted to EPA and TCEQ pursuant to the FFA. The LUC RD will contain implementation and maintenance actions, including periodic inspections. The long-term groundwater and surface water monitoring and MNA performance monitoring plan will also be presented in the RD.¹

The Army will implement, maintain, monitor, report on and enforce land use controls at Army-owned property. The Army shall perform those actions related to land use control activities described in this ROD and in the Remedial Design for the ROD. For portions of the Site subject to land use controls



that are not owned by the Army, the Army will monitor and report on the implementation, maintenance, and enforcement of land use controls, and coordinate with federal, state, and local governments and owners and occupants of properties subject to land use controls. The Army will provide notice of the groundwater and soil (surface and subsurface) contamination and any land use restrictions referenced in the ROD. The Army will send these notices to the federal, state and local governments involved at this site and the owners and occupants of the properties subject to those use restrictions and land use controls. The Army shall provide the initial notice within 90 days of ROD signature. The frequency of subsequent notifications will be described in the Remedial Design for the ROD. The Army remains responsible for ensuring that the remedy remains protective of human health and the environment. The Army will fulfill its responsibility and obligations under CERCLA and the NCP as it implements, maintains, and reviews the selected remedy.¹

Upon transfer of Army-owned property, the Army will provide written notice of the land use controls to the transferee of the groundwater and soil (surface and subsurface) contamination and any land use restrictions referenced in the ROD. Within 15 days of transfer, the Army shall provide EPA and the TCEQ with written notice of the division of implementation, maintenance, and enforcement responsibilities unless such information has already been provided in the LUC RD. The LUC RD will address the procedures to be used by the Army and the transferee to document compliance with the LUCs described in this ROD. In the event property is transferred out of Federal control, the land use controls relating to property and groundwater restrictions shall be recorded in the deed and shall be enforceable by the United States and the state of Texas.¹

U.S. Army and regulators will consult to determine appropriate enforcement actions should there be a failure of a LUC objective at the site after they have been transferred.

The management strategy at LHAAP is to approach each site separately to address human health issues and to approach the sites by sub-area to address ecological risk (Shaw, 2007a). Thus, the implementation of this remedy at LHAAP-29 is independent of any other remedial action at LHAAP to address human health issues. To address ecological risk, LHAAP-29 was grouped with several other sites as part of the Industrial Sub-Area. The final COPECs in soil that require remedial action in the Industrial Sub-Area are 2,4-DNT, 2,6-DNT and 2,4,6-TNT (Shaw, 2010). The remedial actions at LHAAP-29 will be sufficient to remove ecological risks for the sub-area. This management strategy is considered to be endorsed by regulators as evidenced by the regulatory approval of the BERA (Shaw, 2007a) and BERA Addendum (AGEISS, 2014).

1.5 Statutory Determinations

The selected remedy is protective of human health and the environment, complies with Federal and State requirements that are established as applicable or relevant and appropriate requirements (ARARS) for the remedial action, and is cost-effective.

The remedy offers long-term effectiveness through excavation of contaminated soil, thermal treatment of MC DNAPL, flushing, plugging, and abandoning the TNT wastewater and cooling water lines, and the implementation of a LUC, which will minimize the potential risk to the hypothetical future maintenance worker posed by the contaminated soil and groundwater. Evaluation of MNA, including routine monitoring of the attenuation until cleanup levels are met, will document the effectiveness of the selected remedy. In the event that MNA is determined to be ineffective, a contingency remedy consisting of injection of bioremediation amendments to enhance degradation of the groundwater contaminants at selected locations will be developed and implemented.



Development and specific description of the contingency remedy will be presented in a RD/RAWP. The selected remedy is easily and immediately implementable and while it was not the lowest cost alternative, is considered most likely to be effective and successful compared to the other alternatives considered for LHAAP-29.

The thermal treatment (ISTD) component of the selected remedy satisfies the statutory preference for treatment as a principal treatment element of the remedy. The MNA component does not address the statutory preference for treatment to the maximum extent practicable; MNA is a passive remedial action using natural processes.

The selected remedy will reduce the toxicity, mobility, or volume of contaminants in the groundwater through active and passive remedial actions. The high concentrations of MC in the intermediate groundwater at LHAAP-29 have indicated that DNAPL residual source material may be residing in the subsurface and acting as a principal threat in the groundwater.

Because hazardous substances, pollutants, or contaminants will remain at the site above levels that allow for unlimited use and unrestricted exposure, a five-year review will be conducted every 5 years to ensure protection of human health and the environment under CERCLA §121(c), U.S. Code (USC) Title 42 §9621(c). In accordance with Texas Administrative Code (TAC) Title 30 §335.566, a notification will be recorded in Harrison County records restricting land use to nonresidential until it is demonstrated that surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure; that a prohibition of groundwater use (except for environmental monitoring and testing) as a potable source will remain in place until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soil and groundwater allow for unlimited use and unrestricted exposure; and, that the integrity of any current or future remedial or monitoring systems will remain in place until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met. Although the U.S. Army may later pass these procedural responsibilities to the transferee by property transfer agreement, the U.S. Army shall retain ultimate responsibility for remedy integrity per the FFA and CERCLA §121.

1.6 ROD Data Certification Checklist

The following information is included in the Decision Summary section of this ROD. Additional information can be found in the Administrative Record for this site.

- Current and reasonably anticipated future land use assumptions and current and potential future beneficial uses of groundwater as identified in the baseline risk assessment and ROD (**Section 2.6**).
- Potential groundwater use that will be available at the site as a result of the selected remedy (**Section 2.6**).
- COCs and their concentrations (**Section 2.7**).
- Baseline risk represented by the COCs (**Section 2.7**).
- Cleanup levels established for COCs and the basis for these levels (**Sections 2.7.3 and 2.8**).



- Principal threat wastes that will be addressed at this site (**Section 2.11**).
- Key factor(s) that led to selecting the remedy (**Section 2.12**).
- Estimated capital, annual operation and maintenance (O&M), and total present worth costs, discount rate, and the number of years over which the remedy cost estimates are projected (**Section 2.12**).



1.7 Authorizing Signatures

As the lead agency, the U.S. Army issues this ROD for LHAAP-29 which documents the final selected remedy. The undersigned is the appropriate approval authority for this decision.

Thomas E. Lederle

16 Aug 2019

(Date)

Thomas E. Lederle
Division Chief
Base Realignment and Closure Division
Assistant Chief of Staff for Installation
Management
U.S. Army

The United States Environmental Protection Agency approves the selected remedy as provided in the ROD for LHAAP-29.

Wren Stenger

September 19, 2019

(Date)

Wren Stenger
Director
Superfund Division
U.S. Environmental Protection Agency
Region 6



2. Decision Summary

2.1 Site Name, Location, and Description

LHAAP-29, Former TNT Production Area, Group 2
Longhorn Army Ammunition Plant
Karnack, Texas

Comprehensive Environmental Response, Compensation, and Liability Information System
USEPA Identification Number: TX6213820529

Lead Agency: U.S. Army, Department of Defense
Support Agencies: USEPA Region 6, TCEQ

Source of Cleanup Money: U.S. Army, Department of Defense
Site Type: Industrial Facility

The former LHAAP is an inactive, government-owned, formerly contractor operated and maintained, Department of Defense facility located in central east Texas (see **Figure 2-1**) in the northeast corner of Harrison County. LHAAP is approximately 14 miles northeast of Marshall, Texas, and approximately 40 miles west of Shreveport, Louisiana. The former U.S. Army installation occupied 8,416 acres between State Highway 43 at Karnack, Texas, and the southwestern shore of Caddo Lake. The facility can be accessed via State Highways 43 and 134.

LHAAP was placed on the Superfund National Priorities List (NPL) on August 9, 1990. Activities to remediate contamination began in 1990. After its listing on the NPL, the U.S. Army, the USEPA, and the Texas Water Commission (currently known as the TCEQ) entered into a CERCLA §120 FFA for remedial activities at LHAAP. The FFA became effective December 30, 1991. LHAAP operated until 1997 when it was placed on inactive status and classified by the U.S. Army Armament, Munitions, and Chemical Command as excess property. The majority of LHAAP, not including LHAAP-29, has been transferred by the U.S. Army to the U.S. Fish and Wildlife Service (USFWS) for management as the Caddo Lake National Wildlife Refuge.

LHAAP-29, Former TNT Production Area is an 85-acre site located within a heavily wooded section in the western-central portion of LHAAP (**Figure 2-2**). The surface features at LHAAP-29 include the foundations for the former production facilities and the underground pipelines that were originally built for cooling water drainage and TNT wastewater conveyance.

2.2 Site History and Enforcement Activities

2.2.1 History of Site Activities

LHAAP was established in December 1941 with the primary mission of manufacturing TNT. Production of TNT began at Plant 1 in October 1942 and continued through World War II until August 1945, when the facility was placed on standby status until February 1952. In 1952, the LHAAP facility was reactivated with the opening of Plant 2, where pyrotechnic ammunition, such as photoflash bombs, simulators, hand signals, and tracers for 40 millimeter ammunition, were produced until 1956.



In December 1954, a third facility, Plant 3, began production of solid-fuel rocket motors for tactical missiles. Rocket motor production at Plant 3 continued to be the primary operation at LHAAP until 1965 when Plant 2 was reactivated for the production of pyrotechnic and illuminating ammunition. In the years following the Vietnam conflict, LHAAP continued to produce flares and other basic pyrotechnic or illuminating items for the U.S. Department of Defense inventory. From September 1988 to May 1991, LHAAP was also used for the static firing and elimination of Pershing I and II rocket motors in compliance with the Intermediate-Range Nuclear Force Treaty in effect between the United States and the former Union of Soviet Socialist Republics. LHAAP operated until 1997 when it was placed on inactive status and classified by the U.S. Army Armament, Munitions, and Chemical Command as excess property.

LHAAP-29 was used as a TNT manufacturing facility from October 1942 to August 1945. The facility produced approximately 400 million pounds of flake TNT during its operation using six TNT production lines (five active and one standby). The TNT production facility was inactive from August 1945 to 1959. In 1959, most of the buildings and aboveground storage tanks were removed. The debris was burned or flashed at Burning Ground No. 2/Flashing Area (LHAAP-17). Concrete foundations open-top concrete-lined pits, most of the underground utilities, and a network of underground pipelines still remain at the site. Since the end of World War II, the only activity that has been documented to have occurred at LHAAP-29 is the “soak out” or solvent bath of out-of-specification rocket motors. This took place from 1959 to the mid-1970s and involved the use of MC-based industrial solvent at tank 801-F. Waste from this operation was sent to LHAAP-18/24 (Jacobs, 2001).

2.2.2 History of Investigative Activities

As part of the Installation Restoration Program, the U.S. Army began an environmental investigation in 1976 at LHAAP followed by installation wide assessments/investigations that included the following:

- In 1980, U.S. Army Toxic and Hazardous Materials Agency (USATHAMA) conducted a record search to assess the impact of the LHAAP installation activities including usage, storage, treatment, and disposal of toxic and hazardous materials on the environment, and defined conditions that may have adversely affected human health and the environment (USATHAMA, 1980).
- Contamination Survey – In 1982, as part of the LHAAP contamination survey, Environmental Protection Systems collected six groundwater samples for laboratory analyses. Subsequently in 1987, as part of the Resource Conservation and Recovery Act (RCRA) permit application process, and as a continuation of the contamination survey, U.S. Army Environmental Hygiene Agency (USAEHA) identified, described, and evaluated all solid waste management units at LHAAP (USAEHA, 1987). Units requiring further sampling, investigation, and corrective action were delineated.
- RCRA Facility Assessment (RFA) – In 1988, a preliminary RFA was conducted by the U.S. Army (Maley, 1988). Waste at the various sites was characterized, but no samples were collected.
- Several investigations to determine the nature and extent of contamination in the soil, groundwater, surface water, and sediments at LHAAP-29 were conducted and are listed below. Samples were analyzed for (VOCs, semivolatile organic compounds (SVOCs),



metals, explosive compounds, perchlorate, pesticides, polychlorinated biphenyls (PCBs), and/or dioxins/furans, depending on the focus of the investigation.

Figure 2-3 and **Figure 2-4** show the sample locations for all investigations. **Figure 2-5** and **Figure 2-6** show well locations and groundwater elevations for the shallow and intermediate groundwater zones, respectively. For some of the earlier investigations, LHAAP sites were organized into groups, and LHAAP-29 was included in Group 2. The group designation was de-emphasized as the complexities of the individual sites increased.

The following summarizes the investigations at LHAAP-29:

- **Multi-phase investigation of Group 2 sites:** Between 1982 and 1998 numerous investigations were conducted in a phased approach by Jacobs, U.S. Army Corps of Engineers (USACE), and Environmental Protection System. Activities included installation of monitoring wells and analysis of groundwater, surface water, soil, and sediment samples. The results are documented in the RI for Group 2 sites (Jacobs, 2001).
- **Plant-wide perchlorate investigation:** The groundwater investigation was conducted from 2000 through 2002 (Solutions to Environmental Problems, Inc. (STEP), 2005) to delineate perchlorate contamination.
- **Baseline Human Health Risk Assessment:** The BHHRA (Jacobs, 2002) used data from the investigations conducted through 2001, including the plant-wide perchlorate investigation results up to that time. The report concluded that the soil at LHAAP-29 posed a non-carcinogenic hazard and the groundwater posed unacceptable carcinogenic risk and non-carcinogenic hazard to the hypothetical future maintenance worker.
- **Environmental Site Assessment:** Media investigated in 2003 included soil and groundwater (Plexus Scientific Corporation, 2005), although no sampling was conducted at LHAAP-29 for this assessment.
- **Baseline Ecological Risk Assessment:** The BERA (Shaw, 2007a) identified COPECs for the Industrial Sub-Area, which includes LHAAP-29. COPECs for the sub-area are addressed in the remedial actions for LHAAP-29. The evaluation was based on environmental investigations from 1993 to 2004.
- **Data Gaps:** Additional investigations were conducted in 2004 after the BHHRA was finalized to further delineate the extent of groundwater contamination identified during previous sampling events. A new monitoring well was installed and a total of 20 wells sampled in August 2004. The results of the 2004 investigation were presented in the *Data Gaps Investigation* (Shaw, 2007b).

Additional investigations were conducted after the 2004 investigation to further delineate contamination as follows:

- Additional investigations were conducted in December 2004 and February 2005 which included sampling soil near the TNT wastewater line, sediments from cooling water lines, the pump house pond, the outfall ditch, and manholes along the cooling water lines. Water was also sampled at the twelve manholes along the cooling water lines. Additional groundwater monitoring well sampling was conducted in May 2005 (Shaw, 2010).
- Sampling of soil at the foundation of former Wash House 806-D was conducted in February 2005 (USACE, 2005). Additional investigations were conducted between August 2006,



February 2007, and February 2008 to further delineate the extent of contamination, and included the activated persulfate oxidation study report (Shaw, 2010).

- Additional groundwater sampling events occurred in October 2008, January 2009, and June 2009 (Shaw, 2010).
- **Feasibility Study:** The FS (Shaw, 2010) was based on all available results from previous investigations through 2009, and the data collected since the risk assessment was evaluated in the FS. Data not formally submitted was incorporated into the FS. The potential soil-to-groundwater pathway was evaluated for the emerging contaminant perchlorate (found in groundwater) and the explosives posing risks or hazards in soil. The concentrations of these chemicals were compared to their TCEQ soil medium-specific concentrations (MSCs) for industrial use based on groundwater protection (GWP-Ind), which is more stringent than the soil MSC for industrial use based on inhalation, ingestion, and dermal contact (SAI-Ind) (TCEQ, 2006). The data collected from the shallow groundwater sampling indicated that MC concentrations were below Safe Drinking Water Act (SDWA) maximum contaminant level (MCLs), and 1,1-DCE concentrations are above the MCL. Additionally, all VOC groundwater data within the plumes were used to evaluate natural attenuation. The FS identified and evaluated 3 remedial alternatives to address the soil and groundwater contamination in the shallow and intermediate zones.
- **Supplemental Investigation:** A Final PP (U.S. Army, 2011) and Draft ROD were prepared in 2011 based on the RI and other investigation results, and the FS. During further evaluation of the RD requirements for the selected alternative, the U.S. Army determined that additional data were needed to refine the extent of the intermediate zone MC plume and also to collect data to evaluate additional treatment technologies, so a supplemental investigation was conducted in 2014. Additional wells were installed in the intermediate zone to define the extent of the MC plume inferred to be DNAPL, measure aquifer parameters, and evaluate thermal treatment of the MC plume, and in-situ bioremediation potential (AECOM, 2016).
- **Baseline Ecological Risk Assessment Addendum:** After the BERA was completed in 2007, a BERA Addendum was completed (AGEISS, 2014). The results of the re-evaluation indicated that the replacement data collected during the data gaps investigation confirmed the conclusions of the BERA that no explosives compounds in soil should be identified as COPECs in the industrial sub-area.
- **Supplemental FS:** A supplemental FS was prepared in 2017 using the data collected in during the supplemental investigation. A fourth alternative, ISTD, was identified and evaluated to address the MC DNAPL plume in the intermediate groundwater zone (AECOM, 2017). The shallow zone groundwater and soil remediation components of the alternatives did not change from the initial FS.



2.2.3 Site History of CERCLA Enforcement Activities

LHAAP-29 was included in the National Priorities Listing for Longhorn in 1990 and identified as Unit No. 29 in the 1991 FFA. The FS for LHAAP-29 (Shaw, 2010) was issued in April 2010, and the PP (U.S. Army, 2011) was issued in March 2011. A Revised PP was completed in 2018 (U.S. Army, 2018). This ROD follows the Revised PP and precedes the more detailed RD.

2.3 Community Participation

The U.S. Army, USEPA, TCEQ and the LHAAP Restoration Advisory Board (RAB) have provided public outreach to the surrounding community concerning LHAAP-29 and other environmental sites at LHAAP. The outreach program has included fact sheets, media interviews, site visits, invitations to attend quarterly RAB meetings, and public meetings consistent with its public participation responsibilities under Sections 113(k)(2)(B), 117(a), and 121(f)(1)(G) of CERCLA.

The Final Revised PP (U.S. Army, 2018) for the selection of the remedy for LHAAP-29 was released to the Administrative Record and made available to the public for review and comment beginning November 21, 2018. The notice of availability of the Final Revised PP and other related documents in the Administrative Record file was published in *The Shreveport Times* and the *Marshall News Messenger* on November 7, 2018. The newspaper and media notices for the meeting are provided in **Appendix A**. The public comment period for the Revised PP began on November 21, 2018 and ended December 21, 2018. A public meeting was held on December 6, 2018 in a formal format and the meeting recorded by a court reporter. The transcript for the meeting is part of the Administrative Record. The significant comments (oral or written) are addressed in the Responsiveness Summary, which is included in this ROD as **Section 3**.

The previously completed PP (U.S. Army, 2011) was also released to the Administrative Record, and similar public notices and a public meeting were held. Comments received for the 2011 Proposed Plan are also presented in **Section 3**.

The Administrative Record may be found at <http://www.longhornaap.com/> and locally at the information repository maintained at the following location:

Location: Marshall Public Library
 300 S. Alamo
 Marshall, Texas 75670

Business Hours: Monday, Tuesday, Thursday (9:30 AM – 7:30 PM)
 Wednesday and Friday (9:00 AM – 5:30 PM)
 Saturday (9:30 AM – 3:30 PM)

2.4 Scope and Role of Response Action

The response action will prevent potential unacceptable risks associated with exposure to contaminated soil and groundwater in both the shallow and intermediate zones. The removal of source soils will positively impact groundwater by eliminating the potential for the leaching of contaminants from the soil into the groundwater and will remove the contamination that poses a risk to ecological receptors.



Plugging the inlets and outlets of the underground lines with a bentonite slurry mix, including the manholes of the process cooling water lines, will minimize hypothetical future maintenance workers contact with contaminants and prevent water from infiltrating the lines.

The selected action at LHAAP-29 will prevent potential risks associated with exposure to contaminated groundwater. The groundwater at LHAAP is not currently being used as a drinking water source, nor may it be used in the future based on its reasonably anticipated use as a national wildlife refuge. However, when establishing the remedial action objectives (RAOs) for this response action, the U.S. Army has considered the NCP's expectation to return usable groundwater to its potential beneficial uses wherever practicable and has also considered the State of Texas designation of all groundwater as potential drinking water, unless otherwise classified, and consistent with 30 TAC 335.563(h)(1) [background total dissolved solids (TDS) content less than or equal to 10,000 milligrams per liter (mg/L)] and that occurs within a geologic zone that is sufficiently permeable to transmit water to a pumping well in usable quantities.

The U.S. Army intends to return the contaminated shallow and intermediate groundwater zones at LHAAP-29 to their potential beneficial uses, which for the purposes of this ROD is considered to be attainment of the SDWA MCLs to the extent practicable, and consistent with 40 C.F.R. § 300.430(e)(2)(i)(B & C). In the absence of federal drinking water standards, clean-up levels will be based on the Texas Risk Reduction Program (TRRP) Tier 1 Residential Groundwater Protective Concentration Level (PCL) (TRRP Residential Groundwater PCL). The TCEQ soil medium specific concentration (MSC) for industrial use based on groundwater protection (GWP-Ind) is used in accordance with 30 TAC 335.559(g)(2). If a return to potential beneficial uses is not practicable, the NCP expectation is to prevent further migration of the plume, prevent exposure to the contaminated groundwater, and evaluate further risk reduction (40 C.F.R. § 300.430(a)(1)(iii)(F)).

The selected remedial action will also ensure containment of the plume to prevent potential impact to surface water. The selected action will also include groundwater monitoring to demonstrate that the plume is not migrating at levels that present a potential impact to surface water bodies and to verify that contaminant levels are being reduced to cleanup levels when the LUC for groundwater use prohibition may be terminated.

In addition, the selected action includes an active remedial component that will mitigate the principal threat waste. By instituting an ISTD treatment of the intermediate groundwater, this active treatment will be applied to the highest concentration area in the MC groundwater plume and will comply with NCP expectations regarding treatment of affected media where principal threat waste may be considered.

2.5 Site Characteristics

This section of the ROD presents a brief comprehensive overview of the LHAAP-29 site characteristics with respect to the conceptual site model (CSM), physical site features, known or suspected sources of contamination, types of contamination, and affected media. Known or potential routes of contaminant migration are also discussed. Detailed information about the site characteristics can be found in the RI (Jacobs, 2001).

2.5.1 Conceptual Site Model

Figure 2-7 illustrates the human health conceptual site model for LHAAP-29. The model presents the human health pathways that are complete and being considered for remediation. Those



pathways that are likely to be incomplete or have negligible impact are not being considered for remediation. The ecological conceptual model for LHAAP-29 (**Figure 2-8**) is similar to the one presented for human health in terms of the origin and fate and transport mechanisms of the contaminants present at the site. However, only exposure pathways and routes associated with soil are relevant for ecological risk assessment.

Explosive compound releases resulting from the manufacturing process of TNT as well as releases from process tanks and process waste pipelines are the suspected contamination sources at LHAAP-29. Releases from the rocket motor soak-out process that used methylene chloride-based solvents are also contamination sources. The pipelines include the TNT process wastewater gravity line (“red and/or yellow liquor”) that transported the TNT process wastewater to LHAAP-32 for treatment and two cooling water lines (blue water) that transported the cooling water to an outfall ditch. The remaining potential sources of contamination at the site are explosives compounds in stained soils around the foundation of former Buildings 806A and 806-D, isolated perchlorate-containing soils in the northeastern portion of LHAAP-29 at a depth of 8 feet below ground surface (bgs), and TNT-contaminated sediment in the cooling water outfall ditch at a depth of 7 feet bgs.

The red liquor TNT wastewater line was originally installed as a wooden pipeline at a depth of three to five feet bgs and was taken out of service, clear-flushed and abandoned in 1946 (Bate Stamp 001446, RFA, April, 1988). Several trenches were excavated across the wooden line in 1993, and the wood was found to be soft and severely degraded at most locations (Sverdrup Environmental, Inc. [Sverdrup], 1993). Samples collected at that time resulted in high concentrations of TNT in the liquid and sludge (3,500 µg/L and 3,700 mg/kg, respectively), but that data was later deemed unusable for environmental decision making (Jacobs, 2001). The transite TNT wastewater pipeline was added parallel to and north of the wooden line to carry the TNT wastewater. The transite TNT wastewater line has solid residues contaminated with explosives at concentrations above the GWP-Ind.

Two blue cooling water lines, called cooling water line north and cooling water line south, exist at LHAAP-29 and range from 8 to 18 inches in diameter. These are gravity fed lines and are constructed of vitrified clay pipe with manholes. These lines have solid residues contaminated with explosives at concentrations that are above the GWP-Ind MSC (solid residue).

Contamination in the form of explosive compounds, VOCs, perchlorate, and metals from rocket motor washout and TNT production is present in the shallow groundwater zone at LHAAP-29 and poses potential risk to the hypothetical future maintenance worker. In the intermediate groundwater zone, concentrations of VOCs and arsenic are present, with MC posing the highest risk. At monitoring well 29WW16 the MC concentrations in the intermediate zone are approximately half the solubility limit, which indicates a potential for the presence of DNAPL. The horizontal extent of contamination in the shallow and intermediate groundwater zones is presented in **Figures Figure 2-9**, and **Figure 2-11**.

The soil and groundwater at LHAAP-29 may pose an unacceptable risk for the hypothetical future maintenance workers. Even though no impact to surface water from groundwater has been established (Shaw, 2007c); the migration pathway of groundwater contaminants to surface water is being considered for remediation along with soil, soil to groundwater, and future industrial groundwater use.



2.5.2 Overview of the Site

The site boundary of LHAAP-29 comprises approximately 85 acres in the western-central portion of LHAAP. The surface features include the foundations for the former production facilities and the underground pipelines that were originally built for cooling water drainage and TNT wastewater conveyance. The site is currently heavily wooded. Surface runoff is collected by ditches constructed in 1942 when the production facility was built. Surface runoff from the northern part of the site (about 40 percent of the site) enters Goose Prairie Creek located approximately 1,500 feet to the north and east of the site. Surface water runoff in the southern portion of the site (about 60 percent of the site) flows into a tributary of Central Creek located near the southeast portion of the site. Eventually, runoff from the two creeks enters Caddo Lake.

2.5.3 Geology and Hydrogeology

The local geology at LHAAP-29 consists of a silty to clayey sand at the surface that extends from 3 to 10 feet bgs. A clayey silt grading to a silty clay was encountered extending to a depth of 40 feet bgs with an underlying silty to clayey sand. With the exception of boreholes 29WW02, 29WW10, and 29WW11, a sandy silt to silty clay layer was encountered below the sand deposit at depths ranging from 15 to 26 feet bgs. Additional silt, clay and sand layers were encountered with depth in boreholes 29SB52 and 29SB53 that terminated at the top of the Midway formation at depths of 157 and 140 feet bgs, respectively.

There are three groundwater zones at LHAAP-29: shallow, intermediate, and deep. Semi-confining clay or silty clay layers separate the three groundwater zones. The shallow groundwater zone has wells that are screened at two depths (shallow and lower shallow); however, the wells have similar water level elevations and are all considered to be shallow zone wells. The bottom of each of the zones is defined by a continuous or semi-continuous clay layer of varying thickness. The depth of the shallow groundwater zone generally ranges from 17 to 45 feet bgs because of variable ground surface elevations across the site. The intermediate zone is less defined, but its depth has been measured to approximately 88 feet bgs. The deep groundwater zone extends to a depth of approximately 155 feet bgs. Based on the 2007 water levels and historic potentiometric maps, the predominant groundwater flow in the shallow zone is east/southeast and is east/northeast in the intermediate zone. **Figures 2-5** and **2-6** illustrate the groundwater elevations in the shallow zone and intermediate zone, respectively.

2.5.4 Sampling Strategy

Nineteen investigations/sampling events were conducted at LHAAP-29 from 1982 to 2014, as described in **Section 2.2.2**. In the early investigations, soil samples were collected from throughout the site to determine the areas of contamination. Subsequent investigations focused on the areas where contamination was found, performing additional soil, groundwater, and sediment sampling, and installing monitoring wells to delineate the groundwater contamination. Samples from all media were analyzed for various analytes including VOCs, SVOCs, metals, explosives, perchlorate, pesticides, and dioxins/furans. In the intermediate groundwater contaminant plume, groundwater samples were also analyzed for indicators of conditions that promote natural attenuation (biodegradation), such as dissolved oxygen, conductance, pH, oxidation-reduction potential, sulfide, methane, and chloride. During the 2014 investigation, soil resistivity and total organic carbon data were measured to provide additional data to evaluate ISTD technologies, and microcosm testing was performed to evaluate the potential for enhanced in-situ bioremediation (EISB). Aquifer



pumping tests were also conducted to gather site-specific hydraulic conductivity data to support the evaluation of additional treatment technologies.

2.5.5 Nature and Extent of Contamination

Contamination was found in the soil, groundwater (shallow and intermediate zones), and liquid and solid residue remaining in the cooling water and TNT process wastewater underground lines. The COCs are toxic and carcinogenic. Since there is a high cancer risk associated with exposure to groundwater from this region of the intermediate zone, such residual source material may be considered a principal threat waste at LHAAP-29.

Groundwater

Shallow zone groundwater COCs are VOCs (TCE, 1,2-DCA, and daughter products), perchlorate, explosives (2,4-DNT, 2,6-DNT, 2-NT, 3-NT, 4-NT), and metals (arsenic, mercury, nickel, selenium).

The shallow zone plumes for VOCs and perchlorate are shown on **Figure 2-9**. The most recent maximum concentrations of COCs in shallow groundwater were all detected in samples collected from monitoring well 29WW15, and included TCE at 344 µg/L, 1,2-DCA at 5,520 µg/L and perchlorate at 16,800 µg/L. The calculated volume of the perchlorate plume is 4 million gallons. There are 3 shallow zone plumes for explosives as shown on



Figure 2-10. The highest concentration of 2,4-DNT detected was 50.9 µg/L at monitoring well 29WW05. The highest concentration of 2,6-DNT was 239 µg/L at monitoring well 116. The highest concentrations of 2-NT, 3-NT, and 4-NT are 8,140 µg/L, 451 µg/L, and 1,400 µg/L, respectively, also reported at monitoring well 116. The volume of the VOC, perchlorate, and NT plumes are estimated to be approximately 9 million gallons, assuming a total porosity of 0.25 (or 25 percent) with a thickness ranging from 5 to 10 feet.

Intermediate zone groundwater COCs are MC, TCE and daughter products, 1,2-DCA, and arsenic. The intermediate zone plume for VOCs is shown on **Figure 2-11**. The highest concentration of MC detected was 10,300,000 µg/L at monitoring well 29WW16. The most recently reported MC concentration in a 2014 sample collected from this well was 8,260,000 µg/L. Other COCs identified for the intermediate groundwater zone are degradation (daughter) products of TCE that have been non-detect or have not been detected above their SDWA MCLs; however, due to historic results and the high detection limits at 29WW16, it has been assumed that concentrations of 1,2-DCA and TCE still exceed their SDWA MCLs. The calculated volume of contaminated groundwater that exceeds MCLs is 650,000 gallons, assuming a total porosity of 0.40 (or 40 percent) with an average thickness of 15 feet based on drilling observations.

As demonstrated by previous sampling results, the deep groundwater zone at LHAAP-29 is not affected.

Soil

Soil COCs and COPECs are explosives (2,4,6-TNT, 2,4-DNT, 2,6-DNT) and perchlorate. **Figure 2-12** shows the approximate areas of contaminated soil. The maximum 2,4,6-TNT concentration in the soil is 26,000 milligrams per kilogram (mg/kg). Other explosives, 2,4-DNT and 2,6-DNT, have maximum concentrations of 8,000 mg/kg and an estimated concentration of 15 mg/kg, respectively. Additionally, perchlorate has been detected in the soil at a maximum concentration of 8.6 mg/kg. The estimated volume of impacted soil is 3,900 cubic yards in place.

TNT Wastewater Lines

Line solid residue COCs from the transite TNT wastewater line are explosives (2,4,6-TNT, 2,4-DNT, 1,3 dinitrobenzene, 2-amino-4,6-DNT, 4-amino-2,6-DNT). Contaminated explosive solid residue remains within the transite TNT wastewater line at concentrations above the SAI-Ind and GWP-Ind MSCs, but access to the pipe is limited to the inlets and outlets unless the pipe is penetrated. Additionally, the line is buried 3 feet bgs or deeper. The gravity flow portion of the line is approximately 3,000 linear feet. The pressurized portion of the line is approximately 1,000 linear feet. The line is in good condition. **Figure 2-13** shows the TNT wastewater lines at LHAAP-29, soil, sediment, and solid residue sampling results and MSC values.

The wooden TNT wastewater line was previously flushed and abandoned. The results from limited soil samples collected near the line indicate there has not been a release to the surrounding soil above the cleanup levels. The line is buried 3 feet bgs or deeper, but the deteriorated condition of the line and the potential for residual explosives may present a continuing source of groundwater contamination.

Cooling Water Lines

Line solid residue COCs from the vitrified clay cooling water lines are explosives (2,4,6-TNT, 2,4-DNT, 2,6-DNT, 2-amino-4,6-DNT, 4-amino-2,6-DNT). The solid residues from the manholes are



contaminated with explosives at concentrations that are above the GWP-Ind (solid residue). The north and south vitrified clay cooling water lines are accessible through manholes, and the liquid and solid residue contents from the manholes were sampled. There are approximately 5,000 feet of pipe in the main lines, approximately 1,680 linear feet of pipe from each production area to the main line, and 12 manholes. **Figure 2-13** shows the cooling water lines and sampling results of the line contents at LHAAP-29.

2.6 Current and Future Land and Resource Uses

2.6.1 Current and Future Land Uses

LHAAP is located near the unincorporated community of Karnack, Texas. Karnack is a rural community with a population of 775 people. The incorporated community of Uncertain, Texas, population 205, is located to the northeast of LHAAP on the edge of Caddo Lake and is a resort area and an access point to Caddo Lake. The industries in the surrounding area consist of agriculture, timber, oil and natural gas production, and recreation.

LHAAP has been an industrial facility since 1942. Production activities and associated waste management activities continued until the facility was determined to be in excess of the U.S. Army's needs in 1997. The plant area has been relatively dormant since that time. LHAAP is surrounded by a fence (except on the border with Caddo Lake), and current security measures at the LHAAP preclude unlimited public access to areas within the fence. The fence now represents the National Wildlife Refuge boundary. Approved access for hunters is limited.

The reasonably anticipated future use of LHAAP-29 is as part of a national wildlife refuge. This anticipated future use is based on a Memorandum of Agreement (MOA) (U.S. Army, 2004) between the USFWS and the U.S. Army. That MOA documents the transfer process of the LHAAP acreage to USFWS to become the Caddo Lake National Wildlife Refuge and will be used to facilitate a future transfer of LHAAP-29. Presently the Caddo Lake National Wildlife Refuge occupies a little more than 7,100 acres of the 8,416-acre former installation. In accordance with the National Wildlife Refuge System Administration Act of 1966 and its amendments (16 USC 668dd), the land will remain as a national wildlife refuge unless there is a change brought about by an act of Congress, or the land is part of an exchange authorized by the Secretary of the Interior.

2.6.2 Current and Future Surface Water Uses

Streams on LHAAP currently support wildlife and aquatic life. While humans may have limited access to some streams during annual hunts, there is no routine human use of streams on LHAAP. The streams do not carry adequate numbers and size of fish to support either sport or subsistence fishing. During the summer months, the streams cease flowing and/or dry up. The streams flow into Caddo Lake. Caddo Lake is a large recreational area that covers 51 square miles and has a mean depth of 6 feet. The watershed of the lake encompasses approximately 2,700 square miles. It is used extensively for fishing and boating. Caddo Lake is a drinking water supply for multiple cities in Louisiana including Vivian, Oil City, Mooringsport, South Shore, Blanchard, Shreveport, and Bossier City.

The anticipated future uses of the streams and lake are the same as the current uses.



2.6.3 Current and Future Groundwater Uses

Groundwater in the aquifer (250 to 430 feet bgs) near LHAAP is currently used as a drinking water source. The drinking water aquifer should not be confused with the deep zone groundwater, which extends only to a depth of approximately 151 feet bgs. The deep zone groundwater and the drinking water aquifer are distinct from each other and there is no connectivity between the contaminated zone and the drinking water aquifer. There are five active water supply wells near LHAAP that are completed in the drinking water aquifer (**Figure 2-2**). One well is located in and owned by Caddo Lake State Park. The well is completed to a depth of 315 feet bgs and has been in use since 1935. A second well owned by the Karnack Water Supply Corporation services the town of Karnack and is located approximately 0.3 miles northwest of town. This well is completed to approximately 430 feet bgs and has been in use since 1942. The Caddo Lake Water Supply Corporation has three wells located both north and northwest of LHAAP. These wells are identified as Caddo Lake Water Supply Corporation Wells 1, 2, and 3, and all are hydraulically upgradient of LHAAP (Jacobs, 2002). These wells are completed deeper than the deepest zone of contamination at LHAAP. Because of this and the large distance between these wells and LHAAP, water removal from these wells is not expected to affect groundwater flow at the site. In addition, there are several livestock and domestic wells located in the vicinity of LHAAP with depths averaging approximately 250 feet bgs.

Three water supply wells are located within the boundary of LHAAP itself. One well is located at the Fire Station; the second well is located approximately 0.35 miles southwest of the Fire Station. The third well is located north of the USFWS administration building for the Caddo Lake National Wildlife Refuge, near the main entrance to LHAAP. The distances from these water supply wells to LHAAP-29 are approximately 0.4 miles, 0.2 miles, and 0.5 miles, respectively (**Figure 2-2**). The three water supply wells were completed at a depth much greater than the zone of contamination described at LHAAP-29. None of these three wells are currently used for drinking water at LHAAP, although they may supply water for non-potable uses. Two additional wells previously supplied water to the installation, but these have been plugged and abandoned.

Although the anticipated future use of the facility as a wildlife refuge does not include the use of the groundwater at LHAAP-29 as a drinking water source, the State of Texas designates all groundwater as potential drinking water, unless otherwise classified, and consistent with 30 TAC 335.563(h)(1). To be conservative, a hypothetical industrial use scenario was evaluated for risk. The future industrial scenario for LHAAP assumes limited use of groundwater as a drinking water source.

2.7 Summary of Site Risks

The BHHRA and BERA estimate the risks posed by contaminants at the site if no action were taken. These assessments provide the basis for taking action and identify the contaminants and exposure pathways that need to be addressed by the remedial action.

2.7.1 Summary of Human Health Risk Assessment

This section is based on the conclusions presented in the *Final Baseline Human Health and Screening Ecological Risk Assessment for the Group 2 Sites* (Jacobs, 2002), in the *Data Gaps Investigations* (Shaw, 2007b), and in additional data collected in preparation of the *Final Feasibility Study, LHAAP-29* (Shaw, 2010). The risk assessment used data from the investigations conducted through 1998 and the plant-wide perchlorate investigation conducted in 2000. Results from the later investigations conducted through 2014 did not change the overall outcome of the risk assessment



and the discussion of results and risks presented here are as presented in the Baseline HHRA and FS. During the risk assessment, soil and groundwater data were used to calculate the aggregate risk, which was then compared to the USEPA target risk range of 1×10^{-4} to 1×10^{-6} for the excess lifetime carcinogenic risk and to a hazard index (HI) of 1 for non-carcinogenic hazards. If there is no unacceptable risk associated with a medium, and a cleanup level is not exceeded, then the medium is not identified in this ROD for remediation. The CSM that is associated with the risk assessment was introduced in **Section 2.5.1**, and is presented as **Figure 2-7**.

2.7.1.1 Identification of Chemicals of Potential Concern

The BHHRA identified chemicals of potential concern (COPCs) for LHAAP-29 and evaluated the carcinogenic risk and non-carcinogenic hazard for each. **Table 2-1** summarizes the risk assessment data for the COPCs, including minimum and maximum detected concentrations, frequency of detection, and exposure point concentrations (EPCs). Analytical results for various congeners of dioxins and furans are expressed as 2,3,7,8-tetrachlorodibenzo-p-dioxin (TCDD) toxicity equivalence concentration (TEC).

2.7.1.2 Exposure Assessment

The Jacobs risk assessment (Jacobs, 2002) presented the human health risks and hazards to a hypothetical future maintenance worker under an industrial scenario for soil and groundwater.

For soil, reasonable exposure pathways according to the CSM are: incidental ingestion of the surface soil (0 to 2 feet bgs), dermal contact with the surface soil, inhalation of particulates, and inhalation of VOCs from the soil (0 to 7 feet bgs).

For groundwater, reasonable exposure pathways are ingestion of groundwater, dermal contact while showering with contaminated groundwater, and inhalation of VOCs while showering with contaminated groundwater.

2.7.1.3 Toxicity Assessment

The carcinogenic and non-carcinogenic toxicity assessments from the BHHRA are summarized in **Table 2-2** and **Table 2-3**, respectively. The toxicity data assumes that exposure would be chronic to be conservative. Sources for the data include the Integrated Risk Information System (IRIS) and Health Effects Assessment Summary Tables (HEAST).

2.7.1.4 Risk Characterization

Characterization of the carcinogenic risk and non-carcinogenic hazard are summarized in **Table 2-4** and **Table 2-5**, respectively. For carcinogens, risks are generally expressed as the incremental probability of an individual's developing cancer over a lifetime as a result of exposure to the carcinogen. Excess lifetime carcinogenic risk is calculated from the following equation:

$$\text{Risk} = \text{CDI} \times \text{SF}$$

where: risk = unitless probability of an individual developing cancer

CDI = chronic daily intake averaged over 70 years, expressed as milligrams per kilogram per day (mg/kg-day)

SF = slope factor, expressed as (mg/kg-day)⁻¹



These risks are probabilities that usually are expressed in scientific notation. An excess lifetime carcinogenic risk of 1×10^{-6} indicates that an individual experiencing the reasonable maximum exposure estimate has a 1 in 1,000,000 chance of developing cancer as a result of site-related exposure. This is referred to as an “excess lifetime carcinogenic risk” because it would be in addition to the risks of cancer that individuals face from other causes such as smoking or exposure to too much sun. The chance of an individual developing cancer from all other causes has been estimated to be as high as one in three. USEPA’s generally acceptable risk range for site-related exposures is 1×10^{-4} to 1×10^{-6} .

The potential for non-carcinogenic effects is evaluated by comparing an exposure level over a specified time period (e.g., lifetime) with a reference dose (RfD) derived for a similar exposure period. An RfD represents a level that an individual may be exposed to that is not expected to cause any deleterious effect. The ratio of exposure to toxicity is called a hazard quotient (HQ). An $HQ < 1$ indicates that a receptor’s dose of a single contaminant is less than the RfD, and that toxic non-carcinogenic effects from that chemical are unlikely. The HI is generated by adding the HQs for all COCs that affect the same target organ (e.g., liver) or that act through the same mechanism of action within a medium or across all media to which a given individual may reasonably be exposed. An $HI < 1$ indicates that, based on the sum of all HQ’s from different contaminants and exposure routes, toxic non-carcinogenic effects from all contaminants are unlikely. An $HI > 1$ indicates that site-related exposures may present a risk to human health.

The HQ is calculated as follows:

$$\text{Non-carcinogenic HQ} = \text{CDI/RfD}$$

Where: CDI = chronic daily intake
 RfD = reference dose

CDI and RfD are expressed in the same units and represent the same exposure period (e.g., chronic, subchronic, or short-term).

The carcinogenic risks for soil and groundwater are 7.3×10^{-6} and 3.9×10^{-1} , respectively (Jacobs, 2002). The carcinogenic risk for soil is within the USEPA target risk range of 1×10^{-4} to 1×10^{-6} . The HIs for soil and groundwater are 1 and 3,000, respectively, and are above the acceptable HI of < 1 . The carcinogenic risks for groundwater are unacceptable; the non-carcinogenic risk for both soil and groundwater are unacceptable; therefore, the remedial action acts on both the soil and groundwater. Chemicals with a risk greater than 1×10^{-4} in groundwater include TCE, 1,2-DCA, arsenic, 2,4-DNT, 2,6-DNT, MC, chloroform, and 2,3,7,8-TCDD. Chemicals with a HQ greater than one in groundwater include 2-NT, 3-NT, arsenic, nickel, 2,4-DNT, 2,6-DNT, chloroform, perchlorate, TCE, and 1,2-DCA.

The BHHRA included an uncertainty analysis which identified factors that would cause values used in the risk assessment to be over or underestimated. The analysis concluded that the risks and HIs are overestimated, making the BHHRA a conservative evaluation. Additionally, the uncertainty analysis indicated a portion of the noncarcinogenic effects associated with antimony in groundwater at LHAAP-29 may be due to background.

2.7.1.5 Evaluation of COPCs

To further evaluate the occurrence of COPCs, a data gap investigation was conducted (Shaw, 2007b) and additional investigations were conducted when preparing the FS (Shaw, 2010). While



these investigations did not change the overall outcome of the earlier BHHRA, they determined the COPCs to be targeted by the remedial action.

Table 2-6 and **Table 2-7** list the chemicals in groundwater that exceed those values for the carcinogenic risk and HQ greater than 0.1, respectively. There is no carcinogenic risk in soil to the hypothetical maintenance worker. **Table 2-8** lists chemicals in the soil that have an HQ greater than 0.1 for the hypothetical maintenance worker. These tables also summarize the justifications for which of the COPCs should be classified as COCs. COPCs in soil were identified as COCs when they posed a carcinogenic risk above the acceptable range (risk greater than 1×10^{-4}) or when their HQ was greater than 1. COPCs in groundwater were identified as COCs when they posed a carcinogenic risk above the acceptable range (risk greater than 1×10^{-4}), when their HQ was greater than 1, or when the EPC was above the MCL or in the absence of federal drinking water standards, the TRRP Residential Groundwater PCL. Recent data obtained after the BHHRA investigation was used when possible. Based on the comparison of the maximum groundwater concentration since the BHHRA to their associated SDWA MCL or PCL, these COCs have been identified on **Table 2-10** to be of concern in the shallow and intermediate groundwater. **Table 2-10** presents the final list of COCs and all media, along with cleanup levels.

The human health risk assessment, which was based on the reasonably anticipated future use as a national wildlife refuge, does not address unrestricted use. In accordance with 30 TAC 335.566, a notification will be recorded in the Harrison County records stating that the site is suitable for nonresidential use.

2.7.2 Summary of Ecological Risk Assessment

The *Final Installation-Wide Baseline Ecological Risk Assessment* (Shaw, 2007a) and Baseline Ecological Risk Assessment Addendum (AGEISS, 2014) evaluated potential hazards to ecological resources at LHAAP by conducting a screening evaluation to identify initial COPECs in the individual sub-areas and watersheds. The potential of these COPECs to adversely affect communities was evaluated for: (1) organisms that have direct contact with the COPECs (e.g., plants and earthworms growing and living in contaminated soil); and (2) organisms that may be exposed to the chemicals via food chain pathways (e.g., ingestion of an earthworm living in the contaminated soil by a shrew). Potential impacts to invertebrate and plant communities were evaluated by comparing COPEC concentrations to benchmark values available from multiple literature sources. For the food chain exposure assessment, a number of measurement receptors were selected as representative species for the various trophic levels in the food web that could be at risk from contaminants in site media. The measurement receptors that were selected and used in the food chain evaluation included the following:

- Deer Mouse
- Raccoon
- Modified Raccoon (as a surrogate for the Louisiana Black Bear)
- Short-Tailed Shrew
- Red Fox
- Muskrat
- River Otter



- Townsend's Big-Eared Bat
- Common Snapping Turtle
- Bank Swallow
- American Woodcock
- Belted Kingfisher
- Red-Tailed Hawk

A food chain model was developed and used to estimate the total dose for each measurement receptor based on species-specific considerations such as diet, body weight, ingestion rates, etc., using conservative exposure estimates. Ecological hazard estimates were developed based on exposure to all media including soil in a particular sub-area and surface water and sediment from any watersheds present in the sub-areas. Two different soil depths were used for modeling exposure to ecological receptors: surface soil (0 to 0.5 foot) and total soil (0 to 3 feet). Each receptor was assumed to be exposed to one of the two depths based on its life history characteristics (e.g., burrowing animals were assumed to be exposed to total soil). Bioaccumulation of chemicals up the food chain was initially estimated using uptake factors obtained from available literature, and then refined using site-specific data obtained during the BERA. **Figure 2-8** presents the ecological conceptual model, which lays out the exposure pathways for selected species.

Ecological effects quotients (EEQ) were developed for each of the measurement receptors. EEQs are similar to HQs for human health, and are calculated by dividing the total dose that the receptor is exposed to by the toxicity reference value (TRV), which is based on a no-observed adverse effect level (NOAEL) or the lowest-observed adverse effect level concentration. If the EEQ exceeds 1 for a receptor (based on the NOAEL TRV), then that chemical is considered to have a realistic potential to cause adverse ecological impacts, and is identified as a final COPEC that should be addressed either through remediation or further investigation. As discussed in the BERA, there are several important uncertainties associated with the assumptions used in the EEQ process, and it should be noted that EEQs greater than 1 do not necessarily mean that ecological impacts have occurred, or are occurring.

Several sub-areas were established within LHAAP for the BERA. LHAAP-29 falls within the Industrial Sub-Area. For the industrial Sub-Area, four chemicals were selected as final COPECs: cadmium, chromium, zinc, and perchlorate. After that selection, additional sampling data became available and further analysis was performed, leading to the calculation of ecological preliminary cleanup levels (EcoPRGs) for several chemicals in soil. The final COPECs that were initially selected were found to not be of concern and EcoPRGs were calculated for six other chemicals: barium, lead, 2,4-DNT, 2,6-DNT, TNT, and 2,3,7,8-TCDD. Ecological hazards were found to be acceptable for the Industrial Sub-Area that includes LHAAP-29; however, elevated concentrations of NTs (TNT, 2,4-DNT and 2,6-DNT) were identified at one location at the site (Shaw, 2007a). Although NTs were not selected in the Industrial Sub-Area as final COPECs due to low frequency of detection and other considerations, the BERA evaluated measurement receptors and included a spatial analysis at this sample location at LHAAP-29. The results of this analysis identified that the NTs at this location and the adjacent areas may represent a small area of highly elevated concentrations (i.e., a hot spot) that could pose a threat to small-range ecological receptors either through acute toxicity, or as a source area for downgradient surface water transport of contamination (Shaw, 2007b). The EcoPRGs are shown on **Table 2-9**. An Excel spreadsheet analysis was performed by



ranking the detected concentrations of each final COPEC in the Industrial Sub-Area and iteratively re-calculating the 95% upper confidence limit (UCL) on the mean after removing concentrations until the 95% UCL for the Industrial Sub-Area was lower than the EcoPRG. (Note: as discussed in the BERA, the EcoPRG is not a “not to exceed” value for all concentrations; rather, it is a conservative estimate of the average concentration that results in no adverse effects, and as such is equivalent to the 95% UCL of chemical concentrations, rather than to individual sample concentrations.) The order of chemical concentrations was altered to preferentially remove LHAAP-29 samples in order to reduce the ecological risk in the Industrial-Sub Area. It is assumed that the locations associated with these concentrations will be remediated. The outcome of the analysis is included in **Table 2-10** and indicated on **Figure 2-12**.

After the BERA was completed in 2007, additional data review determined that some explosives results used in the BERA were invalid. Additional samples were collected during a data gaps investigation to replace the invalid results and the results were combined with the previously reported useable data and data from samples collected following completion of the BERA to re-evaluate the ecological risks. The results were reported in the BERA Addendum (AGEISS, 2014). The results of the re-evaluation indicated that the replacement data collected during the data gaps investigation confirmed the conclusions of the BERA that no explosives compounds in soil should be identified as COPECs in the industrial sub-area. These results do not change the determination that the areas of elevated NTs at LHAAP-29 might pose a risk to small-range ecological receptors and should be addressed as part of the remedial action.

2.7.3 Basis of Action

The remedial action selected in this ROD is necessary to protect the public health or welfare or the environment from actual or threatened releases of hazardous substances, pollutants, or contaminants into the environment. Actions for the groundwater are necessary to address the potential for human health risks in the unlikely event there is an attempt to use groundwater as a potable water source.

Actions for soil are necessary to address human health risk including the pathway from soil to groundwater and ecological risks. **Table 2-10** presents the COCs and the final cleanup levels for both soil and groundwater with groundwater COCs for the shallow zone and the intermediate zone listed separately, which takes into account both ecological and human receptors.

The SDWA MCL is the cleanup level for the groundwater COCs. In the absence of federal drinking water standards, clean-up levels will be based on TRRP Residential Groundwater PCLs.

2.8 Remedial Action Objectives

The RAOs for LHAAP-29 presented in this ROD for the selected remedy and contingency remedy address contamination associated with the media at the site and take into account the future uses of LHAAP surface waters, land, and groundwater, are:

- Protection of human health by preventing human exposure to the contaminants in the soil, sediment, transite TNT wastewater line, vitrified clay cooling water lines, and groundwater
- Protection of human health and the environment by preventing the migration of contaminants to groundwater and surface water from potential sources in the soil, sediment, and process lines (TNT wastewater and cooling water)



- Protection of human health and the environment by preventing contaminated groundwater from migrating into nearby surface water
- Protection of ecological receptors by preventing exposure to the contaminated soil and sediment
- Return of groundwater to its potential beneficial uses, wherever practicable within a timeframe that is reasonable given the particular circumstances of the site (40 C. F. R. § 300.430(a)(1)(iii)(F)).

The above RAO recognizes USEPA's policy to return all groundwater to beneficial uses, based on non-binding programmatic expectation in the NCP, and is consistent with the NCP regulations requiring the lead agency to establish RAOs specifying contaminants and media of concern, potential exposure pathways, and remediation goals.

Per these RAOs, and consistent with the NCP, groundwater will be returned to its beneficial use. In the absence of federal drinking water standards, the groundwater clean-up level at the Site is the TRRP Residential Groundwater PCL and is protective of human health and the environment.

2.9 Description of Alternatives

Four alternatives (including No Action) were evaluated. This section introduces the remedy components, identifies the common elements and distinguishing features of each alternative, and describes the expected outcomes of each.

2.9.1 Description of Remedy Components

Alternative 1 – No Action

As required by the NCP, the no action alternative provides a comparative baseline against which the action alternatives can be evaluated. Under this alternative, groundwater would be left “as is” without implementing any additional monitoring, containment, removal, treatment, or other mitigating actions. No actions would be implemented to reduce existing or potential future exposure to human and ecological receptors, although natural attenuation would be ongoing.

Estimated Capital Present Worth Cost: \$0

Estimated O&M Present Worth Cost: \$0

Cost Estimate Duration: NA

Estimated Present Worth Cost: \$0

Alternative 2 – Excavation and Off-site Disposal for Soil; Plug Lines; In-situ Chemical Oxidation, MNA and LUCs for Intermediate Zone and Shallow Zone Groundwater

The major components of this alternative include the following:

- Excavation and off-site disposal of impacted soil from LHAAP-29 to protect human and ecological receptors, and to eliminate the potential soil-to-groundwater pathway
- Cooling water and transite TNT process wastewater lines will be flushed with water, inspected, and plugged using a bentonite slurry. Confirmation sampling of the wooden TNT wastewater line during the RD may result in excavation and disposal.



- Injection of chemical oxidant in targeted locations in the intermediate groundwater zone to oxidize organic constituents in the saturated zone and extraction of groundwater to help distribute oxidant with MNA following treatment
- MNA with LTM in the intermediate zone (after chemical oxidation is complete) to reduce groundwater contamination to cleanup levels
- MNA with LTM in the shallow zone to reduce groundwater contamination to cleanup levels
- A contingency remedy to enhance MNA if MNA is found to be ineffective. The contingency remedy would consist of injection of bioremediation amendments to enhance degradation of the groundwater contaminants at selected locations based on data available at the time it is determined MNA is not successful. Details for the contingency remedy would be presented in a RD/RAWP.
- Groundwater monitoring will be conducted to evaluate inorganic COCs. The need to continue groundwater monitoring for this purpose will be evaluated at five year reviews.
- LTM semiannually for 3 years, annually until the next five-year review, then annually until recommended otherwise at the five-year review to evaluate remedy performance and determine if plume conditions remain constant, improve, or worsen. Monitoring will continue until the five-year review demonstrates that cleanup levels are reached
- The LUCs' performance objectives are to prohibit groundwater use (except for environmental testing and monitoring) as a potable source until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met; to restrict land use to nonresidential until it is demonstrated that the surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure; and to maintain the integrity of any current or future remedial or monitoring systems until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10** in groundwater are met.

Estimated Capital Present Worth Cost: \$8,070,000

Estimated O&M Present Worth Cost: \$1,070,000

Cost Estimate Duration: 30 years

Estimated Present Worth Cost: \$9,140,000

Alternative 3 – Excavation and Off-site Disposal of Soil; Plug Lines; Intermediate Zone Groundwater Extraction and Treatment, MNA and LUCs for Intermediate and Shallow Groundwater

The major components of this alternative include the following:

- Excavation and off-site disposal of impacted soil from LHAAP-29 to protect human and ecological receptors, and to eliminate the potential soil-to-groundwater pathway
- Cooling water and transite TNT process wastewater lines will be flushed with water, inspected, and plugged using a bentonite slurry. Confirmation sampling of the wooden TNT wastewater line during the RD may result in excavation and disposal.



- Groundwater extraction to reduce VOC levels throughout the intermediate zone groundwater contaminant plume to favorable conditions for MNA
- MNA with LTM in the intermediate zone (after groundwater extraction) to reduce groundwater contamination to cleanup levels
- MNA with LTM in the shallow zone to reduce groundwater contamination to cleanup levels
- A contingency remedy to enhance MNA if MNA is found to be ineffective. The contingency remedy would consist of injection of bioremediation amendments to enhance degradation of the groundwater contaminants at selected locations based on data available at the time it is determined MNA is not successful. Details for the contingency remedy would be presented in a RD/RAWP.
- Groundwater monitoring will be conducted to evaluate inorganic COCs. The need to continue groundwater monitoring for this purpose will be evaluated at five year reviews.
- LTM semiannually for 3 years, annually until the next five-year review, then annually until recommended otherwise at the five-year review to evaluate remedy performance and determine if plume conditions remain constant, improve, or worsen. Monitoring will continue until the five-year review demonstrates that cleanup levels are reached
- The LUCs' performance objectives are to prohibit groundwater use (except for environmental testing and monitoring) as a potable source until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met; to restrict land use to nonresidential until it is demonstrated that the surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure; and to maintain the integrity of any current or future remedial or monitoring systems until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met.

Estimated Capital Present Worth Cost: \$1,550,000

Estimated O&M Present Worth Cost: \$1,780,000

Cost Estimate Duration: 30 years

Estimated Present Worth Cost: \$3,300,000

Alternative 4 - Excavation and Off-site Disposal for Soil; Plug Lines; ISTD, MNA and LUCs for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater

The major components of this alternative include the following:

- Excavation and off-site disposal of impacted soil from LHAAP-29 to protect human and ecological receptors, and to eliminate the potential soil-to-groundwater pathway
- Cooling water and transite TNT process wastewater lines will be flushed with water, inspected, and plugged using a bentonite slurry. Confirmation sampling of the wooden TNT wastewater line during the RD may result in excavation and disposal.
- One of two ISTD process options, ERH (Alternative 4a) or TCH (Alternative 4b) will be selected during the remedial design phase and implemented to remediate the MC DNAPL plume in the intermediate zone to levels amenable to MNA



- MNA with LTM in the intermediate groundwater zone (after ISTD activities are complete) to reduce groundwater contamination to cleanup levels
- MNA with LTM in the shallow zone to reduce groundwater contamination to cleanup levels
- A contingency remedy to enhance MNA if MNA is found to be ineffective. The contingency remedy would consist of injection of bioremediation amendments to enhance degradation of the groundwater contaminants at selected locations based on data available at the time it is determined MNA is not successful. Details for the contingency remedy would be presented in a RD/RAWP.
- Groundwater monitoring will be conducted to evaluate inorganic COCs. The need to continue groundwater monitoring for this purpose will be evaluated at five year reviews.
- LTM semiannually for 3 years, annually until the next five-year review, then annually until recommended otherwise at the five-year review to evaluate remedy performance and determine if plume conditions remain constant, improve, or worsen. Monitoring will continue until the five-year review demonstrates that cleanup levels are reached
- The LUCs' performance objectives are to prohibit groundwater use (except for environmental testing and monitoring) as a potable source until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met; to restrict land use to nonresidential until it is demonstrated that the surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure; and to maintain the integrity of any current or future remedial or monitoring systems until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met.

Alternative 4a:

Estimated Capital Present Worth Cost: \$3,710,000

Estimated O&M Present Worth Cost: \$1,030,000

Cost Estimate Duration: 30 years

Estimated Total Present Worth Cost: \$4,740,000

Alternative 4b:

Estimated Capital Present Worth Cost: \$4,530,000

Estimated O&M Present Worth Cost: \$1,190,000

Cost Estimate Duration: 30 years

Estimated Total Present Worth Cost: \$5,720,000

2.9.2 Common Elements and Distinguishing Features

Common Elements of Alternative 2, 3, and 4

Common elements of Alternatives 2, 3, and 4 are described below.

Soil Excavation –Contaminated soil would be excavated at LHAAP-29 under Alternatives 2, 3, and 4 to prevent human and ecological receptor exposure to contaminants in the soil and to eliminate the soil-to-groundwater pathway. As part of the RD, confirmation soil samples will be collected along



the north and south cooling water lines as well as the TNT wastewater lines to confirm that leaching has not occurred, which may identify additional soil excavation areas. Disposal will be at a RCRA Subtitle D-permitted landfill.

Process Lines – Transit TNT wastewater line would be flushed, then the inlets and outlets will be inspected and plugged with a bentonite slurry mix or equivalent. The cooling water lines will be evaluated further during the RD in order to base the remedial action on up-to-date data. The lines will be flushed with water, inspected and plugged using a bentonite slurry mix or equivalent. Rinsate water will be containerized and characterized for waste handling. During the RD, samples will be collected from the soil along the deteriorated wooden TNT wastewater line and analyzed for explosives to determine if there are concentrations that represent a potential source for groundwater contamination. If present, above the GWP-Ind, the affected soils may be included in the excavation and disposal activity.

MNA – MNA is a passive remedial action that relies on natural biological, chemical, and physical processes to reduce the mass and concentrations of groundwater COCs under favorable conditions. The natural attenuation evaluation indicates that MNA is a feasible technology for the groundwater at LHAAP-29 (Shaw, 2010). Monitoring activities associated with MNA would confirm the protection of human health and the environment by documenting the return of the groundwater to its potential beneficial use as a drinking water supply, and by documenting reduction of the contaminant mass and protection of surface water through containment of the plume. In all three alternatives, contaminant reduction would occur by MNA alone in the shallow zone. For the intermediate zone, to achieve conditions favorable to MNA, MC would be reduced by chemical oxidation in Alternative 2, groundwater extraction in Alternative 3, and thermal destruction in Alternative 4.

MNA performance monitoring would be conducted quarterly for the first two years. After eight quarterly sampling events, MNA effectiveness will be evaluated. The analytical program would consist of VOCs, including chlorinated compounds and degradation products, nitrotoluenes, methane, ethene, and ethane, among others. The full list of MNA parameters would be developed during the RD phase.

Inspection/Long-Term Groundwater Monitoring – Alternatives 2, 3, and 4 include inspection and long-term groundwater monitoring activities. Monitoring would be continued as required to evaluate the effectiveness of the remedy, to demonstrate compliance with applicable or relevant and appropriate requirements (ARARs) and RAOs, and to support five-year reviews.

LUCs – LUCs would be implemented to support the RAOs. The LUC for groundwater would prevent human exposure to residual groundwater contamination presenting an unacceptable risk to human health and ensure that there is no withdrawal or use of groundwater beneath the sites for anything other than environmental monitoring and testing. The LUC to prohibit groundwater use (except for environmental testing and monitoring) as a potable source would remain until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met; to restrict land use to nonresidential until it is demonstrated that the surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure; and to maintain the integrity of any current or future remedial or monitoring systems until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met.



In addition, within 90 days of signature of this ROD, the Army shall request the Texas Department of Licensing and Regulation to notify well drillers of groundwater use prohibitions based on a preliminary LUC boundary. A LUC Remedial Design (RD) will be finalized as the land use component of the Remedial Design. Within 21 days of the issuance of the ROD, the Army will propose deadlines for completion of the RD Work Plan, RD, and Remedial Action Work Plan. The documents will be prepared and submitted to EPA and TCEQ pursuant to the FFA. The LUC RD will contain implementation and maintenance actions, including periodic inspections. The long-term groundwater and surface water monitoring and MNA performance monitoring will also be presented in the RD. Consistent with the dates presented for these documents, the U.S. Army shall: 1) request the Texas Department of Licensing and Regulation to notify well drillers of the final boundary of groundwater use prohibitions; and 2) notify the Harrison County Courthouse of the LUCs to include a map showing the areas of groundwater and nonresidential use restrictions, and the monitoring system at the site, in accordance with 30 TAC 335.565.

The Army will implement, maintain, monitor, report on and enforce land use controls at Army-owned property. The Army shall perform those actions related to land use control activities described in this ROD and in the Remedial Design for the ROD. For portions of the Site subject to land use controls that are not owned by the Army, the Army will monitor and report on the implementation, maintenance, and enforcement of land use controls, and coordinate with federal, state, and local governments and owners and occupants of properties subject to land use controls. The Army will provide notice of the groundwater and soil (surface and subsurface) contamination and any land use restrictions referenced in the ROD. The Army will send these notices to the federal, state and local governments involved at this site and the owners and occupants of the properties subject to those use restrictions and land use controls. The Army shall provide the initial notice within 90 days of ROD signature. The frequency of subsequent notifications will be described in the Remedial Design for the ROD. The Army remains responsible for ensuring that the remedy remains protective of human health and the environment. The Army will fulfill its responsibility and obligations under CERCLA and the NCP as it implements, maintains, and reviews the selected remedy.

Upon transfer of Army-owned property, the Army will provide written notice of the land use controls to the transferee of the groundwater and soil (surface and subsurface) contamination and any land use restrictions referenced in the ROD. Within 15 days of transfer, the Army shall provide EPA and TCEQ with written notice of the division of implementation, maintenance, and enforcement responsibilities unless such information has already been provided in the LUC RD. The LUC RD will address the procedures to be used by the Army and the transferee to document compliance with the LUCs described in this ROD. In the event property is transferred out of Federal control, the land use controls relating to property and groundwater restrictions shall be recorded in the deed and shall be enforceable by the United States and the state of Texas.

To transfer this property (LHAAP-29), an Environmental Condition of Property (ECP) document would be prepared and the Environmental Protection Provisions from the ECP would be attached to the letter of transfer. The ECP would include the LUCs as part of the Environmental Protection Provisions. The property would be transferred subject to the LUCs identified in the ECP. These restrictions would prohibit or restrict property uses that might result in exposure to the contaminated groundwater (e.g., drilling restrictions) or soil (e.g., residential land use prohibition).

The U.S. Army and regulators will consult to determine appropriate enforcement actions should there be a failure of a LUC objective at these sites after they have been transferred.



Distinguishing Feature of Alternative 2

The distinguishing feature of Alternative 2 compared to Alternatives 3 and 4 is the inclusion of ISCO to treat the MC DNAPL in the intermediate groundwater zone. These actions are described below. Note that the design specifications presented below are used for cost estimating purposes and the final design for the selected remedy will be presented in the Remedial Design document.

In situ chemical oxidation – The components of this action include:

Installation of injection wells. Four intermediate zone injection wells (88 feet bgs) would be installed around existing intermediate monitoring well 29WW16, where MC was detected at the highest concentration. The four wells would be arranged in a square with well 29WW16 in the center of the square. The spacing may be adjusted based on actual field conditions.

Injection of oxidation solution. One pore volume of heat activated (40 degrees Celsius [°C]) combined persulfate and sodium hydroxide solution at 60 grams per liter (g/L) and 15 g/L, respectively, would be injected into four wells while simultaneously extracting groundwater from well 29WW216. Temporary piping would be used for the injection array. The estimated volume is approximately 187,000 gallons of activated persulfate and sodium hydroxide solution to be injected into the subsurface. The solution is estimated to be 94,000 pounds of persulfate reagent and 23,500 pounds of sodium hydroxide. A second round of injection may be required if monitoring indicates COCs are not being effectively reduced from the initial round. For the cost estimate, a second round is assumed. If contaminant concentrations do not decrease as anticipated, the method would be modified.

Simultaneous extraction of groundwater. Well 29WW16, or an equivalent, would be converted to an extraction well. Prior to conversion, a pumping test would be conducted and hydrogeologic parameters would be measured to assess aquifer conditions. Groundwater flow in the vicinity of 29WW16 and the injection wells would be modeled to determine the scope of the modifications needed at 29WW16 and to assess the time required to extract one pore volume.

For estimating purposes, it is assumed the on-site groundwater treatment plant is operating and can handle 187,000 gallons extracted for one pore volume. A temporary piping system would be used to convey the extracted water to three 5,000-gallon on-site storage tanks. The on-site tanks would be interconnected and would be equipped with a high level shut off to the extraction pump. Once every two days, water would be pumped into a tank truck and transported to the LHAAP groundwater treatment plant for treatment and discharge (**Figure 2-14**). A 20-foot by 50-foot gravel pad would be prepared for the tanks, and a 6-inch-layer of gravel would be placed to upgrade the road to the tanks at LHAAP-29. Based on the pumping test results, the quantities of extracted water and best approach to handle the water would be evaluated.

Monitor effectiveness. To monitor the effectiveness of the in situ chemical oxidation, six wells would be monitored biweekly for three sampling events. The six wells would include one new monitoring well, four injection wells, and 29WW16. The effect of the first chemical injection should be evident within a few weeks. It is anticipated that a second injection may be needed after approximately 2 months.

Distinguishing Feature of Alternative 3

The distinguishing feature of Alternative 3 compared to Alternatives 2 and 4 is the inclusion of groundwater extraction to address contamination in the intermediate groundwater zone.



Groundwater extraction is estimated to require 3 years to reduce concentrations to levels amenable to MNA. These actions are described below. Note that the design specifications presented below are used for cost estimating purposes and the final design for the selected remedy will be presented in the Remedial Design document.

Pre-Design Study: This action would begin with a pre-design study. A pump test would be conducted and hydrogeologic parameters would be measured to better design the system. Groundwater flow would be modeled to set performance evaluation parameters and to assess the likely time required for remediation.

Construction: Groundwater contamination in the intermediate zone at LHAAP-29 primarily consists of a MC plume. A minimum of five additional wells (four extraction and one monitoring) are proposed to be installed in the intermediate zone within the region of greatest MC contamination in order to provide a more effective extraction process. Several groundwater monitoring wells are located throughout the site and some of these could also be converted to extract contaminated groundwater if needed. The final number and locations for extraction wells would be determined as part of the pre-design study.

Storage and transport of extracted groundwater: A piping system would be constructed to transport the extracted water from the extraction wells 5,000-gallon storage tanks to be located on-site at LHAAP-29. The tanks would be interconnected and equipped with a high level shut off to the pump. Once every two days or at a determined frequency, the water would be pumped into a tank truck and transported to the existing groundwater treatment plant for treatment and discharge.

Performance Monitoring: During extraction, samples would be collected from the extraction wells and monitoring wells to monitor the effectiveness of the action. During startup of the extraction system (until the system is operating properly), bimonthly sampling would be conducted. Startup is estimated to be approximately six months. After startup, monitoring would be reduced to quarterly for the remaining 2.5 years.

Water Treatment and Discharge: The extracted groundwater from LHAAP-29 would be treated at the LHAAP groundwater treatment plant, which was originally built to treat groundwater containing VOCs and metals extracted from other LHAAP sites. The plant uses air stripping, carbon adsorption, and thermal oxidation. Perchlorate treatment using a fluidized bed reactor was added in April 2001 to the treatment plant. The extracted water from LHAAP-29 would be discharged from the tank truck into the existing 300,000-gallon equalization tank. This tank receives water from other LHAAP sites and is stored in this tank until treatment. After the water is treated, the effluent would be discharged in accordance with plant procedures. The plant presently operates at a fraction of its maximum capacity of 1 to 1.5 million gallons of water per month. The original groundwater treatment plant components have adequate capacity to accommodate the increase in volume that would be introduced to the system when the contaminated groundwater is transported from LHAAP-29 to the plant. The system capacity is limited by effluent storage and discharge rate, and this concern was addressed. Recent mitigating measures include the replacement of the reinjection pipeline to increase the pipe diameter to 4-inches, and the installation of a sprinkler system. The capacity issue would be reevaluated as necessary during the remedial action.

Extraction System: Operation and maintenance would include groundwater extraction system maintenance, groundwater treatment plant operations, and environmental media monitoring. In three years, the extraction wells are anticipated to remove the highest concentrations of VOCs from the groundwater intermediate zone at LHAAP-29, thus reducing the contaminant mass to make



conditions favorable for MNA (estimate assumes 3 years). For MNA, four wells would be selected for use as monitoring wells, and monitoring would be implemented to demonstrate that any remaining VOCs are attenuated by natural processes. During the groundwater extraction operations, the extraction wells would require regular maintenance to prevent fouling of well screens, and the extraction pumps would require routine maintenance and may also require replacement. Cleaning of the pipelines, refurbishing pumps and other maintenance activities would be needed on the groundwater collection and transport system during full-scale operation. O&M costs would include the addition of chemicals, power, and labor; equipment cleaning, tank cleaning, general system maintenance, and replacement; and regulatory monitoring and reporting. O&M activities would also be conducted at the LHAAP plant location as part of the routine plant O&M activities.

Distinguishing Feature of Alternative 4

The distinguishing feature of Alternative 4 compared to Alternatives 2 and 3 is the inclusion of ISTD to remediate the MC DNAPL in the intermediate groundwater zone. Alternative 4a, using ERH as the ISTD process, or 4b using TCH as the ISTD process would be selected during the remedial design phase. Thermal desorption operates by heating the subsurface to effectively and quickly volatilize large quantities of VOCs, including those in the form of non-aqueous phase liquid. Available technologies include ERH, TCH, and steam injection, which are typically coupled with a soil vapor extraction (SVE) system and above ground emission controls to capture and/or destroy volatilized VOCs. Pumping of groundwater or multi-phase extraction (MPE) may also be required to contain the mobilized VOCs in groundwater.

These actions are described below. Note that the design specifications presented below are used for cost estimating purposes and the final design for the selected remedy will be presented in the Remedial Design document. The proposed ERH concept developed for this site and used to develop costs presented in the FS does not include a groundwater extraction component, while the proposed TCH concept incorporated groundwater extraction as part of the MPE component. The TCH conceptual design is significantly more robust than the one proposed for ERH, which may in part explain the difference in total cost between Alternatives 4a and 4b. The actual number of heater wells and the need for groundwater extraction would require further evaluation during the remedial design phase. ISTD for either process option would include extraction of vapors and potentially concurrent groundwater extraction. The need for MPE would be determined during the RD phase. Condensate would be removed from the extracted vapors, which would subsequently be treated in a thermal oxidizer for the destruction of the chlorinated VOCs. Hydrogen chloride generated by combustion of chlorinated compounds would be treated in an acid gas scrubber before discharge to ambient air. If groundwater extraction is required, the extracted water would be combined with the condensate removed from the extracted vapors and transported via truck to the existing groundwater treatment plant (GWTP).

Alternative 4a, In-situ thermal desorption using ERH – ERH delivers electricity through subsurface media via an array of electrodes. The heat generated by electrical resistance typically can raise subsurface temperatures to around the boiling point of water. The steam produced from pore-water serves as a medium to carry out volatilized VOCs for capture via SVE and subsequent ex-situ treatment of extracted vapors. Contaminants are also directly volatilized from unsaturated soil and the applied heat can increase hydrolysis of chlorinated solvents, such as MC, and promote in-situ biological activity. Treatment duration is estimated at 65 – 87 days. The components of the ERH technology for Alternative 4a include:



Installation of electrode borings: An array of electrode borings with co-located vapor extraction wells. The specific depths and layout of the array would be determined during the RD. Temperature monitoring probes would also be installed.

Vapors: The extracted vapor would require treatment to remove VOCs. Although activated carbon is proposed in the vendor's conceptual design, it is assumed that the mass loading would require a thermal oxidizer for treatment of the vapors. The airflow rate is estimated at 320 standard cubic feet per minute (scfm).

Storage, transfer, and treatment of extracted water: Condensate from the co-located vapor extraction wells would be collected and stored in an above-ground tank prior to transfer to the GWTP for treatment and disposal.

Alternative 4b, In-situ thermal desorption using TCH - Thermal conduction is the process of heat flow from a high temperature area to a lower temperature area. TCH involves applying heat and vacuum simultaneously to subsurface media with an array of vertical heater/vacuum wells, thus heating up solids (soil and rock) and liquids (water, air and non-aqueous phase liquids). The heat moves out radially from each thermal well until the heat fronts overlap. Thermal conductivities of subsurface materials, such as sands, silt, and clay show little variability, which leads to highly predictable in-situ heating even in challenging heterogeneous settings. Heating of the wells can either be accomplished by electrically-powered heater coils or using combustion of fuels (e.g. natural gas or propane).

Transport of the vaporized contaminants is improved by the creation of permeability, which results from drying and, if clay is present, shrinking of the soil close to the heaters causing the formation of preferential flow paths that allow capture of the vaporized contaminants. The target temperature for TCH is typically the boiling temperature of the groundwater (e.g. 100 degrees Celsius [°C] at a pressure of 1 atmosphere). The steam produced from the groundwater serves as a medium to carry out volatilized VOCs. In addition, the applied heat can increase hydrolysis and promote in-situ biological activity. Treatment duration estimated at 180 days.

The components of the TCH technology for Alternative 4b include:

Installation of heater wells: An array of heater wells with co-located vapor extraction wells. The specific depths and layout of the array would be determined during the RD. Temperature monitoring probes would also be installed.

Vapor treatment: The extracted vapor would require treatment to remove VOCs. Although activated carbon is proposed in the vendor's conceptual design, it is assumed that the mass loading would require a thermal oxidizer for treatment of the vapors. Assumed airflow rate is 320 scfm

Storage, transfer, and treatment of extracted water: Condensate from the co-located vapor extraction wells would be collected and stored in an above-ground tank prior to transfer to the GWTP for treatment and disposal

2.9.3 Expected Outcomes of Each Alternative

Alternative 1 would allow the site to remain a hazard to human and ecological receptors, since it simply leaves the site as is. Alternatives 2, 3, and 4 provide the same outcome to mitigate exposure to human and ecological receptors by excavation and off-site disposal of the contaminated soil. Soil excavation would also eliminate the potential soil-to-groundwater pathway, preventing further



degradation of groundwater from contaminated soil. Alternatives 2, 3, and 4 would significantly and permanently reduce groundwater contaminant concentrations to the applicable cleanup levels and, therefore, provide long-term effectiveness and permanence. This would be achieved by implementing MNA after the implementation of the initial active treatments from Alternatives 2, 3, and 4 to reduce the VOC contaminant concentrations in the intermediate groundwater zone. These include ISCO for Alternative 2, groundwater extraction (pump and treat) for Alternative 3, and ISTD for Alternative 4. Cleanup levels for the shallow zone should be achieved through MNA in approximately 70 years, and in the intermediate zone the cleanup levels should be achieved through MNA following extraction or active treatment in approximately 90 years for Alternatives 2 and 3, and 5-10 years for Alternatives 4a and 4b. However, considering the lithologic variability, particularly the lateral and vertical change from sand to clay, the times to achieve the cleanup levels may vary by an order of magnitude. The similar outcomes are considered to be attainment of the SDWA MCLs to the extent practicable, and consistent with 40 C.F.R. § 300.430(e)(2)(i)(B & C). If no SDWA MCL has been promulgated for a contaminant, the TRRP Residential Groundwater PCL is used in place of the SDWA MCL. In addition, the monitoring activities associated with MNA would confirm the protection of human health and the environment by documenting the return of the groundwater to its potential beneficial use as a drinking water supply to the extent practicable, given the particular circumstances of the site, by documenting reduction of the contaminant mass and protection of surface water through containment of the plume.

The LUC will remain in place to prohibit groundwater use (except for environmental testing and monitoring) as a potable source until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met; to restrict land use to nonresidential until it is demonstrated that the surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure; and to maintain the integrity of any current or future remedial or monitoring systems until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met.

2.10 Summary of Comparative Analysis

Nine criteria identified in the NCP 40 C.F.R. § 300.430(a)(1)(iii)(F) are used to evaluate the different remediation alternatives individually and against each other in order to select a remedy. This section profiles the relative performance of each alternative against the nine criteria, noting how it compares to the other options under consideration. The nine evaluation criteria are discussed below. **Table 2-11** summarizes the comparative analysis of the alternatives.

2.10.1 Overall Protection of Human Health and the Environment

Overall protection of human health and the environment addresses whether each alternative provides adequate protection of human health and the environment and describes how risks posed through each exposure pathway are eliminated, reduced, or controlled, through treatment, engineering controls, and/or land use controls.

The four alternatives provide varying levels of human health protection. Alternative 1, no action, does not confirm achievement of the RAO for the return of groundwater to its potential beneficial use because there is no monitoring involved. Alternative 1 also provides the least protection of all the



alternatives; it provides no reduction in risks to human health or the environment because no measures would be implemented to eliminate the pathway for human exposure to soil or to the groundwater contamination and potential groundwater impacts to Central Creek and Goose Prairie Creek would not be addressed. Additionally, the soil pathway for ecological receptors would not be addressed.

Alternatives 2, 3, and 4 satisfy the RAOs for LHAAP-29. Alternatives 2, 3, and 4 would remove the contaminated soil and solid residue in lines that pose a hazard, and provide confirmation that human health and the environment would be protected because the monitoring would be conducted to show that MNA is returning the groundwater in the contaminated shallow and intermediate groundwater zones at LHAAP-29 to its potential beneficial use as a drinking water supply to the extent practicable, given the particular circumstances of the site, and to document that the plumes are contained and prevented from impacting surface water at levels that could present a risk to human health and the environment. Furthermore, the LUC for groundwater would protect human health by preventing access to the contaminated groundwater until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soils and groundwater allow for unlimited use and unrestricted exposure. Alternatives 2, 3 and 4 provide treatment of the primary COC, MC, for human health in the intermediate zone.

2.10.2 Compliance with ARARs

Section 121(d) of CERCLA and 40 C.F.R. § 300.430(f)(1)(ii)(B) require that remedial actions at CERCLA sites attain legally applicable or relevant and appropriate Federal and state requirements, standards, criteria, and limitations, which are collectively referred to as “ARARs”, unless such ARARs are waived under CERCLA Section 121(d)(4). The ARARs that pertain to this ROD are discussed in **Section 2.13.2**.

Because contaminated groundwater has the potential to flow into Goose Prairie Creek which flows to Caddo Lake, a drinking water supply, chemical-specific ARARs for surface water consumption are appropriate and relevant. Specifically, Texas surface water quality standards are set forth in 30 TAC 307.6(d)(1), Table 2, for MC, TCE, 1,2-DCA, 1,1-DCE, and VC. For those COCs not listed in Table 2, including perchlorate, cis-1,2-DCE, trans-1,2-DCE, and explosives, the TRRP Residential Groundwater PCLs for those COCs would apply.

Alternative 1 does not comply with chemical-specific ARARs because no remedial action or measures would be implemented. Alternatives 2, 3, and 4 do comply with all chemical-specific ARARs for soil because the contaminated soil above the chemical-specific ARAR would be removed. Alternatives 2, 3, and 4 comply with all chemical-specific ARARs for groundwater because they would return the contaminated shallow and intermediate groundwater zones at LHAAP-29 to their potential beneficial use as drinking water, wherever practicable, which for the purposes of this ROD is considered to be attainment of the relevant and appropriate cleanup levels (SDWA MCLs or TRRP Residential Groundwater PCLs if no SDWA MCL is available) to the extent practicable, and consistent with 40 C.F.R. § 300.430(e)(2)(i)(B & C) and 30 TAC 335.559(d)(2). If a return to potential beneficial uses is not practicable, these alternatives would still meet the NCP expectation to prevent further migration of the plume, prevent exposure to the contaminated groundwater, and evaluate further risk reduction. While Alternative 3 provides a level of overall protection similar to Alternatives 2 and 4, Alternatives 2 and 4 would accelerate the MC cleanup in the intermediate zone. Alternative 4 would achieve the cleanup in the shortest period of time, 5-10 years of MNA after active treatment is completed, compared with 90 years of MNA for Alternatives 2 and 3.



Location-specific and action-specific ARARs would not apply to Alternative 1 since no remedial activities would be conducted. Alternatives 2, 3, and 4 would comply with all action-specific ARARs. There are no location-specific ARARs for Alternatives 2, 3, or 4.

2.10.3 Long-Term Effectiveness and Permanence

Long-term effectiveness and permanence refers to expected residual risk and the ability of a remedy to maintain reliable protection of human health and the environment over time, once clean-up levels have been met. This criterion includes the consideration of residual risk that will remain on-site following remediation and the adequacy and reliability of controls.

Alternative 1 would be the least effective and permanent in the long term because no contaminant source removal or treatment would take place and no measures would be implemented to control exposure risks posed by contaminated site soil, sediment, surface water and groundwater. Alternative 1 would also have no effectiveness and permanence with regards to the contaminated soil, since no soil removal would be conducted.

Removing the source soils and either removing the contaminated groundwater through extraction, or using active treatment to destroy VOCs in the DNAPL plume would provide a permanent solution for the contaminants in affected media.

Alternatives 2, 3, and 4 would provide a moderate degree of long-term effectiveness by removing the source soils and providing restoration of the groundwater by treatment and/or MNA. Alternatives 2 and 4 provide a higher level of effectiveness than Alternative 3 because the intermediate groundwater zone would reach concentrations amenable to natural attenuation in a shorter time frame. Alternative 4 would provide the highest level of effectiveness since the intermediate zone groundwater would reach cleanup levels in the shortest timeframe. Alternative 2 allows the opportunity to evaluate the impact of the in-situ treatment and re-inject if necessary. The impact of the ISTD performed under either of the process options for Alternative 4 would also be evaluated to determine whether additional treatment may be necessary. Alternative 3 is as effective and permanent as Alternatives 2 and 4, but would require more time to reduce concentrations amenable to MNA than Alternatives 2 or 4 and would require a longer period of active operations and maintenance. Alternatives 2 and 4 would significantly reduce initial groundwater contaminant concentrations and thereafter rely on natural attenuation and LUCs until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soils and groundwater allow for unlimited use and unrestricted exposure. Monitoring activities associated with MNA would confirm the protection of human health and the environment by documenting the return of the groundwater to its potential beneficial use as a drinking water supply to the extent practicable, by documenting reduction of the contaminant mass and protection of surface water through containment of the plume.

LUCs would provide a moderate to high degree of effectiveness by limiting land use and preventing exposure to contaminated media. As a federally owned property that will remain so as a refuge after transfer, the ability to achieve permanence and effectiveness is greatly enhanced.

The removal of contaminated solid residue from the transite wastewater line and from the cooling water lines would provide a high degree of long-term effectiveness and permanence by removing the contaminated solid residue from the site.



2.10.4 Reduction of Toxicity, Mobility, or Volume Through Treatment

Reduction of toxicity, mobility, or volume through treatment refers to the anticipated performance of the treatment technologies that may be included as part of a remedy.

Alternative 1 does not employ treatment and would not result in a reduction of toxicity, mobility, or volume of contaminants. Alternative 2 would use excavation, ISCO, and natural attenuation to permanently reduce the mass and concentration of contaminants and, therefore, the toxicity, mobility, and volume of the contaminants. In-situ chemical oxidation is an active treatment process.

Alternative 3 would use excavation, groundwater extraction, and natural attenuation to achieve the same reductions in contamination that are expected from Alternative 2. Groundwater extraction is an active treatment process and would reduce toxicity and volume of the contaminants.

Alternative 4 would use excavation, ISTD, and natural attenuation to permanently reduce the mass and concentration of contaminants and, therefore, the toxicity, mobility, and volume of the contaminants. ISTD is an active treatment process.

Biological activity would generate daughter products that may temporarily increase toxicity or mobility of the contaminant plume. Alternatives 2, 3, and 4 include monitoring so that daughter products would be quantified, documented, and evaluated. The same biological activities would also consume the daughter products, and it is anticipated that these concentrations would be reduced to levels below their associated cleanup levels to return groundwater to its potential beneficial use, wherever practicable.

There is an NCP expectation to use treatment to address principal threat wastes, wherever practicable. Alternatives 2, 3, and 4 satisfy the NCP expectation by including treatment components that address the potential for principal threat wastes associated with the high concentrations of MC in the intermediate groundwater zone.

The soil excavation in Alternatives 2, 3, and 4 would reduce mobility because perchlorate and explosive contaminated soils would be removed from the site and placed in a permitted disposal facility. Toxicity and volume would not be reduced by the excavation portion of the alternatives as the form and quantity of contaminants would not be altered.

The removal of contaminated solid residue from the transite wastewater line and cooling water lines by flushing with water in Alternatives 2, 3, and 4 would reduce mobility because the solid residue would be removed from the site and the rinsate collected, analyzed by the toxicity characteristic leaching procedure (TCLP), and properly disposed. Toxicity and volume would not be reduced since quantity of the contaminants would not be altered.

2.10.5 Short-Term Effectiveness

Short-term effectiveness addresses the period of time needed to implement the remedy and any adverse impacts that may be posed to workers, the community, and the environment during construction and operation of the remedy until cleanup levels are achieved.

Alternative 1 would not involve any remedial measures; therefore, no short-term risk to workers, the community, or the environment would exist. The activities associated with Alternatives 2, 3, and 4 would be protective to the surrounding community from short-term risks except for minimal potential



short-term risks during transport (possible accident when soil is transported off site) of perchlorate and explosive contaminated soil.

Alternatives 2, 3, and 4 would involve potential short-term risks to remediation workers associated with exposure to contaminated groundwater from monitoring and/or operation of drilling/construction equipment, and with exposure to contaminated soil during excavation work.

Alternative 2 has additional short-term risks due to remediation workers handling chemical oxidants and also requires heating of the target zone to 40 degrees C, which would pose similar risks as those posed by the thermal treatment included under Alternative 4, such as potential exposure to high voltage power sources. The thermal treatment component of Alternative 4 has additional potential short-term risks due to potential exposure to high voltage power sources and exposure to hot fluids extracted during ISTD treatment. In addition, workers could be exposed to toxic air emissions during ISTD operations.

Alternative 3 would have short-term risks to the remediation workers associated with exposure during increased operations at the LHAAP groundwater treatment system, which include chemical handling and operation of a high-temperature catalytic oxidizer and in handling contaminated groundwater during extraction, temporary storage on site, and conveyance to the GWTP 1.5 miles away. The implementation of Alternative 3 would require more time than either Alternative 2 or Alternative 4.

Alternatives 2, 3, and 4 include the LUCs as elements of their remedies and would provide almost immediate protection from the contaminated groundwater by prohibiting groundwater use except for environmental monitoring and testing through LUC implementation through a relatively quick implementation period. The time period to achieve groundwater cleanup levels is the most significant difference between Alternative 1 versus Alternatives 2, 3, or 4. Alternatives 2 and 3 are expected to take less time to achieve RAOs than Alternative 1, and either Alternative 4a or 4b would require the shortest time for the intermediate zone groundwater to achieve cleanup levels.

2.10.6 Implementability

Implementability addresses the technical and administrative feasibility of a remedy from design through construction and operation. Factors such as availability of services and materials, administrative feasibility, and coordination with other governmental entities are also considered.

Under Alternative 1, no remedial action would be taken. Therefore, no difficulties or uncertainties would be associated with its implementation. For Alternatives 2, 3, and 4, soil excavation would require extensive coordination between excavation, sampling, transportation and disposal. Plugging of the TNT transite wastewater line and cooling water lines can be conducted without extensive intrusive activities with equipment, services, and materials readily available to conduct the activities for Alternatives 2, 3, and 4. Alternatives 2, 3, and 4 could all be implemented. There are qualified vendors available to implement the ISTD technology. There are also qualified contractors with the capabilities to design and implement ISCO. Installation of utilities, availability of equipment and supplies, and compliance with any local ordinances or other requirements would need to be identified and addressed during the design and construction process.

Alternative 3 would involve the use of a groundwater treatment system which currently exists at LHAAP and is accessible to the site; however, from a technical standpoint the increased duration for extraction would require three years for Alternative 3 compared to six months for Alternative 2 and



Alternative 4b, or approximately 3 months for Alternative 4a. The U.S. Army will be responsible for LTM and enforcement of the LUCs, long-term evaluation of MNA, long-term sampling, and LTM and operation of sampling equipment. Technically, Alternatives 2, 3, and 4 are readily implementable.

Administratively, all of the alternatives are implementable.

2.10.7 Cost

Cost estimates are used in the CERCLA process to eliminate those remedial alternatives that are significantly more expensive than competing alternatives without offering commensurate increases in performance or overall protection of human health or the environment. The cost estimates developed are preliminary estimates with an intended accuracy range of –30 to +50 percent. Final costs would depend on actual labor and material costs, actual site conditions, productivity, competitive market conditions, final scope, final schedule, final engineering design, and other variables.

The cost estimates include capital costs (including fixed-price remedial construction) and long-term O&M costs (post-remediation). Overall present worth costs are developed for each alternative assuming a discount rate of 2.8 percent. The duration used for the estimates is a 30-year period.

The progression of present worth costs from the least expensive alternative to the most expensive alternative is as follows: Alternative 1, Alternative 3, Alternative 4a, Alternative 4b, and Alternative 2. No costs are associated with Alternative 1 because no remedial activities would be conducted.

Alternative 3 has the lowest present worth and capital costs of the active remedial alternatives. The presence of the existing groundwater treatment system at LHAAP greatly reduces the capital costs associated with groundwater extraction in Alternative 3. Alternative 2 has the highest present worth and capital costs primarily due to the activities associated with the injection phase of ISCO. Alternative 4a has a lower present worth than Alternative 4b, however the difference in costs between the ISTD process options may reflect differences between the components included in the proposed designs for ERH and TCH developed for the FS. The proposed ERH concept developed for this site and used to develop costs presented in the FS does not include a groundwater extraction component, while the proposed TCH concept incorporated groundwater extraction as part of the MPE component. The TCH conceptual design is significantly more robust than the one proposed for ECH, which may in part explain the difference in total cost between Alternatives 4a and 4b.

2.10.8 State/Support Agency Acceptance

The USEPA and TCEQ have reviewed the Revised Proposed Plan, which presented Alternative 4 as the preferred alternative. Comments received from the USEPA and TCEQ during the Revised Proposed Plan development have been incorporated. Both agencies concur with the selected remedial action.

2.10.9 Community Acceptance

Community acceptance is an important consideration in the final evaluation of the selected remedy. Verbal comments were received during the public meeting held on December 6, 2018 at the Karnack Community Center. No other comments were received during the 30-day public comment period. The topics of the comments included: excavation and disposal of contaminated soils; requirements for vegetation removal and above ground ecological impacts of the remedial actions; the status of



the wooden wastewater line; prior treatability studies; groundwater plume stability and migration; ISTD effectiveness; and regulatory drivers for remediation. Comment responses were provided and incorporated into the ROD, including describing soil sampling and evaluating the wooden wastewater lines for removal.

Several sets of written public comments were received during the 30-day public comment period and public meeting held for the 2011 PP, and there were several verbal comments from the March 22, 2011 public meeting. Although the selected remedy has changed, most of the comments remain relevant. The relevant topics of the 2011 comments included: excavation of contaminated soils, groundwater treatment plant operation, additional groundwater sampling for DNT isomers, remediation for flushing and plugging the subsurface TNT wastewater line, and plugging the cooling water lines. Comment responses were provided and are summarized in the Responsiveness Summary (**Section 3**).

2.11 Principal Threat Waste

Laboratory results from the groundwater at LHAAP-29 have indicated that residual DNAPL may be residing as a source material in the subsurface. In a phase separate from groundwater, the hazardous contaminant MC is characterized as a highly toxic source material and, thus, a principal threat waste. In accordance with the NCP, treatment alternatives have been evaluated through the remedy selection process. The preferred remedial alternative includes an active remedial component that would mitigate the potential principal threat. By instituting an ISTD treatment of the groundwater, this active treatment would be applied to the highest concentration area in the MC groundwater plume and would comply with the NCP expectations regarding treatment of affected media where principal threat may be considered.

2.12 The Selected Remedy

2.12.1 Summary of Rationale for the Selected Remedy

Alternative 4 (Excavation and Off-site Disposal for Soil; Plug Lines; ISTD, MNA and LUCs for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater) is the selected alternative for LHAAP-29 and is consistent with the intended future use of the site as a national wildlife refuge. Either Alternative 4a, using ERH, or Alternative 4b, using TCH as the ISTD process option would be selected during the RD. The ISTD would rapidly reduce MC concentrations in the intermediate zone to make conditions more amenable for MNA. The selected alternative offers a high degree of long-term effectiveness and can be easily and immediately implemented. This alternative would satisfy the RAOs for the site through the following:

- Contaminated soil and sediment removal with off-site disposal to protect the hypothetical future maintenance worker and ecological receptors from exposure and eliminate the soil-to-groundwater pathway. Additional confirmation soil sampling during the RD may identify additional soil excavation areas, see **Section 2.12.2**.
- Flushing, inspecting, and plugging of the transite TNT wastewater line and the vitrified clay cooling water lines to eliminate potential exposure from residual contamination. The wooden wastewater lines would be evaluated during the RD for excavation and disposal.
- For intermediate groundwater zone: ISTD treatment of the MC DNAPL plume to reduce to levels amenable to MNA.



- MNA was selected as one component of the remedy based on available groundwater evidence as presented in the FS (Shaw, 2010). A tiered approach using three lines of evidence was used to examine the occurrence of natural attenuation. The first line of evidence evaluated reductions in COC concentrations over time and with distance, the second line of evidence evaluated geochemical indicators, while the third line of evidence entailed estimation of natural attenuation rates. Historical decreases in concentrations of chlorinated solvents in individual wells were observed in intermediate groundwater, including the detection of daughter products that suggest the occurrence of complete reductive dechlorination. Concentrations of TCE decreased from 8,800 µg/L to 4,340 µg/L at monitoring well 29WW16 and were completely attenuated at monitoring well 29WW35. Concentrations of MC decreased from 8,770 µg/L to undetectable at 29WW35. These results indicated the intermediate contaminant plume is stable. Thus, natural attenuation was considered feasible for intermediate groundwater, but not as a sole remedy due to the high MC concentrations in groundwater at 29WW16 and vicinity. Additional evaluation would be implemented as part of the MNA component. MNA, together with ISTD, would ultimately restore the intermediate groundwater to attain groundwater cleanup levels. This is anticipated to be completed in approximately 5-10 years. Considering the lithologic variability, particularly the lateral and vertical change from sand to clay, the times to achieve the cleanup levels may vary by an order of magnitude. This approximate timeframe to achieve cleanup levels is considered reasonable based on the anticipated future land use of the site as a national wildlife refuge and the fact that there is no current or anticipated future use of groundwater as a drinking water supply. Thus, MNA is an appropriate component of the remedy for the intermediate groundwater because it would protect human health and the environment, and would document that further reductive dechlorination is occurring within the groundwater plume and that contaminant concentrations are being reduced to attain groundwater cleanup levels.
- For shallow groundwater zone: MNA to reduce contaminant levels to cleanup levels and return groundwater to its potential beneficial uses, and confirm the contaminated groundwater remains localized with minimal migration to protect surface water. MNA was selected as the remedy based on available groundwater evidence as presented in the FS (Shaw, 2010). A tiered approach using three lines of evidence was used to examine the occurrence of natural attenuation. The first line of evidence evaluated reductions in COC concentrations over time and with distance, the second line of evidence evaluated geochemical indicators, while the third line of evidence entailed estimation of natural attenuation rates. Historical decreases in concentrations of perchlorate, explosives, and chlorinated solvents in individual wells were observed in shallow groundwater, including the detection of daughter products that suggest the occurrence of complete reductive dechlorination. These results indicated the shallow contaminant plumes are stable, and monitoring wells formerly with COC concentrations above cleanup levels have attained the cleanup levels in the historical sampling record. Noting some of the highest remaining concentrations; perchlorate decreased from 88,000 µg/L to 16,800 µg/L at monitoring well 29WW15, the explosive o-NT decreased from 18,000 µg/L to 1,230 µg/L at monitoring well 29WW05, and the chlorinated solvent 1,2-DCA decreased from 14,000 µg/L to 5,520 µg/L at 29WW15. Thus, natural attenuation was considered feasible for perchlorate, explosives and chlorinated solvents in shallow groundwater. Additional evaluation would be implemented as part of the MNA component. MNA would ultimately restore the shallow groundwater to attain groundwater cleanup levels. This is anticipated to be completed in approximately 70 years



based on attenuation of 1,2-DCA in 29WW15. Considering the lithologic variability, particularly the lateral and vertical change from sand to clay, the times to MCL may vary by an order of magnitude. This approximate timeframe to achieve cleanup levels is considered reasonable based on the anticipated future land use of the site as a national wildlife refuge and the fact that there is no current or anticipated future use of groundwater as a drinking water supply. Thus, MNA is an appropriate component of the remedy for the shallow groundwater because it would protect human health and the environment and would document that further natural attenuation is occurring within the groundwater plume, and that perchlorate, explosives, and chlorinated solvent concentrations are being reduced to attain groundwater cleanup levels.

- The LUC to prohibit groundwater use (except for environmental testing and monitoring) as a potable source would be implemented to ensure protection of human health by preventing exposure to groundwater until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met. The LUC restricting land use to nonresidential would be implemented until it is demonstrated that the surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure. The LUC to maintain the integrity of any current or future remedial or monitoring systems would be implemented until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met.
- Long-term monitoring and reporting would continue until the cleanup levels are achieved in groundwater to confirm protection of human health by preventing exposure to groundwater until cleanup levels are met.

Five-year reviews will be performed to document that the remedy remains protective of human health and the environment.

Based on information currently available, the U.S. Army believes the selected alternative meets the threshold criteria and provides the best balance of tradeoffs among the other alternatives with respect to the CERCLA §121(b) criteria used to evaluate remedial alternatives. The selected alternative: 1) is protective of human health and the environment; 2) complies with ARARs; 3) is cost-effective; 4) utilizes a permanent solution; and 5) utilizes an active treatment as a principal element. The selected remedy addresses the statutory preference for treatment to the maximum extent possible. As a component of the intermediate groundwater zone, MC is characterized as a highly toxic source material and, thus, a principal threat waste. In accordance with the NCP, treatment alternatives have been evaluated through the remedy selection process. The preferred remedial alternative includes an active remedial component that would mitigate the potential principal threat. By instituting ISTD treatment of the groundwater, this active treatment would be applied to the highest concentration area in the MC groundwater plume and would comply with NCP expectations regarding treatment of affected media where principal threat may be considered.

In the remedial design, the U.S. Army would present details of the soil excavation, selected ISTD process (ERH or TCH), ISTD design, LUC operations and maintenance, groundwater monitoring, flushing TNT transite wastewater line and cooling water lines, evaluating the deteriorated wooden wastewater line for excavation and disposal, plugging and abandoning TNT wastewater and cooling water lines, and MNA remedy implementation for LHAAP-29.



2.12.2 Description of the Selected Remedy

The selected remedy, Alternative 4, was outlined in **Section 2.9**; that description is expanded in the following discussion. The major components of the remedy include:

Soil Excavation. The excavation would remove explosives and perchlorate contaminated soils for off-site disposal at a RCRA Subtitle D-permitted landfill. This action would achieve the following: 1) removal of soil that is a direct risk to the hypothetical future maintenance worker, thereby protecting human health by preventing inhalation, ingestion, and dermal contact with the COCs; 2) removal of contaminated soil that is a potential source of contaminant migration to groundwater; and 3) removal of soil posing a risk to ecological receptors. The cleanup levels are presented in **Table 2-10**. The approximate excavation locations are highlighted on **Figure 2-12**. The removal of soil contamination would be verified by collecting confirmation samples from the walls and floors of the excavation area and submitting them for laboratory analysis for the COCs of interest. Clean borrow soil would be used as needed to backfill the excavations so they can be graded for proper drainage. Additional sampling would be required during the remedial design phase to further define explosives impacts near former Building 812-F and in the cooling water outfall/ditch as part of the remedial design, and confirmation soil samples would be collected adjacent to the North and South Cooling Water lines as well as the wooden and transite TNT wastewater lines to confirm that leaching from the lines has not occurred. Results from the confirmation soil sampling may identify additional areas exceeding the cleanup levels presented in **Table 2-10**, which would require soil excavation.

Plug and Abandon Lines. The transite TNT wastewater line and the cooling water lines would be flushed with water to remove solid residue. After flushing, the lines would be visually inspected to evaluate if there is any remaining residue and/or liquid in the lines. The inspection and closure details would be included in the RD and may include techniques such as sampling of flush water and video camera inspection if there is any uncertainty about the effectiveness of the flushing. The rinsate water would be containerized. During typical flushing operations, the flush water would be sampled, analyzed and screened to TCLP (or the equivalent TCEQ test) to determine disposal. The inlets and outlets of the transite TNT wastewater line would be plugged with a bentonite slurry mix or equivalent. The manholes and outlets of the cooling water lines would then be plugged with a bentonite slurry mix or equivalent. The deteriorated wooden wastewater line would be sampled to determine whether contaminants in the line exceed soil cleanup levels and require excavation and disposal.

In-situ Thermal Desorption for Intermediate Zone VOC Groundwater Plume. Under Alternative 4 the highest concentration area in the MC plume in the Intermediate Zone groundwater would be treated using ISTD with either the process option ERH (Alternative 4a) or TCH (Alternative 4b) to be determined during the remedial design phase. Groundwater extraction may be implemented as part of the in-situ treatment to physically remove mass and to control the hydraulic gradient.

ERH delivers electricity through subsurface media via an array of electrodes. The heat generated by electrical resistance typically can raise subsurface temperatures to around the boiling point of water. The steam produced from pore-water serves as a medium to carry out volatilized VOCs for capture via SVE and subsequent ex-situ treatment of extracted vapors. In addition, the applied heat can increase hydrolysis of chlorinated solvents, such as MC, and promote in-situ biological activity in two ways. First, biological activity is boosted by moderately high temperatures (30 °C) found at the periphery of the heated area during active thermal treatment, and throughout the heated area as it



cools. Second, high temperatures increase the solubility of DNAPL, resulting in an increase in contaminant concentrations in the dissolved form that the microbes are able to use, provided the concentrations of the dissolved COCs are not toxic to the microorganisms.

TCH involves applying heat and vacuum simultaneously to subsurface media with an array of vertical heater/vacuum wells. The wells are typically heated by electrical coils, but can also be heated by fuel (e.g. natural gas or propane) combustion. Heat generated from heating elements/wells is transferred to the subsurface largely through thermal conduction and radiant heating. Similar to ERH, the heat generated by TCH typically raises subsurface temperatures to around the boiling point of groundwater and the steam produced from the groundwater serves as a medium to carry out volatilized VOCs. In addition, the applied heat can increase hydrolysis and promote in-situ biological activity.

Preliminary conceptual designs including the number and spacing of heater wells (TCH) or electrode borings (ERH), associated vapor and/or dual phase extraction systems, and treatment train for extracted vapors and condensate were developed and used for FS costing. The final ISTD process option would be determined during the remedial design phase, and the specific design parameters, including the number and locations of heater wells (TCH) or electrodes (ERH) and associated treatment train components may change as the system design is refined.

The ERH preliminary conceptual design components include:

- **Electrode borings.** Fifteen 12-inch diameter electrode borings would be installed to a depth of 91 feet bgs with co-located vapor extraction wells screened above 48 feet bgs; average electrode spacing would be 18 ft. The electrodes would be used to heat the subsurface to near boiling temperatures to volatilize VOCs from the subsurface, and the vapor extraction wells would extract the steam generated.
- **Temperature monitoring probes.** Four probes, with 10 sensors per probe to monitor subsurface temperatures.
- **Vapor treatment.** Activated carbon or a thermal oxidizer would be used to treat the vapors. The vendor's conceptual design included activated carbon, but the expected mass loading would require a thermal oxidizer to treat the extracted vapor. Airflow rate is estimated at 320 scfm.
- **Condensate collection.** Condensate production is estimated at 0.9 gpm and would be captured and stored in aboveground tanks for transfer to the on-site GWTP.
- **Power control unit.** A 480 volt, 3-phase 700 kilowatt power control unit would be required to supply the required power to the electrodes, extraction and treatment components associated with the system operations.

Figure 2-14 shows the conceptual layout for the network of ERH electrodes. The final layout and number of electrodes would be determined as part of the RD if ERH is the selected ISTD process option.

The TCH preliminary conceptual design components include:

- **Heater borings.** 25 heater borings would be installed to a depth of 91 feet bgs with co-located vapor extraction wells screened at close to the target zone depth. Average electrode spacing would be 18 feet.



- **Multi-phase extraction wells.** 7 multi-phase extraction wells would be installed in 10-inch boreholes to extract vapor and water for treatment.
- **Temperature monitoring probes.** 5 temperature monitoring probes with 3 sensors in each probe would be installed in 6-inch boreholes to monitor subsurface temperatures.
- **Vapor treatment.** Activated carbon or a thermal oxidizer would be used to treat the vapors. The vendor's conceptual design included activated carbon, but the expected mass loading would require a thermal oxidizer to treat the extracted vapor. Airflow rate is estimated at 320 scfm.
- **Water collection and treatment.** The water production rate is estimated at 1-3 gpm and would be captured and stored in aboveground tanks for transfer to the GWTP.
- **Power control unit.** A 480-volt, 3-phase 700 kilowatt power control unit would be required to supply the required power to the heater borings, extraction and treatment components associated with the system operations.

Figure 2-15 shows the conceptual layout for the network of TCH heater borings. The final layout and number of electrodes would be determined as part of the RD if TCH is the selected ISTD process option.

Major components of the MNA remedy include:

- **MNA to return groundwater to its potential beneficial use, wherever practicable given the particular circumstances of the site.** MNA begins following ISTD activities. Historic data suggest that natural attenuation of COCs is occurring at the site; however, additional data collection is necessary to fully evaluate natural attenuation. Monitoring wells would be sampled for eight consecutive quarters to evaluate and confirm the occurrence of natural attenuation in conjunction with historical data. Data from the eight quarterly events would be combined with historic data to evaluate the effectiveness of various natural physical, chemical, and biological processes in reducing contaminant concentrations.
- **Performance objectives to evaluate the MNA remedy performance after two years.** Each of the general performance objectives must be met as indicated below. If MNA is effective, a baseline would be established from the data to that point in time. Specific evaluation criteria would be developed in the RD. A contingency remedy would be developed and implemented to enhance MNA if it is found to be ineffective. If the criteria are not met to illustrate that MNA is an effective remedy, the contingency action would be initiated.

The MNA evaluation would be based on the USEPA lines of evidence (USEPA, 1999) as follows:

- Plume stability (i.e., the plume concentrations are decreasing in the majority of performance wells, and the plume is not expanding in area as demonstrated with compliance wells)
- MNA Process Evaluation demonstrated based on an attenuation rate calculated with empirical performance monitoring data, and MNA Process Demonstration based on the presence of daughter products and bacterial culture counts
- **A contingency remedy to enhance MNA to reach the RAOs if MNA is found to be ineffective.** The area and the elements of the contingency remedy would be selected based on the entire data set available. The contingency remedy would consist of injection of bioremediation amendments to enhance degradation of the groundwater contaminants at



selected locations based on data available at the time it is determined MNA is not successful. Development and specific description of the contingency remedy would be presented in a RD/RAWP.

- **Initiate LTM.** Monitoring would be conducted to evaluate the remedy performance and determine if the plume conditions remain constant, improve or worsen after the baseline is established. LTM would be implemented at a frequency of semiannual for three years, then annually until the next five-year review. The performance monitoring plan would be developed in the RD and would be based on USEPA guidance (USEPA, 2004).
 - Continue LTM annually thereafter until recommended otherwise by the five-year review to evaluate remedy performance and determine if plume conditions remain constant, improve, or worsen. The baseline of the plume for future five-year reviews would be established as part of the MNA evaluation program. The initial LTM plan would be developed in the RD.
 - Groundwater monitoring would be conducted to evaluate inorganic COCs. The need to continue groundwater monitoring for this purpose would be evaluated at five year reviews.
- **Land Use Control.** The LUC objectives include maintaining the integrity of any current or future remedial or monitoring systems, and preventing the use of groundwater contaminated above cleanup levels as a potable water source. The groundwater treatment and MNA remedial components include a groundwater monitoring system that will be used to characterize the condition of the groundwater during the period the groundwater remedy is in place until the groundwater remediation goals are achieved, and to demonstrate achievement of the groundwater remediation goals when the groundwater remedy is complete. As a part of this groundwater remedy, the Army will maintain the remedial and monitoring systems associated with the groundwater remedies until these components of the remedy are no longer needed to achieve cleanup levels, and cleanup levels have been achieved. During the period of operation of the groundwater remedy, if any of the elements of the remedial and groundwater monitoring systems are damaged, destroyed, or become ineffective, they will be repaired or replaced with suitable components to assure that the remedial and groundwater monitoring systems are able to provide data of the quality necessary to determine the progress of and eventual completion of this component of the remedy. The actions to be taken to implement these LUC objectives and requirements will be provided through modifying the “Comprehensive Land Use Control (LUC) Management Plan, Former Longhorn Army Ammunition Plant, Karnack, Texas” and detailed in the LUC RD.
 - The LUC for prohibition of groundwater use (except for monitoring and testing) shall be implemented and shall remain in place at the Site until the COCs (i.e. including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soil and groundwater remaining at the site are reduced below levels that would support unlimited use and unrestricted exposure. A LUC RD will be finalized as the land use component of the Remedial Design. Within 21 days of the issuance of the ROD, the Army will propose deadlines for completion of the RD Work Plan, RD and Remedial Action Work Plan. The documents will be prepared and submitted to the EPA and the TCEQ pursuant to the FFA. The LUC RD will contain implementation and maintenance actions, including periodic inspections. The long-term groundwater and surface water monitoring and MNA performance monitoring plan will also be presented in the RD. The recordation notification for the



- Site which will be filed with Harrison County, will include a description of the LUCs. The preliminary boundary for the groundwater LUC is shown on **Figure 2-16**.
- The LUC restricting land use to nonresidential shall be implemented until it is demonstrated that surface and subsurface soil and groundwater COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) are at levels that allow for unlimited use and unrestricted exposure.
 - The LUC to maintain the integrity of any current or future remedial or monitoring systems will remain in place until the levels of COCs (i.e., including all hazardous substances, pollutants and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in groundwater are met. The LUC to prohibit groundwater use (except for environmental monitoring and testing) as a potable source will remain in place until the levels of COCs (i.e., all hazardous substances, **Table 2-10**) in soil and groundwater allow for unlimited use and unrestricted exposure.

The Army will implement, maintain, monitor, report on and enforce land use controls at Army-owned property. The Army shall perform those actions related to land use control activities described in this ROD and in the Remedial Design for the ROD. For portions of the site subject to land use controls that are not owned by the Army, the Army will monitor and report on the implementation, maintenance, and enforcement of land use controls, and coordinate with federal, state, and local governments and owners and occupants of properties subject to land use controls. The Army will provide notice of the groundwater and soil (surface and subsurface) contamination and any land use restrictions referenced in the ROD. The Army will send these notices to the federal, state and local governments involved at this site, and the owners and occupants of the properties subject to those use restrictions and land use controls. The Army shall provide the initial notice within 90 days of ROD signature. The frequency of subsequent notifications will be described in the Remedial Design for the ROD. The Army remains responsible for ensuring that the remedy remains protective of human health and the environment. The Army will fulfill its responsibility and obligations under CERCLA and the NCP as it implements, maintains, and reviews the selected remedy.

Upon transfer of Army-owned property, the Army will provide written notice of the land use controls to the transferee of the groundwater and soil (surface and subsurface) contamination and any land use restrictions referenced in the ROD. Within 15 days of transfer, the Army shall provide USEPA and TCEQ with written notice of the division of implementation, maintenance, and enforcement responsibilities unless such information has already been provided in the LUC RD. The LUC RD will address the procedures to be used by the Army and the transferee to document compliance with the LUCs described in this ROD. In the event property is transferred out of Federal control, the land use controls relating to property and groundwater restrictions shall be recorded in the deed and shall be enforceable by the United States and the state of Texas.

LUC implementation and maintenance actions will be described in the RD for LHAAP 29. The LUCs would be included in the property transfer documents and a recordation of them filed in the Harrison County Courthouse. The LUC for groundwater will prevent human exposure to groundwater contaminated with chlorinated solvents, explosives, metals, and perchlorate through the prohibition of groundwater use (except for environmental monitoring and testing), restrict land use to nonresidential, require maintenance of the integrity of any current or future remedial or monitoring systems and prevent the use of groundwater contaminated above cleanup levels as a potable water source. In addition, within 90 days of signature of this ROD, the Army shall request the Texas



Department of Licensing and Regulation to notify well drillers of groundwater use prohibitions based on a preliminary LUC boundary. Within 21 days of the issuance of the ROD, the Army will propose deadlines for completion of the RD Work Plan, RD, and Remedial Action Work Plan. The documents will be prepared and submitted to USEPA and TCEQ pursuant to the FFA. The LUC RD will contain implementation and maintenance actions, including periodic inspections. Consistent with the dates presented for these documents, the Army shall: 1) request the Texas Department of Licensing and Regulation to notify well drillers of the final boundary of groundwater use prohibitions; and 2) notify the Harrison County Courthouse of the LUCs to include a map showing the area of groundwater use prohibition at the site, in accordance with 30 TAC 335.565.

Monitoring activities associated with the LUCs will confirm that groundwater is not being used. Long-term operational requirements under this alternative will include maintenance of the LUCs. Groundwater monitoring will demonstrate no migration of the plume and the eventual reduction of contaminants to levels below cleanup levels. The need for continued groundwater monitoring will be evaluated every 5 years during the reviews. Sampling frequency and analytical requirements will be presented as an appendix to the RD for LHAAP-29.

2.12.3 Cost Estimate of the Selected Remedy

Table 2-12 and **Table 2-13** present the present worth analysis of the cost for the selected remedy, Alternative 4a or 4b. The information in the tables is based on the best available information regarding the anticipated scope of the remedial alternative. The quantities used in the estimates are for estimating purposes only. Changes in the cost elements are likely to occur as a result of new information and data collected during the engineering design of the remedial alternative. Changes will be documented in accordance with 40 CFR 300.435(c)(2) in the form of a memorandum in the Administrative Record, an Explanation of Significant Difference (ESD), or a ROD amendment, as necessary. This is an order-of-magnitude engineering cost estimate that is expected to be within -30 to +50 percent of the actual project cost.

The total project present worth, capital, and O&M costs for Alternatives 4a and 4b are shown in **Table 2-12** and **Table 2-13**. For Alternative 4a, the capital costs are \$3.71M, O&M costs are \$1.03M, and total present worth costs are \$4,740M. For Alternative 4b, the capital costs are \$4.53M, O&M costs are \$1.19M, and total present worth costs are \$5.72M. The costs were developed using a discount rate of 2.8%. The O&M costs include evaluation of MNA, maintenance of the LUC, and LTM through Year 30. The LTM would support the required CERCLA five-year reviews.

2.12.4 Expected Outcomes of Selected Remedy

The purpose of this response action is to attain the RAOs stated in **Section 2.8** of this document. **Table 2-10** presents the cleanup levels. The cleanup levels for the COCs in the groundwater are the Federal SDWA MCLs, or in the absence of federal drinking water standards, the cleanup level is the TRRP Residential Groundwater PCL. The cleanup level for the soil is the GWP-Ind MSC. The cleanup levels for the COPECs in soil are the EcoPRGs.

The expected outcome of the selected remedy is that contaminants in soil and groundwater will be reduced to the cleanup levels. Achievement of the cleanup levels (**Table 2-10**) is anticipated to be completed in approximately 70 years based on the MNA for 1,2-DCA in the shallow zone. MNA in the intermediate zone is estimated to take 5-10 years following active treatment of the DNAPL plume. Considering the lithologic variability, particularly the lateral and vertical change from sand to clay, the times to achieve the cleanup levels may vary by an order of magnitude. This approximate



timeframe to achieve cleanup levels is considered reasonable for the anticipated future land use as a national wildlife refuge.

The LUC for the maintenance of the monitoring system will be maintained until the groundwater cleanup levels are achieved. The LUCs for soil and groundwater will be maintained until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) allow for unlimited use and unrestricted exposure. In the short-term (prior to the groundwater achieving cleanup levels), the site will be made part of a national wildlife refuge operated by USFWS, and will continue as such in the long-term (after the groundwater achieves cleanup levels).

In addition, the monitoring activities associated with MNA will confirm the protection of human health and the environment by documenting the return of the groundwater to its potential beneficial use as a drinking water supply, by documenting reduction of the contaminant mass, and protection of surface water through containment of the plume. The LUC for groundwater will prohibit the use of the site's groundwater except for environmental monitoring and testing.

As part of the evaluation of MNA, attenuation rates are computed and evaluated in accordance with the USEPA guidance material (USEPA, 1998). Time-dependent attenuation rate constants and estimated in-well cleanup times are determined based on COC concentration data over time from individual wells assuming first order degradation kinetics. Attenuation rates are calculated for the monitoring wells with the highest concentrations for which the available data allow such a calculation. Attenuation rates are based on the following formula from the USEPA guidance (USEPA, 1998):

$$C = C_0 e^{-kt}$$

where: C = concentration at time t
 C₀ = initial concentration
 k = attenuation rate constant (first order reaction)

2.13 Statutory Determinations

Under CERCLA §121 and the NCP, the U.S. Army must select remedies that are protective of human health and the environment, comply with ARARs (unless a statutory waiver is justified), are cost effective, and utilize permanent solutions and alternative treatment technologies or resource recovery technologies to the maximum extent practicable. In addition, CERCLA includes a preference for remedies that employ treatment that permanently and significantly reduce the volume, toxicity, or mobility of hazardous wastes as a principal element and a bias against off-site disposal of untreated wastes. The following sections discuss how the selected remedy meets the statutory requirements.

2.13.1 Protection of Human Health and the Environment

The selected remedy, Alternative 4, will achieve the RAOs for LHAAP-29. For the protection of human health, the remedial action would remove soil that exceeds the cleanup levels; flush and remove residues from the process lines and properly dispose the rinsate; reduce groundwater COCs with ISTD in the intermediate zone followed by MNA; reduce shallow groundwater COCs with MNA, which would eventually achieve the destruction of the COCs present in the groundwater plumes at



LHAAP-29. Continued maintenance of the LUC for groundwater will prevent human access and exposure to groundwater that poses an unacceptable risk to human health, until COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soils and groundwater, have sufficiently degraded to levels that allow for unlimited use and unrestricted exposure. At LHAAP-29 the evaluation of historical groundwater contaminant trends indicates that natural attenuation processes are occurring at the site. This remedy provides adequate confirmation that human health and the environment are protected because monitoring would be conducted to document the effectiveness of MNA. The monitoring activities associated with MNA will ensure that COCs and by-product (daughter) contaminants in groundwater do not flow to surface water bodies at such levels that ARARs are exceeded. There are currently no surface water impacts identified. The LUCs for soil and groundwater will be maintained until the levels of COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) in soil and groundwater allow for unlimited use and unrestricted exposure.

For the protection of ecological receptors, the remedial action would remove soil at select areas (in addition to those areas excavated for the protection of human health) to address ecological risks. The outcome of the removal is that the soil in the Industrial Sub-Area, which includes LHAAP-29, will satisfy the EcoPRGs.

There are no short-term threats associated with the selected remedy that cannot be readily controlled. In addition, no adverse cross-media impacts are expected from the selected remedy.

2.13.2 Compliance with ARARS

The selected remedy complies with all ARARs. The ARARs are presented below and in **Table 2-14**.

Chemical-Specific ARARs

- **Soil:** There are no federally promulgated chemical-specific ARARs for soil. The State of Texas promulgated cleanup standards under 30 TAC 335, Subchapter S, specifically 30 TAC 335.559 (g)(2) which specifies contaminant concentration limits for nonresidential soil and are used as the chemical-specific ARARs for the site soils. The concentrations represent the non-residential soil-to-groundwater cross-media protection concentrations that must be met to demonstrate that a contaminant in soil does not pose the potential for a future release of leachate in excess of the groundwater concentration considered to be protective for nonresidential worker exposure. It is anticipated that removal of contaminated soils above the Texas standards will prevent further contamination of the groundwater from soil at the site.
- **TNT Wastewater Line and Cooling Water Lines:** The removal of explosive-contaminated solid or liquid residue remaining in the line by flushing and disposing of the rinsate based on TCLP criteria or equivalent TCEQ criteria will prevent any further contamination of the groundwater from any explosive-contaminated residue remaining in the lines, in accordance with 40 C.F.R. § 261.2 and 30 TAC 335.559(g)(2).
- **Surface water:** Section 121(d)(2) of CERCLA states that every remedial action shall require a level of control which at least attains surface water quality criteria established under Sections 304 or 303 of the Clean Water Act of 1972 (CWA) where such goals and criteria are relevant and appropriate under the circumstances of the release or threatened release. Therefore, surface water quality criteria may be ARARs if there is a remedial action that



affects surface water, and measures will be implemented during construction to prevent off-site migration of contaminants to surface waters. In the event of remedy failure resulting in or potentially resulting in a release to surface water, 40 C.F.R. §§ 122, 125, 129, and 130-131, 40 C.F.R. §§ 141.61 and 141.62, and 30 TAC 307.4, 307.6, 307.7, 307.8 and 307.9 are considered potential future ARARs.

- **Groundwater:** Cleanup levels are presented in **Table 2-10**. The cleanup goal for groundwater will be the SDWA MCLs as specified in 40 C.F.R. §§ 141.61 and 141.62, which meet health-based standards and criteria. In the absence of federal drinking water standards, clean-up levels will be based on TRRP Residential Groundwater PCLs.

This alternative will return the contaminated shallow and intermediate groundwater zones at LHAAP-29 to their potential beneficial use as drinking water, wherever practicable, which for the purposes of this ROD is considered to be attainment of the relevant and appropriate SDWA MCLs, and consistent with 40 C. F. R. § 300.430(e)(2)(i)(B&C). If a return to potential beneficial uses is not practicable, this alternative would still meet the NCP expectation to prevent further migration of the plume, prevent exposure to the contaminated groundwater, and evaluate further risk reduction.

Location-Specific ARARs

There are no location-specific ARARs.

Action-Specific ARARs

The selected remedy has potential action-specific ARARs related to the following activities: site preparation, construction, and excavation activities; waste generation, characterization, management, storage, and disposal activities; well construction and abandonment; and water treatment.

- **Site Preparation, Construction, and Excavation Activities:** Certain on-site preparation, construction, and/or excavation activities will be necessary under all remediation actions to prepare the site for remediation, including the soil-moving or site-grading activities. Control of fugitive emissions and storm water runoff during implementation of these activities will be required. Airborne particulate matter resulting from construction or excavation activities is subject to the fugitive dust and opacity limits listed in 30 TAC 111, Subchapter A. No person may cause, suffer, allow, or permit visible emissions from any source to exceed an opacity of 30 percent for any 6-minute period (30 TAC 111.111[a]). Reasonable precautions must also be taken to achieve maximum control of dust to the extent practicable, including the application of water or suitable chemicals or the complete covering of materials (30 TAC 111.143 and 30 TAC 111.145). Texas has also promulgated general nuisance rules for air contaminants mandating that no person shall discharge from any source whatsoever one or more air contaminants, or combinations thereof, in such concentration and of such duration as are or may tend to be injurious to or to adversely affect human health or welfare, animal life, vegetation, or property, or as to interfere with the normal use and enjoyment of animal life, vegetation, or property (30 TAC 101.4). Storm water discharges from construction activities that disturb equal to or greater than one acre of land must comply with the substantive requirements of a USEPA National Pollutant Discharge Elimination System (NPDES) general permit (40 C.F.R. § 122. 26), depending on the amount of acreage disturbed. Substantive requirements include implementation of good construction



management techniques; phasing of large construction projects; minimal clearing; and sediment, erosion, structural, and vegetative controls to mitigate runoff and ensure that discharges meet required parameters.

- Waste and Disposal Activities:** The processes of monitoring, intercepting, or treating contaminated groundwater may generate a variety of primary and secondary waste streams (e.g., soil, personal protective equipment, and dewatering and decontamination fluids). These waste streams are expected to be non-hazardous waste. All solid waste (defined as any solid, liquid, semisolid, or contained gaseous material intended for discard [40 C.F.R. § 261.2]) generated during remedial activities must be appropriately characterized to determine whether it contains RCRA hazardous waste (40 C.F.R. § 262.11; 30 TAC 335.62; 30 TAC 335.503[a][4]; 30 TAC 335.504). All wastes must be managed, stored, treated (if necessary), and disposed in accordance with the ARARs for waste management listed in **Table 2-14** for the particular type of waste stream or contaminants in the waste.
- Well Construction:** The remedial action may involve the placement, use, or eventual plugging and abandonment of some type of groundwater monitoring, injection, and/or extraction wells, either for in-situ treatment or extraction of the contaminated groundwater or for LTM of the groundwater. Available standards for well construction and plugging/abandonment would provide ARARs for such actions and include 30 TAC 331, Subchapters A and H. Specific provisions 30 TAC §331.9(a); 30 TAC §331.10(a); 30 TAC §331.10(d); 30 TAC §331.21; 30 TAC §331.132(a); 30 TAC §331.132(c); 30 TAC §331.132(d)(1); 30 TAC §331.132(d)(4); 30 TAC §331.133(e) apply. Texas has promulgated technical requirements in Chapter 76 of Title 16 of the TAC applicable to construction, operation, and plugging/abandonment of water wells. In particular, 16 TAC 76.1000 (*Locations and Standards of Completion for Wells*), 16 TAC 76.1002 (*Standards for Wells Producing Undesirable Water or Constituents*) (LHAAP-29 contaminated groundwater could be considered “undesirable water” defined pursuant to Section 76.10[36] as “water that is injurious to human health and the environment or water that can cause pollution to land or other waters”), 16 TAC 76.1004 (*Standards for Capping and Plugging of Wells and Plugging Wells that Penetrate Undesirable Water or Constituent Zones*), and 16 TAC 76.1008 (*Pump Installation*) may provide ARARs for the placement, construction, and eventual plugging/abandonment of groundwater injection or extraction wells or the placement and long-term operation of groundwater monitoring wells for proposed groundwater remedial strategies.
- Water treatment:** Contaminated groundwater and wastewaters collected during well drilling or decontamination activities could be transported to the groundwater treatment plant at LHAAP-18/24 for processing, and would subsequently be discharged in compliance with the effluent limits for that plant. Such waters would be characterized, as required, before transport and managed accordingly in compliance with requirements for the type of waste contaminating the water. To assure compliance with the groundwater treatment plant’s discharge limits, the incoming water must meet the waste acceptance criteria for the facility. On-site wastewater treatment units (as defined in 40 C.F.R. § 260.10) that are part of a wastewater treatment facility that is subject to regulation under Section 402 or Section 307(b) of the Clean Water Act of 1972 are not subject to RCRA Subtitle C hazardous waste management standards (40 C.F.R. § 270.1(c)(2)(v) and 40 C.F.R. § 264.1(g)(6).; 30 TAC 335.42[d][1]). The USEPA has clarified that this exemption applies to all tanks, conveyance



systems, and ancillary equipment, including piping and transfer trucks, associated with the wastewater treatment unit (Federal Register Title 53, 34079, September 2, 1988).

2.13.3 Cost-Effectiveness

Table 2-12 and **Table 2-13** present the present worth analysis of the cost estimates for the selected remedy. The information in the tables is based on the best available information regarding the anticipated scope of the remedial alternative. The quantities used in the estimates are for estimating purposes only. Changes in the cost elements are likely to occur as a result of new information and data collected during the engineering design of the remedial alternative. The least expensive alternative to the most expensive alternative (provided no contingencies are implemented) is as

follows: Alternative 1, Alternative 3, Alternative 4a, Alternative 4b, and Alternative 2. No costs are associated with Alternative 1 because no remedial activities would be conducted. Alternative 3 has the lowest present worth and capital costs of the remediation alternatives (Alternatives 2, 3, 4a, and 4b). The present worth cost for Alternative 3 is lower than that of Alternative 2, as it does not involve chemical treatment or construction costs for a groundwater extraction system. However, costs for operation and maintenance are higher for Alternative 3 than that of Alternative 2. Alternative 3 also estimates assume a 3-year duration for extraction; however, the presence of inferred DNAPL and sorbed MC is expected to require extraction for a longer period of time.

Additionally, although Alternative 3 appears to have lower costs than Alternatives 4a or 4b, this ignores that Alternative 3 will take longer to meet the RAOs in the intermediate zone due to the presence of DNAPL and would instead rely on the MNA component and LUCs to protect human health and the environment. Similarly, Alternative 2 is estimated to have the highest cost and has higher uncertainty of ISCO performance in full-scale application compared to bench-scale testing. Since the DNAPL in the intermediate zone may persist longer, MNA and LUCs may need to remain in place longer as well under Alternatives 2 and 3 than under Alternative 4. A comparison between the sub-alternatives 4a (ERH) and 4b (TCH) for treatment of the MC plume in intermediate zone indicates that sub-alternative 4a may have a lower cost; however, the TCH conceptual design under Alternative 4b appears to be more robust.

2.13.4 Utilization of Permanent Solutions and Alternative Treatment (or Resource Recovery) Technologies to the Maximum Extent Practicable

The U.S. Army has determined that the selected remedy represents the maximum extent to which permanent solutions and treatment technologies can be utilized in a practicable manner at the site. Soil excavation would remove impacted soils and groundwater extraction and in-situ treatment would irreversibly reduce groundwater contaminant concentrations in the treated portions of the groundwater plume. Removal of contaminated pipeline solid residue would protect the groundwater from contaminant leaching, and MNA will reduce groundwater contaminants to cleanup levels.

The selected remedy would provide reduction in toxicity, mobility, and volume of the groundwater contaminants via active treatment of the most contaminated areas. The selected remedy would document effectiveness through the confirmation of MNA and the routine monitoring of the attenuation and migration of the contaminants in groundwater.



The selected remedy would provide immediate protection because the LUCs would be implemented quickly. Maintenance of this control would be required until COCs (i.e., including all hazardous substances, pollutants, and contaminants found at the Site at cleanup levels as listed in **Table 2-10**) and by-product (daughter) contaminant concentrations in soil and groundwater that allow for unlimited use and unrestricted exposure.

2.13.5 Preference for Treatment as a Principal Element

The selected remedy would reduce the toxicity, mobility, or volume of contaminants in the groundwater through an active remedial process. By utilizing ISTD as a significant portion of the remedy, the statutory preference for remedies that employ treatment as a principal element is satisfied. In addition, there is a potential principal source threat at LHAAP-29 residing as residual source material in the subsurface. As a component of this groundwater, the hazardous contaminant MC is characterized as a highly toxic source material and, thus, potentially a principal threat waste.

2.13.6 Five-Year Review Requirements

Section 121(c) of CERCLA and NCP 40 C.F.R. § 300.430(f)(5)(iii)(C) provide the statutory and legal bases for conducting five-year reviews. Because this remedy will result in contaminants that remain on site above levels that allow unlimited use and unrestricted exposure, a review will be conducted at least every five years to confirm that the remedy continues to provide adequate protection of human health and the environment.

2.14 Significant Changes from the Proposed Plan

The Revised Proposed Plan was issued for public comments on November 21, 2018 and identified Alternative 4 as the Preferred Alternative, with either Alternative 4a or 4b to be selected during the RD. The U.S. Army reviewed all verbal comments that were discussed during the public meeting (there were no written comments). After careful consideration of the comments, it was determined that no significant changes to the remedy, as originally identified in the Proposed Plan, were necessary or appropriate.



Table 2-1. Summary of Chemicals of Potential Concern and Medium-Specific Exposure Point Concentrations

Scenario Timeframe: Current
Medium: Groundwater
Exposure Medium: Groundwater

Exposure Point	Chemical	Concentration Detected ¹ (mg/L)		Frequency of Detection	Exposure Point Concentration (mg/L)	Statistical Measure
		Minimum	Maximum			
Ingestion, inhalation, dermal contact	Dioxin/Furan					
	2,3,7,8-TCDD TEC	2.82E-09	1.25E-08	---	1.25E-08	maximum
	Explosives					
	2,4-Dinitrotoluene	0.0087	0.530	6/50	5.30E-01	maximum
	2,6-Dinitrotoluene	0.0087	0.530	7/50	5.30E-01	maximum
	2-Amino-4,6-dinitrotoluene	0.0059	0.0059	1/50	5.90E-03	maximum
	2-Nitrotoluene	0.002	4.40	9/50	4.40E+00	maximum
	3-Nitrotoluene	0.0021	0.240	7/50	2.40E-01	maximum
	4-Amino-2,6-dinitrotoluene	0.0059	0.0059	1/50	5.90E-03	maximum
	4-Nitrotoluene	0.0054	2.100	8/50	2.10E+00	maximum
	Metals					
	Aluminum	0.21	130	30/43	1.30E+02	maximum
	Antimony	0.005	0.052	40/68	5.20E-02	maximum
	Arsenic	0.008	0.059	7/68	5.90E-02	maximum
	Barium	0.024	6.5	37/68	6.50E+00	maximum
	Beryllium	0.0005	0.0099	6/43	9.90E-03	maximum
	Cadmium	0.0012	0.00623	2/68	6.23E-03	maximum
	Chromium	0.01	7.6	45/68	7.60E+00	maximum
	Lead	0.00241	0.35	26/68	3.50E-01	maximum
	Manganese	0.022	2.41	42/43	2.41E+00	maximum
	Mercury	0.002	0.003	2/68	3.00E-03	maximum
	Nickel	0.04	8.4	29/68	8.40E+00	maximum
	Selenium	0.006	0.35	6/68	3.50E-01	maximum
	Silver	0.01	0.08	5/68	8.00E-02	maximum
	Strontium	0.2	19	43/43	1.90E+01	maximum
	Thallium	0.0011	0.003	14/68	3.00E-03	maximum
	Vanadium	0.12	0.36	2/43	3.60E-01	maximum
	Non-Metallic Anion					
	Perchlorate	8.00E-03	8.80E+01	13/30	8.80E+01	maximum
	Volatile Organics					
	1,2-Dichloroethane	14	14	1/68	1.40E+01	maximum
	Acetone	0.0058	0.0058	1/68	5.80E-03	maximum
	Bromodichloromethane	0.001	0.0022	3/68	2.20E-03	maximum
Chloroform	0.0012	0.014	7/68	1.40E-02	maximum	
cis-1,2-Dichloroethene	0.0013	0.0013	1/50	1.30E-03	maximum	
Methylene chloride	0.001	6,600	12/68	6.60E+03	maximum	
p-Cymene	0.0029	0.0029	1/43	2.90E-03	maximum	
Trichloroethene	0.0011	1.200	3/68	1.20E+00	maximum	



Table 2-1. Summary of Chemicals of Potential Concern and Medium-Specific Exposure Point Concentrations (continued)

Scenario Timeframe: Current
Medium: Soil
Exposure Medium: Soil (0 to 2 feet below ground surface)

Exposure Point	Chemical	Concentration Detected ¹ (mg/kg)		Frequency of Detection	Exposure Point Concentration (mg/kg)	Statistical Measure
		Minimum	Maximum			
Ingestion, inhalation, dermal contact	Dioxin/Furan					
	2,3,7,8-TCDD TEC	2.63E-07	7.71E-06	---	4.20E-06	95% UCL
	Explosives					
	2,4,6-Trinitrotoluene	3.8	190	5/49	1.90E+02	maximum
	2,4-Dinitrotoluene	0.760	6.2	2/49	6.20E+00	maximum
	2-Amino-4,6-dinitrotoluene	2.6	25	4/49	2.5E+01	maximum
	4-Amino-2,6-dinitrotoluene	1.1	16	3/18	1.60E+01	maximum
	Metals					
	Antimony	1.2	2.5	15/65	1.92E+00	95% UCL
	Mercury	0.12	0.22	3/75	2.20E-01	maximum
	Non-Metallic Anion					
	Perchlorate	2.45E-02	7.03E-02	5/6	7.03E-02	maximum

Notes:

¹ Minimum/maximum detected concentration above the reporting limit

For groundwater, the maximum detected concentrations were used to estimate the exposure point concentration.

For soil, the 95% UCL values were used to estimate the exposure point concentration if the concentration exceeded the average and was below the maximum detected; otherwise, the maximum detected concentration was used to estimate the exposure point concentration.

---: No information available

95% UCL: 95% upper confidence level of the mean

mg/kg: milligrams per kilogram

mg/L: milligrams per liter

TCDD: tetrachlorodibenzo-p-dioxin

TEC: toxicity equivalence concentration

References:

Jacobs Engineering Group, Inc. (Jacobs), 2002, *Baseline Human Health and Screening Ecological Risk Assessment for the Group 2 Sites (Sites 12, 17, 18/24, 29, 32, 49, Harrison Bayou, and Caddo Lake), Longhorn Army Ammunition Plant, Karnack, Texas*, Final, Oak Ridge, TN, August.

Summary of Chemicals of Potential Concern and Medium-Specific Exposure Point Concentrations:

The table presents the chemicals of potential concern (COPCs) and exposure point concentration (EPC) for each (i.e. the concentration used to estimate the exposure and risk from each COPC). The table includes the range of concentrations detected for each COPC, as well as the frequency of detection (i.e., the number of times the chemical was detected in the samples collected at the site), the EPC, and the statistical measure upon which the EPC was based. The COPCs listed are the ones that were quantitatively evaluated for carcinogenic risk and non-carcinogenic hazard in the Baseline Human Health Risk Assessment (Jacobs, 2002).



Table 2-2. Carcinogenic Toxicity Data Summary

Pathway: Ingestion, Dermal Contact

Chemical of Concern	Oral Cancer Slope Factor (mg/kg-day) ⁻¹	Dermal Cancer Slope Factor (mg/kg-day) ⁻¹	Weight of Evidence/ Carcinogen Guideline Description	Source/ Date
Dioxin/Furans				
2,3,7,8-TCDD TEC	1.50E+05	3.00E+05	Not Classified	---
Explosives				
2,4,6-Trinitrotoluene	3.00E-02	5.00E-02	C	TCEQ, 2001
2,4-Dinitrotoluene	6.80E-01	8.00E-01	B2	TCEQ, 2001
2,6-Dinitrotoluene	6.80E-01	8.00E-01	B2	TCEQ, 2001
2-Amino-4,6-dinitrotoluene	1.00E-02	2.00E-02	Not Classified	---
2-Nitrotoluene	NTV	NTV	Not Classified	---
3-Nitrotoluene	NTV	NTV	Not Classified	---
4-Amino-2,6-dinitrotoluene	1.00E-02	2.00E-02	Not Classified	---
4-Nitrotoluene	NTV	NTV	Not Classified	---
Metals				
Aluminum	NTV	NTV	Not Classified	---
Antimony	NTV	NTV	Not Classified	---
Arsenic	1.50E+00	1.58E+00	A	TCEQ, 2001
Barium	NC	NC	D	TCEQ, 2001
Beryllium	NTV	NTV	B1	TCEQ, 2001
Cadmium (water)	NTV	NTV	B1	TCEQ, 2001
Chromium (total)	NC	NC	Not Classified	---
Lead	NTV	NTV	Not Classified	---
Manganese (non-diet)	NC	NC	D	TCEQ, 2001
Mercury	NC	NC	D	TCEQ, 2001
Nickel	NTV	NTV	A	TCEQ, 2001
Selenium	NC	NC	D	TCEQ, 2001
Silver	NC	NC	D	TCEQ, 2001
Strontium	NTV	NTV	Not Classified	---
Thallium	NC	NC	Not Classified	---
Vanadium	NTV	NTV	Not Classified	---
Non-Metallic Anions				
Perchlorate	NTV	NTV	Not Classified	---
Volatile Organics				
1,2-Dichloroethane	9.10E-02	9.10E-02	B2	TCEQ, 2001
Acetone	NC	NC	D	TCEQ, 2001
Bromodichloromethane	6.20E-02	6.33E-02	B2	TCEQ, 2001
Chloroform	6.10E-03	3.05E-02	B2	TCEQ, 2001
cis-1,2-Dichloroethene	NC	NC	D	TCEQ, 2001
Methylene chloride	7.50E-03	7.89E-03	B2	TCEQ, 2001
p-Cymene	NTV	NTV	Not Classified	---
Trichloroethene	1.10E-02	1.10E-02	B2	TCEQ, 2001



Table 2-2. Carcinogenic Toxicity Data Summary (continued)

Pathway: Inhalation

Chemical of Concern	Unit Risk Factor (mg/m ³) ⁻¹	Weight of Evidence/ Carcinogen Guideline Description	Source/Date
Dioxin/Furans			
2,3,7,8-TCDD TEC	3.30E+04	Not Classified	---
Explosives			
2,4,6-Trinitrotoluene	NTV	C	TCEQ, 2001
2,4-Dinitrotoluene	NTV	B2	TCEQ, 2001
2,6-Dinitrotoluene	NTV	B2	TCEQ, 2001
2-Amino-4,6-dinitrotoluene	NTV	Not Classified	---
2-Nitrotoluene	NTV	Not Classified	---
3-Nitrotoluene	NTV	Not Classified	---
4-Amino-2,6-dinitrotoluene	NTV	Not Classified	---
4-Nitrotoluene	NTV	Not Classified	---
Metals			
Aluminum	NTV	Not Classified	---
Antimony	NTV	Not Classified	---
Arsenic	4.30E+00	A	TCEQ, 2001
Barium	NC	D	TCEQ, 2001
Beryllium	2.40E+00	B1	TCEQ, 2001
Cadmium (water)	1.80E+00	B1	TCEQ, 2001
Chromium (total)	NC	Not Classified	---
Lead	NTV	Not Classified	---
Manganese (Non-diet)	NC	D	TCEQ, 2001
Mercury	NC	D	TCEQ, 2001
Nickel	4.80E-01	A	TCEQ, 2001
Selenium	NC	D	TCEQ, 2001
Silver	NC	D	TCEQ, 2001
Strontium	NTV	Not Classified	---
Thallium	NC	Not Classified	---
Vanadium	NTV	Not Classified	---
Non-Metallic Anions			
Perchlorate	NTV	Not Classified	---
Volatile Organics			
1,2-Dichloroethane	2.60E-02	B2	TCEQ, 2001
Acetone	NC	D	TCEQ, 2001
Bromodichloromethane	NTV	B2	TCEQ, 2001
Chloroform	2.30E-02	B2	TCEQ, 2001
cis-1,2-Dichloroethene	NC	D	TCEQ, 2001
Methylene chloride	4.70E-04	B2	TCEQ, 2001
p-Cymene	NTV	Not Classified	---



Table 2-2. Carcinogenic Toxicity Data Summary (continued)

Pathway: Inhalation

Chemical of Concern	Unit Risk Factor (mg/m ³) ⁻¹	Weight of Evidence/ Carcinogen Guideline Description	Source/Date
Volatile Organics (continued)			
Trichloroethene	1.70E-03	B2	TCEQ, 2001

Notes:

--- : No information available

mg/kg-day: milligrams per kilogram per day

mg/m³: milligrams per cubic meter

NC: Chemical not classified as a carcinogen

NTV: no toxicity value available

TCDD: tetrachlorodibenzo-p-dioxin

TEC: toxicity equivalence concentration

Weight of Evidence/Carcinogen Guideline Description:

A - Human carcinogen

B1 - Probable human carcinogen – Indicates that limited human data are available

B2 - Probable human carcinogen – Indicates sufficient evidence in animals and inadequate or no evidence in humans

C - Possible human carcinogen

D - Not classifiable as a human carcinogen

References:

Jacobs Engineering Group, Inc. (Jacobs), 2002, *Baseline Human Health and Screening Ecological Risk Assessment for the Group 2 Sites (Sites 12, 17, 18/24, 29, 32, 49, Harrison Bayou, and Caddo Lake)*, Longhorn Army Ammunition Plant, Karnack, Texas, Final, Oak Ridge, TN, August.

Texas Commission on Environmental Quality (TCEQ), 2001, Update to 1998 Consistency Memorandum. Toxicity Factors Table, 15 March 2001. Medium specific concentrations have been recalculated using updated toxicity values through March 2010.

Summary of Toxicity Assessment:

The table provides carcinogenic risk information which is relevant to the contaminants of potential concern in soil and ground water. The list of chemicals of concern presented here are the ones that were quantitatively evaluated for carcinogenic risk and non-carcinogenic hazard in the Baseline Human Health Risk Assessment (Jacobs, 2002).



Table 2-3. Non-Carcinogenic Toxicity Data Summary

Pathway: Ingestion, Dermal Contact

Chemical of Concern	Chronic/ Subchronic	Oral RfD Value (mg/kg-day)	Dermal RfD (mg/kg-day)	Target Endpoint	Combined Uncertainty/ Modifying Factors	Source/Date
Dioxin/Furans						
2,3,7,8-TCDD TEC	chronic	NTV	NTV	NA	NA	---
Explosives						
2,4,6-Trinitrotoluene	chronic	5.00E-04	3.00E-04	Liver effects	1000/1	USEPA-IRIS, 2001
2,4-Dinitrotoluene	chronic	2.00E-03	1.70E-03	Central nervous system effects	100/1	USEPA-IRIS, 2001
2,6-Dinitrotoluene	chronic	1.00E-03	8.50E-04	Central nervous system effects	3000/1	USEPA-HEAST, 1997
2-Amino-4,6-dinitrotoluene	chronic	1.67E-04	8.33E-05	NA	NA	---
2-Nitrotoluene	chronic	1.00E-02	5.00E-03	Spleen lesions	10000/1	USEPA-HEAST, 1997
3-Nitrotoluene	chronic	1.00E-02	5.00E-03	Spleen lesions	10000/1	USEPA-HEAST, 1997
4-Amino-2,6-dinitrotoluene	chronic	1.67E-04	8.33E-05	NA	NA	---
4-Nitrotoluene	chronic	1.00E-02	5.00E-03	Spleen lesions	10000/1	USEPA-HEAST, 1997
Metals						
Aluminum	chronic	1.00E+00	1.00E-01	NA	NA	---
Antimony	chronic	4.00E-04	6.00E-05	Longevity, blood glucose, and cholesterol	1000/1	USEPA-IRIS, 2001
Arsenic	chronic	3.00E-04	2.85E-04	Skin effects	3/1	USEPA-IRIS, 2001
Barium	chronic	7.00E-02	4.90E-03	Increased kidney weight	3/1	USEPA-IRIS, 2001
Beryllium	chronic	2.00E-03	1.40E-05	Small Intestine	300/1	USEPA-IRIS, 2001
Cadmium (water)	chronic	5.00E-04	1.25E-05	Proteinuria	10/1	USEPA-IRIS, 2001
Chromium (total)	chronic	1.50E+00	1.95E-02	No effects observed	100/10	USEPA-IRIS, 2001
Lead	chronic	NTV	NTV	NA	NA	---
Manganese (non-diet)	chronic	4.70E-02	2.82E-03	Central nervous system effects	1/1	USEPA-IRIS, 2001
Mercury	chronic	3.00E-04	2.10E-05	Autoimmune effects	1000/1	USEPA-IRIS, 2001
Nickel	chronic	2.00E-02	8.00E-04	Decreased Body Weight	300/1	USEPA-IRIS, 2001
Selenium	chronic	5.00E-03	2.50E-03	Skin	3/1	USEPA-IRIS, 2001
Silver	chronic	5.00E-03	2.00E-04	Argyria	3/1	USEPA-IRIS, 2001
Strontium	chronic	6.00E-01	1.20E-01	Rachitic bone	300/1	USEPA-IRIS, 2001
Thallium	chronic	8.00E-05	8.00E-05	Blood	3000/1	USEPA-IRIS, 2001
Vanadium	chronic	7.00E-03	1.82E-04	NA	NA	---



Table 2-3. Non-Carcinogenic Toxicity Data Summary (continued)

Pathway: Ingestion, Dermal Contact

Chemical of Concern	Chronic/ Subchronic	Oral RfD Value (mg/kg-day)	Dermal RfD (mg/kg-day)	Target Endpoint	Combined Uncertainty/ Modifying Factors	Source/Date
Non-Metallic Anions						
Perchlorate	chronic	9.00E-04	9.00E-04	NA	NA	---
Volatile Organics						
1,2-Dichloroethane	chronic	3.00E-02	3.00E-02	NA	NA	---
Acetone	chronic	1.00E-01	8.30E-02	Liver and kidney effects	1000/1	USEPA-IRIS, 2001
Bromodichloromethane	chronic	2.00E-02	1.96E-02	Renal cytomegaly	1000/1	USEPA-IRIS, 2001
Chloroform	chronic	1.00E-02	2.00E-03	Cyst formation in the liver	1000/1	USEPA-IRIS, 2001
cis-1,2-Dichloroethene	chronic	1.00E-02	1.00E-02	Decreased hematocrit and hemoglobin in the blood	3000/1	USEPA-IRIS, 2001
Methylene chloride	chronic	6.00E-02	5.70E-02	Liver toxicity	100/1	USEPA-IRIS, 2001
p-Cymene	chronic	1.00E-01	8.00E-02	NA	NA	---
Trichloroethene	chronic	6.00E-03	6.00E-03	NA	NA	---

Pathway: Inhalation

Chemical of Concern	Chronic/ Subchronic	Inhalation RfC (mg/m ³)	Target Endpoint	Combined Uncertainty/ Modifying Factors	Source/ Date
Dioxin/Furans					
2,3,7,8-TCDD TEC	chronic	NTV	---	---	---
Explosives					
2,4,6-Trinitrotoluene	chronic	0.0001	NA	NA	---
2,4-Dinitrotoluene	chronic	0.00015	NA	NA	---
2,6-Dinitrotoluene	chronic	0.00015	NA	NA	---
2-Amino-4,6-dinitrotoluene	chronic	0.0001	NA	NA	---
2-Nitrotoluene	chronic	0.011	NA	NA	---
3-Nitrotoluene	chronic	0.011	NA	NA	---
4-Amino-2,6-dinitrotoluene	chronic	0.0001	NA	NA	---
4-Nitrotoluene	chronic	0.011	NA	NA	---
Metals					
Aluminum	chronic	0.0035	NA	NA	---
Antimony	chronic	0.0005	Pulmonary toxicity, chronic interstitial inflammation	300/1	USEPA-IRIS, 2001
Arsenic	chronic	NTV	---	---	---
Barium	chronic	0.00049	Fetus, developmental effects	1000/1	USEPA-HEAST, 1997



Table 2-3. Non-Carcinogenic Toxicity Data Summary (continued)

Pathway: Inhalation

Chemical of Concern	Chronic/ Subchronic	Inhalation RfC (mg/m ³)	Target Endpoint	Combined Uncertainty/ Modifying Factors	Source/ Date
Metals (continued)					
Beryllium	chronic	0.00002	Beryllium sensitization and progression to Chronic Beryllium Disease	10/1	USEPA-IRIS, 2001
Cadmium (water)	chronic	0.0002	NA	NA	---
Chromium (total)	chronic	0.0001	NA	NA	---
Lead	chronic	NTV	---	---	---
Manganese (non-diet)	chronic	0.00005	Impairment of neurobehavioral function	1000/1	USEPA-IRIS, 2001
Mercury	chronic	0.0003	Hand tremor, memory loss	30/1	USEPA-IRIS, 2001
Nickel	chronic	0.0002	Respiratory effects	NA	ATSDR, 1997
Selenium	chronic	0.0002	NA	NA	---
Silver	chronic	0.00001	NA	NA	---
Strontium	chronic	NTV	---	---	---
Thallium	chronic	0.0001	NA	NA	---
Vanadium	chronic	0.00005	NA	NA	---
Non-Metallic Anions					
Perchlorate	chronic	NTV	---	---	---
Volatile Organics					
1,2-Dichloroethane	chronic	0.005	NA	NA	---
Acetone	chronic	0.59	NA	NA	---
Bromodichloromethane	chronic	NTV	---	---	---
Chloroform	chronic	0.000301	NA	NA	---
cis-1,2-Dichloroethene	chronic	0.793	NA	NA	---
Methylene chloride	chronic	3	Liver toxicity	100/1	USEPA-HEAST, 1997
p-Cymene	chronic	0.3	NA	NA	---
Trichloroethene	chronic	NTV	---	---	---

Notes:

---: No information for a compound with no toxicity value (NTV)

IRIS: Integrated Risk Information System, USEPA

mg/kg-day: milligrams per kilogram per day

mg/m³: milligrams per cubic meter

NA: Information not available

NTV: No toxicity value available

RfC: Reference concentration

RfD: Reference dose

TCDD: tetrachlorodibenzo-p-dioxin

TEC: toxicity equivalence concentration

References:

Agency for Toxic Substances and Disease Registry (ATSDR), 1997, Minimal Risk Levels (MRLs) for Hazardous Substances.

Jacobs Engineering Group, Inc. (Jacobs), 2002, *Baseline Human Health and Screening Ecological Risk Assessment for the Group 2 Sites (Sites 12, 17, 18/24, 29, 32, 49, Harrison Bayou, and Caddo Lake)*, Longhorn Army Ammunition Plant, Karnack, Texas, Final, Oak Ridge, TN, August.

USEPA-HEAST, 1997. Health Effects Summary Table (HEAST). FY 1995, Annual Office of Emergency and Remedial Response. Washington, D.C. EPA/340/R-95-036.

USEPA-IRIS, 2001. Integrated Risk Information System (IRIS). United States Environmental Protection Agency Online Database for Toxicity Information on Hazardous Chemicals, 2001.

Summary of Toxicity Assessment:

This table provides non-carcinogenic risk information relevant to the contaminants of concern in both soil and ground water. The list of chemicals of potential concern presented here are the ones that were quantitatively evaluated for carcinogenic risk and non-carcinogenic hazard in the Baseline Human Health Risk Assessment (Jacobs, 2002). The uncertainty factor and modifying factor are used in the development of a reference dose. The uncertainty factor adjusts results from dose-response studies in animals to make them applicable to humans. The modifying factor is used to account for uncertainties in the available toxicity data from which the reference dose is derived. In the risk assessment, the reference doses and concentrations were for the chronic case, to be conservative.



Table 2-4. Risk Characterization Summary – Carcinogens

Scenario Timeframe: Future
 Receptor Population: Maintenance Worker
 Receptor Age: Adult

Medium	Exposure Medium	Exposure Point	Chemical of Concern	Carcinogenic Risk			
				Ingestion	Inhalation	Dermal	Exposure Routes Total
Ground-water	Ground-water	Ingestion or exposure through showering	Dioxin/Furan				
			2,3,7,8-TCDD TEC	6.5E-06	NE	5.4E-05	6.1E-05
			Explosive				
			2,4,6-Trinitrotoluene	ND	ND	ND	NA
			2,4-Dinitrotoluene	1.3E-03	NE	NE (Kp<=0.01)	1.3E-03
			2,6-Dinitrotoluene	1.3E-03	NE	NE (Kp<=0.01)	1.3E-03
			2-Amino-4,6-dinitrotoluene	2.1E-07	NE	NE (Kp<=0.01)	2.1E-07
			2-Nitrotoluene	NTV	NTV	NTV	NA
			3-Nitrotoluene	NTV	NTV	NTV	NA
			4-Amino-2,6-dinitrotoluene	2.1E-07	NE	NE (Kp<=0.01)	2.1E-07
			4-Nitrotoluene	NTV	NTV	NTV	NA
			Metals				
			Aluminum	NTV	NE	NE (Kp<=0.01)	NA
			Antimony	NTV	NE	NE (Kp<=0.01)	NA
			Arsenic	3.1E-04	NE	NE (Kp<=0.01)	3.1E-04
			Barium	NC	NE	NE (Kp<=0.01)	NA
			Beryllium	NTV	NE	NE (Kp<=0.01)	NA
			Cadmium (water)	NTV	NE	NE (Kp<=0.01)	NA
			Chromium (total)	NC	NE	NE (Kp<=0.01)	NA
			Lead	NTV	NE	NE (Kp<=0.01)	NA
			Manganese	NC	NE	NE (Kp<=0.01)	NA
			Mercury	NC	NE	NE (Kp<=0.01)	NA
			Nickel	NTV	NE	NE (Kp<=0.01)	NA
			Selenium	NC	NE	NE (Kp<=0.01)	NA
			Silver	NC	NE	NE (Kp<=0.01)	NA
			Strontium	NTV	NE	NE (Kp<=0.01)	NA
			Thallium	NC	NE	NE (Kp<=0.01)	NA
			Vanadium	NTV	NE	NE (Kp<=0.01)	NA
			Non-Metallic Anion				
			Perchlorate	NTV	NE	NE (Kp<=0.01)	NA
			Volatile Organics				
			1,2-Dichloroethane	4.5E-03	2.2E-02	2.0E-03	2.9E-02
			Acetone	NC	NC	NE (Kp<=0.01)	NA
			Bromodichloro-methane	4.8E-07	NTV	2.5E-07	7.2E-07
			Chloroform	3.0E-07	2.0E-05	1.2E-06	2.1E-05
			Cis-1,2-Dichloroethene	NC	NC	NE (Kp<=0.01)	NA
			Methylene chloride	1.7E-01	1.9E-01	NE (Kp<=0.01)	3.6E-01
			p-Cymene	NTV	NTV	NTV	NA
			Trichloroethene	4.6E-05	1.2E-04	6.2E-05	2.3E-04
Groundwater risk total =							3.9E-01



Table 2-4. Risk Characterization Summary – Carcinogens (continued)

Scenario Timeframe: Future
 Receptor Population: Maintenance Worker
 Receptor Age: Adult

Medium	Exposure Medium	Exposure Point	Chemical of Concern	Carcinogenic Risk			
				Ingestion	Inhalation	Dermal	Exposure Routes Total
Soil (0 to 2 feet)	Soil and particulates	Incidental Ingestion, inhalation of particulates, and dermal contact	Dioxin/Furan				
			2,3,7,8-TCDD TEC	2.2E-07	7.3E-12	8.4E-08	3.0E-07
			Explosive				
			2,4,6-Trinitrotoluene	2.0E-06	NTV	2.1E-06	4.1E-06
			2,4-Dinitrotoluene	1.5E-06	NTV	1.1E-06	2.6E-06
			2,6-Dinitrotoluene	ND	ND	ND	NA
			2-Amino-4,6-dinitrotoluene	8.7E-08	NTV	1.1E-07	2.0E-07
			2-Nitrotoluene	ND	ND	ND	NA
			3-Nitrotoluene	ND	ND	ND	NA
			4-Amino-2,6-dinitrotoluene	5.6E-08	NTV	7.2E-08	1.3E-07
			4-Nitrotoluene	ND	ND	ND	NA
			Metals				
			Aluminum	ND	ND	ND	NA
			Antimony	NTV	NTV	NTV	NA
			Arsenic	ND	ND	ND	NA
			Barium	ND	ND	ND	NA
			Beryllium	ND	ND	ND	NA
			Cadmium (water)	ND	ND	ND	NA
			Chromium (total)	ND	ND	ND	NA
			Lead	ND	ND	ND	NA
			Manganese (non-diet)	ND	ND	ND	NA
Mercury	NC	NC	NC	NA			
Nickel	ND	ND	ND	NA			
Selenium	ND	ND	ND	NA			
Silver	ND	ND	ND	NA			
Strontium	ND	ND	ND	NA			
Thallium	ND	ND	ND	NA			
Vanadium	ND	ND	ND	NA			
Non-Metallic Anion							
Perchlorate	NTV	NTV	NTV	NA			
Volatile Organics							
1,2-Dichloroethane	ND	ND	ND	NA			
Acetone	ND	ND	ND	NA			
Bromodichloro-methane	ND	ND	ND	NA			
Chloroform	ND	ND	ND	NA			



Table 2-4. Risk Characterization Summary – Carcinogens (continued)

Scenario Timeframe: Future
 Receptor Population: Maintenance Worker
 Receptor Age: Adult

Medium	Exposure Medium	Exposure Point	Chemical of Concern	Carcinogenic Risk			
				Ingestion	Inhalation	Dermal	Exposure Routes Total
			Cis-1,2-Dichloroethene	ND	ND	ND	NA
Volatile Organics (continued)							
			Methylene chloride	ND	ND	ND	NA
			p-Cymene	ND	ND	ND	NA
			Trichloroethene	ND	ND	ND	NA
Soil risk total =							7.3E-06
Total risk (soil and groundwater) =							3.9E-01

Notes:

Kp	Dermal permeability coefficient
NA	Not applicable
NC	Not classified as a carcinogen
ND	Not detected in associated media or not selected as a chemical of potential concern
NE	Not evaluated through this exposure pathway. Chemical is not identified as volatile.
NE(Kp<=0.01)	Based on USEPA Region 6 guidance, chemicals of potential concern with a Kp<=0.01 were not evaluated for dermal contact while showering (USEPA, 1995)
NTV	No toxicity value available
TCDD	Tetrachlorodibenzo-p-dioxin
TEC	Toxicity equivalence concentration

References:

U.S. Environmental Protection Agency (USEPA), 1989, *Risk Assessment Guidance for Superfund, Vol. I: Human Health Evaluation Manual, (Part A)*, EPA/540/1-89/002, December.

USEPA, *Supplemental Region VI Risk Assessment Guidance, May 5, 1995.*

Summary of Risk Characterization:

The table provides risk estimates for the significant routes of exposure at LHAAP-29. These risk estimates are based on a reasonable maximum exposure and were developed by taking into account various conservative assumptions about the frequency and duration of a hypothetical future maintenance worker's exposure to soil and groundwater, as well as the toxicity of the chemicals of concern. The total risk from exposure to contaminated soil and groundwater at this site is estimated to be 3.9×10^{-01} . A risk below 1×10^{-4} is generally considered to be acceptable (USEPA, 1989). The total groundwater risk is unacceptable.



Table 2-5. Risk Characterization Summary – Non-Carcinogens

Scenario Timeframe: Future
 Receptor Population: Maintenance Worker
 Receptor Age: Adult

Medium	Exposure Medium	Exposure Point	Chemical of Concern	Target End-point	Non-Carcinogenic Hazard Quotient			
					Ingestion	Inhalation	Dermal	Exposure Routes Total
Ground-water	Ground-water	Ingestion or exposure through showering	Dioxin/Furan					
			2,3,7,8-TCDD TEC	NA	NTV	NE	NTV	NA
			Explosive					
			2,4,6-Trinitrotoluene	Liver effects	ND	ND	ND	NA
			2,4-Dinitrotoluene	NA	2.6E+00	NE	NE (Kp<=0.01)	2.6E+00
			2,6-Dinitrotoluene	NA	5.2E+00	NE	NE (Kp<=0.01)	5.2E+00
			2-Amino-4,6-dinitrotoluene	NA	3.5E-01	NE	NE (Kp<=0.01)	3.5E-01
			2-Nitrotoluene	Spleen lesions	4.3E+00	6.8E+01	4.1E-01	7.3E+01
			3-Nitrotoluene	Spleen lesions	2.3E-01	3.7E+00	2.2E-02	4.0E+00
			4-Amino-2,6-dinitrotoluene	NA	3.5E-01	NE	NE (Kp<=0.01)	3.5E-01
			4-Nitrotoluene	Spleen lesions	2.1E+00	3.3E+01	1.9E-01	3.5E+01
			Metals					
			Aluminum	NA	1.3E+00	NE	NE (Kp<=0.01)	1.3E+00
			Antimony	Longevity, blood glucose, and cholesterol	1.3E+00	NE	NE (Kp<=0.01)	1.3E+00
			Arsenic	Skin effects	1.9E+00	NE	NE (Kp<=0.01)	1.9E+00
			Barium	Fetus, developmental effects, increased kidney weight	9.1E-01	NE	NE (Kp<=0.01)	9.1E-01
			Beryllium	Beryllium sensitization and progression to Chronic Beryllium Disease	4.8E-02	NE	NE (Kp<=0.01)	4.8E-02
			Cadmium(water)	Proteinuria	1.2E-01	NE	NE (Kp<=0.01)	1.2E-01
			Chromium (total)	Proteinuria	5.0E-02	NE	NE (Kp<=0.01)	5.0E-02
			Lead	Gastrointestinal	NTV	NE	NE (Kp<=0.01)	NA



Table 2-5. Risk Characterization Summary – Non-Carcinogens (continued)

Scenario Timeframe: Future
 Receptor Population: Maintenance Worker
 Receptor Age: Adult

Medium	Exposure Medium	Exposure Point	Chemical of Concern	Target End-point	Non-Carcinogenic Hazard Quotient				
					Ingestion	Inhalation	Dermal	Exposure Routes Total	
			Metals (continued)						
			Manganese (non-diet)	CNS	5.0E-01	NE	NE (Kp<=0.01)	5.0E-01	
			Mercury	CNS	9.8E-02	NE	NE (Kp<=0.01)	9.8E-02	
			Nickel	Respiratory effects, decreased body weight	4.1E+00	NE	NE (Kp<=0.01)	4.1E+00	
			Selenium	NA	6.8E-01	NE	NE (Kp<=0.01)	6.8E-01	
			Silver	Argyria	1.6E-01	NE	NE (Kp<=0.01)	1.6E-01	
			Strontium	Rachitic bone	3.1E-01	NE	NE (Kp<=0.01)	3.1E-01	
			Thallium	Blood	3.7E-01	NE	NE (Kp<=0.01)	3.7E-01	
			Vanadium	NA	5.0E-01	NE	NE (Kp<=0.01)	5.0E-01	
			Non-Metallic Anion						
			Perchlorate	---	9.6E+02	NE	NE (Kp<=0.01)	9.6E+02	
			Volatile Organics						
			1,2-Dichloroethane	NA	4.6E+00	4.8E+02	2.1E+00	4.9E+02	
			Acetone	NA	5.7E-04	1.7E-03	NE (Kp<=0.01)	2.3E-03	
			Bromodichloromethane	---	1.1E-03	NTV	5.6E-04	1.6E-03	
			Chloroform	NA	1.4E-02	8.0E+00	5.4E-02	8.0E+00	
			cis-1,2-Dichloroethene	NA	1.3E-03	2.8E-04	NE (Kp<=0.01)	1.6E-03	
			Methylene chloride	Decreased hematocrit and hemoglobin in the blood	1.1E+03	3.8E+02	NE (Kp<=0.01)	1.5E+03	
			p-Cymene	NA	2.8E-04	1.7E-03	2.7E-04	2.2E-03	
			Trichloroethene	Liver and kidney effects	2.0E+00	NTV	2.6E+00	4.6E+00	
			Groundwater Hazard Index Total =						3.0E+03



Table 2-5. Risk Characterization Summary – Non-Carcinogens (continued)

Scenario Timeframe: Future
 Receptor Population: Maintenance Worker
 Receptor Age: Adult

Medium	Exposure Medium	Exposure Point	Chemical of Concern	Target Endpoint	Non-Carcinogenic Hazard Quotient			
					Ingestion	Inhalation	Dermal	Exposure Routes Total
Soil (0 to 2 feet)	Soil and particulates	Incidental ingestion, inhalation of particulates, dermal contact	Dioxin/Furan					
			2,3,7,8-TCDD TEC	NA	NTV	NTV	NTV	NA
			Explosive					
			2,4,6-Trinitrotoluene	Liver effects	3.7E-01	2.8E-04	4.0E-01	7.7E-01
			2,4-Dinitrotoluene	IA	3.0E-03	6.1E-06	2.3E-03	5.3E-03
			2,6-Dinitrotoluene	IA	ND	ND	ND	NA
			2-Amino-4,6-dinitrotoluene	NA	1.5E-01	3.7E-05	1.9E-01	3.3E-01
			2-Nitrotoluene	Spleen lesions	ND	ND	ND	NA
			3-Nitrotoluene	Spleen lesions	ND	ND	ND	NA
			4-Amino-2,6-dinitrotoluene	NA	9.4E-02	2.4E-05	1.2E-01	2.1E-01
			4-Nitrotoluene	Spleen lesions	ND	ND	ND	NA
			Metals					
			Aluminum	NA	ND	ND	ND	NA
			Antimony	longevity, blood glucose, and cholesterol	4.7E-03	5.7E-07	2.0E-03	6.7E-03
Arsenic	skin effects	ND	ND	ND	NA			
Barium	retus, developmental effects, increased kidney weight	ND	ND	ND	NA			
Beryllium	beryllium sensitization and progression to chronic beryllium disease	ND	ND	ND	NA			
Cadmium (water)	proteinuria	ND	ND	ND	NA			
Chromium (total)	proteinuria	ND	ND	ND	NA			
Lead	gastrointestinal	ND	ND	ND	NA			
Manganese (non-diet)	CNS	ND	ND	ND	NA			
Mercury	CNS	7.2E-04	1.1E-07	6.6E-04	1.4E-03			
Nickel	Respiratory effects, decreased body weight	ND	ND	ND	NA			
Selenium	NA	ND	ND	ND	NA			



Table 2-5. Risk Characterization Summary – Non-Carcinogens (continued)

Scenario Timeframe: Future
Receptor Population: Maintenance Worker
Receptor Age: Adult

Medium	Exposure Medium	Exposure Point	Chemical of Concern	Target Endpoint	Non-Carcinogenic Hazard Quotient			
					Ingestion	Inhalation	Dermal	Exposure Routes Total
			Metals (continued)					
			Silver	Argyria	ND	ND	ND	NA
			Strontium	Rachitic bone	ND	ND	ND	NA
			Thallium	Blood	ND	ND	ND	NA
			Vanadium	NA	ND	ND	ND	NA
			Non-Metallic Anion					
			Perchlorate	---	7.6E-05	NTV	4.9E-06	8.1E-05
			Volatile Organics					
			1,2-Dichloroethane	NA	ND	ND	ND	NA
			Acetone	NA	ND	ND	ND	NA
			Bromodichloromethane	---	ND	ND	ND	NA
			Chloroform	NA	ND	ND	ND	NA
			cis-1,2-Dichloroethene	NA	ND	ND	ND	NA
			Methylene chloride	Decreased hematocrit and hemoglobin in the blood	ND	ND	ND	NA
			p-Cymene	NA	ND	ND	ND	NA
			Trichloroethene	Liver and kidney effects	ND	ND	ND	NA
Soil Hazard Index Total =								1.3E+00
Hazard Index Total (soil and groundwater) =								3.0E+03

Notes:

CNS central nervous system
Kp Dermal permeability coefficient
NA Not applicable
ND Not detected in associated media or not selected as a chemical of potential concern
NE Not evaluated through this exposure pathway. Chemical is not identified as a volatile.
NE (Kp<=0.01) Based on USEPA Region 6 guidance, chemicals of potential concern with a Kp<=0.01 were not evaluated for dermal contact while showering (USEPA, 1995)
NTV No toxicity value
TCDD Tetrachlorodibenzo-p-dioxin
TEC Toxicity equivalence concentration

References:

U.S. Environmental Protection Agency (USEPA), 1989, *Risk Assessment Guidance for Superfund, Vol. I: Human Health Evaluation Manual, (Part A)*, EPA/540/1-89/002, December.

Summary of Risk Characterization:

The table provides hazard quotients (HQs) for each route of exposure and the hazard index (sum of hazard quotients) for all routes of exposure for LHAAP-29. The Risk Assessment Guidance for Superfund (USEPA, 1989) states that, generally, a hazard index (HI) greater than 1 indicates the potential for adverse non-carcinogenic effects. The estimated HI for groundwater is 3.0E+03 and for soil is 1. Both values are unacceptable and indicate that the potential for adverse non-carcinogenic effects could occur from exposure to contaminants in those mediums.



Table 2-6. Chemicals Contributing to Carcinogenic Risk in Groundwater

Chemical	Baseline Risk Assessment			Data Since Risk Assessment			Comparison Levels		Retained as COC ?
	Cancer Risk Groundwater ^a	EPC (µg/L)	Well	Maximum (µg/L)	Well	Adjusted Risk	MCL (µg/L)	TRRP Tier 1 Residential Groundwater PCLs (µg/L)	
Methylene chloride	3.6×10^{-1}	6,600,000	29WW16	10,300,000	29WW16	5.6×10^{-1}	5	5	Yes, 1
1,2-Dichloroethane	2.9×10^{-2}	14,000	29WW15	<12,500	29WW16	--	5	5	Yes, 1
2,4-Dinitrotoluene	1.3×10^{-3}	530	29WW20	50.9 32.4	29WW05 29WW20	1.2×10^{-4}	--	1.3	Yes, 2
2,6-Dinitrotoluene	1.3×10^{-3}	530	29WW20	239 112	116 29WW20	5.9×10^{-4}	--	1.3	Yes, 2
Arsenic	3.1×10^{-4}	59	29WW25	141	29WW25	7.4×10^{-4}	10	10	Yes, 5
Trichloroethene	2.3×10^{-4}	1,200	29WW15	<12,500	29WW16	--	5	5	Yes, 1
2,3,7,8-TCDD	6.1×10^{-5}	1.25×10^{-5}	29WW03	NR	--	--	3.0×10^{-5}	--	No, 4
Chloroform	2.1×10^{-5}	14	29WW21	9.75 ND	29WW15 29WW21	1.5×10^{-5}	80 ^b	1,000	No, 4

Notes and Abbreviations:

1. Identified as COC because most recent maximum concentration is above the Safe Drinking Water Act (SDWA) MCL.
2. Identified as COC because carcinogenic risk is $>10^{-4}$.
3. Excluded because detections are isolated.
4. Excluded because EPC is below the SDWA MCL.
5. Identified as a COC subject to further verification.

^a From Baseline Risk Assessment Table C-71 (Jacobs, 2002)

^b SDWA MCL for total trihalomethanes was used for chloroform.

µg/L	micrograms per liter
COC	contaminant of concern
EPC	exposure point concentration
MCL	Safe Drinking Water Act maximum contaminant level
NR	not resampled for this constituent since Baseline Risk Assessment



Table 2-7. Chemicals with Hazard Quotient Greater than 0.1 in Groundwater

Chemical	Baseline Risk Assessment			Data Since Risk Assessment			Comparison Levels		Retained as COC ?
	Hazard Quotient Groundwater ^a	EPC ^a (µg/L)	Well	Maximum (µg/L)	Well	Adjusted Hazard Quotient	MCL (µg/L)	TRRP Tier 1 Residential Groundwater PCLs (µg/L)	
Methylene chloride	1,500	6,600,000	29WW16	7,110,000	29WW16	1600	5		Yes, 1
Perchlorate	960	88,000	29WW15	16,800	29WW15	180		17	Yes, 2
1,2-Dichloroethane	490	14,000	29WW15	5,520	29WW15	190	5		Yes, 1
4-Nitrotoluene (p-)	35	2,100	29WW20	1,400 374	116 29WW20	23		57	Yes, 2
Chloroform	8.0	14	29WW21	9.75 ND	29WW15 29WW21	5.6	80 ^b		No, 3
2-Nitrotoluene (o-)	7.3	4,400	116	8,140	116	14		4.1	Yes, 2
2,6-Dinitrotoluene	5.2	530	29WW20	239 112	116 29WW20	2.3		1.3	Yes, 2
Trichloroethene	4.6	1,200	29WW15	344	29WW15	1.3	5		Yes, 1
Nickel	4.1	8,400	29WW11	3,190 40	29WW07 29WW11	1.6 <0.1		490	Yes, 9
3-Nitrotoluene (m-)	4.0	240	29WW05	451 123	116 29WW05	7.5		240	Yes, 2
2,4-Dinitrotoluene	2.6	530	29WW20	50.9 32.4	29WW05 29WW20	0.33		1.3	Yes, 5
Arsenic	1.9	59	29WW25	141	29WW25	4.5	10		Yes, 9
Aluminum	1.3	130,000	115	713	29WW08 ^c	<0.1		24,000	No, 6
Antimony	1.3	52	29WW09	1.45	29WW08	<0.1	6		No, 7
Barium	0.91	6,500	116	1,100 48.5 J	115 116	0.15 <0.1	2,000		No, 6
Selenium	0.68	350	118	75.3	118	0.15	50		Yes, 9
Manganese	0.50	2,410	115	1,310	114 ^c	0.27		1,100	No, 4
Vanadium	0.50	360	115	7.5 J	29WW04 ^c	<0.1		44	No, 8
Thallium	0.37	3.0	29WW03	0.339 J	29WW25 ^c	<0.1	2		No, 7



Table 2-7. Chemicals with Hazard Quotient Greater than 0.1 in Groundwater

Chemical	Baseline Risk Assessment			Data Since Risk Assessment			Comparison Levels		Retained as COC ?
	Hazard Quotient Groundwater ^a	EPC ^a (µg/L)	Well	Maximum (µg/L)	Well	Adjusted Hazard Quotient	MCL (µg/L)	TRRP Tier 1 Residential Groundwater PCLs (µg/L)	
2-Amino-4,6-dinitrotoluene	0.35	5.9	29WW05	ND	29WW05	-		4.1	No, 8
4-Amino-2,6-dinitrotoluene	0.35	5.9	29WW05	16.3	29WW05	0.97		4.1	Yes, 1
Strontium	0.31	19,000	119	NR	-	-		15,000	Yes, 1
Silver	0.16	80	29WW09	ND	All wells resampled ^c	-		120	No, 8
Cadmium	0.12	6.23	119	1.2 1.12	115 116	< 0.1 < 0.1	5		No.6

Notes and Abbreviations:

1. Identified as COC because EPC is above the Safe Drinking Water Act (SDWA) MCL or TRRP Tier 1 Residential Groundwater PCL
2. Identified as COC because HQ is > 1.0
3. Excluded because EPC is below the SDWA MCL
4. Excluded because EPC is below the 95% UTL value for Manganese of 7,820 µg/L from *Final Evaluation of Perimeter Well Data for Use as Groundwater Background (Shaw, 2007)* and HQ is <1.0.
5. Already identified as a COC due to carcinogenic risk (**Table 2-2**)
6. More recent sample results indicate lower concentrations of chemical, reducing HQ to <1.0
7. More recent sample results indicate lower concentrations of chemical below the MCL
8. Excluded because EPC and/or most recent maximum is below the TRRP Tier 1 Residential Groundwater PCL and HQ is <1.0
9. Identified as a COC subject to further verification.

^a From Baseline Risk Assessment Table C-68 (Jacobs, 2002)

^b SDWA MCL for total trihalomethanes was used for chloroform

^c Well with maximum in Baseline Risk Assessment was dry in most recent sampling event and the identified well has the most recent maximum

µg/L	micrograms per liter
COC	contaminant of concern
EPC	exposure point concentration
HQ	hazard quotient
J	estimated; the analyte was positively identified; the concentration is estimated
MCL	Safe Drinking Water Act maximum contaminant level

ND not detected in associated media or not selected as a chemical of potential concern

NR chemical not resampled in most recent sampling event



Table 2-8. Chemicals with Hazard Quotient Greater than 0.1 in Soil

Chemical	Baseline Risk Assessment			Data Through 2008			Retained as COC ?
	Soil Hazard Quotient ^a	EPC (mg/kg)	Soil Sample Location	Adjusted Hazard Quotient ^b	Maximum (mg/kg)	Soil Sample Location	
2,4,6-Trinitrotoluene	0.77	190	29SD13	105	26,000	29SD46	Yes, 2
2-Amino-4,6-dinitrotoluene	0.33	25	29SD13	0.63	48	29SD46	No, 1
4-Amino-2,6-dinitrotoluene	0.21	16	29SD13	0.21	16	29SD13	No, 1
2,4-Dinitrotoluene	0.0053	6.2	29SB15	6.8	8,000	29SD46	Yes, 3
Perchlorate ^c	8.1×10^{-5}	0.0703	Max from BHHRA Table 3-66	0.0099	8.6	29SB86	Yes, 4

Notes and Abbreviations:

1. Not identified as contaminant of concern (COC) because HQ is less than 1.0.
2. Identified as COC because risk assessment HQ is almost 1 and most recent sample concentration is greater than the SAI-Ind GWP-Ind.
3. Identified as COC because EPC is above the SAI-Ind and GWP-Ind values and Hazard Quotient is greater than 1.0.
4. Identified as COC because contaminant is COC in groundwater and exceeds the GWP-Ind.

^a HQ from Baseline Risk Assessment Table C-68 (Jacobs, 2002)

^b calculated HQ based on the most recent maximum concentration.

^c Even though HI <0.1, listed because recent maximum concentration is greater than EPC

BHHRA	Baseline Human Health Risk Assessment
EPC	Exposure Point Concentration from Baseline Risk Assessment (Jacobs, 2002)
GWP-Ind	Soil medium-specific concentration for industrial use based on groundwater protection
HQ	hazard quotient
mg/kg	milligrams per kilogram.
SAI-Ind	Soil medium-specific concentration for industrial use based on inhalation, ingestion, and dermal contact



Table 2-9. Chemicals in Soil Compared to EcoPRGs

Chemical	SS EcoPRG ^a (mg/kg)	TS EcoPRG ^a (mg/kg)	Maximum ^b (mg/kg)	Retained as Contaminant of Potential Ecological Concern?
2,4,6-Trinitrotoluene	6.1	4.7	26,000	Yes
2,4-Dinitrotoluene	—	12	8,000	Yes
2,6-Dinitrotoluene	2.7	6.8	15	Yes

Notes and Abbreviations:

^a From Baseline Ecological Risk Assessment Table 16-1 (Shaw, 2007b).

^b Maximum soil concentrations from samples collected in the upper 3 feet of soil at 29SD46 collected (Shaw, 2007a)

EcoPRG Ecological Preliminary Cleanup Level

mg/kg milligrams per kilogram.

SS surface soil from 0-0.5 feet (applicable to deer mouse)

TS total soil from 0-3 feet (applicable to short-tailed shrew)



Table 2-10. Cleanup Levels at LHAAP-29

Medium	Chemical of Concern	Cleanup Level
Shallow Zone Groundwater		MCL (µg/L)
	Trichloroethene	5
	1,2-Dichloroethane	5
	1,1-Dichloroethene*	7
	cis-1,2-Dichloroethene*	70
	trans-1,2-Dichloroethene*	100
	Vinyl chloride*	2
	Arsenic	10
	Mercury	2
	Selenium	50
		TRRP Tier 1 Residential Groundwater PCL - (µg/L)
	2,4-Dinitrotoluene	1.3
	2,6-Dinitrotoluene	1.3
	2-Nitrotoluene	4.1
	3-Nitrotoluene	240
	4-Nitrotoluene	57
	Perchlorate	17
	Nickel	490
Intermediate Zone Groundwater		MCL (µg/L)
	Methylene chloride	5
	Trichloroethene	5
	1,2-Dichloroethane	5
	1,1-Dichloroethene*	7
	cis-1,2-Dichloroethene*	70
	trans-1,2-Dichloroethene*	100
	Vinyl chloride*	2
	Arsenic	10
Soil		GWP-Ind (mg/kg)
	2,4,6-Trinitrotoluene	4.7 ^a 5.1 ^b
	2,4-Dinitrotoluene	0.042
	2,6-Dinitrotoluene	0.042
	Perchlorate**	7.2



Table 2-10. Cleanup Levels at LHAAP-29 (continued)

Medium	Chemical of Concern	Cleanup Level
Transite TNT Wastewater Line		GWP-Ind (mg/kg)
Solid Residue	1,3-Dinitrobenzene	1
	2,4,6-Trinitrotoluene	5.1
	2,4-Dinitrotoluene	0.042
	2-amino-4,6-Dinitrotoluene	1.7
	4-amino-2,6-Dinitrotoluene	1.7
Cooling Water Drain Line		GWP-Ind (mg/kg)
Solid Residue	2,4,6-Trinitrotoluene	5.1
	2,4-Dinitrotoluene	0.042
	2,6-Dinitrotoluene	0.042
	2-amino-4,6-Dinitrotoluene	1.7
	4-amino-2,6-Dinitrotoluene	1.7

Notes:

* Trichloroethene daughter products

** Potential COC in soil due to high perchlorate concentration in groundwater

^a applies to 0-3 feet below ground surface

^b applies from 3 feet below ground surface to groundwater interface

GWP-Ind Texas Commission on Environmental Quality soil medium specific concentration for industrial use based on groundwater protection

µg/L micrograms per liter

MCL Safe Drinking Water Act (SDWA) maximum contaminant level

mg/kg milligrams per kilogram

MSC medium-specific concentration



Table 2-11. Comparative Analysis of Alternatives

Criteria	Alternative 1 No Action	Alternative 2 Excavation and off-site disposal for soil; plug lines; in situ chemical oxidation, MNA and LUCs for intermediate zone groundwater; and MNA and LUCs for shallow zone groundwater	Alternative 3 Excavation and off-site disposal for soil; plug lines; groundwater extraction, MNA and LUCs for groundwater	Alternative 4a Excavation and Offsite Disposal for Soil; Plug Lines; Electrical Resistance Heating (ERH) , MNA and LUCs for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater	Alternative 4b Excavation and Offsite Disposal for Soil; Plug Lines; Thermal Conduction Heating (TCH) and MNA for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater
Overall protection of human health and the environment	No protection. Does not achieve RAOs.	Achieves RAOs. Protection of human health and environment provided by soil removal and remediation of groundwater COCs to cleanup levels	Achieves RAOs. Protection of human health and environment provided by soil removal and remediation of groundwater COCs to cleanup levels.	Achieves RAOs in shallow soil/sediment, shallow zone groundwater and intermediate zone groundwater. LUCs can be removed upon completion of active treatment and post-treatment MNA.	Achieves RAOs in shallow soil/sediment, shallow zone groundwater and intermediate zone groundwater. LUCs can be removed upon completion of active treatment and post-treatment MNA.
Compliance with ARARs	No compliance with chemical-specific ARARs.	Complies with all ARARs.	Complies with all ARARs.	Complies with all ARARs.	Complies with all ARARs.



Criteria	Alternative 1 No Action	Alternative 2 Excavation and off-site disposal for soil; plug lines; in situ chemical oxidation, MNA and LUCs for intermediate zone groundwater; and MNA and LUCs for shallow zone groundwater	Alternative 3 Excavation and off-site disposal for soil; plug lines; groundwater extraction, MNA and LUCs for groundwater	Alternative 4a Excavation and Offsite Disposal for Soil; Plug Lines; Electrical Resistance Heating (ERH) , MNA and LUCs for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater	Alternative 4b Excavation and Offsite Disposal for Soil; Plug Lines; Thermal Conduction Heating (TCH) and MNA for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater
Long-term effectiveness and permanence	Is not effective at protection of human health and the environment and does not provide permanence.	<p>Soil removal is a permanent remedy for shallow soil. Excavation of soil is effective long-term and permanent as contamination would be removed from the site and placed in a permitted landfill. In situ chem-ox for intermediate zone should be effective and permanent; however, uncertainty exists concerning the effectiveness of in situ treatment for reducing groundwater contaminant concentrations to cleanup levels. Treatability and pilot studies would be required to further assess the effectiveness of this treatment method and a pre-design would be required to determine the optimum extraction technique configuration. Evaluation of natural attenuation suggests that contaminants are degrading naturally. MNA sampling would be conducted to confirm its effectiveness. Flushing and plugging the process lines would effectively remove any remaining contaminants and there would be no remaining potential for release to surrounding soils. Land use controls would be effective and reliable so long as they are maintained.</p>	<p>Soil removal is a permanent remedy for shallow soil. Excavation of soil is effective long-term and permanent as contamination would be removed from the site and placed in a permitted landfill. Groundwater extraction should be effective and permanent for intermediate zone, based on the efficiency exhibited by the current groundwater treatment system. A pre-design study would be required to determine the optimum extraction technique/configuration. Evaluation of natural attenuation suggests that contaminants are degrading naturally. MNA sampling would be conducted to confirm its effectiveness. Flushing and plugging the process lines would effectively remove any remaining contaminants and there would be no remaining potential for release to surrounding soils. Land use controls would be effective and reliable so long as they are maintained.</p>	<p>Soil removal is a permanent remedy for shallow soil. Excavation of soil is effective long-term and permanent as contamination would be removed from the site and placed in a permitted landfill. MNA has been demonstrated to be effective in managing residual contamination in shallow zone groundwater. ERH is expected to be effective on DNAPL in intermediate zone, MNA demonstrated to be effective in reducing concentrations in intermediate zone groundwater outside of DNAPL zone. Flushing and plugging the process lines would effectively remove any remaining contaminants and there would be no remaining potential for release to surrounding soils. LUCs would remain in place until MNA is completed.</p>	<p>Soil removal is a permanent remedy for shallow soil. Excavation of soil is effective long-term and permanent as contamination would be removed from the site and placed in a permitted landfill. MNA has been demonstrated to be effective in managing residual contamination in shallow zone groundwater. ERH is expected to be effective on DNAPL in intermediate zone, MNA demonstrated to be effective in reducing concentrations in intermediate zone groundwater outside of DNAPL zone. Flushing and plugging the process lines would effectively remove any remaining contaminants and there would be no remaining potential for release to surrounding soils. LUCs would remain in place until MNA is completed.</p>



Table 2-11. Comparative Analysis of Alternatives (continued)

Criteria	Alternative 1 No Action	Alternative 2 Excavation and off-site disposal for soil; plug lines; in situ chemical oxidation, MNA and LUCs for intermediate zone groundwater; and MNA and LUCs for shallow zone groundwater	Alternative 3 Excavation and off-site disposal for soil; plug lines; groundwater extraction, MNA and LUCs for groundwater	Alternative 4a Excavation and Offsite Disposal for Soil; Plug Lines; Electrical Resistance Heating (ERH) , MNA and LUCs for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater	Alternative 4b Excavation and Offsite Disposal for Soil; Plug Lines; Thermal Conduction Heating (TCH) and MNA for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater
Reduction of toxicity, mobility, or volume through treatment	No reduction.	Provides permanent and irreversible reduction of intermediate zone. Provides active reduction of toxicity and volume of groundwater contaminants through biological degradation component of MNA.	Extraction and treatment of contaminated groundwater intermediate zone reduces toxicity, mobility, and volume of groundwater contaminants in this area outside of natural processes. Provides active reduction of toxicity and volume of groundwater contaminants through biological degradation component of MNA.	Reduced toxicity, mobility, and volume through excavation, in situ hydrolysis of DNAPL and DNAPL removal followed by ex-situ destruction, and MNA.	Reduced toxicity, mobility, and volume through excavation, in situ hydrolysis of DNAPL and DNAPL removal followed by ex-situ destruction, and MNA.
Short-term effectiveness	No short-term impacts.	Greater potential for impacts to the community or hypothetical future maintenance worker through off-site transportation of contaminated soil. Release to environment can be controlled during construction.	Greater potential for impacts to the community or hypothetical future maintenance worker through off-site transportation of contaminated soil. Release to environment can be controlled during construction.	Greater potential for impacts to the community or LHAAP workers through off-site transportation of excavated soil. Release to environment can be controlled during construction. Potential for impacts to workers from exposure to hot fluids and high voltage power during ERH application. Duration of MNA and LUCs expected to be 5-10 years following active remediation.	Greater potential for impacts to the community or LHAAP workers through off-site transportation of excavated soil. Release to environment can be controlled during construction. Potential for impacts to workers from exposure to hot fluids during TCH application. Duration of MNA and LUCs expected to be 5-10 years following active remediation.
Implementability	Inherently implementable.	Implementable, but uncertainty exists whether in situ chemical oxidation would lower contaminant concentrations to cleanup levels. Specialized knowledge required for implementation. Use of on-site storage tanks may limit storage capacity. A groundwater treatment system is already operating at LHAAP.	Implementable. Use of on-site storage tanks may limit storage capacity. A groundwater treatment system is already operating at LHAAP. Potential exists for limited groundwater recovery which may affect ability of system to remove contaminants to cleanup levels. A pre-design study would be required.	Soil excavation readily implemented with standard Earthmoving equipment. ERH has been proven to be effective on DNAPL and within low hydraulic conductivity zones.	Soil excavation readily implemented with standard Earthmoving equipment. TCH has been proven to be effective on DNAPL and within low hydraulic conductivity zones.
Cost* (present worth)					



Table 2-11. Comparative Analysis of Alternatives (continued)

Criteria	Alternative 1 No Action	Alternative 2 Excavation and off-site disposal for soil; plug lines; in situ chemical oxidation, MNA and LUCs for intermediate zone groundwater; and MNA and LUCs for shallow zone groundwater	Alternative 3 Excavation and off-site disposal for soil; plug lines; groundwater extraction, MNA and LUCs for groundwater	Alternative 4a Excavation and Offsite Disposal for Soil; Plug Lines; Electrical Resistance Heating (ERH) , MNA and LUCs for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater	Alternative 4b Excavation and Offsite Disposal for Soil; Plug Lines; Thermal Conduction Heating (TCH) and MNA for Intermediate Zone Groundwater; MNA and LUCs for Shallow Zone Groundwater
Capital	\$0	\$2,109,000	\$1,360,000	\$3,710,000	\$4,530,000
O&M	\$0	\$919,000	\$1,558,000	1,030,000	1,190,000
Total	\$0	\$3,028,000	\$2,918,000	\$4,740,000	\$5,720,000
State Acceptance	This criterion will be evaluated in the Proposed Plan after state agency comments are provided.				
Community Acceptance	This criterion will be evaluated in the Proposed Plan after community comments are provided.				

Notes and Abbreviations:

* Costs have been rounded to nearest \$1,000

ARAR	applicable or relevant and appropriate requirement
COC	contaminant of concern
LHAAP	Longhorn Army Ammunition Plant
LUC	land use controls
MNA	monitored natural attenuation
O&M	operation and maintenance
RAO	remedial action objective



Table 2-12. Remediation Cost Table Selected Remedy (Alternative 4a) Present Worth Analysis

Year	Capital Costs		Excavation & Pipeline Flushing	Operation and Maintenance Costs			Present Value (NPV)		
	Design, Permitting, Construction Management, Sub G&A	Electrical Resistance Heating (ERH)		MNA/LUC	O&M GWTP (condensate treatment)	Total O&M	Discount Rate 2.8%	Capital	O&M
Estimate Base Year	2010 (FS)	2014	2010 (FS)	2010 (FS)	2010 (FS)				
Escalation Factor *	1.14	1.02	1.14	1.14	1.14				
1	814,530	1,443,282	670,076	252,638	23,005	275,643	NPV	3,710,000	1,030,000
2		799,659		127,777		127,777			
3		0		66,717		66,717		Total NPV	\$4,740,000
4		0		58,711		58,711			
5		0		115,041		115,041			
6		0		40,147		40,147			
7		0		40,147		40,147			
8		0		40,147		40,147			
9		0		40,147		40,147			
10		0		88,471		88,471			
11		0				0			
12		0				0			
13		0				0			
14		0				0			
15		0		89,078					
16		0				0			
17		0				0			
18		0				0			
19		0				0			
20		0		89,078					
21		0				0			
22		0				0			
23		0				0			



Table 2-12. Remediation Cost Table Selected Remedy (Alternative 4a) Present Worth Analysis (continued)

Year	Capital Costs		Excavation & Pipeline Flushing	Operation and Maintenance Costs			Present Value (NPV)		
	Design, Permitting, Construction Management, Sub G&A	Electrical Resistance Heating (ERH)		MNA/LUC	O&M GWTP (condensate treatment)	Total O&M	Discount Rate 2.8%	Capital	O&M
24		0				0			
25		0		89,078					
26		0				0			
27		0				0			
28		0				0			
29		0				0			
30		0		89,078					
	814,530	2,242,941	670,076	1,225,755	23,005	1,248,761			

*Escalation based on construction cost index published by RS Means (<https://www.rsmeansonline.com/references/unit/refpdf/hci.pdf>)

Notes:

LUC	land use control
MNA	monitored natural attenuation
NPV	net present value
O&M	operation & maintenance
VOC	volatile organic compound

Major assumptions are as described below. Quantities and assumptions are for cost estimating purposes only.

Capital costs include: excavation evaluation, excavation and disposal activities, flow tests, engineering support, and construction management. The soil is assumed to be classified as nonhazardous for disposal purposes.

Monitoring costs are based on the assumption that sampling is conducted at five shallow zone wells and three intermediate zone wells, with one quality control sample in each zone. In the shallow zone, monitoring begins 6 months into Year 2 when groundwater extraction ends and MNA begins. The sampling frequency is quarterly for 2 years, then semiannually for 3 years, then annually for Years 7 through 10, and finally every 5 years (Years 15, 20, 25, and 30). Analysis of the shallow zone groundwater is for VOCs and perchlorate. In the intermediate zone, monitoring begins at the start of Year 1 when MNA begins. The sampling frequency is quarterly for 2 years (Years 1 and 2), then semiannually for 3 years (Years 3 through 5), then annually for Years 6 through 10, and finally every 5 years (Years 15, 20, 25, and 30). Analysis of the intermediate zone groundwater is for VOCs.

The discount rate of 2.8% is based on the Office of Management and Budget Circular No. A-94, January 2008.

The information in this cost estimate summary table is based on the best available information regarding the anticipated scope of the remedial alternative. Changes in the cost elements are likely to occur as a result of new information and data collected during the engineering design of the remedial alternative. Changes will be documented in accordance with 40 CFR 300.435(c)(2) in the form of a memorandum in the Administrative Record, an Explanation of Significant Difference (ESD), or a ROD amendment, as necessary. This is an order-of-magnitude engineering cost estimate that is expected to be within +50 to -30 percent of the actual project cost.



Table 2-13. Remediation Cost Table Selected Remedy (Alternative 4b) Present Worth Analysis

Year	Capital Costs		Excavation & Pipeline Flushing	Operation and Maintenance Costs			Present Value (NPV)		
	Design, Permitting, Construction Management, Sub G&A	Thermal Conduction Heating (TCH)		MNA/LUC	O&M GWTP (condensate treatment)	Total O&M	Discount Rate 2.8%	Capital	O&M
Estimate Base Year	2010 (FS)	2014	2010 (FS)	2010 (FS)	2010 (FS)				
Escalation Factor *	1.14	1.02	1.14	1.14	1.14				
1	997,004	1,889,115	670,076	250,927	175,509	470,436	NPV	4,530,000	1,190,000
2		1,003,601		126,415		126,415			
3		0		66,266		66,266		Total NPV	\$5,720,000
4		0		66,266		66,266			
5		0		114,262		114,262			
6		0		39,875		39,875			
7		0		39,875		39,875			
8		0		39,875		39,875			
9		0		39,875		39,875			
10		0		87,871		88,871			
11		0				0			
12		0				0			
13		0				0			
14		0				0			
15		0		88,475		88,475			
16		0				0			
17		0				0			
18		0				0			
19		0				0			
20		0		88,475		88,475			
21		0				0			
22		0				0			
23		0				0			
24		0				0			



Table 2-13. Remediation Cost Table Selected Remedy (Alternative 4b) Present Worth Analysis (continued)

Year	Capital Costs		Excavation & Pipeline Flushing	Operation and Maintenance Costs			Present Value (NPV)		
	Design, Permitting, Construction Management, Sub G&A	Thermal Conduction Heating (TCH)		MNA/LUC	O&M GWTP (condensate treatment)	Total O&M	Discount Rate 2.8%	Capital	O&M
25		0		88,475		88,475			
26		0				0			
27		0				0			
28		0				0			
29		0				0			
30		0		88,475		88,475			
	997,004	2,892,716	670,076	1,225,406	175,509	1,404,915			

*Escalation based on construction cost index published by RS Means (<https://www.rsmeansonline.com/references/unit/refpdf/hci.pdf>)

Notes:

LUC	land use control
MNA	monitored natural attenuation
NPV	net present value
O&M	operation & maintenance
VOC	volatile organic compound

Major assumptions are as described below. Quantities and assumptions are for cost estimating purposes only.

Capital costs include: excavation evaluation, excavation and disposal activities, flow tests, engineering support, and construction management. The soil is assumed to be classified as nonhazardous for disposal purposes.

Monitoring costs are based on the assumption that sampling is conducted at five shallow zone wells and three intermediate zone wells, with one quality control sample in each zone. In the shallow zone, monitoring begins 6 months into Year 2 when groundwater extraction ends and MNA begins. The sampling frequency is quarterly for 2 years, then semiannually for 3 years, then annually for Years 7 through 10, and finally every 5 years (Years 15, 20, 25, and 30). Analysis of the shallow zone groundwater is for VOCs and perchlorate. In the intermediate zone, monitoring begins at the start of Year 1 when MNA begins. The sampling frequency is quarterly for 2 years (Years 1 and 2), then semiannually for 3 years (Years 3 through 5), then annually for Years 6 through 10, and finally every 5 years (Years 15, 20, 25, and 30). Analysis of the intermediate zone groundwater is for VOCs.

The discount rate of 2.8% is based on the Office of Management and Budget Circular No. A-94, January 2008.

The information in this cost estimate summary table is based on the best available information regarding the anticipated scope of the remedial alternative. Changes in the cost elements are likely to occur as a result of new information and data collected during the engineering design of the remedial alternative. Changes will be documented in accordance with 40 CFR 300.435(c)(2) in the form of a memorandum in the Administrative Record, an Explanation of Significant Difference (ESD), or a ROD amendment, as necessary. This is an order-of-magnitude engineering cost estimate that is expected to be within +50 to -30 percent of the actual project cost.



Table 2-14. Description of ARARs for Selected Remedy

Citation	Activity or Prerequisite/Status	Requirement
Soil		
TCEQ Texas Risk Reduction Rules 30 TAC 335.558 and 30 TAC 335.559(g)(2)	Ensures adequate protection of human health and the environment from potential exposure to contaminants associated with releases – relevant and appropriate for remediation of contaminated soil and contaminated pipeline solid residue for cross-media contamination pathways such as soil to groundwater and for hypothetical future maintenance workers.	Non-residential (industrial) soils shall conform to the non-residential soil-to-groundwater cross media protection concentration. Non-residential (industrial) soils shall conform to the non-residential soil-to-groundwater cross media protection concentration MSC (GWP-Ind) values for 2,4,6-TNT for soils less than 3 ft bgs and for soils deeper than 3 ft bgs; 2,4-DNT and 2,6-DNT and Perchlorate in accordance with 30 TAC 335.559(g)(2) and as listed in Table 2-10 of this report. The concentration of contamination in soil and pipeline solid residue shall not exceed the non-residential soil-to-groundwater protection MSC (GWP-Ind) for the COCs listed in Table 2-10 . COCs in soil and pipeline solid residue COCs include 2,4,6-TNT, 2,4-DNT, 2,6-DNT, 1,3-DNB, 2-amino-4,6-DNT and 4-amino-2,6-DNT.
Groundwater		
Federal Safe Drinking Water Act (SDWA) MCLs 40 C. F. R. §§ 141.61 and 141.62	Applicable to drinking water for a public water system— relevant and appropriate for water that could potentially be used for human consumption.	Must not exceed SDWA MCLs for water designated as a current or potential source of drinking water. The MCLs for organic contaminants TCE, MC, 1,2-DCE, 1,2-DCA; 1,1-DCE; cis-1,2-DCE; trans-1,2-DCE; and VC are provided in 40 C. F. R. § 141.61(a) and the MCLs for inorganic contaminants arsenic; mercury; and, selenium are provided in 40 C. F. R. § 141.62 (b) and Table 2-10 of this report.
General Site Preparation, Construction, and Excavation Activities		
Opacity Standard 30 TAC 111.111(a)(8)(A)	Fugitive emissions from land-disturbing activities (e.g., excavation, construction)— applicable .	Visible emissions shall not be permitted to exceed opacity of 30% for any 6-minute period from any source.
Air Contaminants – General Nuisance Rules 30 TAC 101.4	Emissions of air contaminants— applicable .	No person shall discharge from any source whatsoever one or more air contaminants or combinations thereof, to exceed an opacity of 30 percent for any 6-minute period as are or may tend to be injurious to or to adversely affect human health or welfare, animal life, vegetation, or property, or as to interfere with the normal use and enjoyment of animal life, vegetation, or property.
Fugitive Particulate Matter Standard 30 TAC 111.145	Fugitive emissions from land-disturbing activities (e.g., excavation, construction)— applicable .	No person may cause, suffer, allow, or permit a structure, road, street, alley or parking area to be constructed, altered, repaired, or demolished, or land to be cleared without taking at least the following precautions to achieve control of dust emissions: <ul style="list-style-type: none"> • Use of water or of suitable oil or chemicals for control of dust in the demolition of structures, in construction operations, in work performed on a road, street, alley, or parking area, or in the clearing of land; and • Use of adequate methods to prevent airborne particulate matter during sandblasting of structures or similar operations
Storm Water Runoff Controls 40 C.F.R. § 122.26	Storm water discharges associated with construction activities— applicable to disturbances of equal to or greater than 1 acre of land.	Specific to areas of excavation of contaminated soil. Good construction management techniques, phasing of construction projects, minimal clearing, and sediment, erosion, structural, and vegetative controls shall be implemented to mitigate storm water run-on/runoff.



Table 2-14. Description of ARARs for Selected Remedy (continued)

Citation	Activity or Prerequisite/Status	Requirement
Waste Generation, Management, and Storage		
Characterization of Solid Waste 40 C.F.R. § 262.11 30 TAC 335.62 30 TAC 335.504 30 TAC 335.503(a)(4)	Generation of solid waste, as defined in 30 TAC 335.1— applicable .	<p>Must determine whether the generated solid waste is RCRA hazardous waste by using prescribed testing methods or applying generator knowledge based on information regarding material or process used. If the waste is determined to be hazardous, it must be managed in accordance with 40 C. F. R. § 262–268.</p> <p>After making the hazardous waste determination as required, if the waste is determined to be nonhazardous, the generator shall then classify the waste as Class 1, Class 2, or Class 3 (as defined in Section 335.505 through Section 335.507) using one or more of the methods listed in Section 335.503(a)(4) and Section 335.508 and manage the waste in accordance with the requirements of Chapter 335 of the TAC for industrial solid waste.</p>
Characterization of Hazardous Waste 40 C.F.R. §.264.13(a)(1); 40 C.F.R. § 268.7 30 TAC 335.504(3) 30 TAC 335.509 30 TAC 335.511	Generation of a RCRA hazardous waste for treatment, storage, or disposal— applicable if hazardous waste is generated (e.g., PPE).	<p>Must obtain a detailed chemical and physical analysis of a representative sample of the waste(s) that at a minimum contains all the information that must be known to treat, store, or dispose of the waste in accordance with 40 C. F. R. §264 and 268.</p> <p>Must also determine whether the waste is restricted from land disposal under 40 C. F. R. § 268 et seq. by testing in accordance with prescribed methods or use of generator knowledge of waste.</p>
Requirements for Temporary Storage of Hazardous Waste in Accumulation Areas 40 C. F. R. § 262.34(a) and (c)(1) 30 TAC 335.69(a) and (d)	On-site accumulation of 55 gallons or less of RCRA hazardous waste for 90 days or less at or near the point of generation— applicable if hazardous waste is generated (e.g., PPE) and stored in an accumulation area.	<p>Remedial activities derived waste (from monitoring and treating contaminated groundwater) is expected for this facility. A generator may accumulate hazardous waste at the facility provided that</p> <ul style="list-style-type: none"> • Waste is placed in containers that comply with 40 C. F. R. § 264.171 to 264.173 (Subpart I); and • Container is marked with the words “hazardous waste”; or • Container may be marked with other words that identify the contents.
Requirements for the Use and Management of Containers 40 C. F. R. § 264.171–264.173 30 TAC 335.69(e) 30 TAC 335.152(a)(7)	On-site storage/treatment of RCRA hazardous waste in containers for greater than 90 days— applicable if hazardous waste is generated (e.g., PPE) and is stored in containers.	Design and operating standards of 40 C. F. R. § 264.175(c) and 40 C. F. R. § 264.171, §.264.172, and §.264.173(a) and (b) must be met for the use and management of hazardous waste in containers.
Wells		
Well Construction Standards—Monitoring or Injection Wells 16 TAC 76.1000	Construction of water wells— applicable to construction of new monitoring or injection wells, if needed.	Injection wells shall be completed in accordance with the technical requirements of Section 76.1000, as appropriate. Substantive requirements applicable to the injection wells will be adhered to.



Table 2-14. Description of ARARs for Selected Remedy (continued)

Citation	Activity or Prerequisite/Status	Requirement
Wells (continued)		
Class V Injection Wells 30 TAC §331.9(a); 30 TAC §331.10(a); 30 TAC §331.10(d); 30 TAC §331.21; 30 TAC §331.132(a); 30 TAC §331.132(c); 30 TAC §331.132(d)(1); 30 TAC §331.132(d)(4); 30 TAC §331.133(e)	Installation, operation, and closure of injection wells for in situ chemical oxidation fall in the category of Class V Injection Wells— relevant and appropriate.	Injection wells shall be constructed to the required specifications for isolation casing, surface completion, prevention of commingling, and confinement of undesirable groundwater to its zone of origin. Closure shall be accomplished by removing all of the removable casing and the entire well shall be pressure filled via a tremie pipe with cement from bottom to the land surface, or closure shall be performed by the alternative method for Class V Wells completed in zones of undesirable groundwater. Groundwater concentrations at time of well closure will determine the appropriate method of abandonment. Substantive requirements applicable to the injection wells will be adhered to.
Well Construction Standards— Extraction Wells 16 TAC 76.1000(a) and (c) through (h) 16 TAC 76.1002(a) through (c) 16 TAC 76.1008(a) through (c)	Construction of water wells— applicable to construction of extraction (recovery) wells.	Substantive requirements applicable to extraction (recovery) wells will be adhered to. Wells shall be completed in accordance with the technical requirements of Section 76.1000, as appropriate. Water wells completed to produce undesirable water shall be cased to prevent the mixing of water or constituent zones. The annular space between the casing and the wall of the borehole shall be pressure grouted with cement or bentonite grout to the land surface. Bentonite grout may not be used if a water zone contains chloride water above 1500 parts per million (ppm) or if hydrocarbons are present. Wells producing undesirable water or constituents shall be completed in such a manner that will not allow undesirable fluids to flow onto the land surface. During installation of a water well pump, installer shall make a reasonable effort to maintain integrity of groundwater and to prevent contamination by elevating the pump column and fittings, or by other means suitable under the circumstances. Pump shall be constructed so that no unprotected openings into the interior of the pump or well casing exist.
Treatment/Disposal		
Disposal of Wastewater (e.g., contaminated groundwater, dewatering fluids, decontamination liquids) 40 C. F. R. § 268.1(c)(4)(i) 30 TAC 335.431(c)	RCRA-restricted characteristically hazardous waste intended for disposal— applicable if extracted groundwater is determined to be RCRA characteristically hazardous.	Appropriate and relevant in the event of a spill. Disposal is not prohibited if such wastes are managed in a treatment system subject to regulation under Section 402 of the CWA that subsequently discharges to waters of the United States.
Closure		



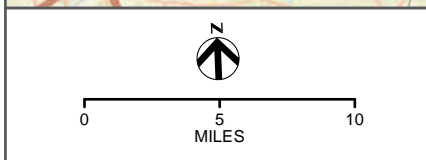
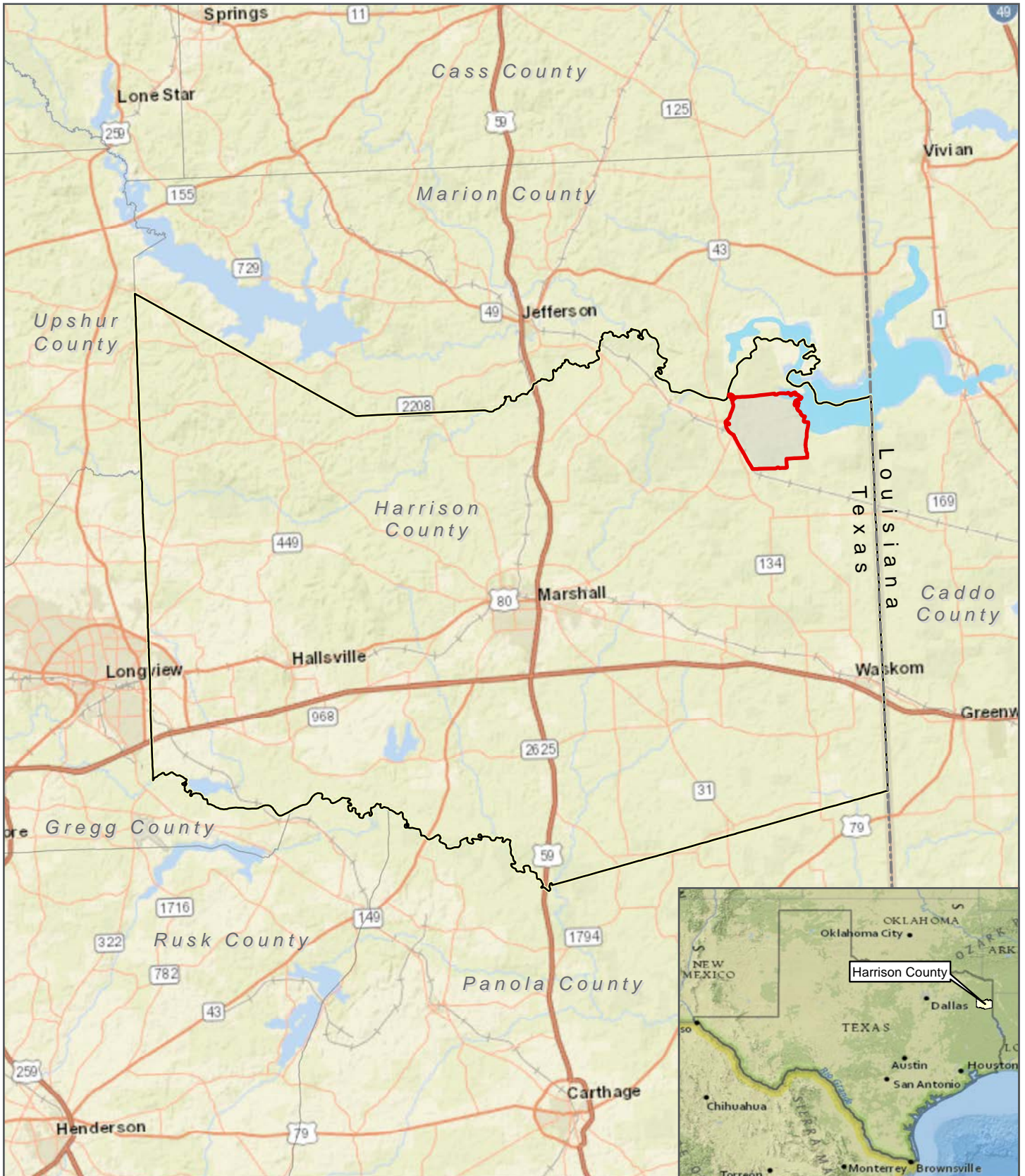
Table 2-14. Description of ARARs for Selected Remedy (continued)

Citation	Activity or Prerequisite/Status	Requirement
<p>Standards for Plugging Wells that Penetrate Undesirable Water or Constituent Zones</p> <p>16 TAC 76.1004(a) through (c)</p>	<p>Plugging and abandonment of wells—applicable to plugging and closure of monitoring and/or extraction wells.</p>	<p>If a well is abandoned, all removable casing shall be removed and the entire well pressure filled via a tremie pipe with cement from bottom up to the land surface. In lieu of this procedure, the well shall be pressure-filled via a tremie tube with bentonite grout of a minimum 9.1 lb/gal weight followed by a cement plug extending from land surface to a depth of not less than 2 feet. Undesirable water or constituents or the freshwater zone(s) shall be isolated with cement plugs.</p>

Abbreviations:

ARAR	<i>applicable or relevant and appropriate requirement</i>
bgs	<i>below ground surface</i>
C. F. R.	<i>Code of Federal Regulations</i>
CWA	<i>Clean Water Act of 1972</i>
USEPA	<i>U.S. Environmental Protection Agency</i>
FR	<i>Federal Register</i>
lb/gal	<i>pound per gallon</i>
LHAAP	<i>Longhorn Army Ammunition Plant</i>
MCL	<i>maximum contaminant level</i>

MCLG	<i>maximum contaminant level goal</i>
MSC	<i>medium-specific concentration</i>
%	<i>percent</i>
PPE	<i>personal protective equipment</i>
ppm	<i>part per million</i>
RCRA	<i>Resource Conservation and Recovery Act of 1976</i>
SDWA	<i>Safe Drinking Water Act</i>
TAC	<i>Texas Administrative Code</i>
TCEQ	<i>Texas Commission on Environmental Quality</i>

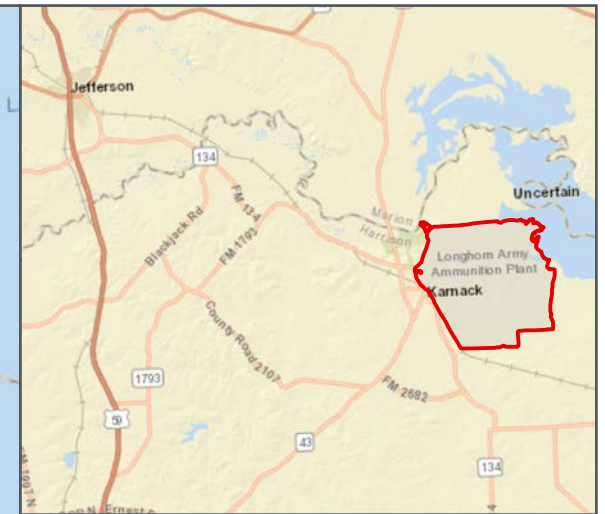
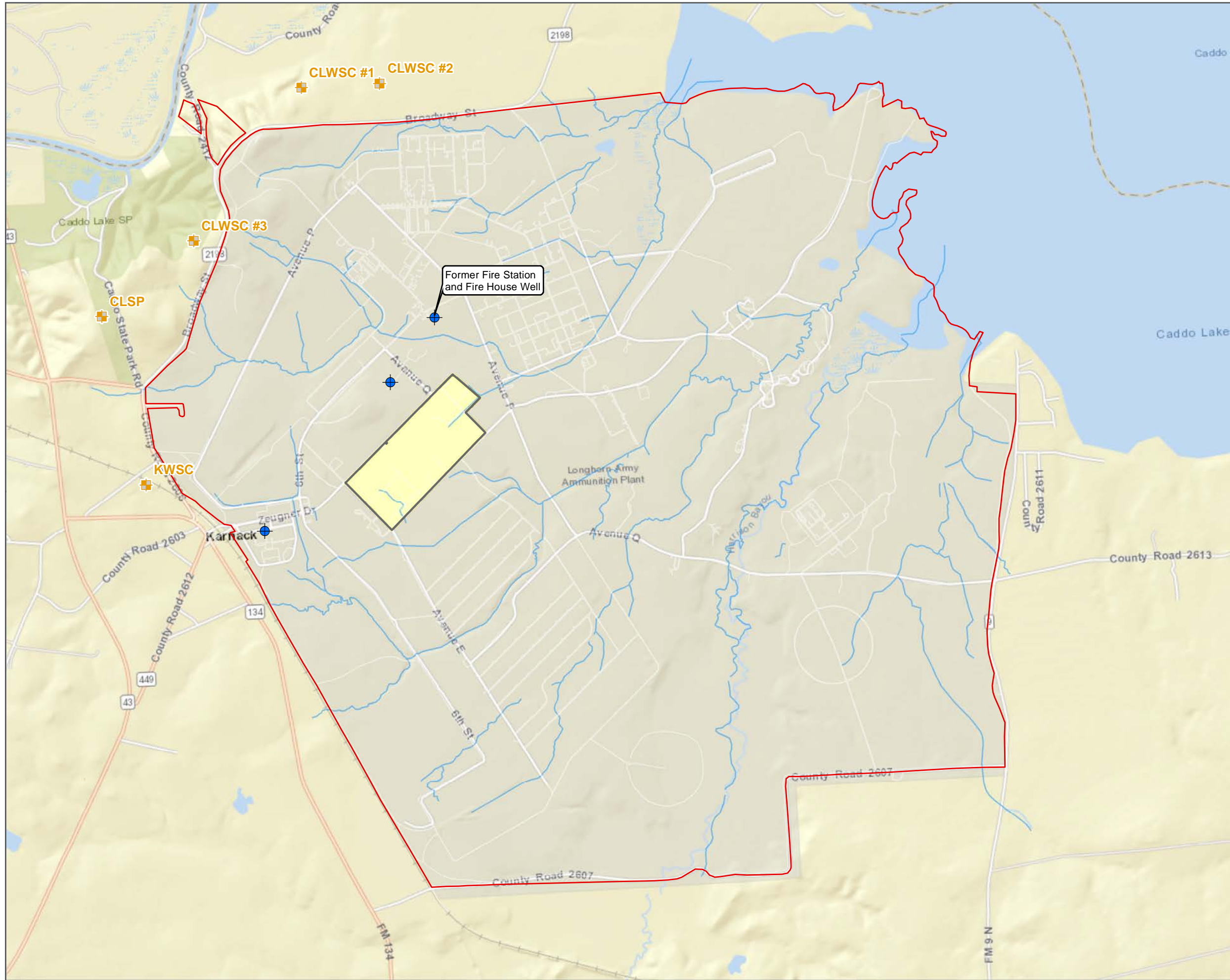


LEGEND	
	LHAAP Installation Boundary
	Harrison County
	State Boundary

SITE LOCATION
 LONGHORN ARMY
 AMMUNITION PLANT
 KARNACK, TEXAS



DATA SOURCES: AECOM, 2017, Draft Final Feasibility Study Addendum LHAAP-29, Former TNT Production Area, Group 2 Longhorn Army Ammunition Plant, Karnack, Texas. August
 DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.



LEGEND

- Public Water Supply Well Locations
- Potable Water Wells
- Roads
- Streams
- Site LHAAP-29
- LHAAP Installation Boundary

DATA SOURCES: AECOM, 2016, Draft Final Remedial Investigation Addendum LHAAP-29, Former TNT Production Area, Group 2 Longhorn Army Ammunition Plant, Karnack, Texas. July. ESRI

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

SITE VICINITY MAP

LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

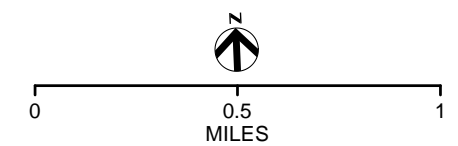
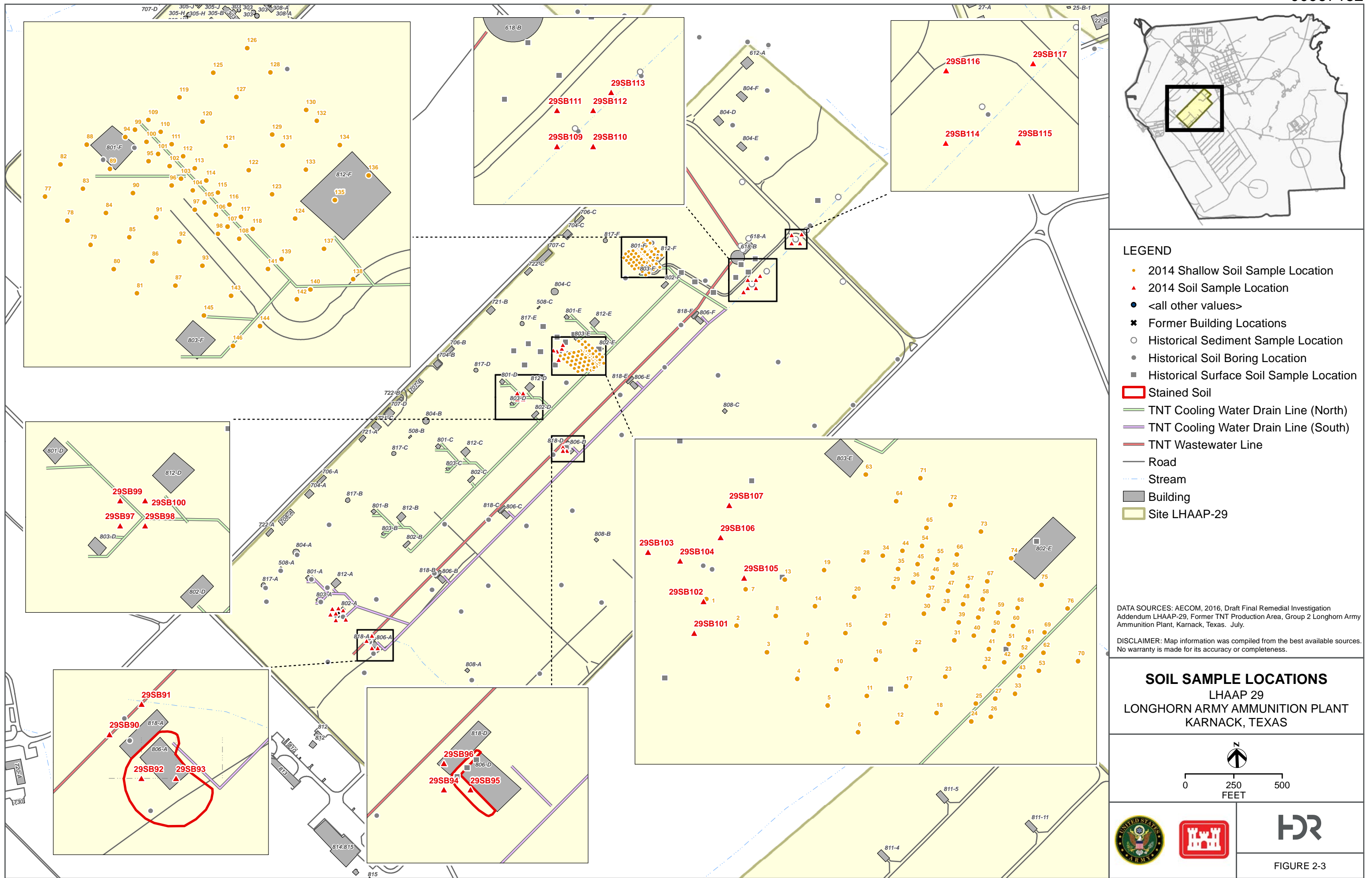


FIGURE 2-2



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Figure 2-3. LHAAP-29 Soil Sample Locations



LEGEND

- 2014 Shallow Soil Sample Location
- ▲ 2014 Soil Sample Location
- <all other values>
- ✖ Former Building Locations
- Historical Sediment Sample Location
- Historical Soil Boring Location
- Historical Surface Soil Sample Location
- ▭ Stained Soil
- TNT Cooling Water Drain Line (North)
- TNT Cooling Water Drain Line (South)
- TNT Wastewater Line
- Road
- Stream
- Building
- Site LHAAP-29

DATA SOURCES: AECOM, 2016, Draft Final Remedial Investigation Addendum LHAAP-29, Former TNT Production Area, Group 2 Longhorn Army Ammunition Plant, Karnack, Texas. July.

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

SOIL SAMPLE LOCATIONS
LHAAP 29
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

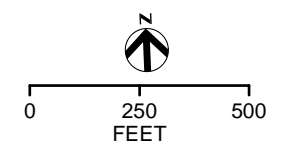
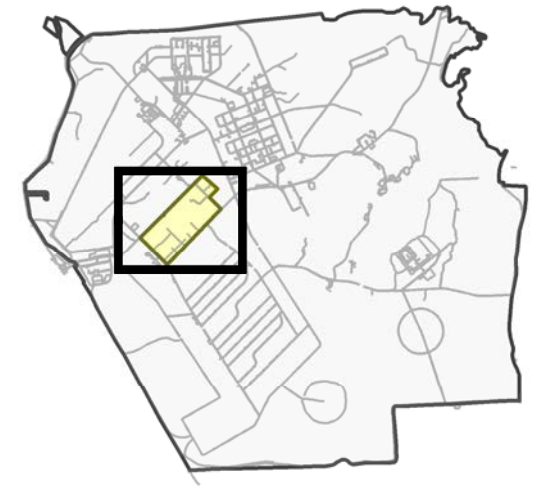
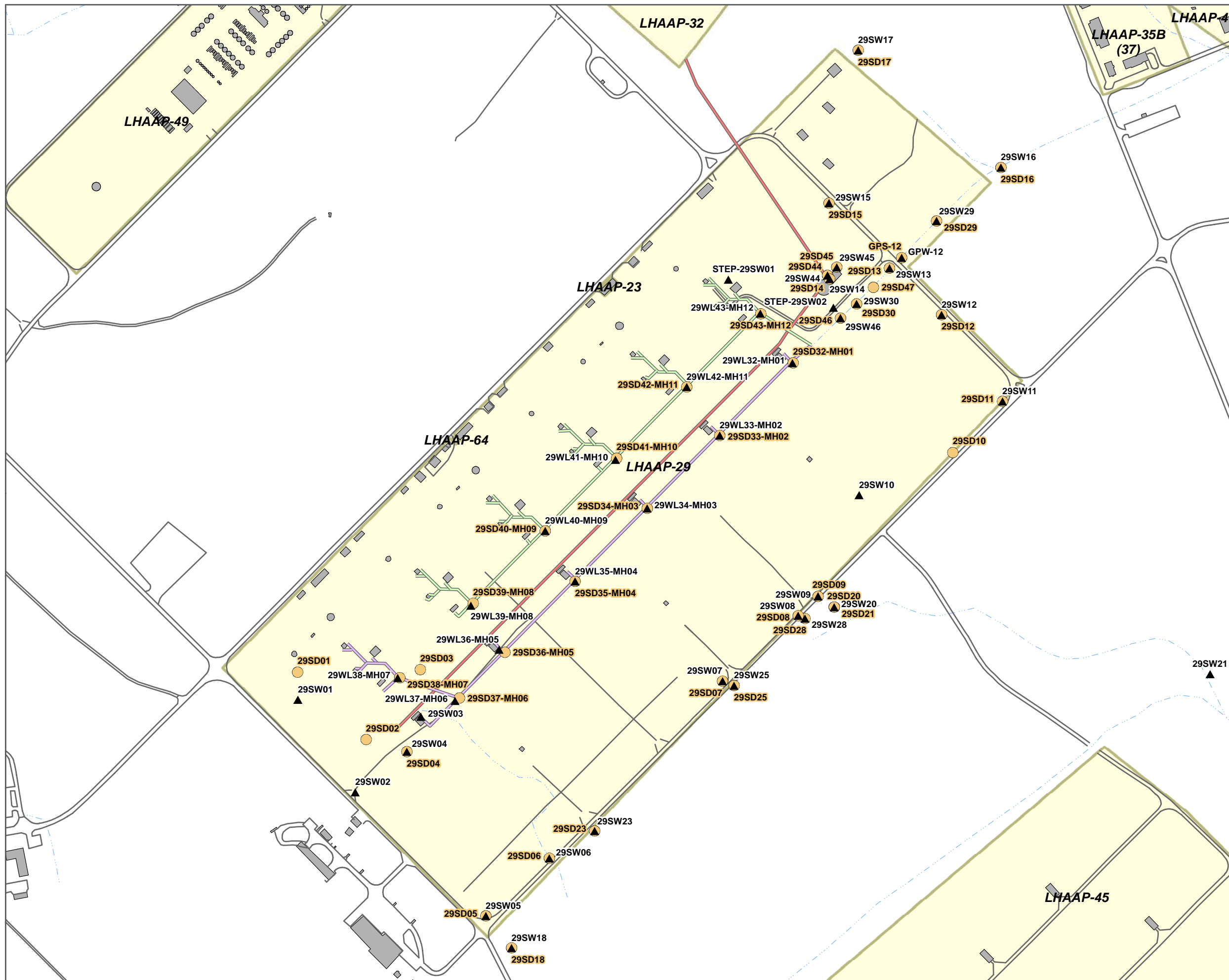


FIGURE 2-3



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Figure 2-4. LHAAP Surface Water and Sediment Sample Locations



LEGEND

- ▲ Surface Water Sample Location
- Sediment Sample Location
- TNT Cooling Water Drain Line (North)
- TNT Cooling Water Drain Line (South)
- TNT Wastewater Line
- - - Stream
- Road
- Building
- Site LHAAP-29

DATA SOURCES: Shaw, 2010, Final Feasibility Study, LHAAP-29, Former TNT Production Area, Group 2, Longhorn Army Ammunition Plant, Karnack, Texas, December.

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

**SURFACE WATER AND
SEDIMENT SAMPLE LOCATIONS**
LHAAP 29
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

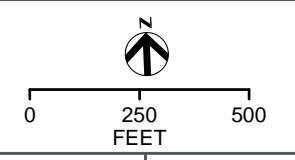


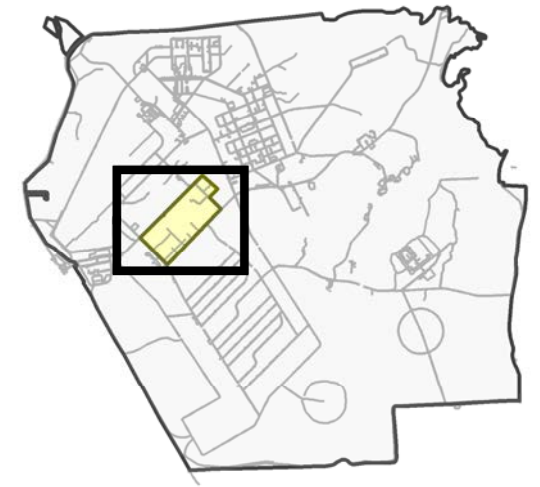
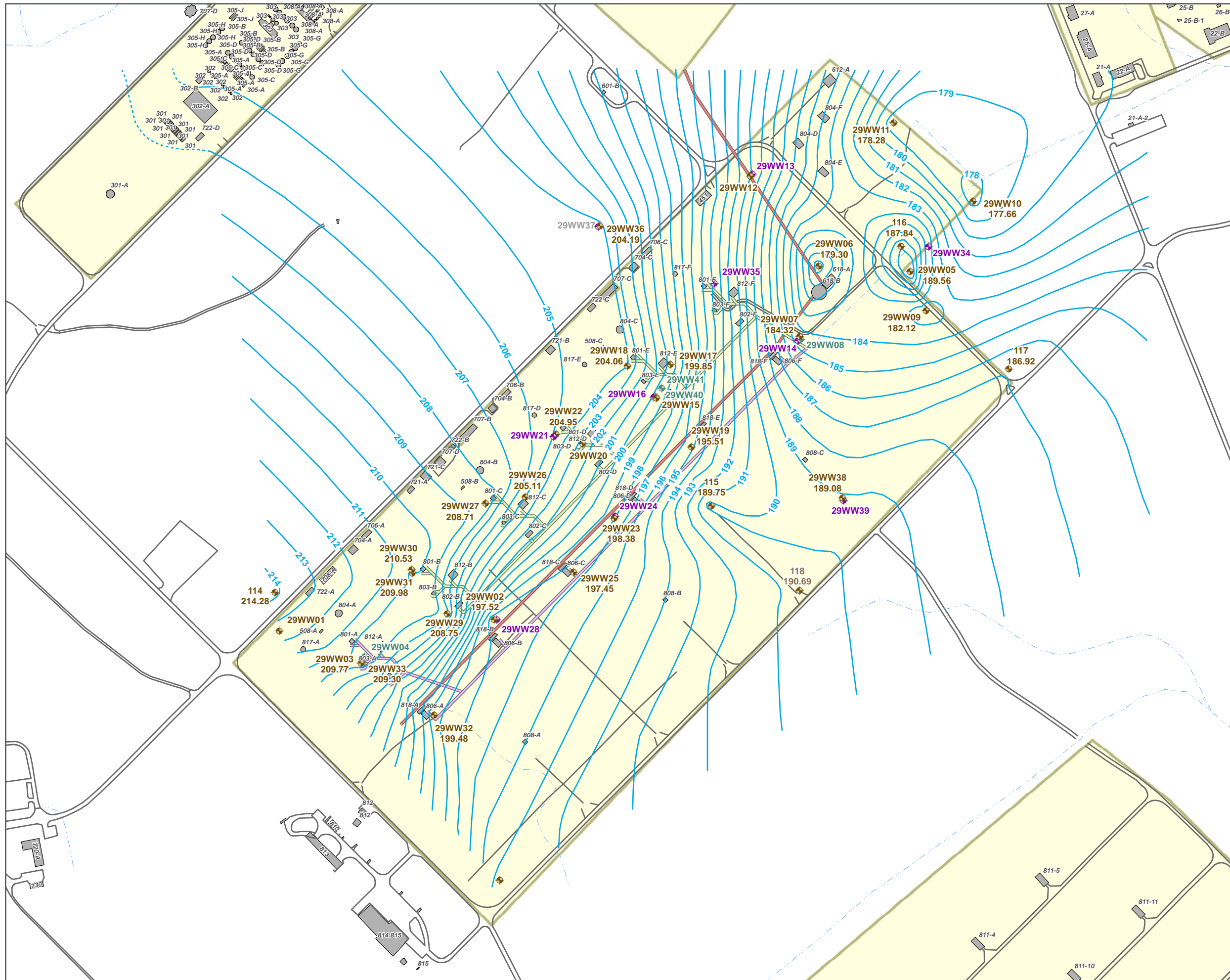
FIGURE 2-4

PATH: J:\2018\18-025_LHAAP_SITES29_47_18AND24_ROD_(WERNER)7.2_WORK_IN_PROGRESS\MAP_DOCS\DRAWING\LHAAP29\LHAAP_29_ROD_SURFACEWATER_SEDIMENTSAMPLE_LOCATIONS.MXD - USER: KLOFGREN - DATE: 5/17/2019



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Figure 2-5. LHAAP-29 Shallow Zone Wells and Groundwater Elevations



- LEGEND**
- Shallow or Lower Shallow Monitoring Well
 - Intermediate Monitoring Well
 - Deep Monitoring Well
 - Groundwater Elevation Contour (Dashed Where Inferred)
 - TNT Cooling Water Drain Line (North)
 - TNT Cooling Water Drain Line (South)
 - TNT Wastewater Line
 - Stream
 - Road
 - Building
 - Site LHAAP-29

NOTE:
Groundwater contours are based on data collected November 29, 2007 through December 3, 2007.

DATA SOURCES: AECOM, 2016, Draft Final Remedial Investigation Addendum LHAAP-29, Former TNT Production Area, Group 2 Longhorn Army Ammunition Plant, Karnack, Texas. July.

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

GROUNDWATER ELEVATION MAP (SHALLOW ZONE)
LHAAP 29
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

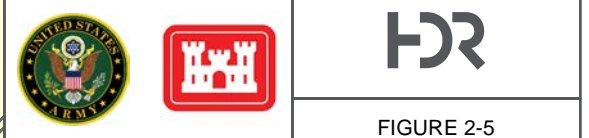
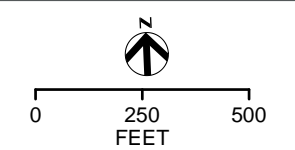
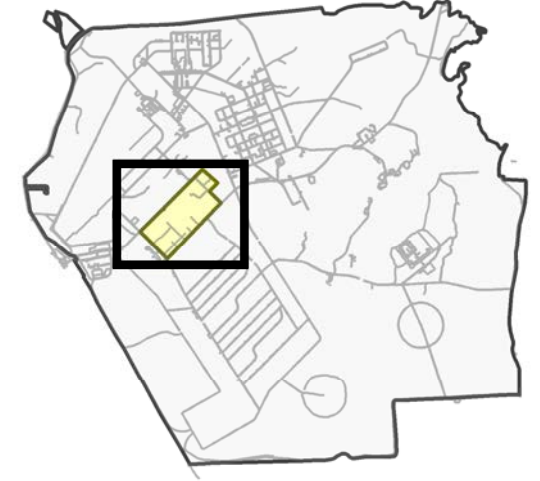
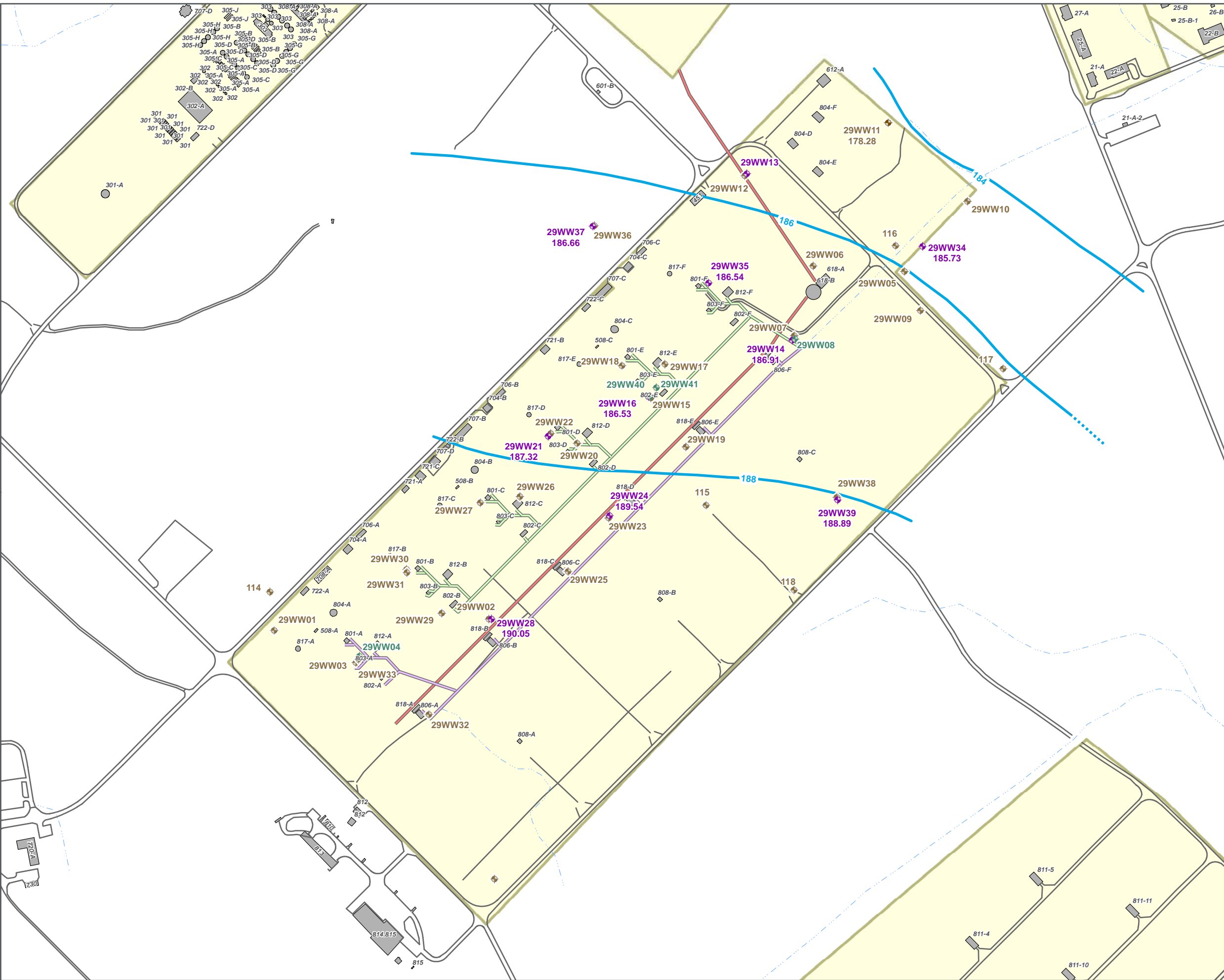


FIGURE 2-5

PATH: J:\2018\18-025_LHAAP_SITES29_47_18AND24_ROD_(WERNER)7.2_WORK_IN_PROGRESS\MAP_DOCS\DRAWING\LHAAP29\LHAAP_29_ROD_GW_ELEVATION_SHALLOW.MXD - USER: KLOFGREN - DATE: 5/17/2019



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LEGEND

- Shallow or Lower Shallow Monitoring Well
- Intermediate Monitoring Well
- Deep Monitoring Well
- Groundwater Elevation Contour (Dashed Where Inferred)
- TNT Cooling Water Drain Line (North)
- TNT Cooling Water Drain Line (South)
- TNT Wastewater Line
- Stream
- Road
- Building
- Site LHAAP-29

NOTE:
Groundwater contours are based on data collected November 29, 2007 through December 3, 2007.

DATA SOURCES: AECOM, 2016, Draft Final Remedial Investigation Addendum LHAAP-29, Former TNT Production Area, Group 2 Longhorn Army Ammunition Plant, Karnack, Texas. July.

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

**GROUNDWATER ELEVATION MAP
(INTERMEDIATE ZONE)**
LHAAP 29
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS

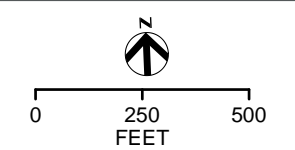


FIGURE 2-6



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Figure 2-7. LHAAP-29 Human Health Conceptual Site Model

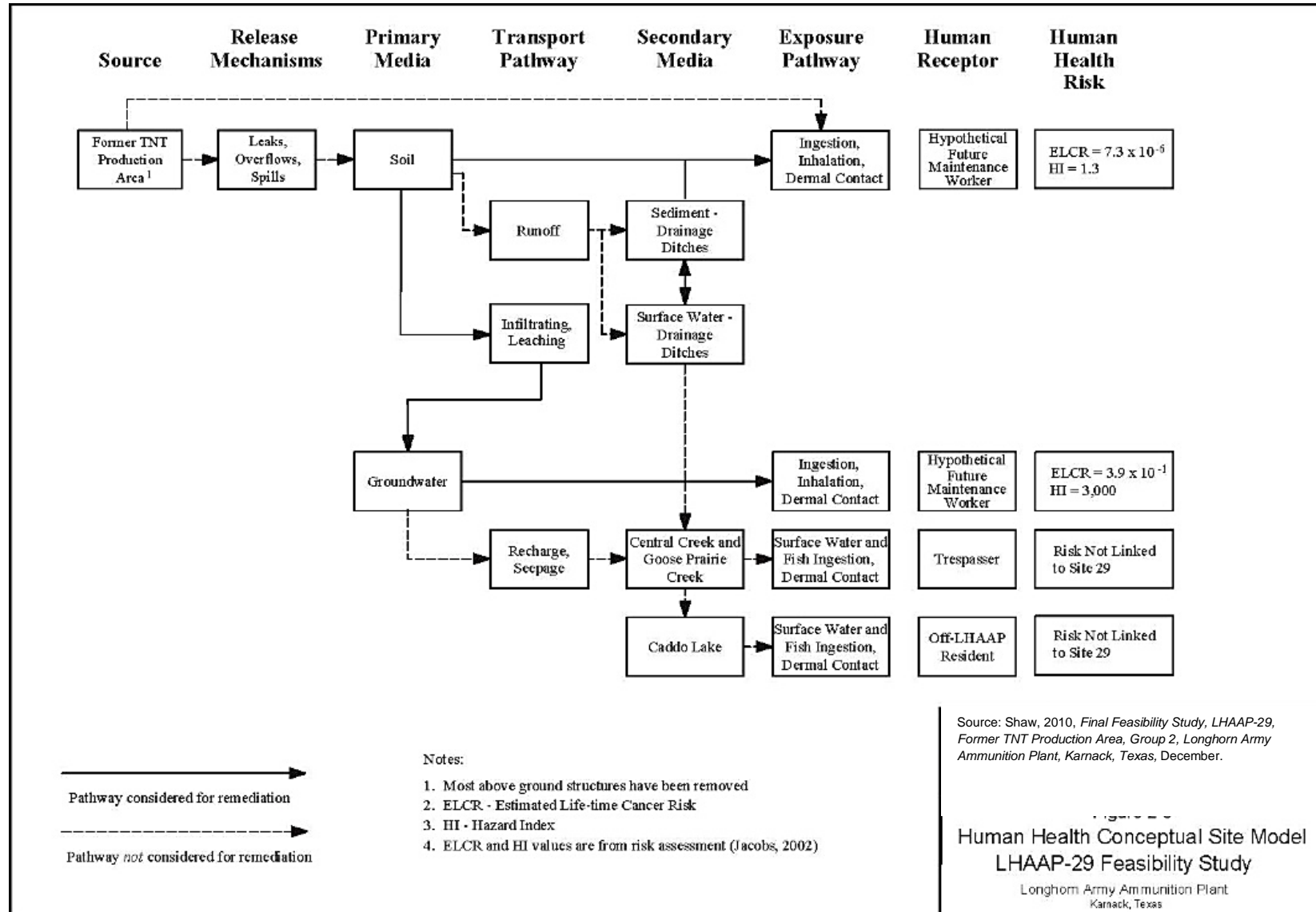
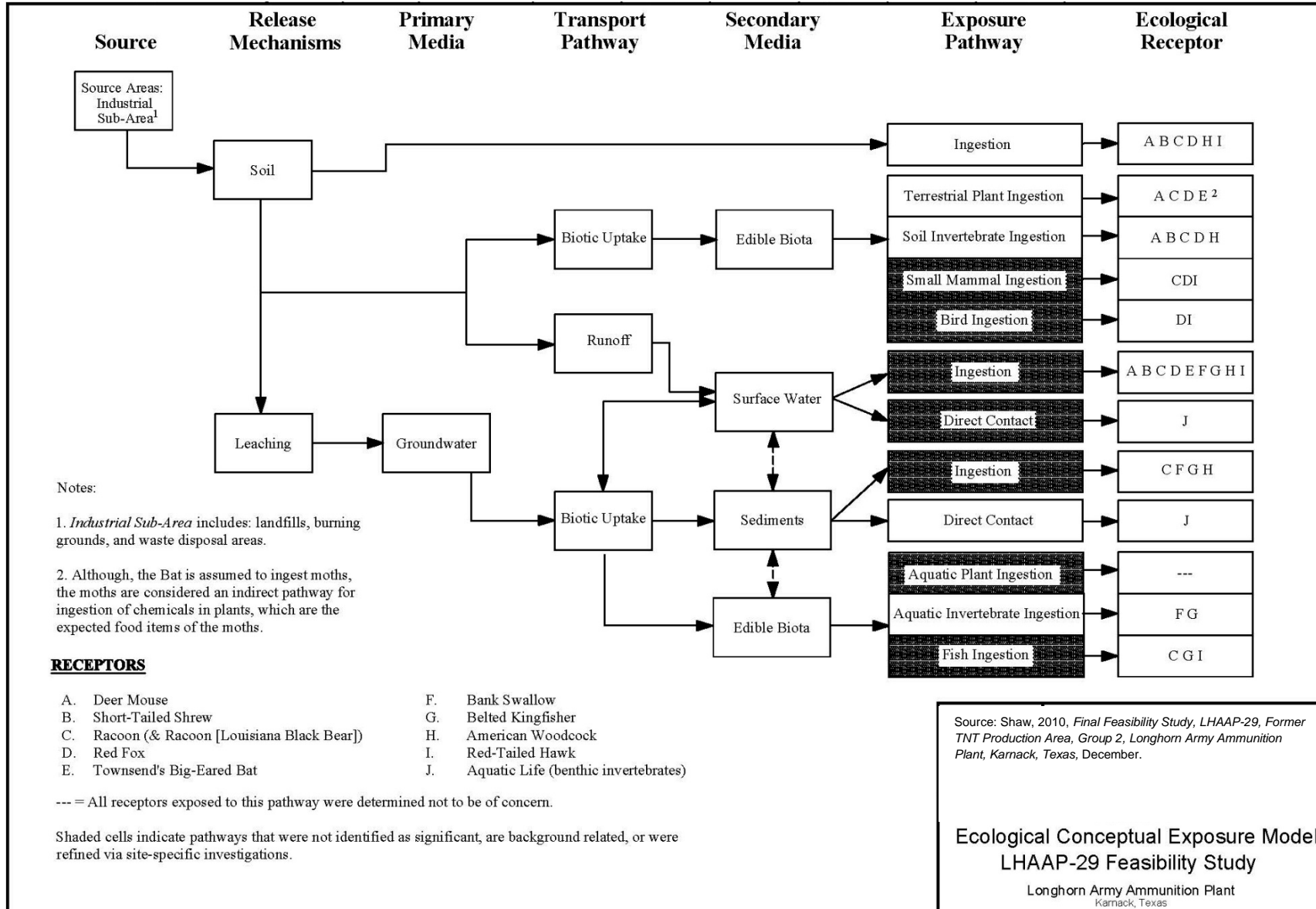
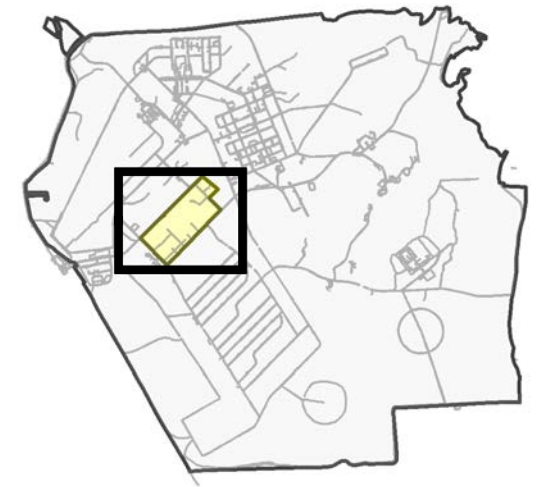




Figure 2-8. LHAAP-29 Ecological Conceptual Exposure Model





LEGEND

- Shallow Groundwater Monitoring Well (S)
- Intermediate Groundwater Monitoring Well (I)
- Deep Groundwater Monitoring Well (D)

Plume in Shallow Groundwater

- 1,2-Dichloroethane (MCL= 5 µg/L)
- Perchlorate (TRRP Tier 1 Residential Groundwater PCL = 17 ug/L)
- Trichloroethene (MCL= 5 µg/L)
- Stream
- Road
- Building
- Site LHAAP-29

NOTES:

- All perchlorate concentrations in the intermediate and deep zones are below the TRRP Tier 1 Residential Groundwater PCL of 17 ug/L
- All concentrations are reported in micrograms per liter (µg/L).
- Analytes are non detect unless labeled on map.
- MCL = Maximum Contaminant Level.

DATA SOURCES: HDR, 2018, Draft Revised Proposed Plan for LHAAP-29 Former TNT Production Area Group 2 Longhorn Army Ammunition Plant, August.

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

VOCS AND PERCHLORATE IN SHALLOW ZONE GROUNDWATER
 LHAAP 29
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS

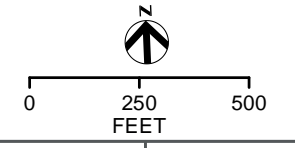


FIGURE 2-9

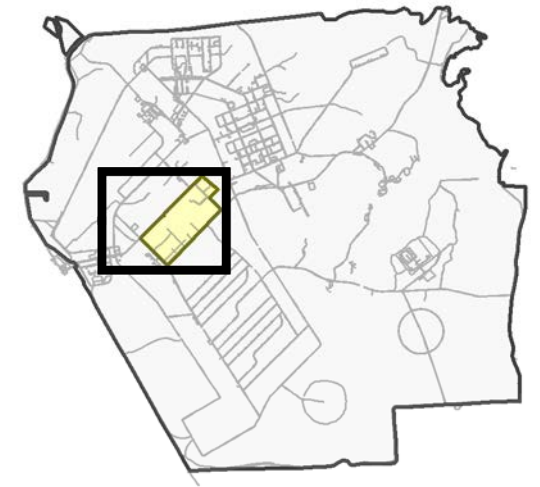
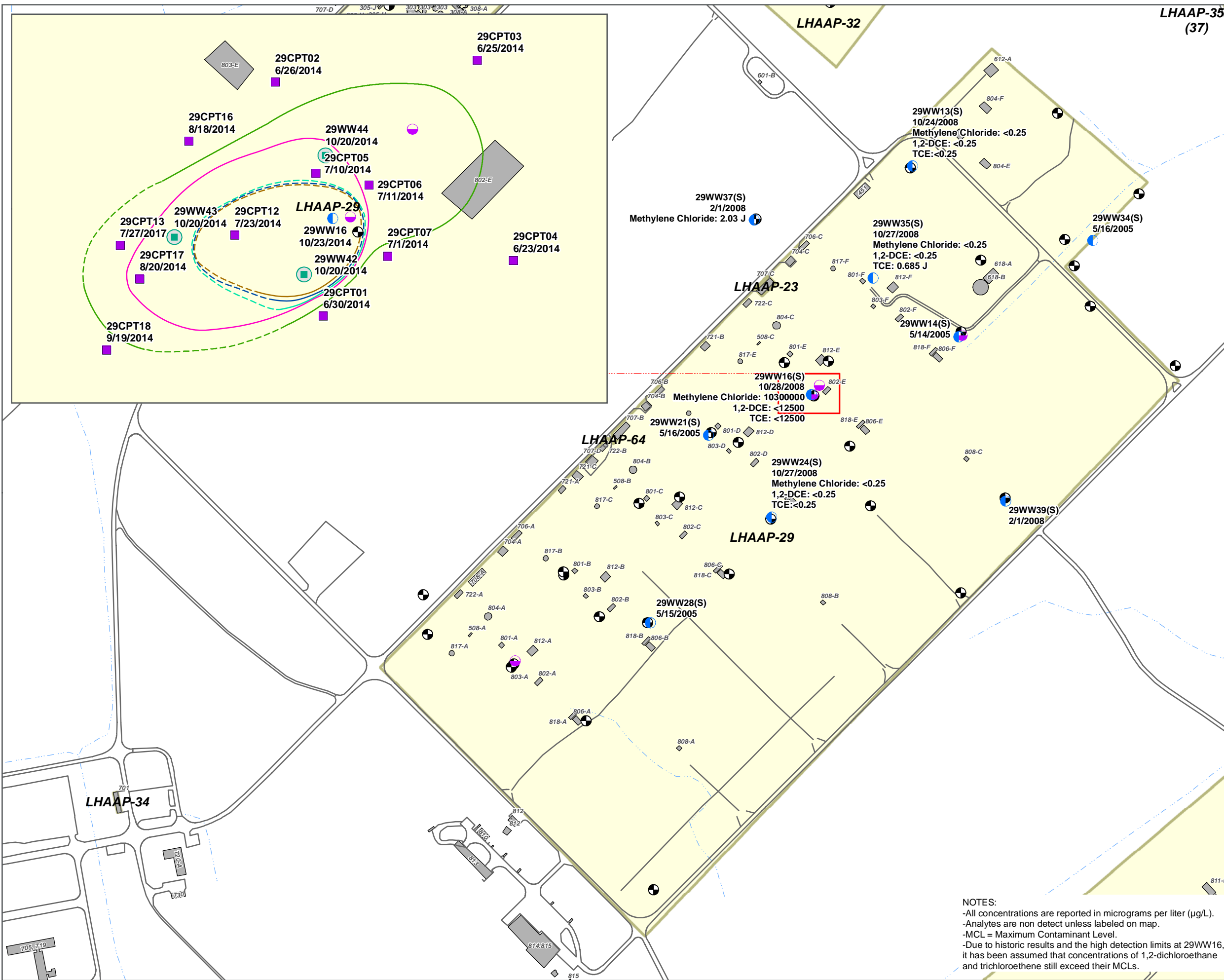


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Figure 2-11. VOCs in Intermediate Zone Groundwater



LEGEND

- Shallow Groundwater Monitoring Well (S)
- Intermediate Groundwater Monitoring Well (I)
- Deep Groundwater Monitoring Well (D)
- Intermediate Zone Well (2014)
- Intermediate Zone Soil Boring (2014)

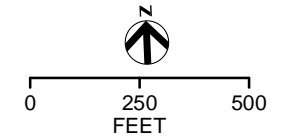
Intermediate Groundwater Contours (Dashed Where Inferred), Contour Concentration

- Trichloroethene (MCL = 5 µg/L)
- 1,1- DCE (MCL = 7 µg/L)
- cis - 1,2 - DCE (MCL = 70 µg/L)
- Vinyl Chloride (MCL = 2 µg/L)
- MC (MCL = 5 µg/L)
- Stream
- Road
- Building
- Site LHAAP-29

DATA SOURCES: HDR, 2018, Draft Revised Proposed Plan for LHAAP-29 Former TNT Production Area Group 2 Longhorn Army Ammunition Plant, August.

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

VOCs IN INTERMEDIATE ZONE GROUNDWATER
 LHAAP 29
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS



NOTES:

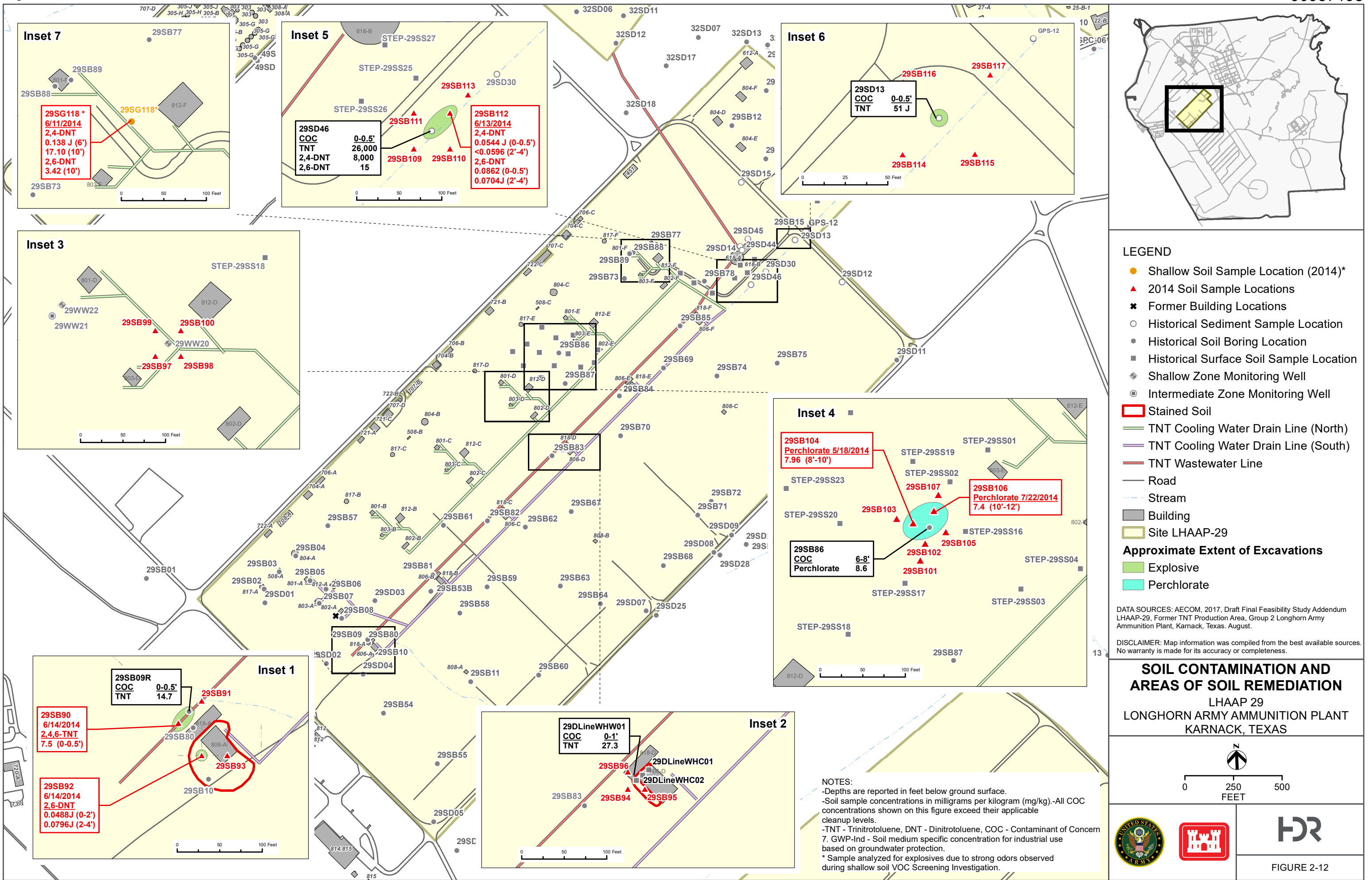
- All concentrations are reported in micrograms per liter (µg/L).
- Analytes are non detect unless labeled on map.
- MCL = Maximum Contaminant Level.
- Due to historic results and the high detection limits at 29WW16, it has been assumed that concentrations of 1,2-dichloroethane and trichloroethene still exceed their MCLs.

FIGURE 2-11



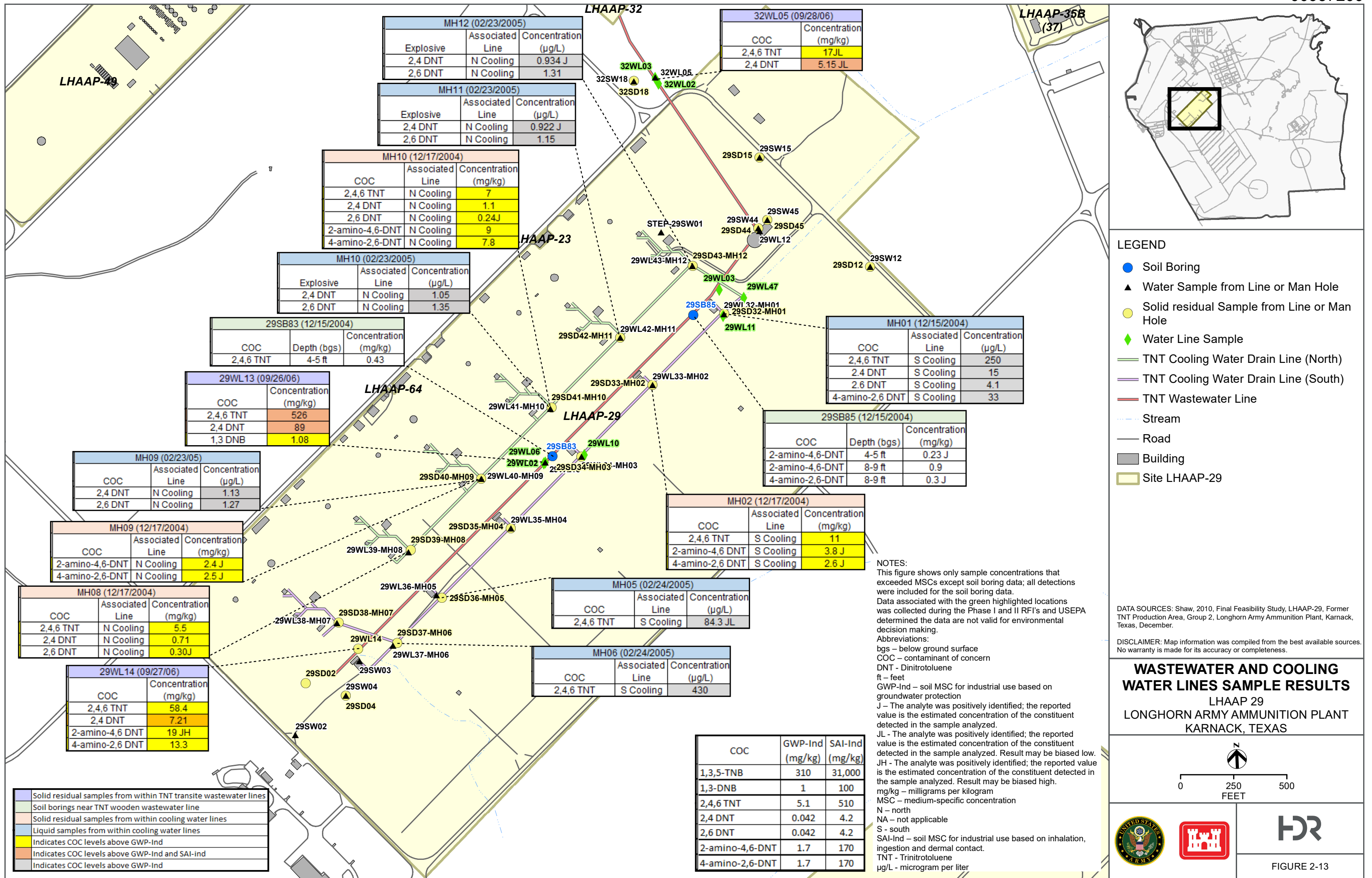
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Figure 2-12. Soil Contamination and Areas of Soil Remediation



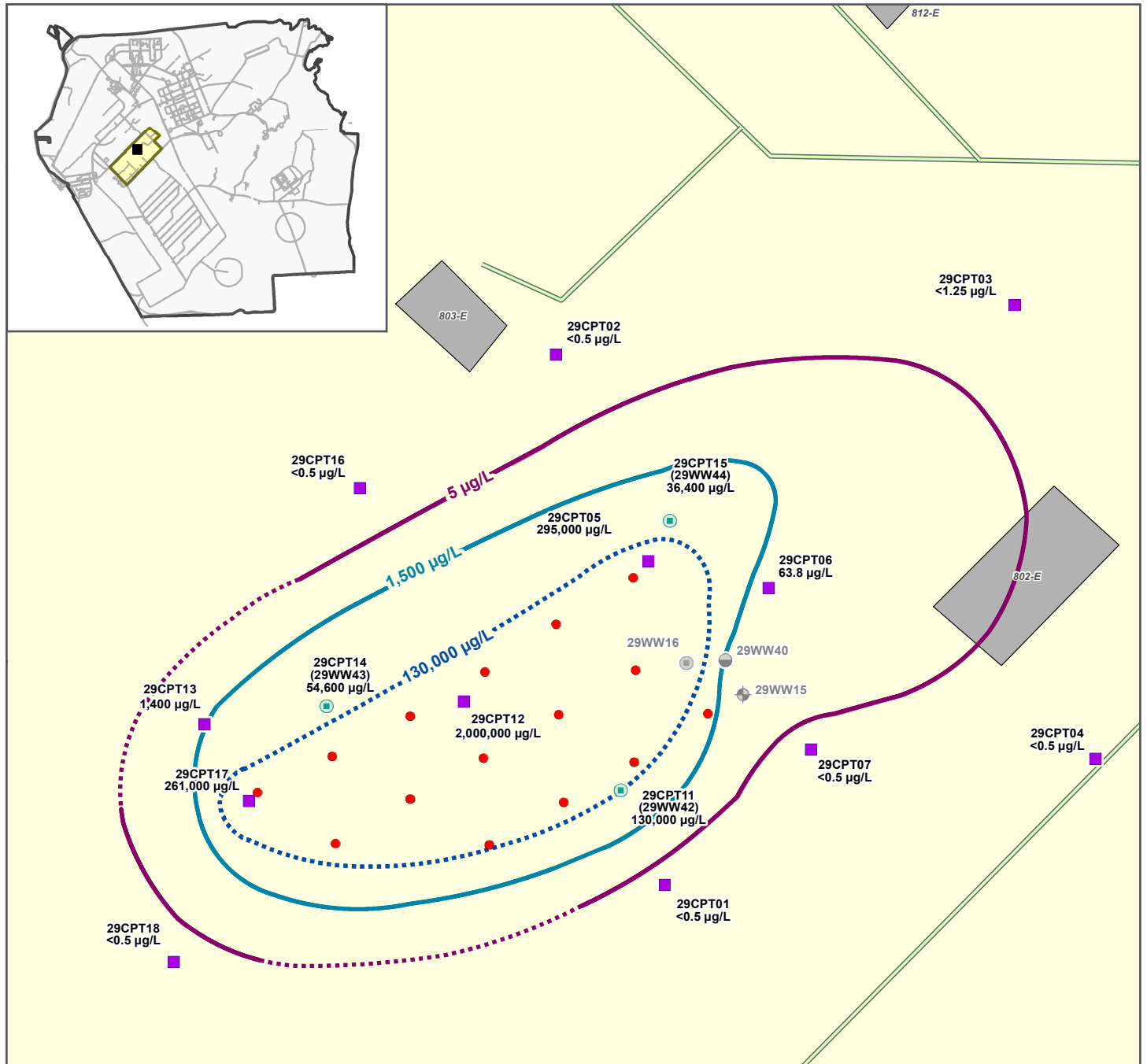


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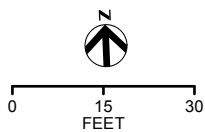


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LEGEND

- Electrode Boring with Co-Located SVE Well
- Intermediate Zone Soil Boring (2014)
- Intermediate Zone Well (2014)
- Deep Zone Well (2006)
- Intermediate Zone Well (1998)
- Shallow Zone Well (1998)
- ⋯ Inferred MC DNAPL Zone Contour (130,000 µg/L ~ 1% Solubility)
- MC Isoconcentration Contour (MCL = 5 µg/L) (Dashed Where Inferred)
- Approximate Outline of MC Exceeding 1,500 µg/L
- TNT Cooling Water Drain Line (North)
- Buildings
- Site LHAAP-29



Notes:
 µg/L = Concentrations in micrograms per liter.
 < = Concentration below laboratory reporting limit.
 DNAPL = Dense Non-Aqueous Phase Liquid
 MC = Methylene Chloride

CONCEPTUAL LAYOUT OF ELECTRICAL RESISTANCE HEATING (ERH) WELL NETWORK

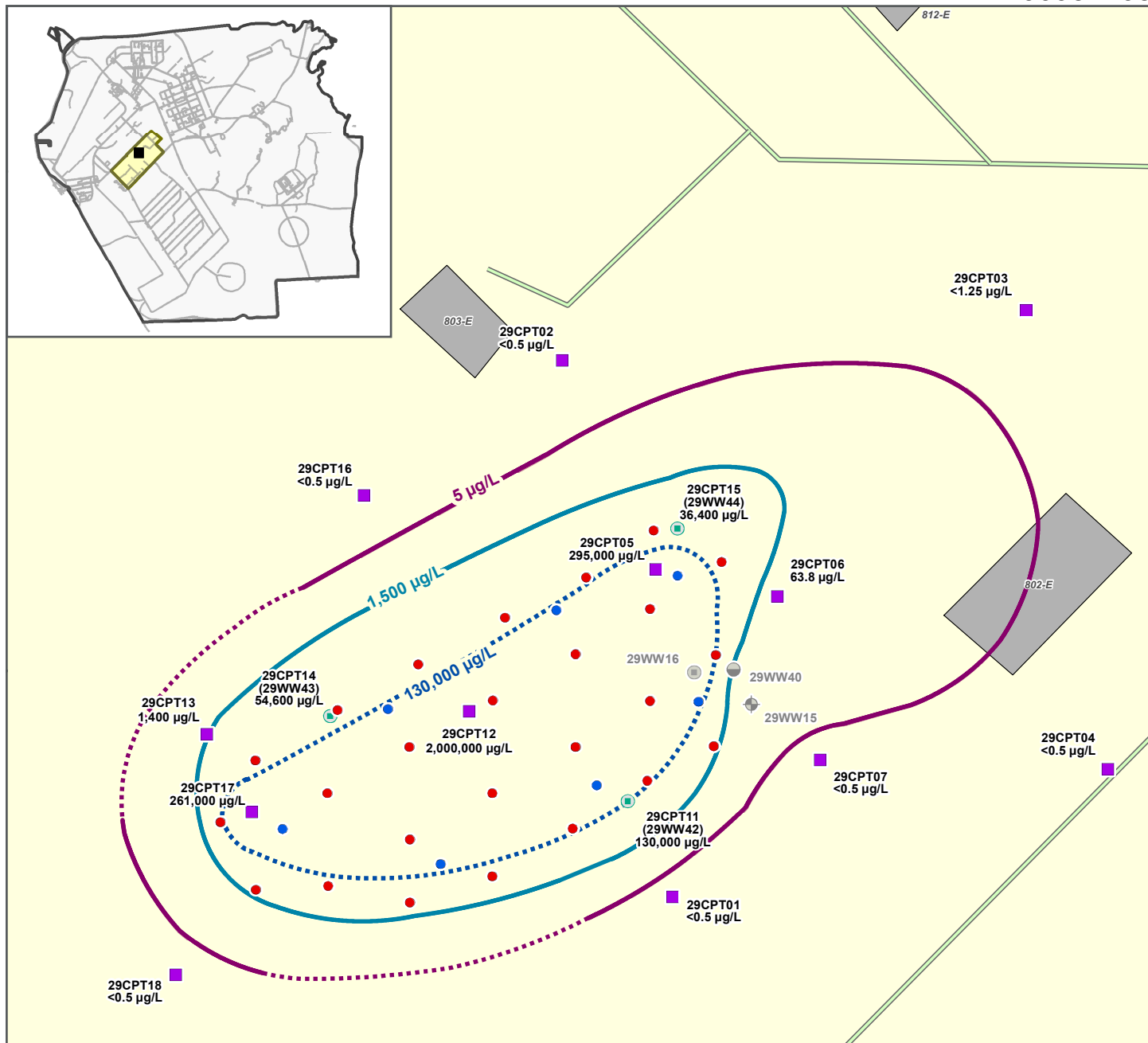
LHAAP 29
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS



DATA SOURCES:

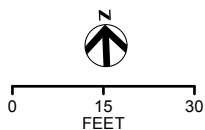
DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

FIGURE 2-14



LEGEND

- Heater Boring with Co-Located SVE Well (Proposed)
- Multi-phase Extraction Well
- Intermediate Zone Soil Boring (2014)
- Intermediate Zone Well (2014)
- Deep Zone Well (2006)
- Intermediate Zone Well (1998)
- Shallow Zone Well (1998)
- Inferred MC DNAPL Zone Contour (130,000 µg/L ~ 1% Solubility)
- MC Isoconcentration Contour (MCL = 5 µg/L) (Dashed Where Inferred)
- Approximate Outline of MC Exceeding 1,500 µg/L
- TNT Cooling Water Drain Line (North)
- Buildings
- Site LHAAP-29



Notes:
 µg/L = Concentrations in micrograms per liter.
 < = Concentration below laboratory reporting limit.
 DNAPL = Dense Non-Aqueous Phase Liquid
 MC = Methylene Chloride

CONCEPTUAL LAYOUT OF THERMAL CONDUCTION HEATING (TCH) WELL NETWORK

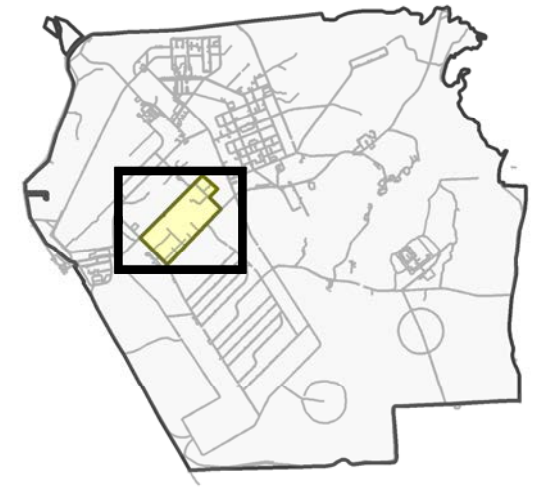
LHAAP 29
 LONGHORN ARMY AMMUNITION PLANT
 KARNACK, TEXAS



DATA SOURCES:

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

FIGURE 2-15



LEGEND

- Preliminary Land Use Control Boundary
- TNT Cooling Water Drain Line
- TNT Cooling Water Drain Line
- TNT Wastewater Line
- Stream
- Road
- Building
- Site LHAAP-29

DISCLAIMER: Map information was compiled from the best available sources. No warranty is made for its accuracy or completeness.

**PRELIMINARY
LAND USE CONTROL BOUNDARY
LHAAP 29
LONGHORN ARMY AMMUNITION PLANT
KARNACK, TEXAS**

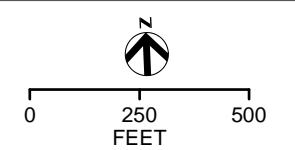


FIGURE 2-16



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3. Responsiveness Summary

The Responsiveness Summary serves three purposes. First, it provides the U.S. Army, USEPA, and TCEQ with information about community concerns with the preferred alternative at LHAAP-29 as presented in the Revised Proposed Plan. Second, it shows how the public's comments were considered in the decision-making process for selection of the remedy. Third, it provides a formal mechanism for the U.S. Army to respond to public comments. Two public comment periods and public meetings were held, one for the LHAAP-29 PP and one for the Revised PP. Responsiveness summaries for both meetings are provided.

The U.S. Army, USEPA, and TCEQ provide information regarding LHAAP-29 through public meetings, the Administrative Record for the facility, and announcements published in the Shreveport Times and Marshall News Messenger newspapers. **Section 2.3** discusses community participation on LHAAP-29, including the dates for the public comment period, the date, location, and time of the public meetings, and the location of the Administrative Record. The following documents related to community involvement were added to the Administrative Record for the two comment periods and public meetings:

- Transcript of the public meeting held on March 22, 2011
- Presentation slides from the March 22, 2011 public meeting
- Written questions and comments from the public during the public comment period pertinent to the revised PP, and the U.S. Army response to those comments are provided in **Section 3.1**.
- Transcript of the public meeting held for the Revised Proposed Plan on December 6, 2018
- Presentation slides from the December 6, 2018 public meeting
- Verbal questions and comments (no written questions provided) from the public during the public comment period, and the U.S. Army response to those comments are provided in **Section 3.1**.

3.1 Stakeholder Issues and Lead Agency Responses

This section responds to significant issues raised by stakeholders including the public and community groups that were received in written or verbal form.

2018 Revised Proposed Plan Comments

No written comments were received on the Revised Proposed Plan. Verbal comments and questions were discussed and addressed during the public meeting on December 6, 2018 and are summarized below.

Question/comment: Will the contaminated soil remain at the site?

Response: The estimated 3,900 cubic yards of contaminated soil will be excavated and disposed at a permitted landfill. The Landfill will be selected as part of the remedial design process.

Question/comment: Are they going to fill in the excavation areas?



Response: *What is typically done in these situations is to backfill with clean soil and re-vegetate the area. The specific details will be developed as part of the remedial design, which hasn't happened yet.*

Question/comment: The information presented states the wooden wastewater line has been flushed and abandoned. I would think the wood has rotted by now. Does it still exist, and is there anything more to be done there?

Response: *As part of the Remedial Design, the soil along the wooden TNT wastewater lines will be sampled to confirm that leaching from the lines has not occurred. Results from the confirmation soil sampling may identify additional areas exceeding the cleanup levels, which would require soil excavation and disposal. Additionally, as part of the Remedial Design, the deteriorated wooden wastewater line will be sampled to determine whether contaminants in the line exceed soil cleanup levels and require excavation and disposal.*

Question/comment: Does the cooling water line made of clay still exist? Has it been flushed?

Response: *Yes, the clay line will be flushed and the water sampled.*

Question/comment: I guess they're expecting that there could be some holes in that old clay pipe, so they're going to check it with a camera. And if they see a hole, then they'll check the soil around where that hole was. Because I'm wondering how aren't these falling apart and disintegrated over time.

Response: *The lines haven't disintegrated. They will be flushed and inspected, and if any breaks are found, soil in the vicinity will be sampled to determine if contaminants have leached into the soil.*

Question/comment: There was a process where they were trying microorganisms out in the area near the fire station about three or four years ago. Did that prove to be successful in any way?

Response: *We've performed a treatability study at the old Chemlab. It's different than what we are doing now. We do enhanced bioremediation at two areas, and will begin in a third area soon. We've had good success with this process in these areas. The treatability study at Site 37 was not successful. That bioremediation at LHAAP-37 was aerobic as opposed to anaerobic process, which has been used at the other three plumes. For Site 29, the contaminant is methylene chloride, which is almost at the DNAPL phase (that is a separate phase and not dissolved in groundwater). The concentrations are so high that it is toxic to the microorganisms and they can't degrade the methylene chloride. They can survive around the edges of the plume but not in the main portion. The data and posters provided show that concentrations decrease rapidly as you get further from the center of the plume where the DNAPL is present and indicates that degradation is probably occurring.*

One thing to note about the shallow and intermediate plumes at Site 29 is that they appear to be stable and not migrating due to the geology. The contaminant plumes appear to be contained by clay lenses. This is fairly unusual but in this case is helpful.

Question/comment: *In heating up the ground that deep is that going to harm the animals or vegetation?*



Response: *We don't expect the soil at or near the ground surface to be heated up to temperatures that could harm animals or vegetation. The heating will occur in the subsurface in the contaminated zone at 45 feet below ground surface. The heater wells or electrodes will be placed at that depth so that only the main part of the plume will be targeted to be heated to the highest temperatures (up to 140 degrees F). There will be a halo of warmed up soil and groundwater around the primary area though. That is considered possibly beneficial to help with destruction of VOCs in the shallower groundwater zone, but we're not sure just how much that might happen or the extent of the warming to shallower depths.*

Question/comment: Are we going to see clearing of that land and taking trees and dirt and everything out for these heating wells?

Response: *Much of the surface area was cleared of low-growing vegetation for the previous investigations and well installation. Some regrowth has occurred and there will have to be some vegetation or tree removal in specific areas that will be excavated, for well installation, and equipment access. That will need to be determined during the remedial design phase. We will try to minimize the amount of environmental damage. We were discussing how to do the work with minimal impact with the regulators and USFWS but we still have to balance that with the need to remove the contamination.*

Question/comment: Are you going to protect this area using fencing or other means?

Response: *During remedial activities, there will be an exclusion zone established to keep people out for safety reasons. One of the gates might need to be closed for a time while there is work going on.*

Question/comment: Is thermal desorption new, is it a proven practice?

Response: *This technology has been around for five to ten years. Because we have DNAPL at depth and there aren't many options to clean it up, we're anticipating that this approach will be able to vaporize the DNAPL so we can remove it.*

Question/comment: Earlier you mentioned that deep groundwater, beyond 88 feet is not contaminated, that this DNAPL is in the intermediate zone?

Response: *That is correct. There are monitoring wells in the deep zone that confirm the contamination does not extend to that zone.*

Question/comment: Once the decision has been made, will you go out with another contractor or will this company be the one to do the work?

Response: *We will have a separate remedial design and potentially a separate remedial action contract.*

Question/comment: This contamination has been here so long, hasn't the damage been done? Is the fear of migrating what is urging us to want to remove it completely? Not a lot has been damaged so why are we doing all this and planning to spend millions of dollars for this?

Response: *At other sites there is a visible risk to surface water. At this site the DNAPL in the intermediate groundwater is actually considered a principal threat waste because it will continue to*



contaminate groundwater. The EPA regulations require that a principal threat waste must be remediated. In addition, even though the plumes aren't moving, there are Federal and State of Texas regulatory requirement to return groundwater to beneficial uses whether we're using it or not, so remedial action is still required. Before 2009 if the groundwater wasn't being used you could let it go, but EPA's rules changed with regard to groundwater and now it is required to return all groundwater to its beneficial use. We have a small area of soil that is an ecological risk that must be remediated.

Question/comment: You've said you don't know what future uses [of groundwater] might be and that scares me. We went through a long process for this to become a refuge and now you're saying there is an opportunity for someone to come in and take the property and use it for another use?

Response: *That is not the intent at all. There are only two ways the property could come out of refuge property, a congressional act or some type of transfer to another federal agency. There is very limited potential for this to happen and is not anticipated. Everything that's being done has the anticipated use as a refuge and the regulatory requirement with regard to returning groundwater to beneficial uses is the driver for the remedial action planning. There are sites throughout the nation where this is required so Longhorn is not being required to do more than other sites.*

Question/comment: The Caddo Lake Institute has a third party contaminant expert who reviews these documents and he finds this to be a good plan. His largest question is the depth and extent of the DNAPL and that has been answered. I wanted to share with the community that he found this to be a good plan.

Response: *Thank You*

2011 Proposed Plan Comments

Question/comment: Remedial Alternatives 2 and 3 includes Excavation and Offsite Disposal of Soil, but no amount is given. How many acres within site 29 are we talking about? How deep would the excavation go? How many cubic yards are proposed to be removed from site 29?

Response: *Based on comparison of concentrations versus cleanup levels, several excavation locations are identified to various depths. Confirmation samples will be collected to verify the cleanup levels are met. Excavation will continue until the cleanup levels are attained in both the vertical (floor) and horizontal (walls) directions. Thus, the actual excavation size and depth may vary from the proposed excavations.*

The proposed excavation areas to mitigate human health risk are:

- *Area around 29SD46 – explosives contaminated soil with the approximate dimensions of 120 feet by 20 feet to a depth of 1 foot and an estimated total volume of 90 cubic yards (cy).*
- *Area around 29SB86 – perchlorate contaminated soil with approximate dimensions of 100-foot diameter circle to a depth of 10 feet and an estimated volume of approximately 2,900 cy.*

The proposed excavation areas to mitigate ecological risk by removal of explosives contaminated soil are:



- Area around 29SB08 (former Building 802-A) – approximate dimensions of a 60-foot diameter circular area and an estimated volume of approximately 200 cy.
- Area around 29DLineWHW01 – stained soil around former Building 806-D and former Building 806-A, with an estimated volume of approximately 30 cy.
- Area around 29SD13, 29SB15, and GPS-12 (cooling water ditch north of Avenue D), 150 feet by 20 feet, approximately 440 cy.

In total, the areas are less than a half-acre with a volume of approximately 3,900 cy.

Question/comment: It is a poor plan for the Army to waste money flushing and plugging the old TNT wastewater, and cooling water lines at site 29 in hope it removes the dangerous chemicals contaminating the groundwater. This is not a cure for the problem. Given the high concentration levels of dangerous chemicals listed, these lines should be dug up and disposed of. Any contaminated soil adjacent to the lines caused by leakage from the lines should be dug up as well and disposed of. By doing this, it for sure resolves the problem by preventing further contamination to groundwater.

Response: *The contamination in the lines has the potential to cause contamination in the groundwater only if there is no action and the contamination remains in the lines and the line deteriorates to allow water to infiltrate or the contamination to come into contact with the soil. Currently, there is no evidence that the explosive residue in the lines has caused the isolated shallow explosive groundwater plumes, as they are not located adjacent to the lines. Explosive soil contamination that poses a potential to leach into the groundwater was identified in the surface soil. (These areas are proposed for excavation as described in the previous response.) The samples collected from the subsurface soil adjacent to the TNT wastewater lines do not have explosive concentrations exceeding the groundwater protection standard, or do not indicate contamination that poses a potential to leach into the groundwater.*

Additionally, as part of the RD, confirmation soil samples will be collected adjacent to the North and South Cooling Water lines as well as the TNT wastewater lines to confirm that leaching from the lines has not occurred.

Over time, the lines may eventually deteriorate or break down. After line flushing, no contamination (no solid or liquid residue) will remain in the lines; therefore, there will be no groundwater infiltration and transport of contaminants as the lines deteriorate. The lines will remain buried below ground and since the lines do not contain any liquid or soil residue, they cannot leak and cannot be a conduit for contaminating the surrounding soil or groundwater from past operations at the site.

Question/comment: Several comments have been received regarding the mitigation of the TNT transite process wastewater line and the two cooling water lines (North and South) as follows:

- All Lines –
 - What is the procedure to plug the lines? Plugging the inlets and outlets to prevent water from infiltrating and transporting may not be correct because the lines may develop leaks, as the seals will eventually fail.



- What is the length of the lines, diameter, how deep they are buried, and about how much it would cost to remove or excavate and dispose of the lines off site?
- TNT process wastewater line – How will it be determined whether high concentrations of contaminants remain in the TNT wastewater line after it is flushed? How will lines be sampled if not accessible by a manhole?
- Cooling water lines –
 - For the RD evaluation only two locations along the northern cooling water line will be sampled. This is not sufficient to characterize contamination along the entire northern line. High concentrations of contaminants have been found at other locations along the northern line. Most of the line has not been sampled because it is not accessible through manholes.
 - The southern cooling water line will not be sampled for the RD evaluation even though high concentrations of contaminants have been found in the line. As with the northern line, most of the southern line has not been sampled because it is not accessible through manholes.
 - If the cooling water lines are flushed, the Army has not explained how it will determine whether high concentrations of contaminants remain in the lines after flushing.

Response: *To clearly describe the lines and the approach to the selected remedy, it must be understood that there are two distinctly different processes that generated the liquid flowing through these lines and remaining residue. Thus, a discussion of each line to address the above comments is presented below:*

TNT Process Wastewater Line – *The TNT wastewater line carried away process wastewater from the washing process during TNT production (also known as “red water, yellow water, red liquor and yellow liquor”). The TNT wastewater line carried wash water from the process and is expected to have some solids in it (5 to 15%). The wastewater was pumped and treated at the wastewater treatment plant (LHAAP-32). This line was originally installed as a wooden line before the transite line was installed. Historical documentation (Bate Stamp 001446, RCRA FA, April 1988) indicated a wooden TNT line (a.k.a. red water line) was used for a short time before the transite line was put into service. The wooden line was “clear-flushed” in 1946 and abandoned. It was determined that no further action was necessary for this line (Bate Stamp 001446). During investigation activities, the wooden TNT wastewater line was found 5 feet south of the transite TNT process wastewater line. The wooden line was cut to allow sampling of the contents. Although the sample results indicated the presence of explosives, the data was later determined by EPA to be unsuitable for environmental decision-making. Because the line is deteriorated and has been cut in multiple locations, the line cannot be flushed. Instead, the line contents and surrounding soil will be sampled and then excavated and disposed offsite if required.*

The TNT transite process wastewater line is approximately 3 feet bgs. The gravity flow portion of the line is approximately 3,000 linear feet. The pressurized portion of the line is approximately 500 linear feet, and the lines are in good condition. The diameter of the TNT transite process wastewater line along the entire line is not known but should range from 8 inches to 18 inches based on the original



design of the wooden stave line. There are no manholes in the transite line, and the line will be cut at select locations for any additional sampling and the implementation of the remedy.

For the investigation, the gravity fed portion of the line was cut at two locations and a thick viscous residue was observed. Samples were collected at these two locations. Samples were also collected at two points along the pressurized portion of the line that extends from the former pump house location to LHAAP-32, located north of LHAAP-29. Since explosives were identified in the viscous residue in the gravity fed portion of the line, it is assumed that this residue remains throughout the line and the samples were representative.

Thus, the selected remedy is to flush the line with potable water. The exact procedure will be developed in the RD, but generally the flushing will be conducted in segments since the line was broken during the investigation phase and the line is no longer a single continual run of pipe. Visual verification that the residue is not sticking to the sides of the pipe will also be conducted. The flush water will be containerized, sampled for waste characterization, and properly treated or disposed. Flushing will ensure that no residual material is left in the lines. Thus, even if the pipes break in the future, there is no contamination remaining to leach. After flushing, the inlets and outlets of the TNT wastewater line will be plugged with a bentonite slurry mix or cementitious grout to stop future infiltration. The procedures for the plugging activities will be included in the RD.

The transite TNT wastewater line is a combination of cement and asbestos that, when disturbed, can become friable and pose additional risk. Exposure to friable asbestos requires special handling and disposal requirements, increasing risks to workers through exposure by the process of digging the lines out of the ground (causing the material to become friable), handling the pieces onsite and during the subsequent transportation and disposal. The location of the line remaining in place will be surveyed and filed with the County. Since the TNT wastewater line is asbestos, the county notification will provide information on the existence and location of the transite line in order to avoid disturbing the line in the event future excavation or other activities are being planned in the area. Additionally, the depth of the line is deeper than the 2 feet bgs, the depth used for industrial use. Since no residue would remain in the lines after flushing, no contamination would remain in the line or have the potential to contaminate the surrounding soil or groundwater even if the line would deteriorate. Additionally, the removal of the line will impact a large area of the site that is currently covered with mature trees and native vegetation.

Cooling Water Lines – The TNT manufacturing process generated a lot of heat; cold water was used to cool the reaction equipment. It flowed over the equipment and down a drain into the cooling water drain lines. Thus, the cooling water lines carried water and would not have carried solids. There were two lines, the North and South Cooling Water (a.k.a. blue cooling water lines) lines, that were gravity fed lines constructed of vitrified clay pipe with manholes. The cooling water from six production plants flowed into the two main collection lines (North and South). The main cooling water lines are approximately 5,000 feet, with approximately 280 feet of smaller line at each of the six production plants. The lines are approximately 8 feet bgs and range from 8 to 18 inches in diameter. The lines drain into a ditch along 16th Street which eventually flows into Goose Prairie Creek.

Small amounts of solid sediment residue and water were found during the investigations of the pipeline. Samples collected from some of the manholes detected explosives in both the solid residue and water at concentrations above the cleanup levels. These lines only carried water, and the solids found in the manholes during investigations are expected to be from open inlets after demolition of



the facility and through the open manholes. The manholes are low spots in the lines where several lines intersect and solid residue would collect at these locations. It stands to reason that minimal solid residue is expected to be found in the actual lines between the manholes. An attempt will be made to collect additional solid residue samples from the manholes during the remedial action.

For the selected remedy, each manhole and outlet will be inspected prior to flushing. The exact procedure will be developed in the RD, but generally the flushing will be conducted between two manholes. The rinsate water will be containerized, transported, sampled, analyzed, and treated at the onsite groundwater treatment plant or appropriately disposed off-site based on the explosives concentrations. After flushing of the lines, there would be no solid residue remaining in the pipe to conduct additional sampling of the solids; however, the flush water will be tested. Thus, even if the pipes break in the future, there is no contamination remaining to leach. The manholes will then be plugged with a bentonite slurry mix or cement grout. The procedures for the plugging activities will be included in the RD.

Question/comment: The Army claims that "...soil samples collected near the line indicate there has not been a release to the surrounding soil." This is incorrect. Sample collected from soil borings along the line were found to contain TNT, 2-amino-4,6-DNT, and 4-amino-2,6-DNT.

Response: *The commenter is referencing the samples collected near the wooden TNT process wastewater line, which is 3 feet bgs. Soil samples were collected in 2004 near the line 4 to 5 feet bgs to evaluate the potential of contaminants leaching from the line and contaminating the surrounding soil. Table 2-6 from the FS summarizes the detected results. Even though samples were found to contain TNT, 2-amino-4,6-DNT, and 4-amino-1,6-DNT, the concentrations are less than both the SAI-Ind and GWP-Ind. Sampling of the wooden TNT process wastewater line and associated soil will take place during the remedial design phase, and if the results indicate unacceptable levels of explosives are present, excavation and disposal off-site of the material will take place during the remedial action phase.*

Question/comment: *The Army estimates that it will take 90 years for natural attenuation to reduce contaminant concentrations acceptable levels. It is not reasonable to propose a plan that could require the maintenance of LUCs for almost a century.*

Response: *All alternatives evaluated have a long time for restoration. Given the nature of the residual contaminants that are present at LHAAP-29, the length of time that will be required to achieve cleanup levels would be long for any of the remedial alternatives, whether treatment, migration control, or source control by removal. The reasonably anticipated future use of the site is as a national wildlife refuge (i.e. Caddo Lake National Wildlife Refuge). Once the property is transferred into the refuge system, the property must be kept as a National Wildlife Refuge unless there is an act of Congress which removes the parcel or the land is exchanged in accordance with the National Wildlife Refuge System Administration Act of 1966 and the National Wildlife Refuge System Act Amendments of 1974. This proposed transfer as a national wildlife refuge, which by its very nature includes physical access and use restrictions, is subject to control and continual inspection by Refuge personnel. Also, the property is intended to remain under ownership and management of a federal government agency. The LUC will restrict access to the groundwater for purposes other than environmental testing until cleanup levels are met. Additionally, access of groundwater through well installations requires a permit from the Texas Department of Licensing and Regulation or Texas Water District authority. The department will be provided a copy of the county*



recordation that indicates the location of contaminated groundwater at the site and associated restriction.

Question/comment: The extremely high concentration of methylene chloride in the intermediate groundwater zone indicates that DNAPL may be present. If so, it may be very difficult to clean up the groundwater. The Army does not appear to have developed any plans to deal with DNAPL. The Army should explain how it will deal with any DNAPL that may be present.

Response: *The current active remedy to conduct in-situ thermal desorption will reduce contaminant concentrations even if DNAPL is present.*

Question/comment: High concentrations of arsenic, mercury, and selenium exist in the shallow groundwater zone. High concentrations of arsenic exist in the deep groundwater zone. The metals are not subject to natural attenuation by biodegradation and the Army has not explained how they will be cleaned up. The Army should explain how it will cleanup these metals.

Response: *The elevated concentrations of metals were suspected to be a result of sampling methodology and/or turbid samples which could falsely elevate the metals concentration result due to excess solids in the sample. Subsequent sampling and redevelopment of some of the wells resulted in lowered concentrations of some metals.*

The arsenic sample collected from the deep well had high aluminum concentrations; the geochemical relationship between the high aluminum in the same sample as the arsenic indicates arsenic may be naturally occurring. Mercury detections are intermittent and appear to be isolated and the calculated Hazard Index for mercury is less than 1, based on the Final Baseline Human Health and Screening Ecological Risk Assessment for Group 2 Sites, from Jacobs Engineering Corp., 2002.

The extent of arsenic and mercury in groundwater will be assessed site-wide during the remedial design. Selenium was only detected above the MCL in one shallow well and the concentrations have fluctuated over the years. The most recent concentration has an associated adjusted hazard quotient value of 0.15. Selenium will be added as a contaminant of concern.

Question/comment: The current distribution of groundwater contaminants at site 29 is not well known. This is because the Army has not sampled many of the monitor wells (18 of 47) since 2005, and has not sampled any monitor well since 2009. In addition, many of the most recent sample analyses are incomplete. No wells have been analyzed for explosives or perchlorate since February 2007.

The only way to ensure that the current distribution of contaminants is known is to sample all the monitor wells at the site. They should be sampled for solvents (e.g., methylene chloride, TCE), explosives (e.g., TNT, 2,4-DNT), metals (e.g., arsenic, mercury), and perchlorate. The Army should do this before it completes the design of the groundwater cleanup plan.

Response: *The analytical suites selected for investigations were based on the past operations that could cause potential contamination at LHAAP-29. Initial investigations had several analytical suites which were reduced to refine the nature and extent of those chemicals that were detected above screening levels. During the remedial design phase, additional data will be gathered to verify plume boundaries, develop monitoring networks, and determine concentrations for the treatment.*



Question/comment: The Army's proposed cleanup plan does not address high concentrations of dangerous chemicals in surface water. The Army should either 1) explain why it is unnecessary to cleanup surface water, or 2) prepare a plan to clean it up.

Response: *The surface water sample collected at 29SW46 has the highest contamination and is located at the cooling water ditch outfall. Both surface water and sediments at this location were sampled and concentrations in the surface water and sediments were above the groundwater MSC for residential use and GWP-Ind levels for explosives, respectively. This location is the outfall of the cooling water lines and is collocated with the high sediment levels which will be removed as part of the selected remedy. The next sample location downgradient of 29SW46 did not show high concentrations of contaminants in the surface water. Thus, the action to remove residual contamination from the cooling water lines (so water cannot infiltrate them and carry contamination into the ditch) and to remove the contaminated sediment where the surface water sample was collected will mitigate impacts to surface water along the cooling water ditch. Also, as part of this action, surface water monitoring will be conducted downgradient of this mitigated area.*

Question/comment: Major components of the cleanup are yet to be determined. These include the evaluation of the cooling water lines, and details of: the plan to monitor metals near the in-situ oxidation area, the soil excavation plan, the groundwater monitoring plan, and MNA implementation. These components will be presented in the RD, which will be completed after the Record of Decision is published. Given the importance these components, the Army should make the RD available for public review and comment as soon as possible.

Response: *The public will be provided with updates on remedial design and remedial action status through the RAB meetings and any concerns can be addressed through this forum. The RAB meetings provide the forum for dissemination of information and discussion. The RD will include performance objectives, schedule and other design criteria and will follow established regulatory guidance for the components of the remedy. The RD will also become part of the Administrative Record and will be available to the public at the Marshall Public Library, the public repository for the LHAAP Administrative Record.*

Question/comment: Recommend expanding environmental testing to include all six isomer of DNT (2,4-DNT, 2,6-DNT, 3,4-DNT, 2,3-DNT, 2,5-DNT and 3,5-DNT), site characterization and evaluation of human health risk should incorporate and consider all six isomers of DNT in all media, and corresponding enforceable remedial goals should be established for the four minor isomers of DNT for soils, sediments, groundwater, and surface water. In the event the presence of minor isomers of DNT is confirmed, there may be potential impacts to aquatic ecosystems that should be addressed.

Response: *The selected remedy will address and monitor the explosive contaminated shallow groundwater and surface water. At this time, there are no Federal or State of Texas promulgated screening levels for DNT isomers, other than for 2,4-DNT and 2,6-DNT. However, as part of the CERCLA process, the statutory five-year reviews will evaluate the effectiveness of the remedy, including any changes in applicable or relevant and appropriate requirements (ARARs) concerning DNT isomers, and would recommend implementation of other measures if needed.*



3.2 Technical and Legal Issues

This section is used to expand on technical and legal issues. However, there are no issues of that nature beyond the technical issues already discussed in **Section 3.1**.



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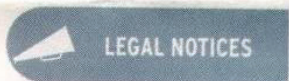


Appendix A
Public Notice Affidavits



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PUBLIC NOTICE

THE UNITED STATES ARMY INVITES PUBLIC COMMENT ON THE REVISED PROPOSED PLAN FOR ENVIRONMENTAL SITE LHAAP-29, FORMER TNT PRODUCTION AREA LONGHORN ARMY AMMUNITION PLANT, TEXAS PUBLIC MEETING ON December 6, 2018 AT THE KARNACK COMMUNITY CENTER, KARNACK, TX

The U.S. Army is the lead agency for environmental response actions at Longhorn Army Ammunition Plant. In partnership with the U.S. Environmental Protection Agency Region 6 (USEPA), the lead Oversight Agency, and Texas Commission on Environmental Quality, the Supporting Agency, the U.S. Army has developed the Revised Proposed Plan for site LHAAP-29, Former TNT Production Area. Although the Revised Proposed Plan identifies the preferred remedy for the site, the U.S. Army welcomes the public's review and comments. Beginning on November 21, 2018, copies of the Revised Proposed Plan and supporting documentation will be available for public review at the Marshall Public Library, 300 S. Alamo, Marshall, Texas, 75670 and on the LHAAP website at http://www.longhornaap.com/LHAAP-29. The public comment period is November 21, 2018, through December 21, 2018. The public meeting will be held on Thursday, December 6, 2018 at the Karnack Community Center, Karnack, TX beginning at 6:00 PM and ending at 7:30 PM. The Karnack Community Center is located at Highway 134 and Spur 449 near the front gate of the Caddo Lake National Wildlife Refuge. Questions, comments, and responses on the Revised Proposed Plan will be recorded by a court reporter during the public meeting. Written comments will be accepted throughout the public comment period.

Longhorn Army Ammunition Plant (LHAAP) is an inactive, government-owned, formerly contractor-operated industrial facility located in central-east Texas in the northeastern corner of Harrison County. The installation occupies nearly 8,416 acres between State Highway 43 at Karnack, Texas, and the western shore of Caddo Lake. LHAAP was established in December 1941 near the beginning of World War II for the manufacture of trinitrotoluene. Other past industrial operations at the installation included the production of secondary explosives, rocket motor propellants, and various pyrotechnics. LHAAP was found to have actual and potential releases of hazardous substances or pollutants or contaminants associated with past operations, and it was added to the National Priorities List (NPL) in 1990. LHAAP-29, former TNT Production Area, encompasses approximately 85 acres in the west-central portion of LHAAP. The site was used as a TNT manufacturing facility from October 1942 to August 1945. The facility produced approximately 400 million pounds of flake TNT during its operation using six TNT production lines. Since the end of World War II, the only activity that has been documented to have occurred at LHAAP 29 is the "soak out" or solvent bath of out-of-specification rocket motors. This took place from 1959 to the mid-1970s and involved the use of a methylene chloride-based industrial solvent.

The Revised Proposed Plan for LHAAP-29 addresses contamination in soil, process lines, and groundwater at LHAAP-29. The full list of alternatives evaluated is: 1) No action; 2) Excavation and off-site disposal of soil, flushing and plugging of process lines, monitored natural attenuation (MNA) for shallow groundwater, in situ chemical oxidation followed by MNA for intermediate groundwater, and land use controls (LUCs); 3) Excavation and offsite disposal of soil; flushing and plugging of process lines, groundwater extraction for intermediate groundwater, MNA for shallow groundwater, and LUCs; 4) Excavation and offsite disposal of soil, flushing and plugging of process lines, in situ thermal desorption for intermediate groundwater, MNA for shallow groundwater, and LUCs. Based on available information, the preferred remedy is Alternative 4, which would remove contaminated soil from LHAAP-29 with off-site disposal; flush and plug process lines thereby eliminating potential for contaminants to leach into groundwater; reduce groundwater contamination throughout the intermediate groundwater contaminant plume via in situ thermal desorption and MNA; use MNA for shallow groundwater to assure protection of human health and the environment by documenting that the contaminated groundwater remains localized and contaminant concentrations are being reduced to cleanup levels; and implement LUCs to protect human health by preventing exposure to contaminated media.

For further information or to submit written comments, contact: Dr. Rose M. Zeiler, Longhorn Army Ammunition Plant, P.O. Box 220, Ratcliff, Arkansas, 72951; phone number 479-635-0110 or email rose.m.zeiler.civ@mail.mil.

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INVITES PUBLIC COMMENT ON THE REVISED
PROPOSED PLAN FOR ENVIRONMENTAL SITE
LHAAP-29, FORMER TNT PRODUCTION AREA
LONGHORN A

Notice published in the Times on 11/07/18

(Signed) *M. Scuin*

Melanie C Altz 11/7/2018
(Notary)



PUBLIC NOTICE

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added to the National Priorities List (NPL) in 1990. LHAAP-29, former TNT Production Area, encompasses approximately 85 acres in the west-central portion of LHAAP. The site was used as a TNT manufacturing facility from October 1942 to August 1945. The facility produced approximately 400 million pounds of flake TNT during its operation using six TNT production lines. Since the end of World War II, the only activity that has been documented to have occurred at LHAAP 29 is the "soak out" or solvent bath of out-of-specification rocket motors. This took place from 1959 to the mid-1970s and involved the use of a methylene chloride-based industrial solvent.

The Revised Proposed Plan for LHAAP-29 addresses contamination in soil, process lines, and groundwater at LHAAP-29. The full list of alternatives evaluated is: 1) No action; 2) Excavation and off-site disposal of soil, flushing and plugging of process lines, monitored natural attenuation (MNA) for shallow groundwater, in situ chemical oxidation followed by MNA for intermediate groundwater, and

land use controls (LUCs); 3) Excavation and offsite disposal of soil; flushing and plugging of process lines, groundwater extraction for intermediate groundwater, MNA for shallow groundwater, and LUCs; 4) Excavation and offsite disposal of soil, flushing and plugging of process lines, in situ thermal desorption for intermediate groundwater, MNA for shallow groundwater, and LUCs. Based on available information, the preferred remedy is Alternative 4, which would remove contaminated soil from LHAAP-29 with off-site disposal; flush and plug process lines thereby eliminating potential for contaminants to leach into groundwater; reduce groundwater contamination throughout the intermediate groundwater contaminant plume via in situ thermal desorption and MNA; use MNA for shallow groundwater to assure protection of human health and the environment by documenting that the contaminated groundwater remains localized and contaminant concentrations are being reduced to cleanup levels; and implement LUCs to protect human health by preventing exposure to contaminated media. For further information or to submit written comments, contact: Dr. Rose M. Zeiler, Longhorn Army Ammunition Plant, P.O. Box 220, Ratcliff, Arkansas, 72951; phone number 479-635-0110 or email rose.m.zeiler.civ@mail.mil.

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