

Cover Page

Order ID : Q3164

Project ID : AEC-2025-0013 CSC 4151 Coram

Client : ATG-GREENVILLE AEC

Lab Sample Number

Q3164-01
Q3164-02
Q3164-03

Client Sample Number

DA-1
DA-2
DA-3

I certify that the data package is in compliance with the terms and conditions of the contract, both technically and for completeness, for other than the conditions detailed above. Release of the data contained in this hard copy data package has been authorized by the laboratory manager or his designee, as verified by the following signature.

Signature : _____

Date: 9/27/2025

NYDOH CERTIFICATION NO - 11376

NJDEP CERTIFICATION NO - 20012

DATA REPORTING QUALIFIERS- ORGANIC

For reporting results, the following “ Results Qualifiers” are used:

Value	If the result is a value greater than or equal to the detection limit, report the value
U	Indicates the compound was analyzed for but was not detected. Report the minimum detection limit for the sample with the U, i.e. “10 U”. This is not necessarily the instrument detection limit attainable for this particular sample based on any concentration or dilution that may have been required.
ND	Indicates the analyte was analyzed for, but not detected
J	Indicates an estimated value. This flag is used: (1) When estimating a concentration for a tentatively identified compound (library search hits, where a 1:1 response is assumed.) (2) When the mass spectral data indicated the identification, however the result was less than the specified detection limit greater than zero. If the detection limit was 10ug/L and a concentration of 3 ug/L was calculated report as 3 J. This flag is used when similar situation arise on any organic parameter i.e. Pest, PCB and others.
B	Indicates the analyte was found in the blank as well as the sample report as “12 B”.
E	Indicates the analyte ‘s concentration exceeds the calibrated range of the instrument for that specific analysis.
D	This flag identifies all compounds identified in an analysis at a secondary dilution factor.
P	This flag is used for Pesticide/PCB target analyte when there is >25% difference for detected concentrations between the two GC columns. The lower of the two values is reported on Form 1 and flagged with a “P”.
N	This flag indicates presumptive evidence of a compound. This is only used for tentatively identified compounds (TICs), where the identification is based on a mass spectral library search. It applies to all TIC results. For generic characterization of a TIC, such as chlorinated hydrocarbon, the flag is not used.
A	This flag indicates that a Tentatively Identified Compound is a suspected aldol-condensation product.
Q	Indicates the LCS did not meet the control limits requirements

APPENDIX A

QA REVIEW GENERAL DOCUMENTATION

Project #: Q3164

Completed

For thorough review, the report must have the following:

GENERAL:

Are all original paperwork present (chain of custody, record of communication,airbill, sample management lab chronicle, login page)

✓

Check chain-of-custody for proper relinquish/return of samples

✓

Is the chain of custody signed and complete

✓

Check internal chain-of-custody for proper relinquish/return of samples /sample extracts

✓

Collect information for each project id from server. Were all requirements followed

✓

COVER PAGE:

Do numbers of samples correspond to the number of samples in the Chain of Custody on login page

✓

Do lab numbers and client Ids on cover page agree with the Chain of Custody

✓

CHAIN OF CUSTODY:

Do requested analyses on Chain of Custody agree with form I results

✓

Do requested analyses on Chain of Custody agree with the log-in page

✓

Were the correct method log-in for analysis according to the Analytical Request and Chain of Custody

✓

Were the samples received within hold time

✓

Were any problems found with the samples at arrival recorded in the Sample Management Laboratory Chronicle

✓

ANALYTICAL:

Was method requirement followed?

✓

Was client requirement followed?

✓

Does the case narrative summarize all QC failure?

✓

All runlogs and manual integration are reviewed for requirements

✓

All manual calculations and /or hand notations verified

✓

QA Review Signature: KETAN PATEL

Date: 09/27/2025

LAB CHRONICLE

OrderID:	Q3164	OrderDate:	9/22/2025 2:37:00 PM
Client:	ATG-GREENVILLE AEC	Project:	AEC-2025-0013 CSC 4151 Coram
Contact:	Staci Bruce	Location:	J22,VOA Lab

LabID	ClientID	Matrix	Test	Method	Sample Date	Prep Date	Anal Date	Received
Q3164-01	DA-1	Water	VOC-BTEX	8260-Low	09/18/25		09/24/25	09/22/25
Q3164-02	DA-2	Water	VOC-BTEX	8260-Low	09/18/25		09/25/25	09/22/25
Q3164-03	DA-3	Water	VOC-BTEX	8260-Low	09/18/25		09/24/25	09/22/25



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Hit Summary Sheet
SW-846

SDG No.: Q3164

Client: ATG-GREENVILLE AEC

Sample ID	Client ID	Matrix	Parameter	Concentration	C	MDL	RDL	Units
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Client ID:

0

Total Voc :

Total Concentration:



QC SUMMARY

Surrogate Summary

SDG No.: **Q3164**

Client: **ATG-GREENVILLE AEC**

Analytical Method: **SW8260-Low**

Lab Sample ID	Client ID	Parameter	Spike	Result	Recovery (%)	Qual	Limits (%)	
							Low	High
Q3164-01	DA-1	1,2-Dichloroethane-d4	50	52.3	105		74	125
		Dibromofluoromethane	50	40.0	80		75	124
		Toluene-d8	50	44.2	88		86	113
		4-Bromofluorobenzene	50	40.7	81		77	121
Q3164-02	DA-2	1,2-Dichloroethane-d4	50	56.3	113		74	125
		Dibromofluoromethane	50	51.6	103		75	124
		Toluene-d8	50	47.6	95		86	113
		4-Bromofluorobenzene	50	44.9	90		77	121
Q3164-03	DA-3	1,2-Dichloroethane-d4	50	51.1	102		74	125
		Dibromofluoromethane	50	40.0	80		75	124
		Toluene-d8	50	44.0	88		86	113
		4-Bromofluorobenzene	50	39.6	79		77	121
VV0924WBL01	VV0924WBL01	1,2-Dichloroethane-d4	50	62.2	124		74	125
		Dibromofluoromethane	50	54.6	109		75	124
		Toluene-d8	50	44.1	88		86	113
		4-Bromofluorobenzene	50	48.0	96		77	121
VV0924WBS02	VV0924WBS02	1,2-Dichloroethane-d4	50	49.0	98		74	125
		Dibromofluoromethane	50	50.0	100		75	124
		Toluene-d8	50	49.5	99		86	113
		4-Bromofluorobenzene	50	52.0	104		77	121
VV0924WBSD0	VV0924WBSD02	1,2-Dichloroethane-d4	50	49.5	99		74	125
		Dibromofluoromethane	50	50.1	100		75	124
		Toluene-d8	50	50.7	101		86	113
		4-Bromofluorobenzene	50	55.0	110		77	121
VV0925WBL01	VV0925WBL01	1,2-Dichloroethane-d4	50	60.1	120		74	125
		Dibromofluoromethane	50	53.3	107		75	124
		Toluene-d8	50	47.3	95		86	113
		4-Bromofluorobenzene	50	44.1	88		77	121
VV0925WBS01	VV0925WBS01	1,2-Dichloroethane-d4	50	45.1	90		74	125
		Dibromofluoromethane	50	47.8	96		75	124
		Toluene-d8	50	47.9	96		86	113
		4-Bromofluorobenzene	50	51.2	102		77	121

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.:	<u>Q3164</u>	Analytical Method:	<u>SW8260-Low</u>
Client:	<u>ATG-GREENVILLE AEC</u>	Datafile :	<u>VV039150.D</u>

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Low	Limits High	RPD
VV0924WBS02	Benzene	20	20.1	ug/L	101			82	109	
	Toluene	20	21.8	ug/L	109			82	110	
	Ethyl Benzene	20	19.9	ug/L	100			83	109	
	m/p-Xylenes	40	43.2	ug/L	108			82	110	
	o-Xylene	20	20.4	ug/L	102			83	109	

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.:	<u>Q3164</u>	Analytical Method:	<u>SW8260-Low</u>
Client:	<u>ATG-GREENVILLE AEC</u>	Datafile :	<u>VV039165.D</u>

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Low	Limits High	RPD
VV0924WBSD02	Benzene	20	18.8	ug/L	94	7		82	109	15
	Toluene	20	20.0	ug/L	100	9		82	110	16
	Ethyl Benzene	20	19.2	ug/L	96	4		83	109	16
	m/p-Xylenes	40	40.4	ug/L	101	7		82	110	15
	o-Xylene	20	19.8	ug/L	99	3		83	109	20

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.:	<u>Q3164</u>	Analytical Method:	<u>SW8260-Low</u>
Client:	<u>ATG-GREENVILLE AEC</u>	Datafile :	<u>VV039171.D</u>

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Low	Limits High	RPD
VV0925WBS01	Benzene	20	19.9	ug/L	100			82	109	
	Toluene	20	21.5	ug/L	108			82	110	
	Ethyl Benzene	20	20.0	ug/L	100			83	109	
	m/p-Xylenes	40	43.1	ug/L	108			82	110	
	o-Xylene	20	21.0	ug/L	105			83	109	

VOLATILE METHOD BLANK SUMMARY

Client ID

VV0924WBL01

Lab Name: Alliance

Contract: ATGG01

Lab Code: ACE

SDG NO.: Q3164

Lab File ID: VV039148.D

Lab Sample ID: VV0924WBL01

Date Analyzed: 09/24/2025

Time Analyzed: 13:09

GC Column: DB-624UI ID: 0.18 (mm)

Heated Purge: (Y/N) N

Instrument ID: MSVOA_V

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES, MS AND MSD:

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
VV0924WBS02	VV0924WBS02	VV039150.D	09/24/2025
DA-1	Q3164-01	VV039160.D	09/24/2025
DA-3	Q3164-03	VV039162.D	09/24/2025
VV0924WBSD02	VV0924WBSD02	VV039165.D	09/24/2025

COMMENTS: _____

VOLATILE METHOD BLANK SUMMARY

Client ID

VV0925WBL01

Lab Name: Alliance

Contract: ATGG01

Lab Code: ACE

SDG NO.: Q3164

Lab File ID: VV039170.D

Lab Sample ID: VV0925WBL01

Date Analyzed: 09/25/2025

Time Analyzed: 11:01

GC Column: DB-624UI ID: 0.18 (mm)

Heated Purge: (Y/N) N

Instrument ID: MSVOA_V

THIS METHOD BLANK APPLIES TO THE FOLLOWING SAMPLES, MS AND MSD:

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
VV0925WBS01	VV0925WBS01	VV039171.D	09/25/2025
DA-2	Q3164-02	VV039173.D	09/25/2025

COMMENTS: _____



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VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name: Alliance

Contract: ATGG01

Lab Code: ACE

SDG NO.: Q3164

Lab File ID: VV039134.D

BFB Injection Date: 09/22/2025

Instrument ID: MSVOA_V

BFB Injection Time: 10:15

GC Column: DB-624UI ID: 0.18 (mm)

Heated Purge: Y/N N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	19.6
75	30.0 - 60.0% of mass 95	51.9
95	Base Peak, 100% relative abundance	100
96	5.0 - 9.0% of mass 95	7.1
173	Less than 2.0% of mass 174	1.1 (1.6) 1
174	50.0 - 100.0% of mass 95	73.3
175	5.0 - 9.0% of mass 174	5.5 (7.5) 1
176	95.0 - 101.0% of mass 174	70.4 (96) 1
177	5.0 - 9.0% of mass 176	4.4 (6.3) 2

1-Value is % mass 174

2-Value is % mass 176

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

CLIENT ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDICC010	VSTDICC010	VV039137.D	09/22/2025	12:26
VSTDICC020	VSTDICC020	VV039138.D	09/22/2025	12:48
VSTDICCC050	VSTDICCC050	VV039139.D	09/22/2025	13:26
VSTDICC100	VSTDICC100	VV039140.D	09/22/2025	13:49
VSTDICC005	VSTDICC005	VV039142.D	09/22/2025	15:37
VSTDICC001	VSTDICC001	VV039143.D	09/22/2025	16:35



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VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name: Alliance

Contract: ATGG01

Lab Code: ACE

SDG NO.: Q3164

Lab File ID: VV039145.D

BFB Injection Date: 09/24/2025

Instrument ID: MSVOA_V

BFB Injection Time: 09:52

GC Column: DB-624UI ID: 0.18 (mm)

Heated Purge: Y/N N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	19.6
75	30.0 - 60.0% of mass 95	51.7
95	Base Peak, 100% relative abundance	100
96	5.0 - 9.0% of mass 95	6.7
173	Less than 2.0% of mass 174	1.4 (1.9) 1
174	50.0 - 100.0% of mass 95	77.1
175	5.0 - 9.0% of mass 174	5.2 (6.7) 1
176	95.0 - 101.0% of mass 174	73.5 (95.3) 1
177	5.0 - 9.0% of mass 176	4.9 (6.6) 2

1-Value is % mass 174

2-Value is % mass 176

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

CLIENT ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDCCC050	VSTDCCC050	VV039146.D	09/24/2025	10:49
VV0924WBL01	VV0924WBL01	VV039148.D	09/24/2025	13:09
VV0924WBS02	VV0924WBS02	VV039150.D	09/24/2025	14:01
DA-1	Q3164-01	VV039160.D	09/24/2025	18:07
DA-3	Q3164-03	VV039162.D	09/24/2025	18:52
VV0924WBSD02	VV0924WBSD02	VV039165.D	09/24/2025	19:59



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VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name: Alliance

Contract: ATGG01

Lab Code: ACE

SDG NO.: Q3164

Lab File ID: VV039167.D

BFB Injection Date: 09/25/2025

Instrument ID: MSVOA_V

BFB Injection Time: 09:14

GC Column: DB-624UI ID: 0.18 (mm)

Heated Purge: Y/N N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	16.2
75	30.0 - 60.0% of mass 95	49.9
95	Base Peak, 100% relative abundance	100
96	5.0 - 9.0% of mass 95	6.3
173	Less than 2.0% of mass 174	1.3 (1.7) 1
174	50.0 - 100.0% of mass 95	72.3
175	5.0 - 9.0% of mass 174	5.7 (7.9) 1
176	95.0 - 101.0% of mass 174	69.9 (96.6) 1
177	5.0 - 9.0% of mass 176	4.4 (6.3) 2

1-Value is % mass 174

2-Value is % mass 176

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

CLIENT ID	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDCCC050	VSTDCCC050	VV039168.D	09/25/2025	09:59
VV0925WBL01	VV0925WBL01	VV039170.D	09/25/2025	11:01
VV0925WBS01	VV0925WBS01	VV039171.D	09/25/2025	11:24
DA-2	Q3164-02	VV039173.D	09/25/2025	12:10

VOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name:	<u>Alliance</u>	Contract:	<u>ATGG01</u>
Lab Code:	<u>ACE</u>	SDG NO.:	<u>Q3164</u>
Lab File ID:	<u>VV039146.D</u>	Date Analyzed:	<u>09/24/2025</u>
Instrument ID:	<u>MSVOA_V</u>	Time Analyzed:	<u>10:49</u>
GC Column:	<u>DB-624UI</u>	ID:	<u>0.18</u> (mm)
		Heated Purge:	(Y/N) <u>N</u>

	IS1 AREA #	RT #	IS2 AREA #	RT #	IS3 AREA #	RT #
12 HOUR STD	558154	4.60	907128	5.53	923608	8.78
UPPER LIMIT	1116310	5.1	1814260	6.032	1847220	9.277
LOWER LIMIT	279077	4.1	453564	5.032	461804	8.277
EPA SAMPLE NO.						
DA-1	739770	4.60	1401751	5.53	1290012	8.78
DA-3	755839	4.60	1427337	5.53	1322483	8.78
VV0924WBL01	463770	4.60	858892	5.53	831675	8.77
VV0924WBS02	492318	4.60	795602	5.53	784865	8.78
VV0924WBSD02	545364	4.60	903558	5.53	885424	8.78

IS1 = Pentafluorobenzene
IS2 = 1,4-Difluorobenzene
IS3 = Chlorobenzene-d5

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.

VOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name:	<u>Alliance</u>	Contract:	<u>ATGG01</u>
Lab Code:	<u>ACE</u>	SDG NO.:	<u>Q3164</u>
Lab File ID:	<u>VV039146.D</u>	Date Analyzed:	<u>09/24/2025</u>
Instrument ID:	<u>MSVOA_V</u>	Time Analyzed:	<u>10:49</u>
GC Column:	<u>DB-624UI</u>	ID:	<u>0.18</u> (mm)
		Heated Purge:	(Y/N) <u>N</u>

	IS4 AREA #	RT #				
12 HOUR STD	551620	11.172				
UPPER LIMIT	1103240	11.672				
LOWER LIMIT	275810	10.672				
EPA SAMPLE NO.						
DA-1	589916	11.17				
DA-3	607726	11.17				
VV0924WBL01	379013	11.18				
VV0924WBS02	468522	11.17				
VV0924WBSD02	510716	11.17				

IS4 = 1,4-Dichlorobenzene-d4

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.

VOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: Alliance Contract: ATGG01
 Lab Code: ACE SDG NO.: Q3164
 Lab File ID: VV039168.D Date Analyzed: 09/25/2025
 Instrument ID: MSVOA_V Time Analyzed: 09:59
 GC Column: DB-624UI ID: 0.18 (mm) Heated Purge: (Y/N) N

	IS1 AREA #	RT #	IS2 AREA #	RT #	IS3 AREA #	RT #
12 HOUR STD	605297	4.60	933338	5.53	901987	8.77
UPPER LIMIT	1210590	5.096	1866680	6.029	1803970	9.273
LOWER LIMIT	302649	4.096	466669	5.029	450994	8.273
EPA SAMPLE NO.						
DA-2	514758	4.60	932817	5.54	926008	8.78
VV0925WBL01	480047	4.60	864912	5.53	870930	8.78
VV0925WBS01	557017	4.60	898902	5.54	871912	8.78

IS1 = Pentafluorobenzene
 IS2 = 1,4-Difluorobenzene
 IS3 = Chlorobenzene-d5

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.

VOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: Alliance Contract: ATGG01
 Lab Code: ACE SDG NO.: Q3164
 Lab File ID: VV039168.D Date Analyzed: 09/25/2025
 Instrument ID: MSVOA_V Time Analyzed: 09:59
 GC Column: DB-624UI ID: 0.18 (mm) Heated Purge: (Y/N) N

	IS4 AREA #	RT #				
12 HOUR STD	535710	11.172				
UPPER LIMIT	1071420	11.672				
LOWER LIMIT	267855	10.672				
EPA SAMPLE NO.						
DA-2	438853	11.18				
VV0925WBL01	413592	11.18				
VV0925WBS01	508947	11.17				

IS4 = 1,4-Dichlorobenzene-d4

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.



SAMPLE DATA

Report of Analysis

Client:	ATG-GREENVILLE AEC	Date Collected:	09/18/25
Project:	AEC-2025-0013 CSC 4151 Coram	Date Received:	09/22/25
Client Sample ID:	DA-1	SDG No.:	Q3164
Lab Sample ID:	Q3164-01	Matrix:	Water
Analytical Method:	8260D	% Solid:	0
Sample Wt/Vol:	5 Units: mL	Final Vol:	5000 uL
Soil Aliquot Vol:	uL	Test:	VOC-BTEX
GC Column:	DB-624UI ID : 0.18	Level :	LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039160.D	1	09/24/25 18:07	VV092425

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	0.15	U	0.15	1.00	ug/L
108-88-3	Toluene	0.14	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	0.13	U	0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	0.24	U	0.24	2.00	ug/L
95-47-6	o-Xylene	0.12	U	0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	52.3		74 - 125	105%	SPK: 50
1868-53-7	Dibromofluoromethane	40.0		75 - 124	80%	SPK: 50
2037-26-5	Toluene-d8	44.2		86 - 113	88%	SPK: 50
460-00-4	4-Bromofluorobenzene	40.7		77 - 121	81%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	740000	4.6			
540-36-3	1,4-Difluorobenzene	1400000	5.532			
3114-55-4	Chlorobenzene-d5	1290000	8.776			
3855-82-1	1,4-Dichlorobenzene-d4	590000	11.172			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039160.D
 Acq On : 24 Sep 2025 18:07
 Operator : SY/MD
 Sample : Q3164-01
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 17 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 DA-1

Quant Time: Sep 25 08:21:14 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

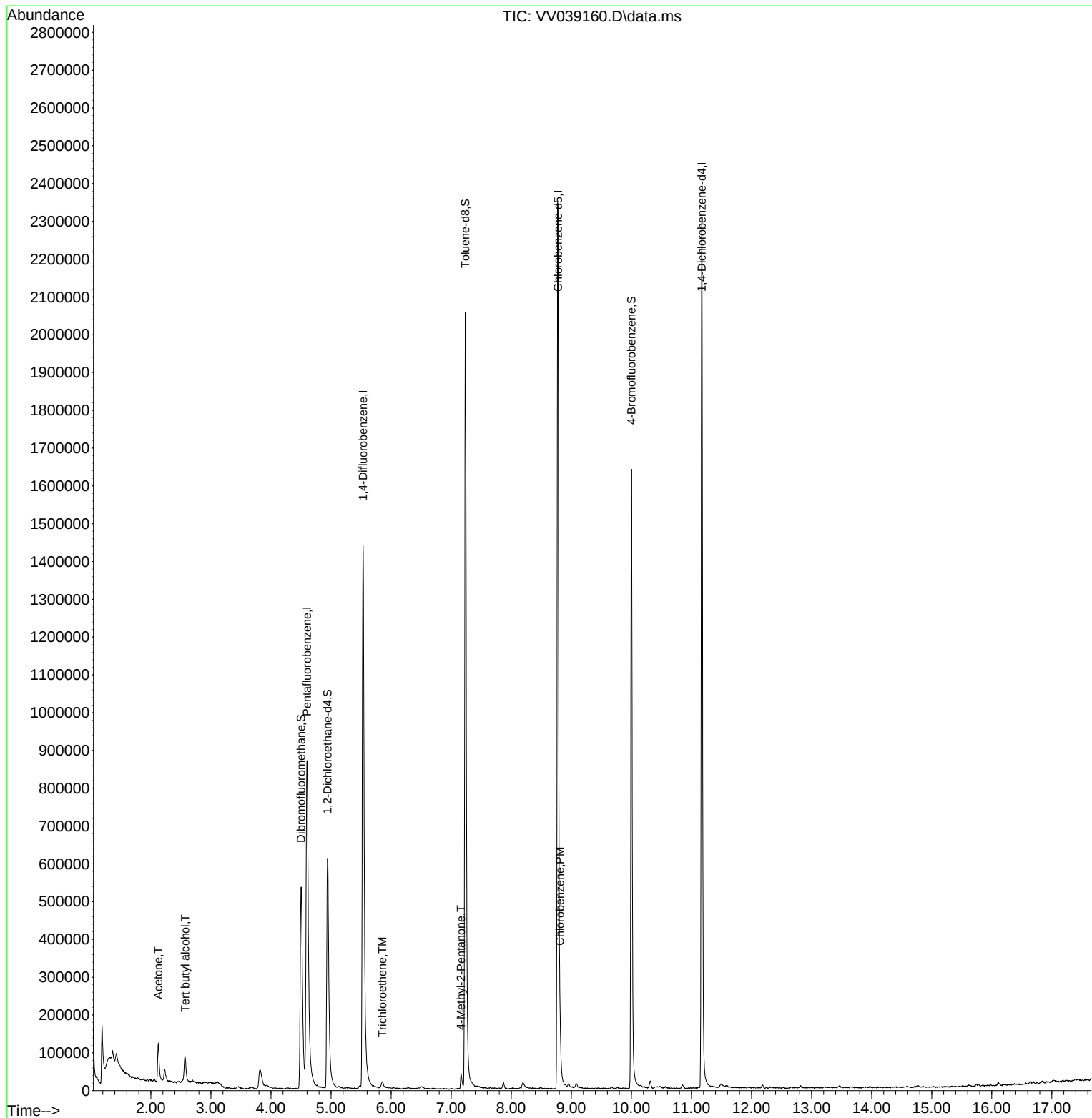
Internal Standards						
1) Pentafluorobenzene	4.600	168	739770	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	1401751	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	1290012	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	589916	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.940	65	560410	52.342	ug/l	0.00
Spiked Amount 50.000	Range 81 - 118		Recovery = 104.680%			
35) Dibromofluoromethane	4.503	113	451140	40.034	ug/l	0.00
Spiked Amount 50.000	Range 80 - 119		Recovery = 80.060%			
50) Toluene-d8	7.236	98	1463568	44.221	ug/l	0.00
Spiked Amount 50.000	Range 89 - 112		Recovery = 88.440%#			
62) 4-Bromofluorobenzene	10.001	95	554931	40.728	ug/l	0.00
Spiked Amount 50.000	Range 85 - 114		Recovery = 81.460%#			
Target Compounds						Qvalue
11) Tert butyl alcohol	2.571	59	42660	18.852	ug/l #	83
16) Acetone	2.124	43	125333	13.774	ug/l	97
44) Trichloroethene	5.854	130	6464	0.505	ug/l	97
51) 4-Methyl-2-Pentanone	7.165	43	24301	1.310	ug/l	98
65) Chlorobenzene	8.808	112	97239	2.706	ug/l	97

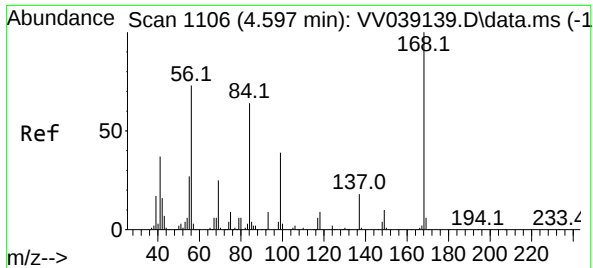
(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039160.D
Acq On : 24 Sep 2025 18:07
Operator : SY/MD
Sample : Q3164-01
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 17 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
DA-1

Quant Time: Sep 25 08:21:14 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

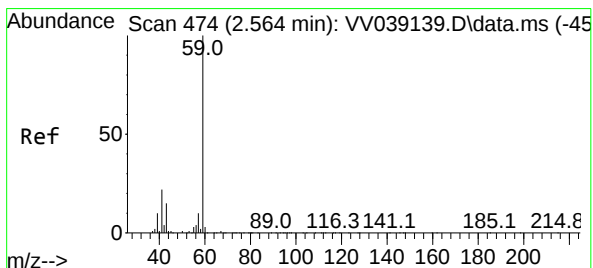
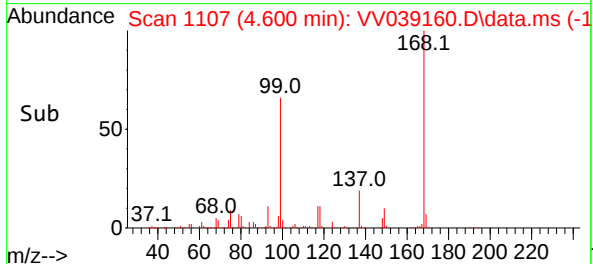
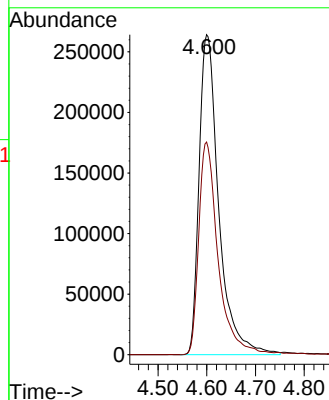
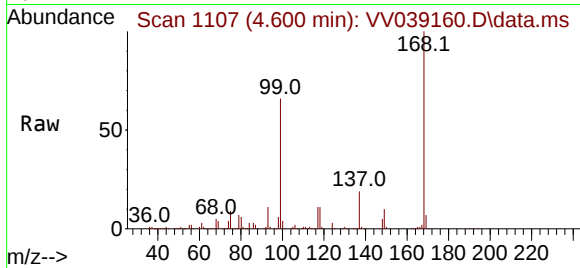




#1
Pentafluorobenzene
Concen: 50.000 ug/l
RT: 4.600 min Scan# 1106
Delta R.T. 0.003 min
Lab File: VV039160.D
Acq: 24 Sep 2025 18:07

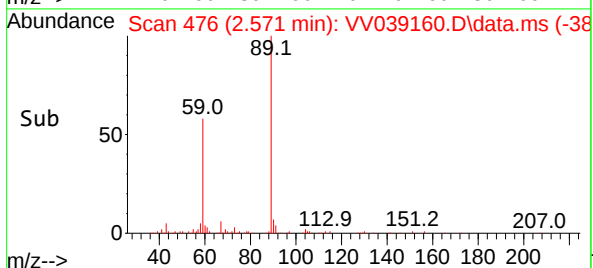
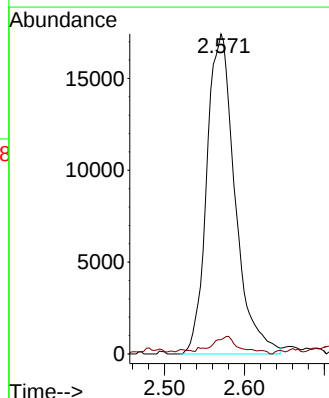
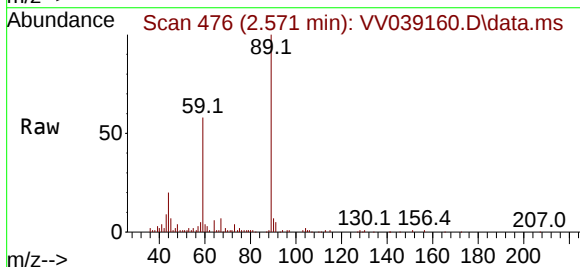
Instrument :
MSVOA_V
ClientSampleId :
DA-1

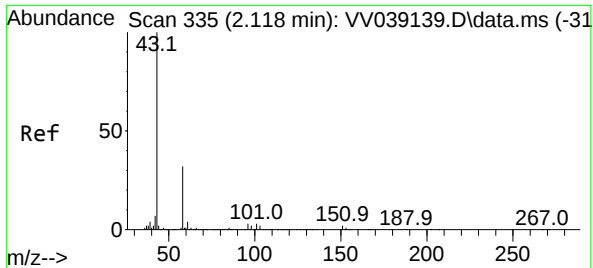
Tgt Ion:168 Resp: 739770
Ion Ratio Lower Upper
168 100
99 66.5 49.6 74.4



#11
Tert butyl alcohol
Concen: 18.852 ug/l
RT: 2.571 min Scan# 476
Delta R.T. 0.006 min
Lab File: VV039160.D
Acq: 24 Sep 2025 18:07

Tgt Ion: 59 Resp: 42660
Ion Ratio Lower Upper
59 100
57 3.4 7.8 11.8#

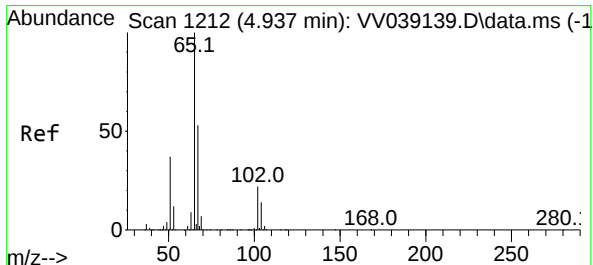
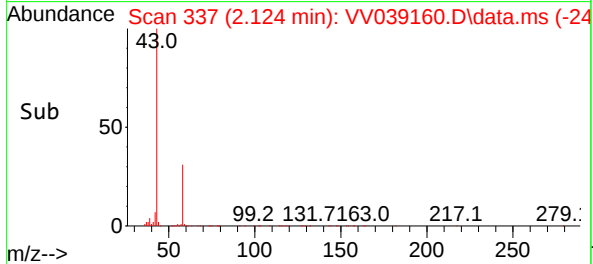
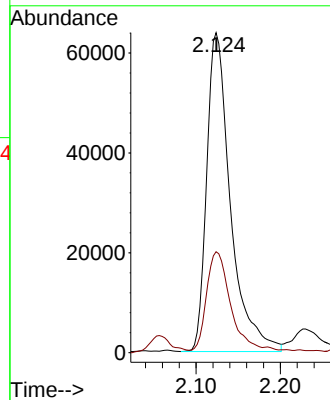
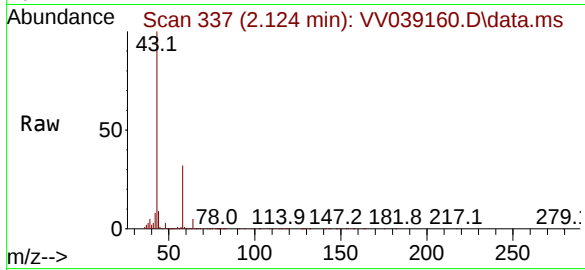




#16
Acetone
Concen: 13.774 ug/l
RT: 2.124 min Scan# 31
Delta R.T. 0.006 min
Lab File: VV039160.D
Acq: 24 Sep 2025 18:07

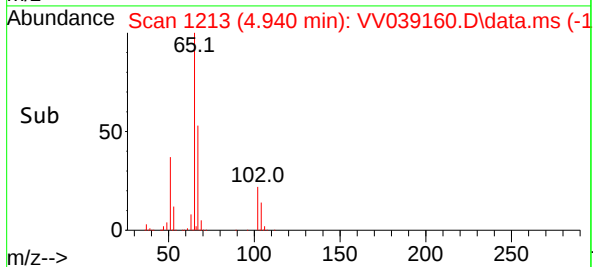
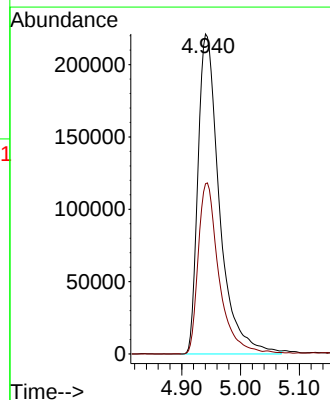
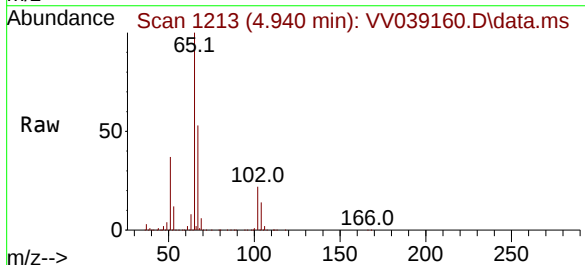
Instrument :
MSVOA_V
ClientSampleId :
DA-1

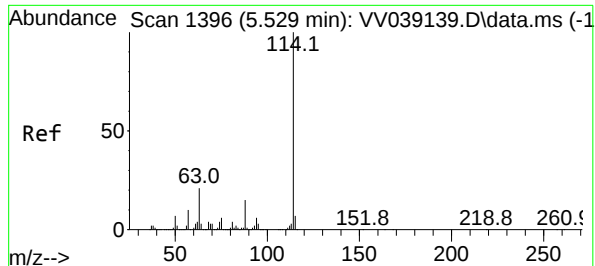
Tgt Ion: 43 Resp: 125333
Ion Ratio Lower Upper
43 100
58 30.8 25.8 38.6



#33
1,2-Dichloroethane-d4
Concen: 52.342 ug/l
RT: 4.940 min Scan# 1213
Delta R.T. 0.003 min
Lab File: VV039160.D
Acq: 24 Sep 2025 18:07

Tgt Ion: 65 Resp: 560410
Ion Ratio Lower Upper
65 100
67 51.3 0.0 107.0





#34

1,4-Difluorobenzene

Concen: 50.000 ug/l

RT: 5.532 min Scan# 114

Delta R.T. 0.003 min

Lab File: VV039160.D

Acq: 24 Sep 2025 18:07

Instrument :

MSVOA_V

ClientSampleId :

DA-1

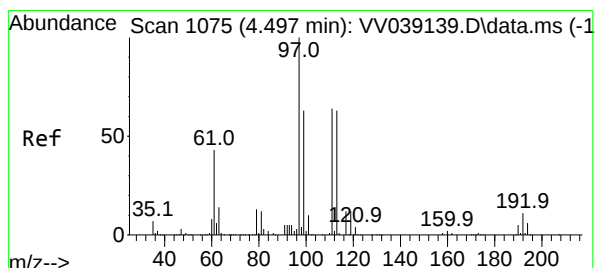
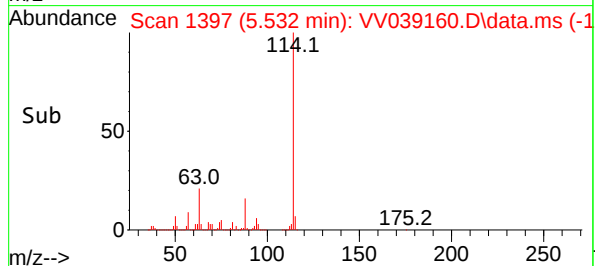
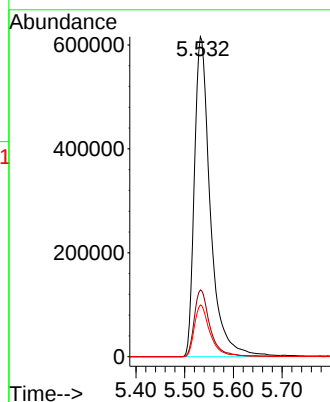
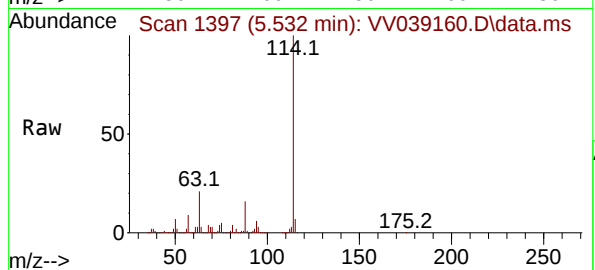
Tgt Ion:114 Resp: 1401751

Ion Ratio Lower Upper

114 100

63 20.8 0.0 42.6

88 16.1 0.0 30.8



#35

Dibromofluoromethane

Concen: 40.034 ug/l

RT: 4.503 min Scan# 1077

Delta R.T. 0.006 min

Lab File: VV039160.D

Acq: 24 Sep 2025 18:07

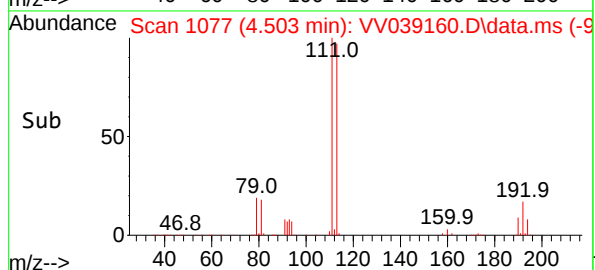
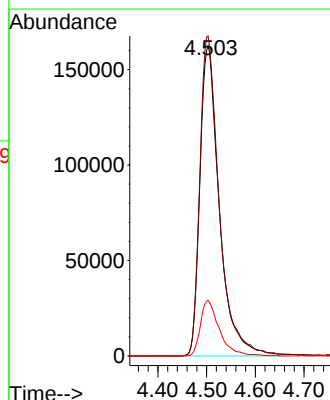
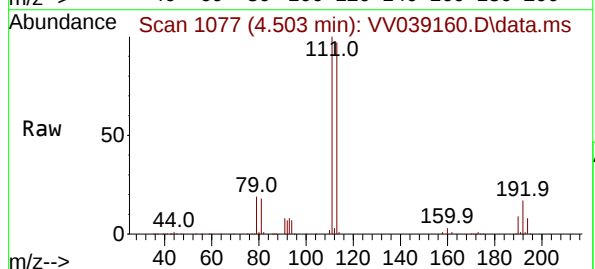
Tgt Ion:113 Resp: 451140

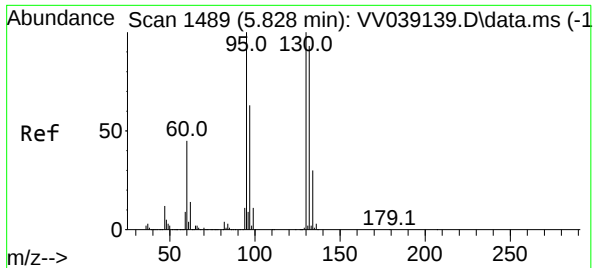
Ion Ratio Lower Upper

113 100

111 102.6 82.4 123.6

192 17.2 13.8 20.8





#44

Trichloroethene

Concen: 0.505 ug/l

RT: 5.854 min Scan# 1489

Delta R.T. 0.026 min

Lab File: VV039160.D

Acq: 24 Sep 2025 18:07

Instrument :

MSVOA_V

ClientSampleId :

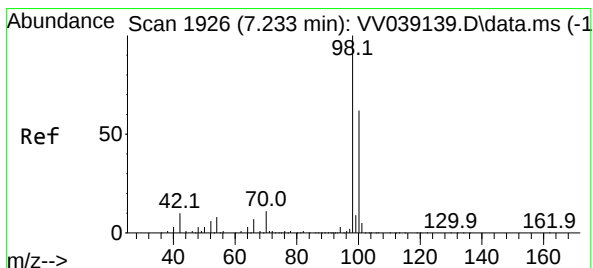
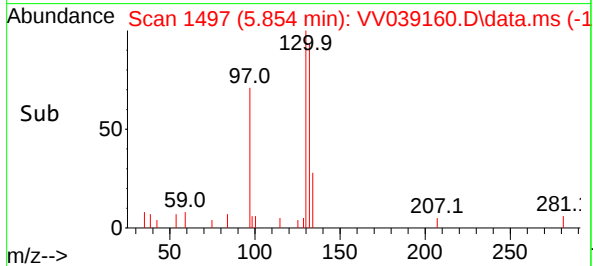
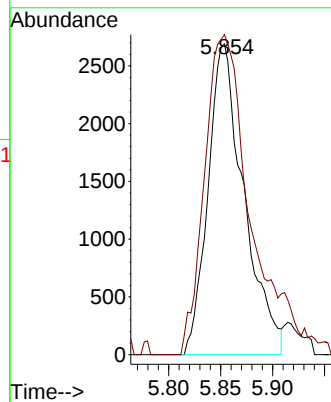
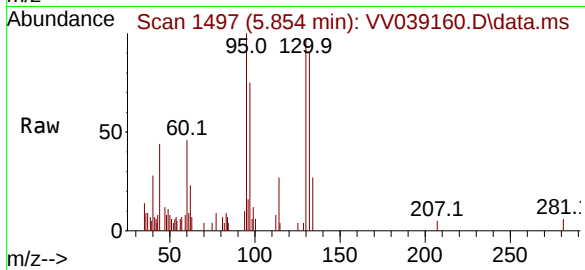
DA-1

Tgt Ion:130 Resp: 6464

Ion Ratio Lower Upper

130 100

95 103.0 0.0 199.2



#50

Toluene-d8

Concen: 44.221 ug/l

RT: 7.236 min Scan# 1927

Delta R.T. 0.003 min

Lab File: VV039160.D

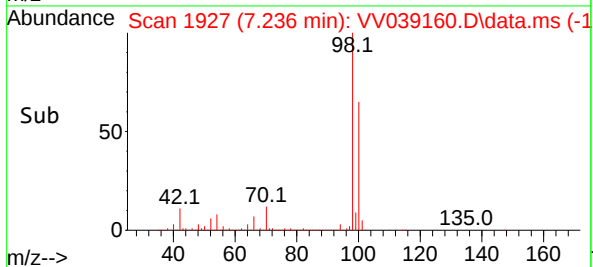
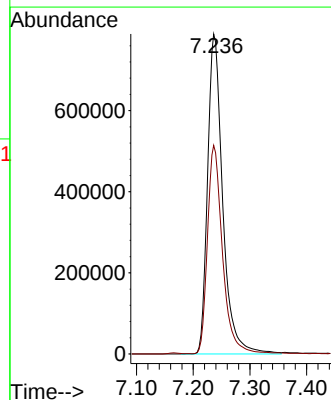
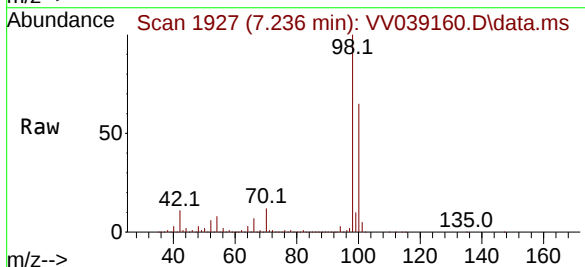
Acq: 24 Sep 2025 18:07

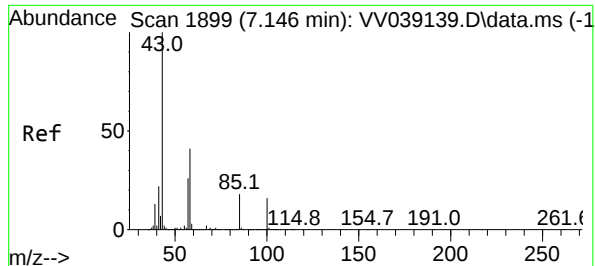
Tgt Ion: 98 Resp: 1463568

Ion Ratio Lower Upper

98 100

100 64.8 52.2 78.2

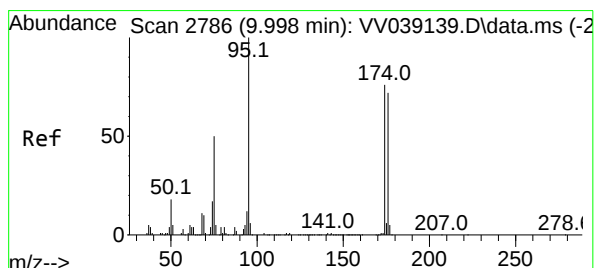
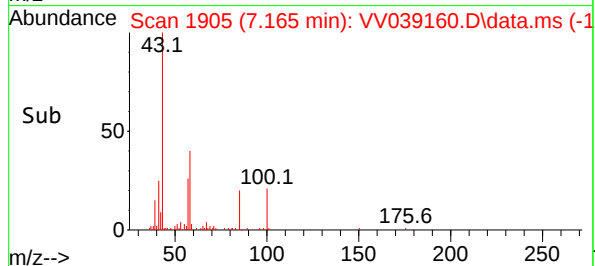
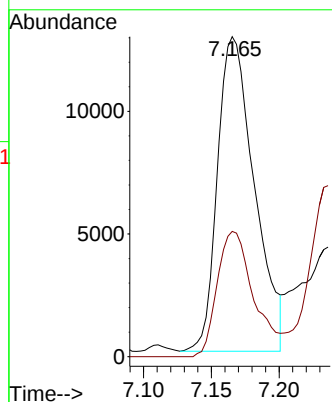
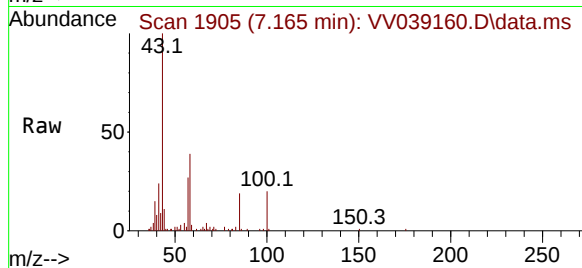




#51
4-Methyl-2-Pentanone
Concen: 1.310 ug/l
RT: 7.165 min Scan# 1905
Delta R.T. 0.019 min
Lab File: VV039160.D
Acq: 24 Sep 2025 18:07

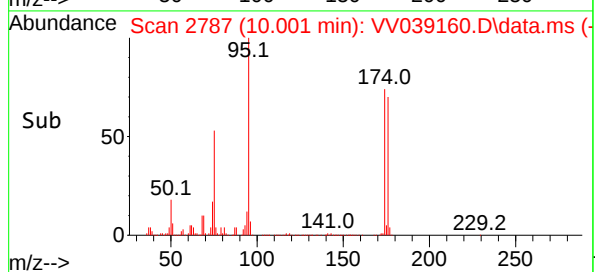
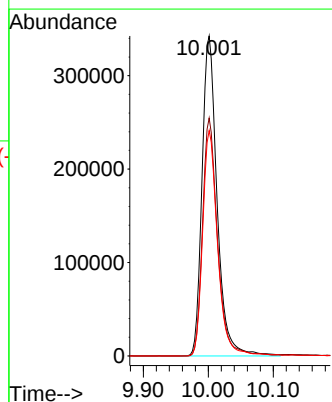
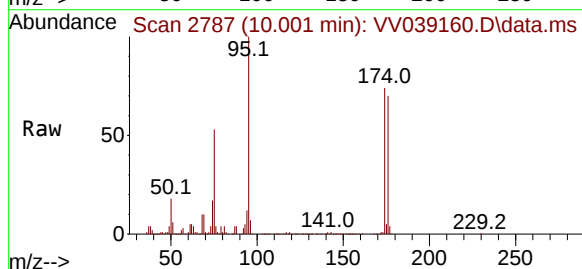
Instrument :
MSVOA_V
ClientSampleId :
DA-1

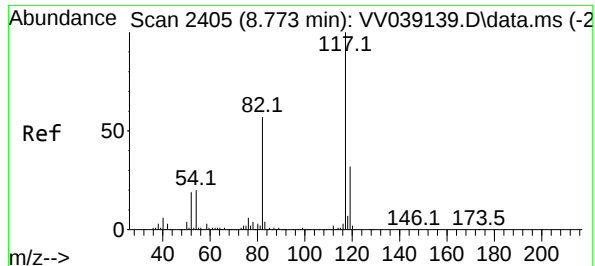
Tgt Ion: 43 Resp: 24301
Ion Ratio Lower Upper
43 100
58 39.4 32.3 48.5



#62
4-Bromofluorobenzene
Concen: 40.728 ug/l
RT: 10.001 min Scan# 2787
Delta R.T. 0.003 min
Lab File: VV039160.D
Acq: 24 Sep 2025 18:07

Tgt Ion: 95 Resp: 554931
Ion Ratio Lower Upper
95 100
174 73.8 0.0 150.4
176 70.1 0.0 144.2





#63

Chlorobenzene-d5

Concen: 50.000 ug/l

RT: 8.776 min Scan# 2405

Delta R.T. 0.003 min

Lab File: VV039160.D

Acq: 24 Sep 2025 18:07

Instrument :

MSVOA_V

ClientSampleId :

DA-1

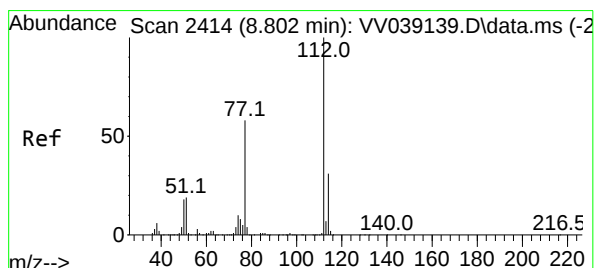
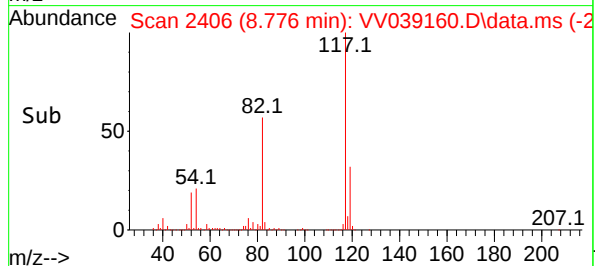
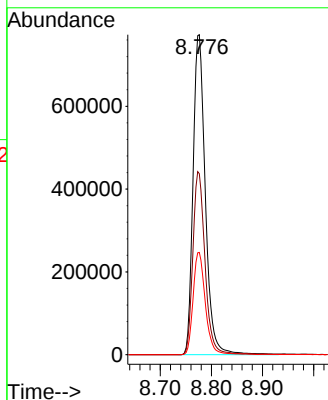
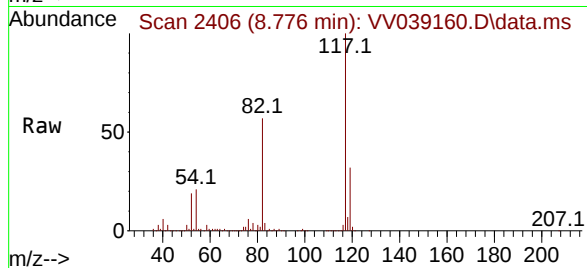
Tgt Ion:117 Resp: 1290012

Ion Ratio Lower Upper

117 100

82 56.6 45.4 68.2

119 32.0 25.6 38.4



#65

Chlorobenzene

Concen: 2.706 ug/l

RT: 8.808 min Scan# 2416

Delta R.T. 0.006 min

Lab File: VV039160.D

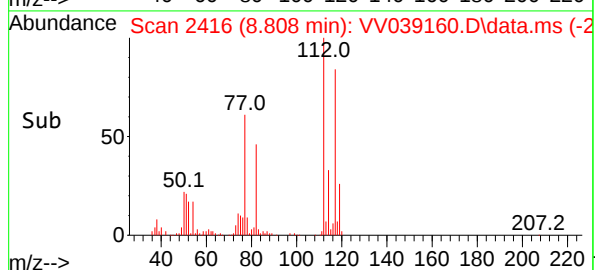
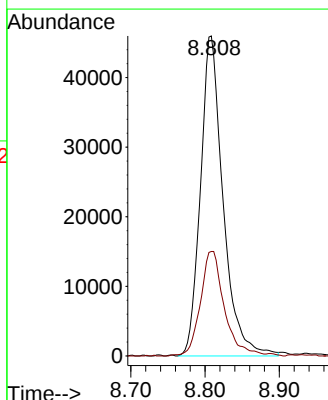
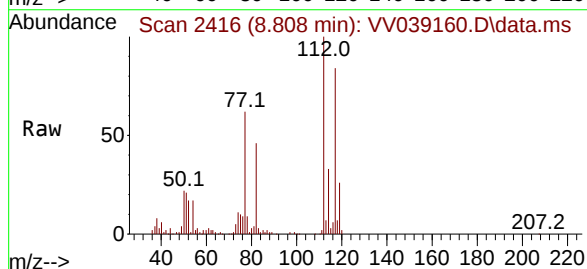
Acq: 24 Sep 2025 18:07

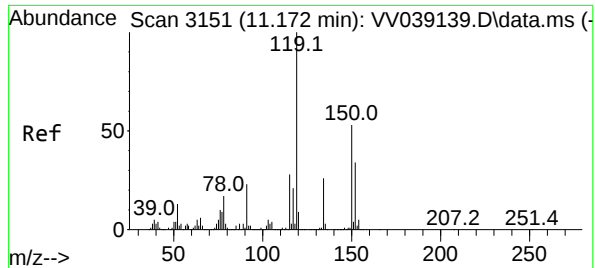
Tgt Ion:112 Resp: 97239

Ion Ratio Lower Upper

112 100

114 32.3 24.6 37.0





#72

1,4-Dichlorobenzene-d4

Concen: 50.000 ug/l

RT: 11.172 min Scan# 31

Delta R.T. -0.000 min

Lab File: VV039160.D

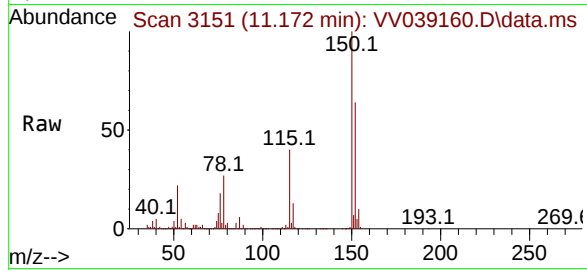
Acq: 24 Sep 2025 18:07

Instrument :

MSVOA_V

ClientSampleId :

DA-1



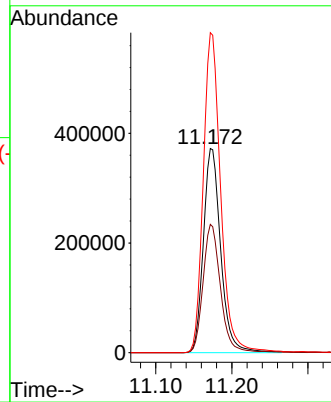
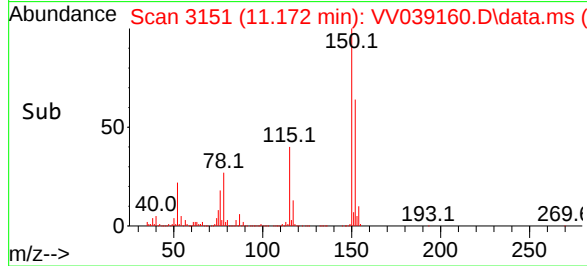
Tgt Ion:152 Resp: 589916

Ion Ratio Lower Upper

152 100

115 63.4 44.8 134.4

150 157.5 0.0 353.8



Report of Analysis

Client:	ATG-GREENVILLE AEC		Date Collected:	09/18/25	
Project:	AEC-2025-0013 CSC 4151 Coram		Date Received:	09/22/25	
Client Sample ID:	DA-2		SDG No.:	Q3164	
Lab Sample ID:	Q3164-02		Matrix:	Water	
Analytical Method:	8260D		% Solid:	0	
Sample Wt/Vol:	5	Units: mL	Final Vol:	5000	uL
Soil Aliquot Vol:		uL	Test:	VOC-BTEX	
GC Column:	DB-624UI	ID : 0.18	Level :	LOW	
Prep Method :					

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039173.D	1	09/25/25 12:10	VV092525

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	0.15	U	0.15	1.00	ug/L
108-88-3	Toluene	0.14	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	0.13	U	0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	0.24	U	0.24	2.00	ug/L
95-47-6	o-Xylene	0.12	U	0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	56.3		74 - 125	113%	SPK: 50
1868-53-7	Dibromofluoromethane	51.6		75 - 124	103%	SPK: 50
2037-26-5	Toluene-d8	47.6		86 - 113	95%	SPK: 50
460-00-4	4-Bromofluorobenzene	44.9		77 - 121	90%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	515000	4.603			
540-36-3	1,4-Difluorobenzene	933000	5.535			
3114-55-4	Chlorobenzene-d5	926000	8.776			
3855-82-1	1,4-Dichlorobenzene-d4	439000	11.175			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039173.D
 Acq On : 25 Sep 2025 12:10
 Operator : SY/MD
 Sample : Q3164-02
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 7 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 DA-2

Quant Time: Sep 26 01:24:03 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

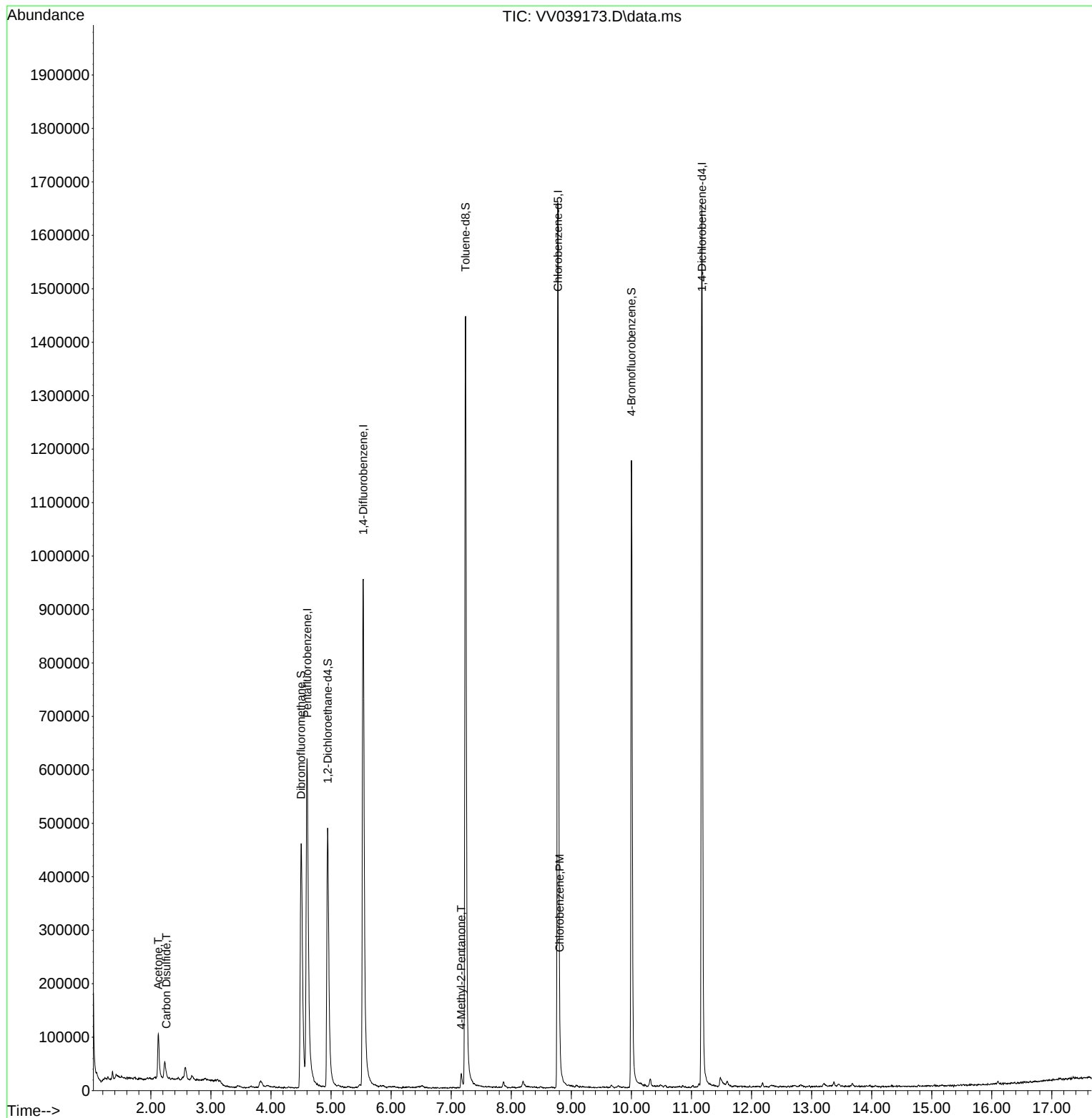
Internal Standards						
1) Pentafluorobenzene	4.603	168	514758	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.535	114	932817	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	926008	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.175	152	438853	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.944	65	419629	56.325	ug/l	0.00
Spiked Amount 50.000	Range 81 - 118		Recovery = 112.660%			
35) Dibromofluoromethane	4.503	113	386984	51.604	ug/l	0.00
Spiked Amount 50.000	Range 80 - 119		Recovery = 103.200%			
50) Toluene-d8	7.240	98	1048304	47.596	ug/l	0.00
Spiked Amount 50.000	Range 89 - 112		Recovery = 95.200%			
62) 4-Bromofluorobenzene	10.001	95	406703	44.855	ug/l	0.00
Spiked Amount 50.000	Range 85 - 114		Recovery = 89.720%			
Target Compounds					Qvalue	
16) Acetone	2.124	43	108214	18.172	ug/l	99
17) Carbon Disulfide	2.259	76	6286	0.265	ug/l	97
51) 4-Methyl-2-Pentanone	7.169	43	16756	1.358	ug/l	91
65) Chlorobenzene	8.805	112	5564	0.216	ug/l	96

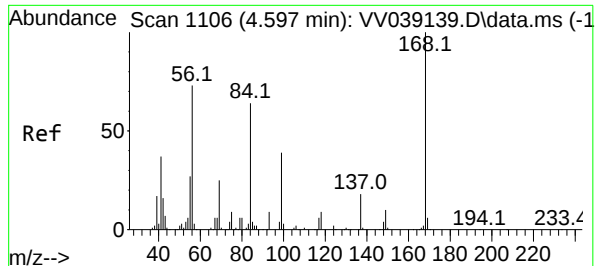
(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
Data File : VV039173.D
Acq On : 25 Sep 2025 12:10
Operator : SY/MD
Sample : Q3164-02
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 7 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
DA-2

Quant Time: Sep 26 01:24:03 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

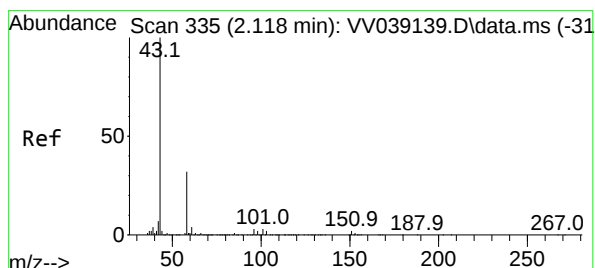
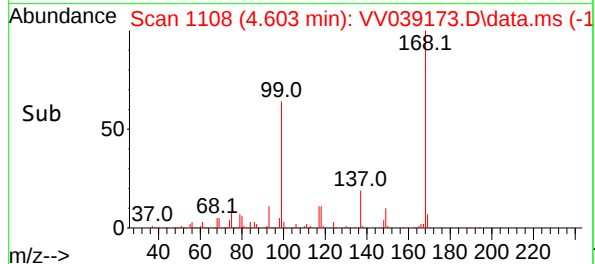
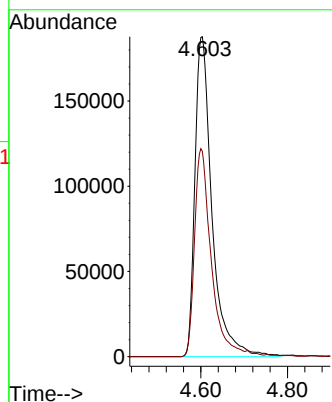
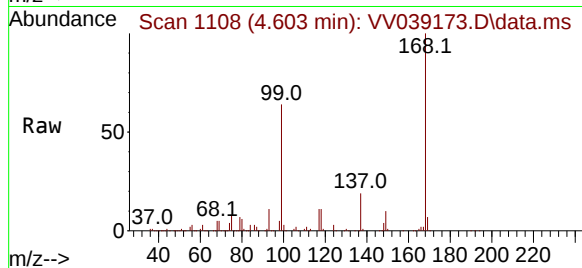




#1
Pentafluorobenzene
Concen: 50.000 ug/l
RT: 4.603 min Scan# 1106
Delta R.T. 0.006 min
Lab File: VV039173.D
Acq: 25 Sep 2025 12:10

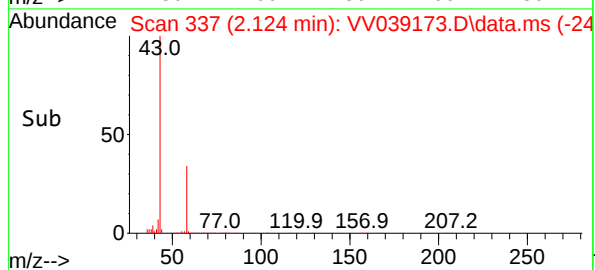
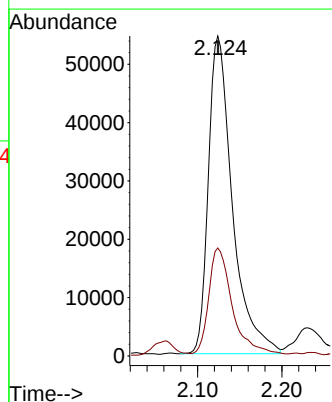
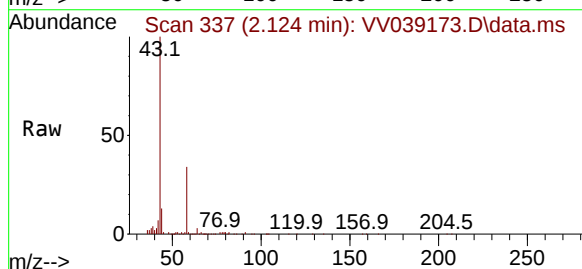
Instrument :
MSVOA_V
ClientSampleId :
DA-2

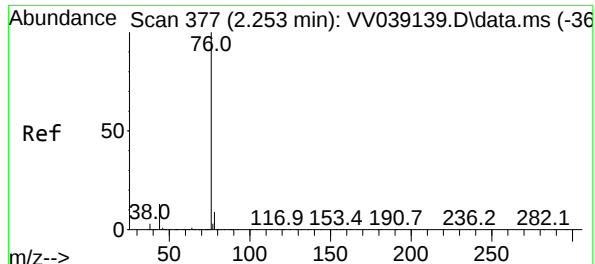
Tgt Ion:168 Resp: 514758
Ion Ratio Lower Upper
168 100
99 64.2 49.6 74.4



#16
Acetone
Concen: 18.172 ug/l
RT: 2.124 min Scan# 337
Delta R.T. 0.006 min
Lab File: VV039173.D
Acq: 25 Sep 2025 12:10

Tgt Ion: 43 Resp: 108214
Ion Ratio Lower Upper
43 100
58 33.0 25.8 38.6





#17

Carbon Disulfide

Concen: 0.265 ug/l

RT: 2.259 min Scan# 31

Delta R.T. 0.006 min

Lab File: VV039173.D

Acq: 25 Sep 2025 12:10

Instrument :

MSVOA_V

ClientSampleId :

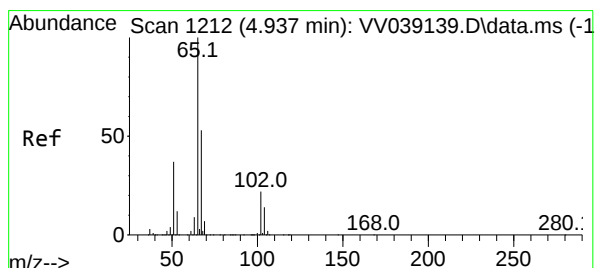
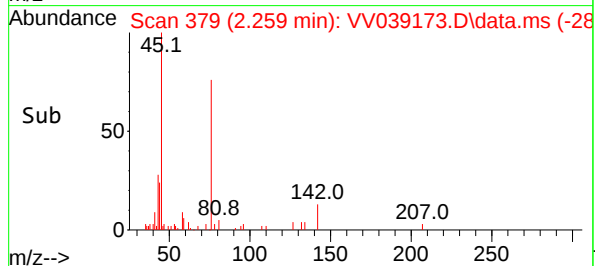
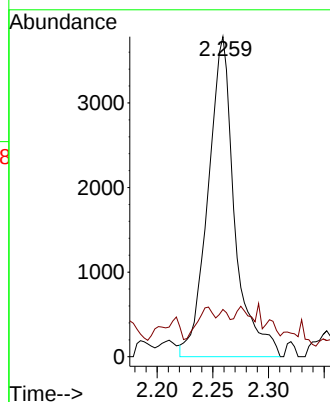
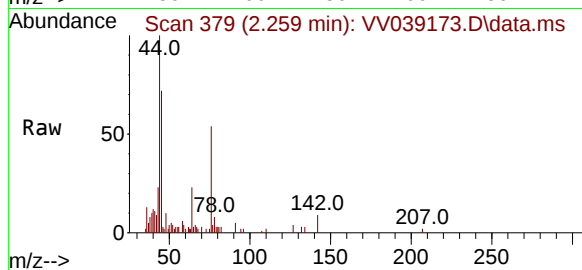
DA-2

Tgt Ion: 76 Resp: 6286

Ion Ratio Lower Upper

76 100

78 8.2 7.4 11.0



#33

1,2-Dichloroethane-d4

Concen: 56.325 ug/l

RT: 4.944 min Scan# 1214

Delta R.T. 0.006 min

Lab File: VV039173.D

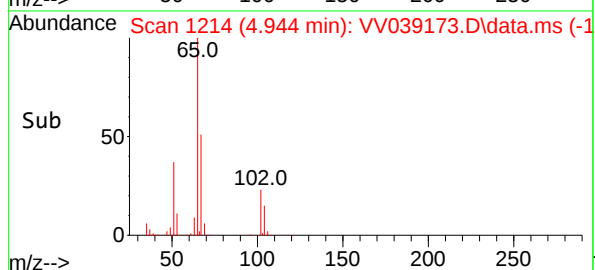
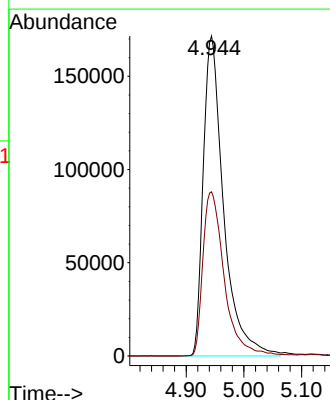
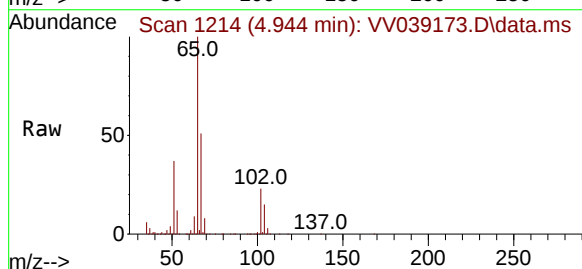
Acq: 25 Sep 2025 12:10

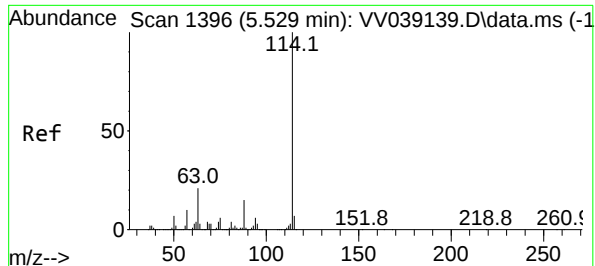
Tgt Ion: 65 Resp: 419629

Ion Ratio Lower Upper

65 100

67 54.1 0.0 107.0





#34

1,4-Difluorobenzene

Concen: 50.000 ug/l

RT: 5.535 min Scan# 11

Delta R.T. 0.006 min

Lab File: VV039173.D

Acq: 25 Sep 2025 12:10

Instrument :

MSVOA_V

ClientSampleId :

DA-2

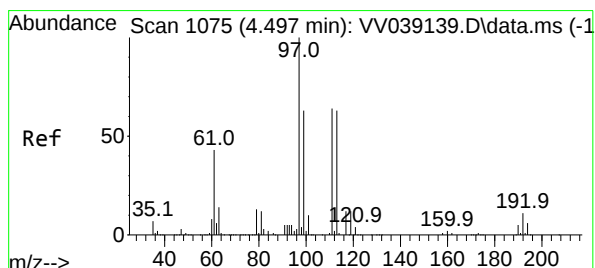
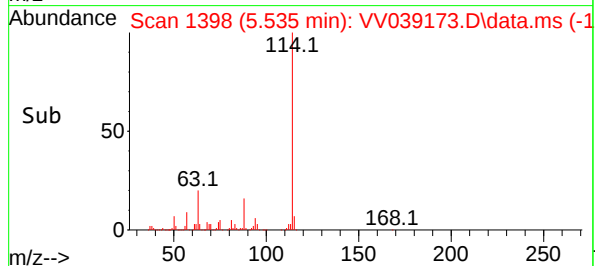
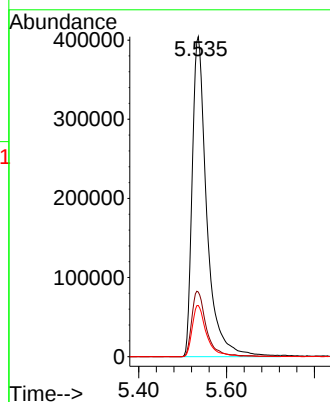
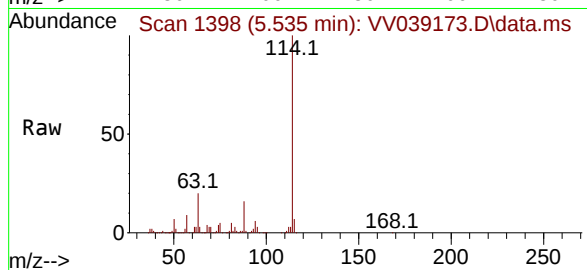
Tgt Ion:114 Resp: 932817

Ion Ratio Lower Upper

114 100

63 20.2 0.0 42.6

88 16.0 0.0 30.8



#35

Dibromofluoromethane

Concen: 51.604 ug/l

RT: 4.503 min Scan# 1077

Delta R.T. 0.006 min

Lab File: VV039173.D

Acq: 25 Sep 2025 12:10

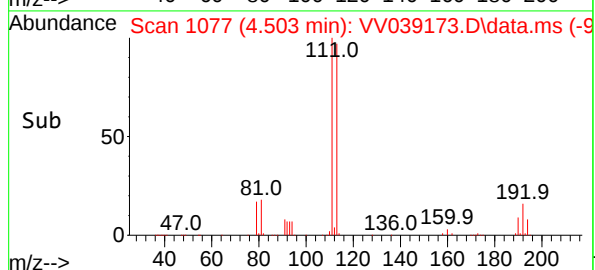
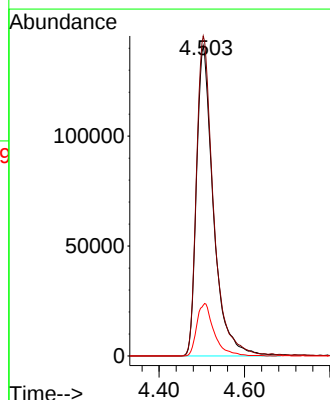
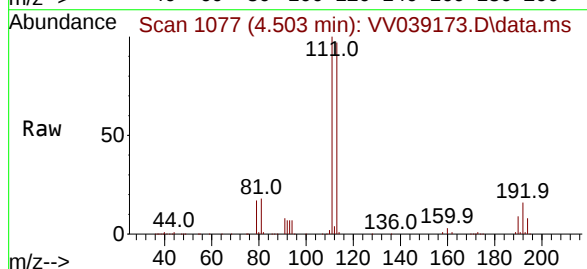
Tgt Ion:113 Resp: 386984

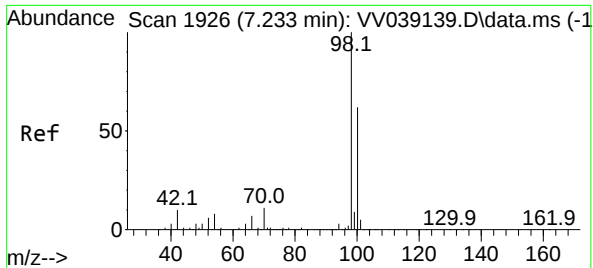
Ion Ratio Lower Upper

113 100

111 103.4 82.4 123.6

192 17.3 13.8 20.8

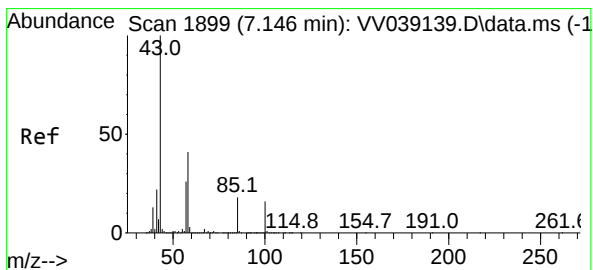
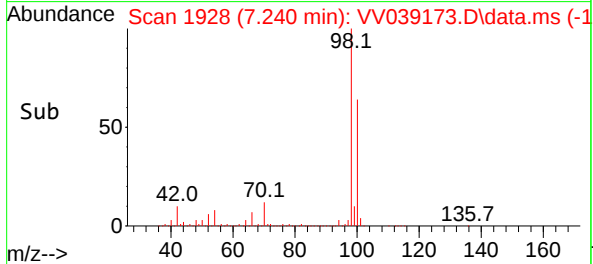
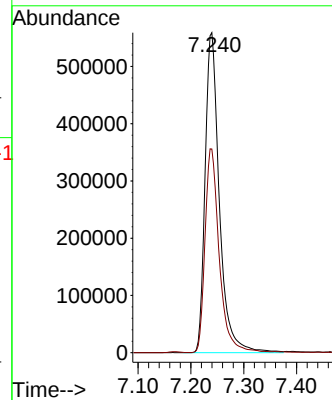
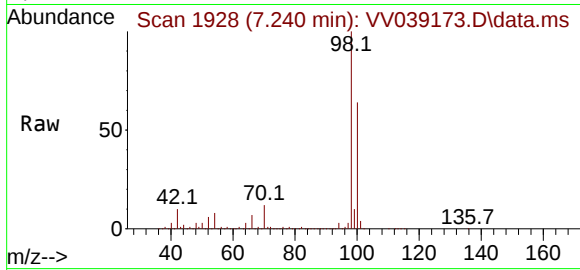




#50
Toluene-d8
Concen: 47.596 ug/l
RT: 7.240 min Scan# 1926
Delta R.T. 0.006 min
Lab File: VV039173.D
Acq: 25 Sep 2025 12:10

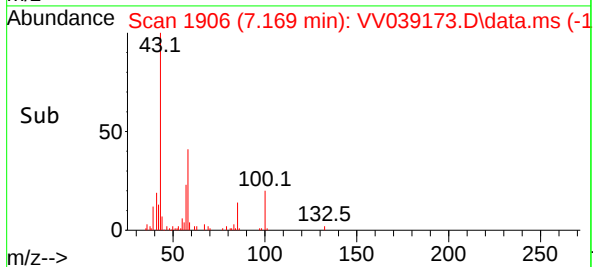
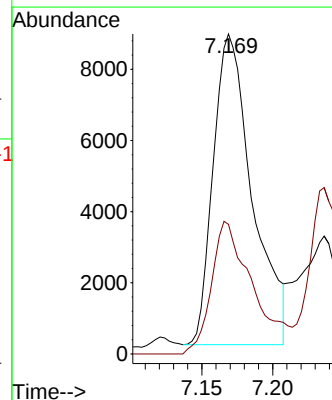
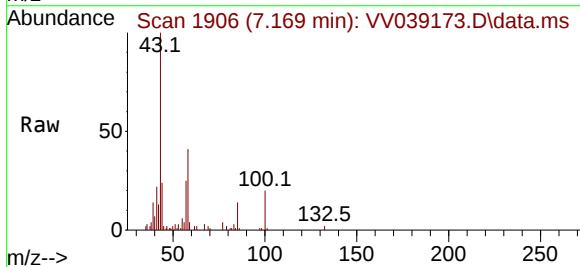
Instrument :
MSVOA_V
ClientSampleId :
DA-2

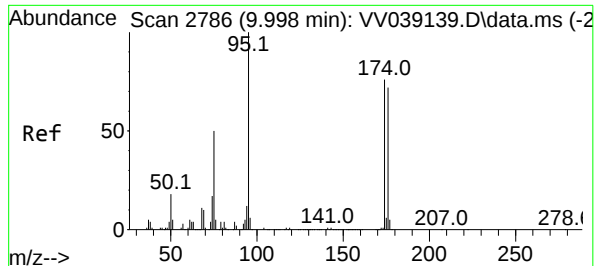
Tgt Ion: 98 Resp: 1048304
Ion Ratio Lower Upper
98 100
100 63.4 52.2 78.2



#51
4-Methyl-2-Pentanone
Concen: 1.358 ug/l
RT: 7.169 min Scan# 1906
Delta R.T. 0.023 min
Lab File: VV039173.D
Acq: 25 Sep 2025 12:10

Tgt Ion: 43 Resp: 16756
Ion Ratio Lower Upper
43 100
58 45.8 32.3 48.5





#62

4-Bromofluorobenzene

Concen: 44.855 ug/l

RT: 10.001 min Scan# 21

Delta R.T. 0.003 min

Lab File: VV039173.D

Acq: 25 Sep 2025 12:10

Instrument :

MSVOA_V

ClientSampleId :

DA-2

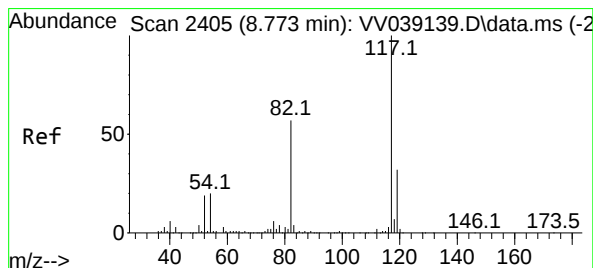
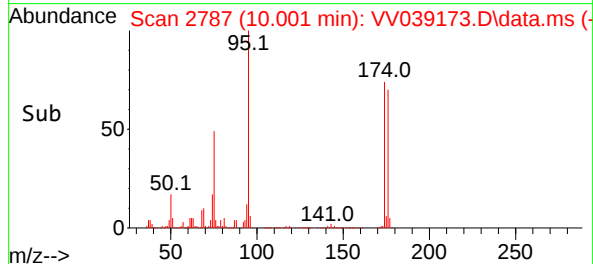
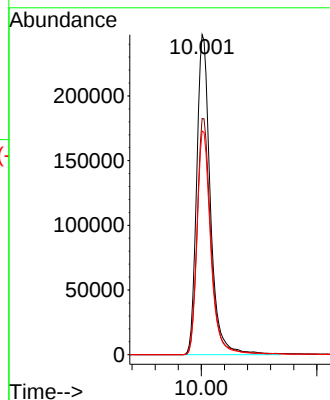
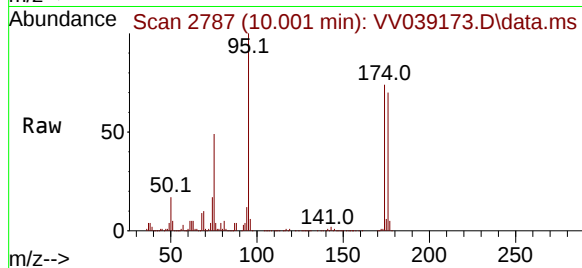
Tgt Ion: 95 Resp: 406703

Ion Ratio Lower Upper

95 100

174 74.1 0.0 150.4

176 69.8 0.0 144.2



#63

Chlorobenzene-d5

Concen: 50.000 ug/l

RT: 8.776 min Scan# 2406

Delta R.T. 0.003 min

Lab File: VV039173.D

Acq: 25 Sep 2025 12:10

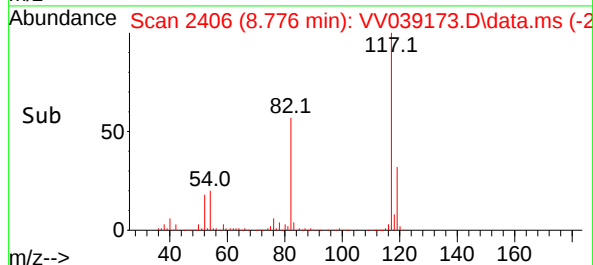
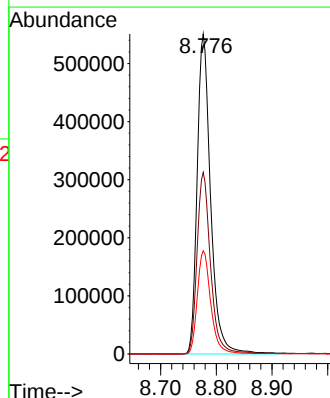
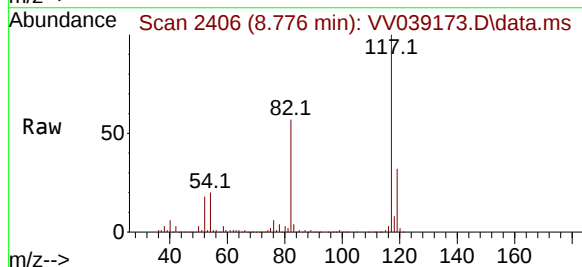
Tgt Ion: 117 Resp: 926008

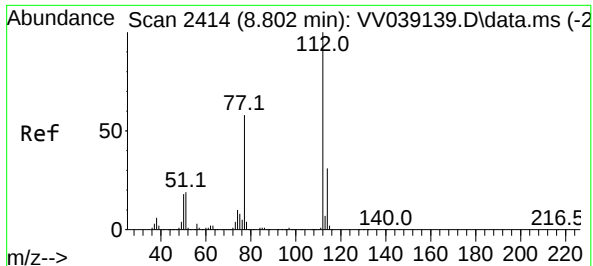
Ion Ratio Lower Upper

117 100

82 56.7 45.4 68.2

119 32.3 25.6 38.4





#65

Chlorobenzene

Concen: 0.216 ug/l

RT: 8.805 min Scan# 2414

Delta R.T. 0.003 min

Lab File: VV039173.D

Acq: 25 Sep 2025 12:10

Instrument :

MSVOA_V

ClientSampleId :

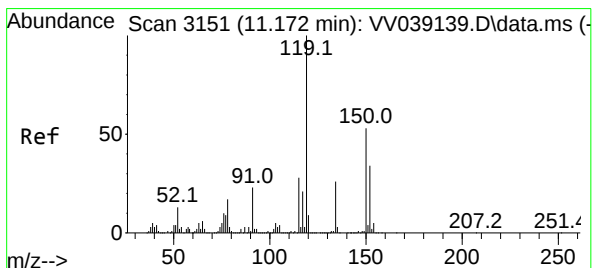
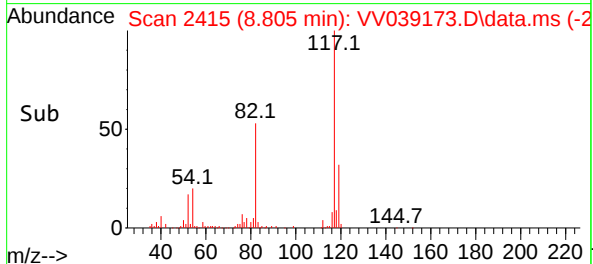
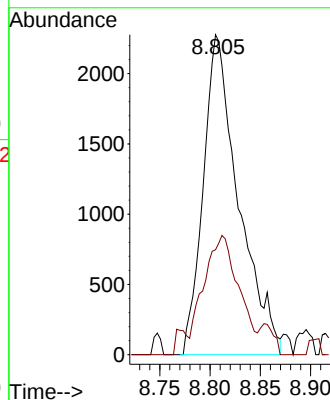
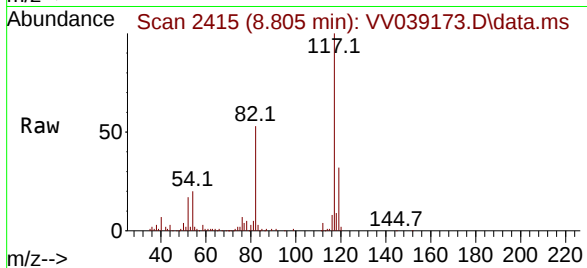
DA-2

Tgt Ion:112 Resp: 5564

Ion Ratio Lower Upper

112 100

114 32.8 24.6 37.0



#72

1,4-Dichlorobenzene-d4

Concen: 50.000 ug/l

RT: 11.175 min Scan# 3152

Delta R.T. 0.003 min

Lab File: VV039173.D

Acq: 25 Sep 2025 12:10

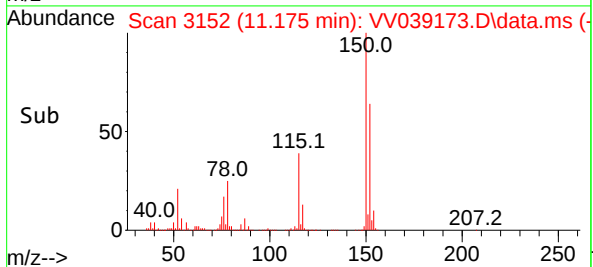
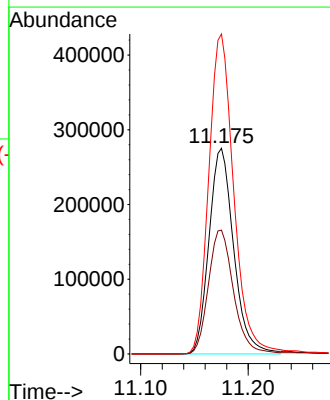
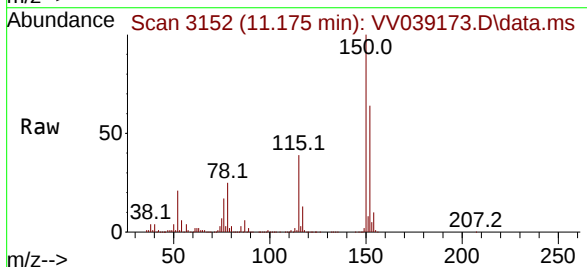
Tgt Ion:152 Resp: 438853

Ion Ratio Lower Upper

152 100

115 61.9 44.8 134.4

150 158.9 0.0 353.8



Report of Analysis

Client:	ATG-GREENVILLE AEC			Date Collected:	09/18/25	
Project:	AEC-2025-0013 CSC 4151 Coram			Date Received:	09/22/25	
Client Sample ID:	DA-3			SDG No.:	Q3164	
Lab Sample ID:	Q3164-03			Matrix:	Water	
Analytical Method:	8260D			% Solid:	0	
Sample Wt/Vol:	5	Units:	mL	Final Vol:	5000	uL
Soil Aliquot Vol:			uL	Test:	VOC-BTEX	
GC Column:	DB-624UI	ID :	0.18	Level :	LOW	
Prep Method :						

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039162.D	1	09/24/25 18:52	VV092425

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	0.15	U	0.15	1.00	ug/L
108-88-3	Toluene	0.14	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	0.13	U	0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	0.24	U	0.24	2.00	ug/L
95-47-6	o-Xylene	0.12	U	0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	51.1		74 - 125	102%	SPK: 50
1868-53-7	Dibromofluoromethane	40.0		75 - 124	80%	SPK: 50
2037-26-5	Toluene-d8	44.1		86 - 113	88%	SPK: 50
460-00-4	4-Bromofluorobenzene	39.6		77 - 121	79%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	756000	4.6			
540-36-3	1,4-Difluorobenzene	1430000	5.532			
3114-55-4	Chlorobenzene-d5	1320000	8.776			
3855-82-1	1,4-Dichlorobenzene-d4	608000	11.172			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039162.D
 Acq On : 24 Sep 2025 18:52
 Operator : SY/MD
 Sample : Q3164-03
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 19 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 DA-3

Quant Time: Sep 25 08:22:22 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T. QIon		Response	Conc	Units	Dev(Min)

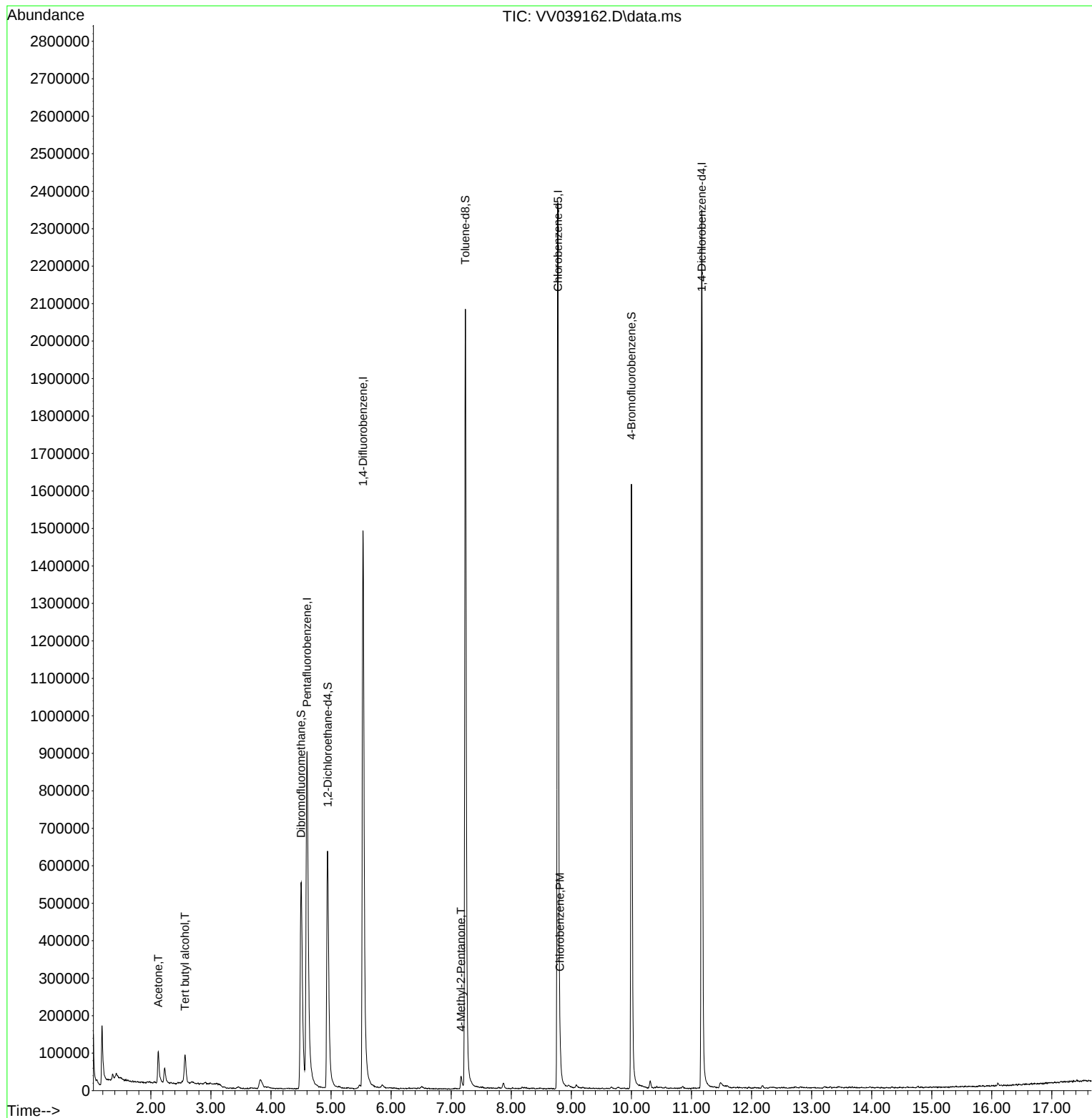
Internal Standards						
1) Pentafluorobenzene	4.600	168	755839	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	1427337	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	1322483	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	607726	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.944	65	559230	51.121	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	102.240%
35) Dibromofluoromethane	4.503	113	458688	39.974	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	79.940%#
50) Toluene-d8	7.236	98	1484627	44.053	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	88.100%#
62) 4-Bromofluorobenzene	10.001	95	549383	39.599	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	79.200%#
Target Compounds						
11) Tert butyl alcohol	2.568	59	53242	23.028	ug/l	Qvalue # 90
16) Acetone	2.124	43	107662	10.872	ug/l	96
51) 4-Methyl-2-Pentanone	7.162	43	24286	1.286	ug/l	92
65) Chlorobenzene	8.808	112	45385	1.232	ug/l	100

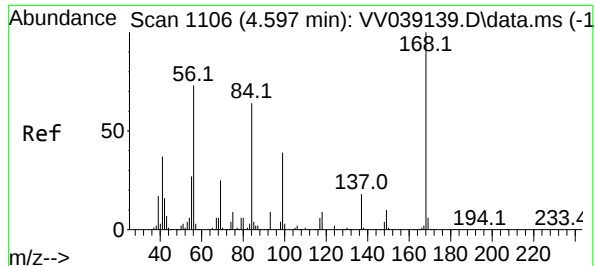
(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039162.D
Acq On : 24 Sep 2025 18:52
Operator : SY/MD
Sample : Q3164-03
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 19 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
DA-3

Quant Time: Sep 25 08:22:22 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

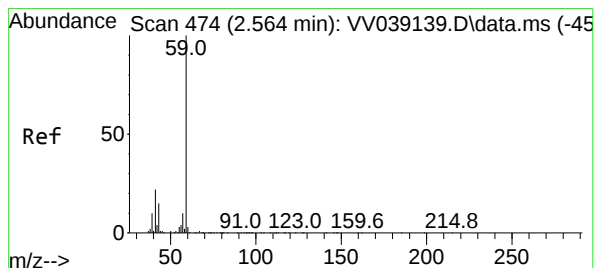
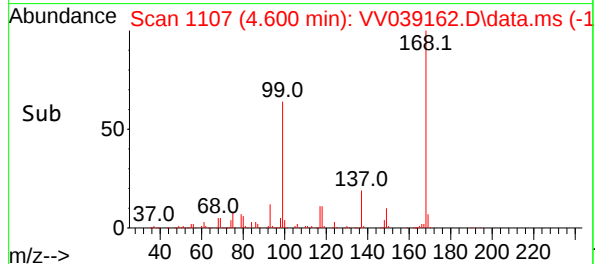
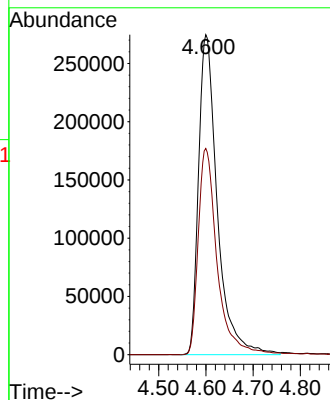
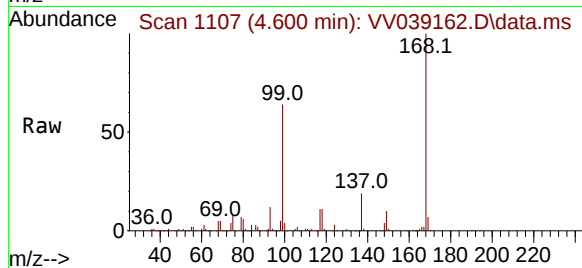




#1
Pentafluorobenzene
Concen: 50.000 ug/l
RT: 4.600 min Scan# 1106
Delta R.T. 0.003 min
Lab File: VV039162.D
Acq: 24 Sep 2025 18:52

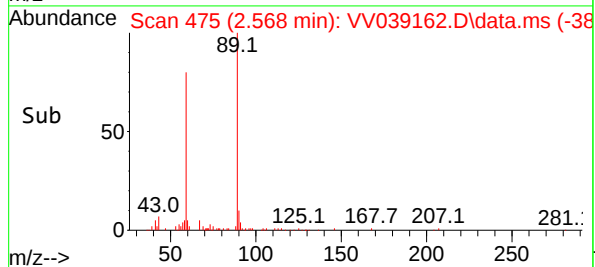
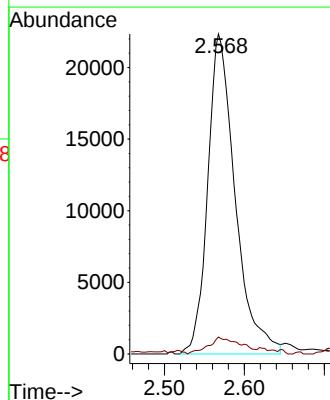
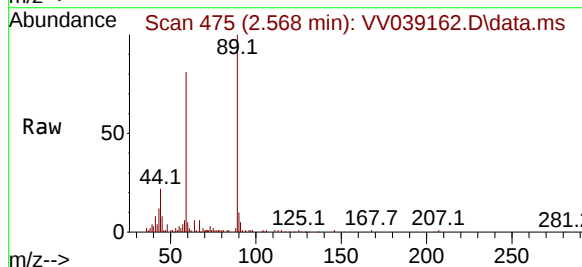
Instrument :
MSVOA_V
ClientSampleId :
DA-3

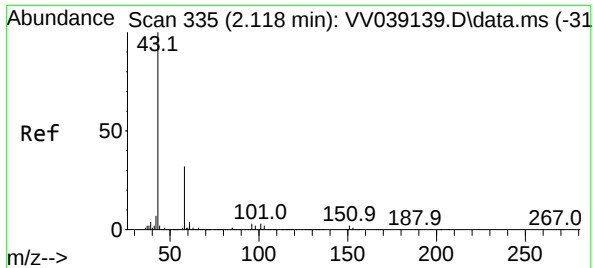
Tgt Ion:168 Resp: 755839
Ion Ratio Lower Upper
168 100
99 64.4 49.6 74.4



#11
Tert butyl alcohol
Concen: 23.028 ug/l
RT: 2.568 min Scan# 475
Delta R.T. 0.003 min
Lab File: VV039162.D
Acq: 24 Sep 2025 18:52

Tgt Ion: 59 Resp: 53242
Ion Ratio Lower Upper
59 100
57 6.0 7.8 11.8#

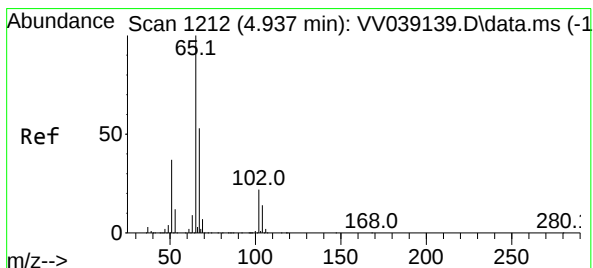
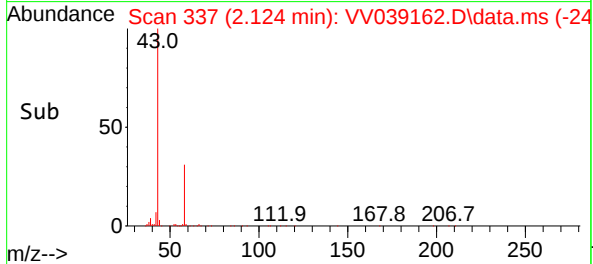
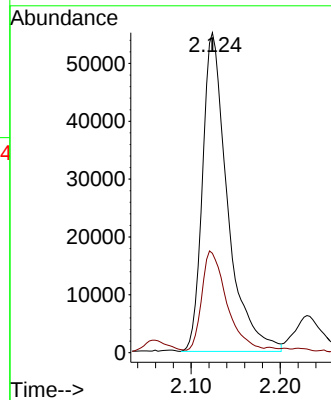
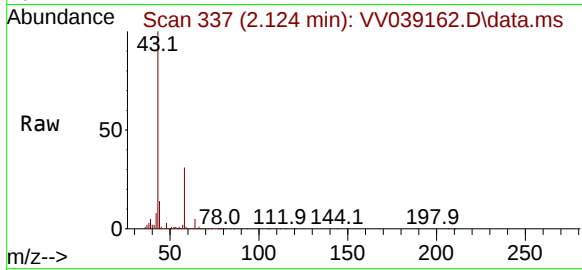




#16
Acetone
Concen: 10.872 ug/l
RT: 2.124 min Scan# 31
Delta R.T. 0.006 min
Lab File: VV039162.D
Acq: 24 Sep 2025 18:52

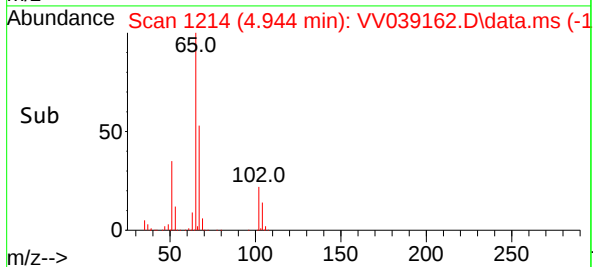
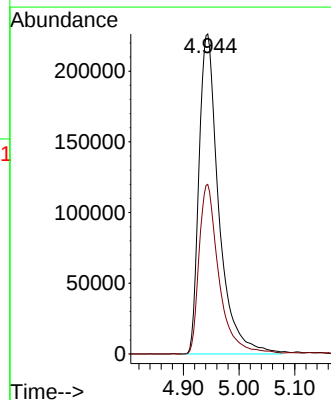
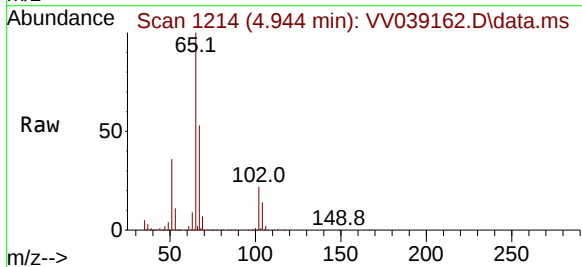
Instrument :
MSVOA_V
ClientSampleId :
DA-3

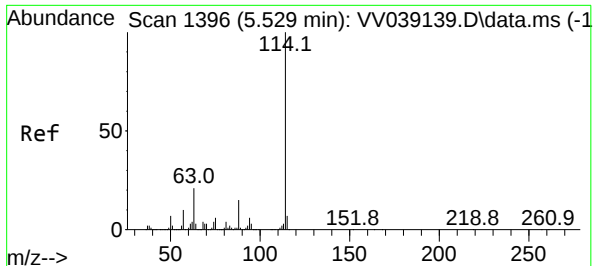
Tgt Ion: 43 Resp: 107662
Ion Ratio Lower Upper
43 100
58 30.2 25.8 38.6



#33
1,2-Dichloroethane-d4
Concen: 51.121 ug/l
RT: 4.944 min Scan# 1214
Delta R.T. 0.006 min
Lab File: VV039162.D
Acq: 24 Sep 2025 18:52

Tgt Ion: 65 Resp: 559230
Ion Ratio Lower Upper
65 100
67 53.2 0.0 107.0





#34

1,4-Difluorobenzene

Concen: 50.000 ug/l

RT: 5.532 min Scan# 11

Delta R.T. 0.003 min

Lab File: VV039162.D

Acq: 24 Sep 2025 18:52

Instrument :

MSVOA_V

ClientSampleId :

DA-3

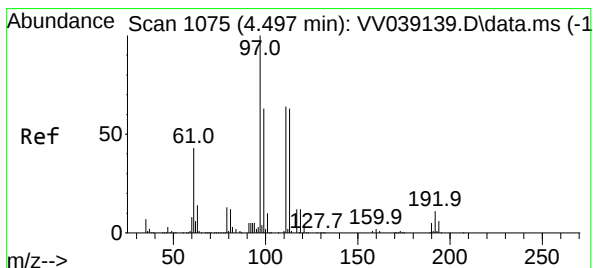
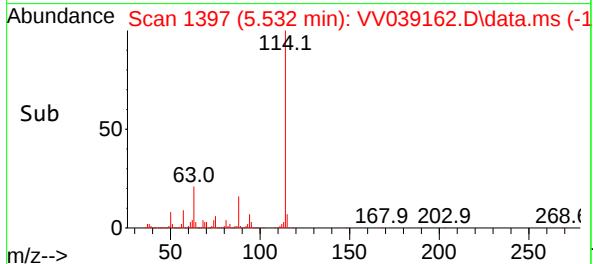
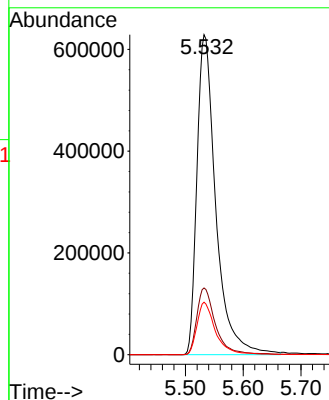
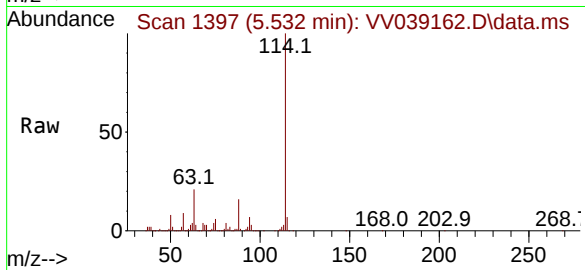
Tgt Ion:114 Resp: 1427337

Ion Ratio Lower Upper

114 100

63 20.9 0.0 42.6

88 16.4 0.0 30.8



#35

Dibromofluoromethane

Concen: 39.974 ug/l

RT: 4.503 min Scan# 1077

Delta R.T. 0.006 min

Lab File: VV039162.D

Acq: 24 Sep 2025 18:52

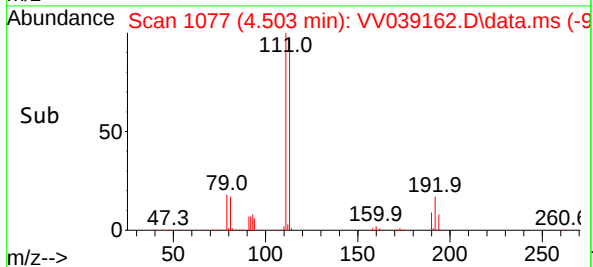
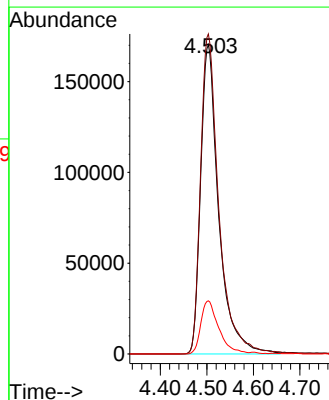
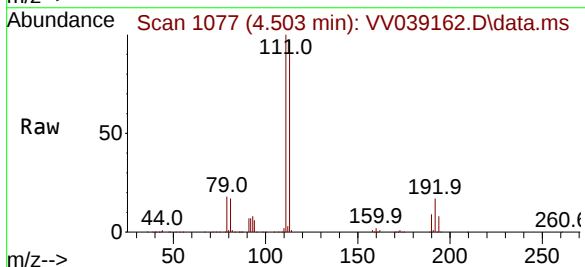
Tgt Ion:113 Resp: 458688

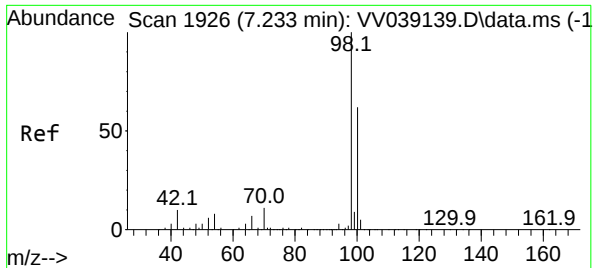
Ion Ratio Lower Upper

113 100

111 104.3 82.4 123.6

192 17.1 13.8 20.8





#50

Toluene-d8

Concen: 44.053 ug/l

RT: 7.236 min Scan# 1926

Delta R.T. 0.003 min

Lab File: VV039162.D

Acq: 24 Sep 2025 18:52

Instrument :

MSVOA_V

ClientSampleId :

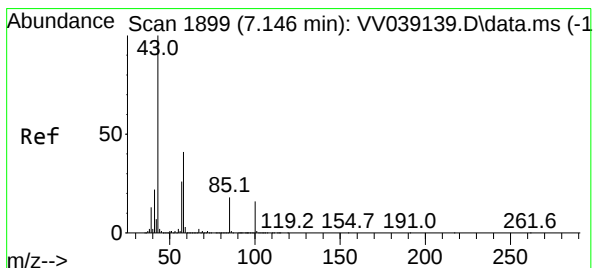
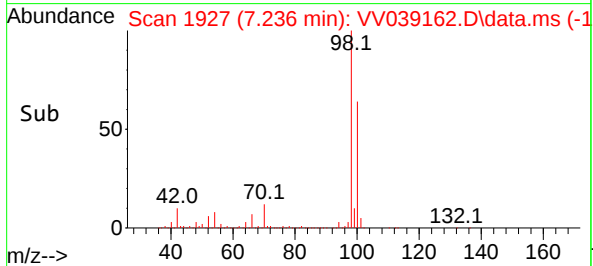
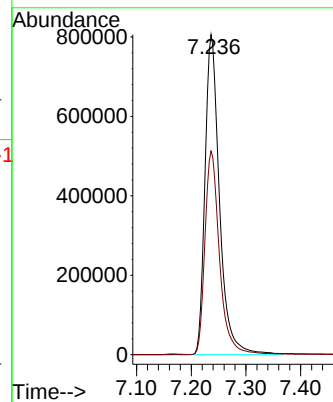
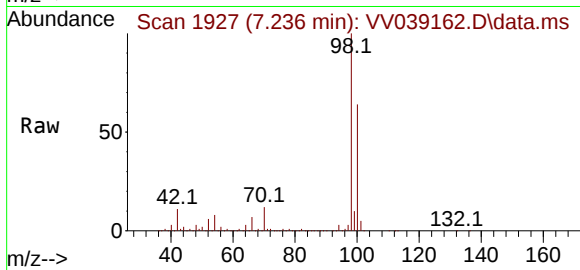
DA-3

Tgt Ion: 98 Resp: 1484627

Ion Ratio Lower Upper

98 100

100 63.5 52.2 78.2



#51

4-Methyl-2-Pentanone

Concen: 1.286 ug/l

RT: 7.162 min Scan# 1904

Delta R.T. 0.016 min

Lab File: VV039162.D

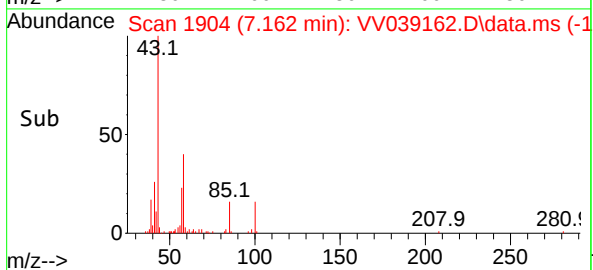
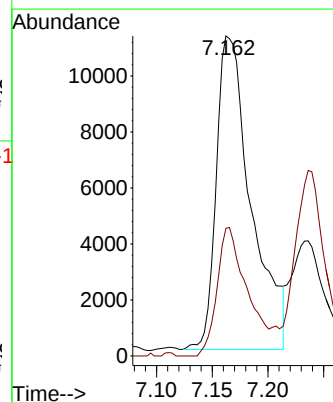
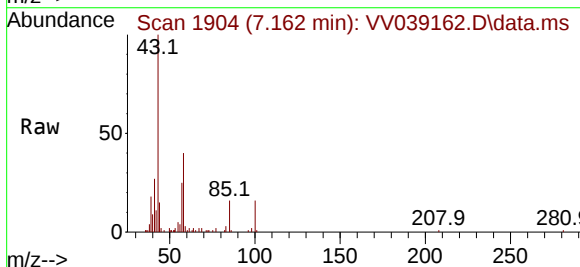
Acq: 24 Sep 2025 18:52

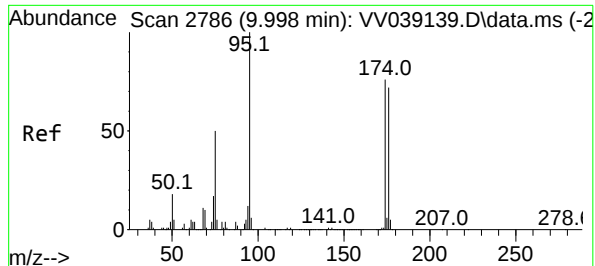
Tgt Ion: 43 Resp: 24286

Ion Ratio Lower Upper

43 100

58 35.4 32.3 48.5





#62

4-Bromofluorobenzene

Concen: 39.599 ug/l

RT: 10.001 min Scan# 21

Delta R.T. 0.003 min

Lab File: VV039162.D

Acq: 24 Sep 2025 18:52

Instrument :

MSVOA_V

ClientSampleId :

DA-3

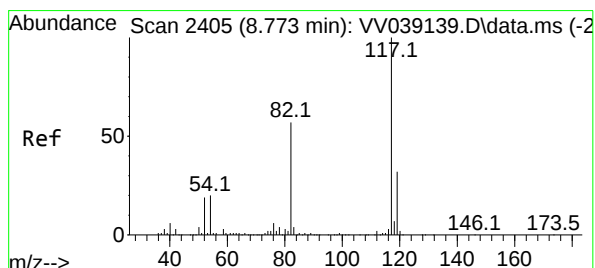
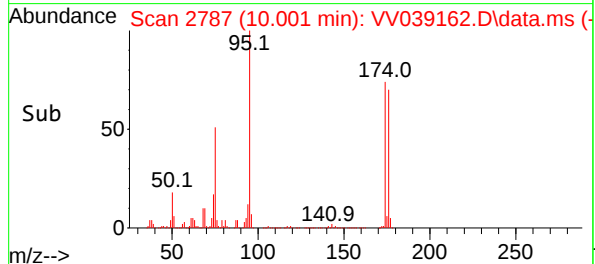
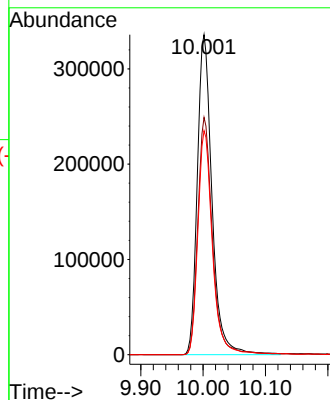
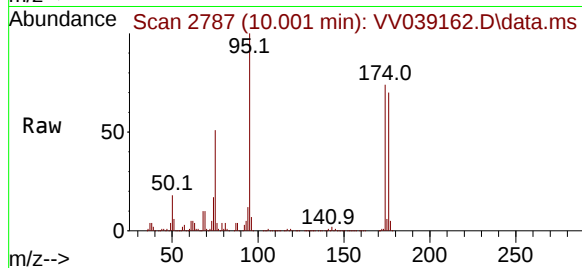
Tgt Ion: 95 Resp: 549383

Ion Ratio Lower Upper

95 100

174 74.3 0.0 150.4

176 69.1 0.0 144.2



#63

Chlorobenzene-d5

Concen: 50.000 ug/l

RT: 8.776 min Scan# 2406

Delta R.T. 0.003 min

Lab File: VV039162.D

Acq: 24 Sep 2025 18:52

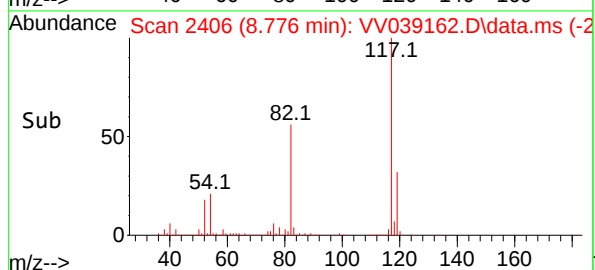
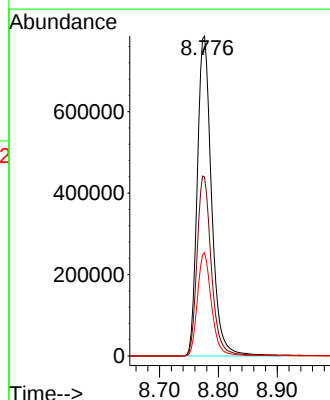
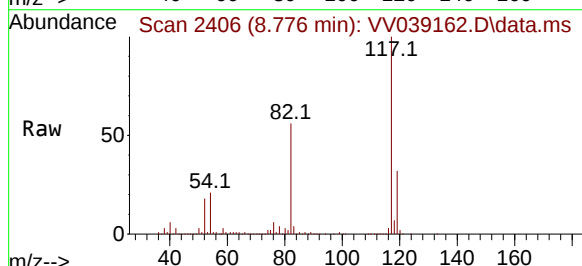
Tgt Ion: 117 Resp: 1322483

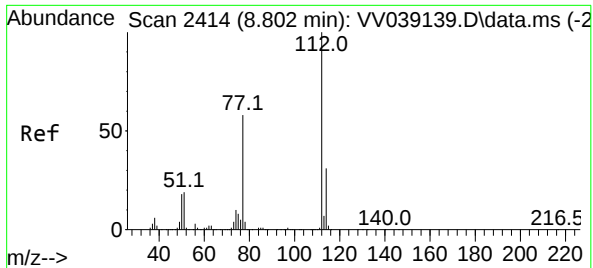
Ion Ratio Lower Upper

117 100

82 56.0 45.4 68.2

119 32.2 25.6 38.4





#65

Chlorobenzene

Concen: 1.232 ug/l

RT: 8.808 min Scan# 2414

Delta R.T. 0.006 min

Lab File: VV039162.D

Acq: 24 Sep 2025 18:52

Instrument :

MSVOA_V

ClientSampleId :

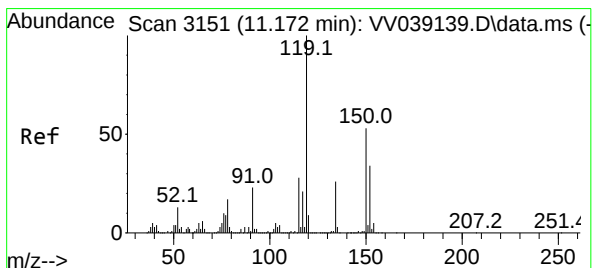
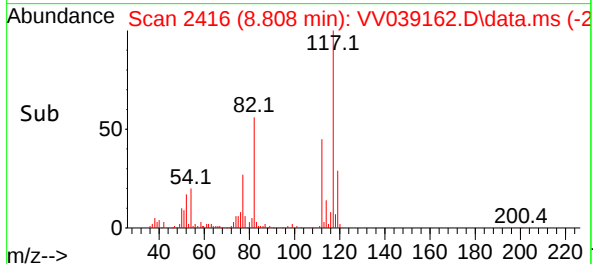
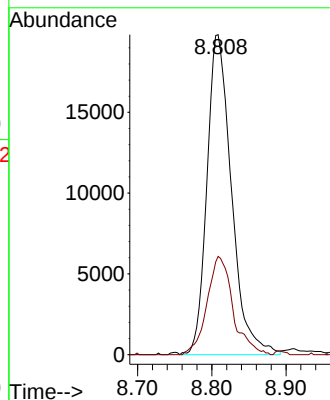
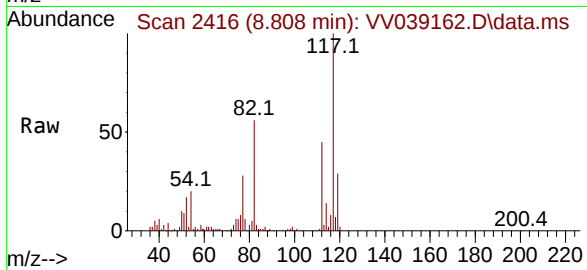
DA-3

Tgt Ion:112 Resp: 45385

Ion Ratio Lower Upper

112 100

114 30.7 24.6 37.0



#72

1,4-Dichlorobenzene-d4

Concen: 50.000 ug/l

RT: 11.172 min Scan# 3151

Delta R.T. -0.000 min

Lab File: VV039162.D

Acq: 24 Sep 2025 18:52

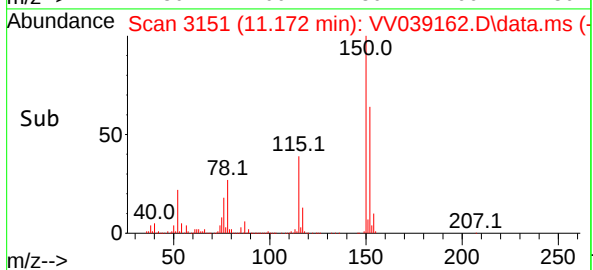
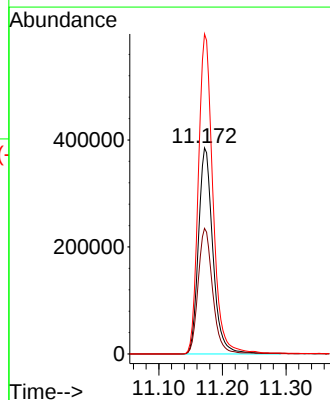
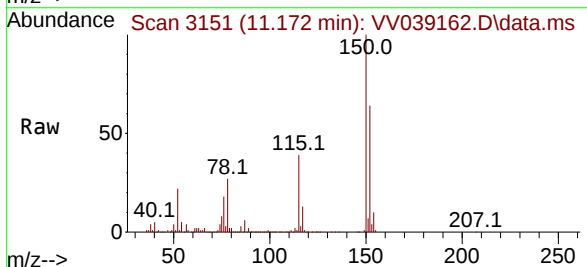
Tgt Ion:152 Resp: 607726

Ion Ratio Lower Upper

152 100

115 60.9 44.8 134.4

150 156.9 0.0 353.8





CALIBRATION SUMMARY



284 Sheffield Street, Mountainside, New Jersey 07092, Phone : 908 789 8900,
Fax : 908 789 8922

VOLATILE ORGANICS INITIAL CALIBRATION DATA

Lab Name: Alliance

Contract: ATGG01

Lab Code: ACE

SDG No.: Q3164

Instrument ID: MSVOA_V

Calibration Date(s): 09/22/2025 09/22/2025

Heated Purge: (Y/N) N

Calibration Time(s): 12:26 16:35

GC Column: DB-624UI ID: 0.18 (mm)

LAB FILE ID:		RRF010 = VV039137.D		RRF020 = VV039138.D		RRF050 = VV039139.D		
		RRF100 = VV039140.D		RRF005 = VV039142.D		RRF001 = VV039143.D		
COMPOUND	RRF010	RRF020	RRF050	RRF100	RRF005	RRF001	RRF	% RSD
Benzene	2.077	1.978	1.990	1.931	1.963	1.936	1.979	2.7
Toluene	1.211	1.199	1.229	1.214	1.130	0.901	1.147	10.9
Ethyl Benzene	2.028	2.143	2.382	2.418	1.953	1.725	2.108	12.5
m/p-Xylenes	0.820	0.882	0.949	0.938	0.713	0.580	0.814	17.7
o-Xylene	0.703	0.742	0.846	0.880	0.635	0.518	0.720	18.7
1,2-Dichloroethane-d4	0.773	0.663	0.646	0.618	0.919		0.724	17.1
Dibromofluoromethane	0.435	0.382	0.377	0.360	0.454		0.402	10.1
Toluene-d8	1.166	1.092	1.136	1.097	1.411		1.181	11.2
4-Bromofluorobenzene	0.481	0.464	0.507	0.501	0.477		0.486	3.6

* Compounds with required minimum RRF and maximum %RSD values.
All other compounds must meet a minimum RRF of 0.010.
RRF of 1,4-Dioxane = Value should be divide by 1000.

Method Path : Z:\voasrv\HPCHEM1\MSVOA_V\Method\
 Method File : 82V092225W.M
 Title : SW846 8260
 Last Update : Wed Sep 24 08:08:08 2025
 Response Via : Initial Calibration

Calibration Files

1 =VV039143.D 5 =VV039142.D 10 =VV039137.D 20 =VV039138.D 50 =VV039139.D 100 =VV039140.D

	Compound	1	5	10	20	50	100	Avg	%RSD
1) I	Pentafluorobenzene	-----ISTD-----							
2) T	Dichlorodifluo...	0.934	0.772	0.848	0.757	0.740	0.699	0.791	10.78
3) P	Chloromethane	0.946	0.889	0.949	0.814	0.770	0.729	0.850	10.90
4) C	Vinyl Chloride	1.026	0.932	1.007	0.856	0.836	0.801	0.910	10.27#
5) T	Bromomethane		0.655	0.631	0.548	0.508	0.524	0.573	11.50
6) T	Chloroethane	0.915	0.606	0.662	0.568	0.533	0.500	0.631	23.83
7) T	Trichlorofluor...	1.546	1.365	1.394	1.253	1.193	1.151	1.317	11.15
8) T	Diethyl Ether	0.572	0.513	0.505	0.461	0.447	0.435	0.489	10.50
9) T	1,1,2-Trichlor...	0.939	0.804	0.839	0.732	0.702	0.686	0.784	12.28
10) T	Methyl Iodide		0.638	0.772	0.742	0.786	0.793	0.746	8.50
11) T	Tert butyl alc...		0.166	0.169	0.157	0.137	0.136	0.153	10.26
12) CM	1,1-Dichloroet...	0.873	0.789	0.804	0.721	0.674	0.663	0.754	10.87#
13) T	Acrolein		0.025	0.027	0.027	0.026	0.027	0.027	4.01
14) T	Allyl chloride	1.523	1.434	1.388	1.283	1.243	1.208	1.347	9.06
15) T	Acrylonitrile	0.587	0.522	0.507	0.462	0.436	0.427	0.490	12.38
16) T	Acetone	0.712	0.569	0.537	0.458	0.449	0.414	0.523	20.84
17) T	Carbon Disulfide	2.670	2.462	2.453	2.181	2.081	1.986	2.306	11.43
18) T	Methyl Acetate	1.198	1.097	1.146	1.036	0.983	0.973	1.072	8.46
19) T	Methyl tert-bu...	2.591	2.556	2.547	2.285	2.233	2.202	2.402	7.50
20) T	Methylene Chlo...	1.987	1.113	0.930	0.848	0.779	0.742	1.066	44.05
21) T	trans-1,2-Dich...	0.959	0.835	0.846	0.755	0.724	0.696	0.803	12.11
22) T	Diisopropyl ether	2.659	2.560	2.757	2.530	2.520	2.433	2.576	4.44
23) T	Vinyl Acetate	2.243	2.313	2.393	2.241	2.204	2.128	2.254	4.04
24) P	1,1-Dichloroet...	1.942	1.659	1.669	1.466	1.388	1.329	1.576	14.42
25) T	2-Butanone	0.554	0.570	0.604	0.584	0.586	0.580	0.579	2.90
26) T	2,2-Dichloropr...	1.906	1.393	1.364	1.275	1.239	1.204	1.397	18.57
27) T	cis-1,2-Dichlo...	1.001	0.832	0.908	0.842	0.868	0.860	0.885	7.06
28) T	Bromochloromet...	0.743	0.525	0.836	0.753	0.679	0.637	0.696	15.49
29) T	Tetrahydrofuran	0.335	0.350	0.376	0.372	0.369	0.374	0.363	4.57
30) C	Chloroform	1.880	1.746	1.752	1.570	1.490	1.433	1.645	10.60#
31) T	Cyclohexane		1.407	1.325	1.222	1.245	1.254	1.291	5.85
32) T	1,1,1-Trichlor...	1.590	1.513	1.562	1.321	1.298	1.263	1.424	10.26
33) S	1,2-Dichloroet...		0.919	0.773	0.663	0.646	0.618	0.724	17.13
34) I	1,4-Difluorobenzene	-----ISTD-----							
35) S	Dibromofluorom...		0.454	0.435	0.382	0.377	0.360	0.402	10.10
36) T	1,1-Dichloropr...	0.650	0.536	0.624	0.604	0.620	0.612	0.608	6.33
37) T	Ethyl Acetate	0.825	0.722	0.715	0.740	0.714	0.710	0.737	5.96
38) T	Carbon Tetrach...	0.890	0.736	0.742	0.710	0.703	0.675	0.743	10.25
39) T	Methylcyclohexane	0.622	0.680	0.701	0.720	0.823	0.857	0.734	12.15
40) TM	Benzene	1.936	1.963	2.077	1.978	1.990	1.931	1.979	2.69
41) T	Methacrylonitrile	0.185	0.215	0.273	0.292	0.330	0.342	0.273	22.92
42) TM	1,2-Dichloroet...	0.740	0.693	0.725	0.694	0.666	0.643	0.693	5.20
43) T	Isopropyl Acetate	0.869	0.933	1.032	1.029	1.069	1.100	1.005	8.71
44) TM	Trichloroethene	0.435	0.460	0.471	0.456	0.463	0.454	0.457	2.64
45) C	1,2-Dichloropr...	0.465	0.471	0.509	0.519	0.502	0.486	0.492	4.40#
46) T	Dibromomethane	0.396	0.381	0.392	0.367	0.368	0.354	0.376	4.40
47) T	Bromodichlorom...	0.829	0.767	0.832	0.764	0.753	0.725	0.778	5.54
48) T	Methyl methacr...	0.491	0.356	0.409	0.449	0.504	0.526	0.456	14.15
49) T	1,4-Dioxane	0.009	0.005	0.007	0.007	0.007	0.008	0.007	17.72
50) S	Toluene-d8		1.411	1.166	1.092	1.136	1.097	1.181	11.21
51) T	4-Methyl-2-Pen...	0.459	0.613	0.715	0.736	0.732	0.714	0.661	16.48
52) CM	Toluene	0.901	1.130	1.211	1.199	1.229	1.214	1.147	10.93#
53) T	t-1,3-Dichloro...	0.548	0.645	0.701	0.696	0.728	0.753	0.678	10.83
54) T	cis-1,3-Dichlo...	0.595	0.704	0.803	0.777	0.811	0.803	0.749	11.37
55) T	1,1,2-Trichlor...	0.553	0.524	0.534	0.502	0.494	0.480	0.514	5.31
56) T	Ethyl methacry...	0.507	0.510	0.569	0.603	0.716	0.761	0.611	17.43

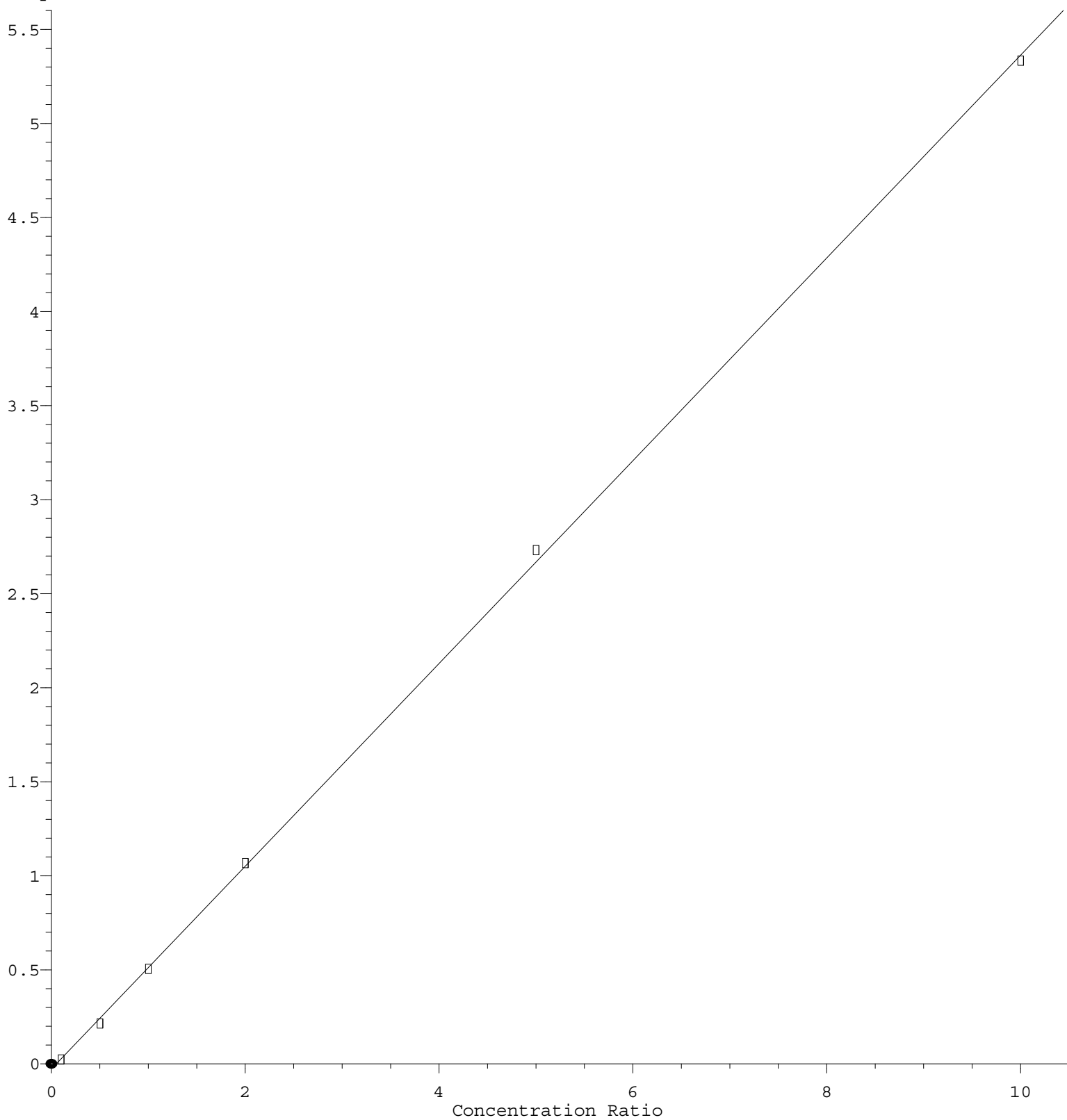
Method Path : Z:\voasrv\HPCHEM1\MSVOA_V\Method\
 Method File : 82V092225W.M

57)	T	1,3-Dichloropr...	0.797	0.764	0.861	0.827	0.820	0.802	0.812	4.02
58)	T	2-Chloroethyl ...	0.136	0.196	0.264	0.304	0.353	0.351	0.267	32.62
59)	T	2-Hexanone	0.231	0.429	0.505	0.534	0.546	0.533	0.463	26.18
60)	T	Dibromochlorom...	0.620	0.598	0.605	0.570	0.567	0.558	0.586	4.20
61)	T	1,2-Dibromoethane	0.484	0.535	0.556	0.538	0.529	0.514	0.526	4.70
62)	S	4-Bromofluorob...	0.477	0.481	0.464	0.507	0.501	0.486		3.62
63)	I	Chlorobenzene-d5	-----ISTD-----							
64)	T	Tetrachloroethene	0.415	0.431	0.435	0.412	0.412	0.407	0.419	2.74
65)	PM	Chlorobenzene	1.428	1.417	1.403	1.368	1.383	1.358	1.393	1.99
66)	T	1,1,1,2-Tetrac...	0.543	0.504	0.520	0.488	0.488	0.477	0.503	4.84
67)	C	Ethyl Benzene	1.725	1.953	2.028	2.143	2.382	2.418	2.108	12.53#
68)	T	m/p-Xylenes	0.580	0.713	0.820	0.882	0.949	0.938	0.814	17.68
69)	T	o-Xylene	0.518	0.635	0.703	0.742	0.846	0.880	0.720	18.67
70)	T	Styrene	0.828	1.038	1.239	1.381	1.553	1.537	1.263	22.78
71)	P	Bromoform	0.397	0.385	0.396	0.384	0.396	0.400	0.393	1.68
72)	I	1,4-Dichlorobenzen...	-----ISTD-----							
73)	T	Isopropylbenzene	3.226	3.296	3.564	3.638	3.939	4.096	3.627	9.49
74)	T	N-amyl acetate	1.031	1.279	1.425	1.506	1.673	1.764	1.447	18.47
75)	P	1,1,2,2-Tetrac...	1.975	1.684	1.600	1.461	1.425	1.389	1.589	13.83
76)	T	1,2,3-Trichlor...	1.375	1.201	1.136	1.048	1.049	1.023	1.138	11.73
77)	T	Bromobenzene	0.978	0.927	0.948	0.919	0.925	0.949	0.941	2.34
78)	T	n-propylbenzene	3.873	3.967	4.330	4.442	4.899	4.986	4.416	10.44
79)	T	2-Chlorotoluene	2.114	2.585	2.752	2.688	2.833	2.883	2.643	10.59
80)	T	1,3,5-Trimethy...	2.462	2.697	2.924	3.141	3.387	3.475	3.014	13.12
81)	T	trans-1,4-Dich...	0.427	0.478	0.451	0.495	0.521	0.474		7.77
82)	T	4-Chlorotoluene	2.381	2.877	3.160	3.216	3.338	3.413	3.064	12.48
83)	T	tert-Butylbenzene	2.875	2.757	2.886	2.881	3.222	3.413	3.006	8.43
84)	T	1,2,4-Trimethy...	2.332	2.542	2.767	3.100	3.378	3.440	2.926	15.47
85)	T	sec-Butylbenzene	3.605	3.608	3.823	3.969	4.364	4.550	3.987	9.88
86)	T	p-Isopropyltol...	2.874	2.994	3.237	3.351	3.682	3.772	3.318	10.85
87)	T	1,3-Dichlorobe...	1.975	1.943	1.931	1.836	1.857	1.882	1.904	2.83
88)	T	1,4-Dichlorobe...	2.522	2.139	2.070	1.951	1.934	1.919	2.089	10.97
89)	T	n-Butylbenzene	2.646	2.816	3.034	3.141	3.613	3.763	3.169	13.88
90)	T	Hexachloroethane	1.044	0.841	0.808	0.740	0.728	0.742	0.817	14.63
91)	T	1,2-Dichlorobe...	1.935	1.797	1.751	1.652	1.738	1.802	1.779	5.26
92)	T	1,2-Dibromo-3-...	0.476	0.333	0.299	0.283	0.285	0.302	0.330	22.39
93)	T	1,2,4-Trichlor...	1.028	0.935	0.928	0.963	1.076	1.189	1.020	9.87
94)	T	Hexachlorobuta...	0.683	0.530	0.512	0.474	0.480	0.501	0.530	14.65
95)	T	Naphthalene	3.054	2.665	2.888	3.114	3.814	4.244	3.296	18.32
96)	T	1,2,3-Trichlor...	1.113	0.901	0.974	0.973	1.107	1.191	1.043	10.58

(#) = Out of Range

2-Hexanone

Response Ratio



$$\text{Response} = 5.390\text{e-}001 * \text{Amt} - 2.507\text{e-}002$$

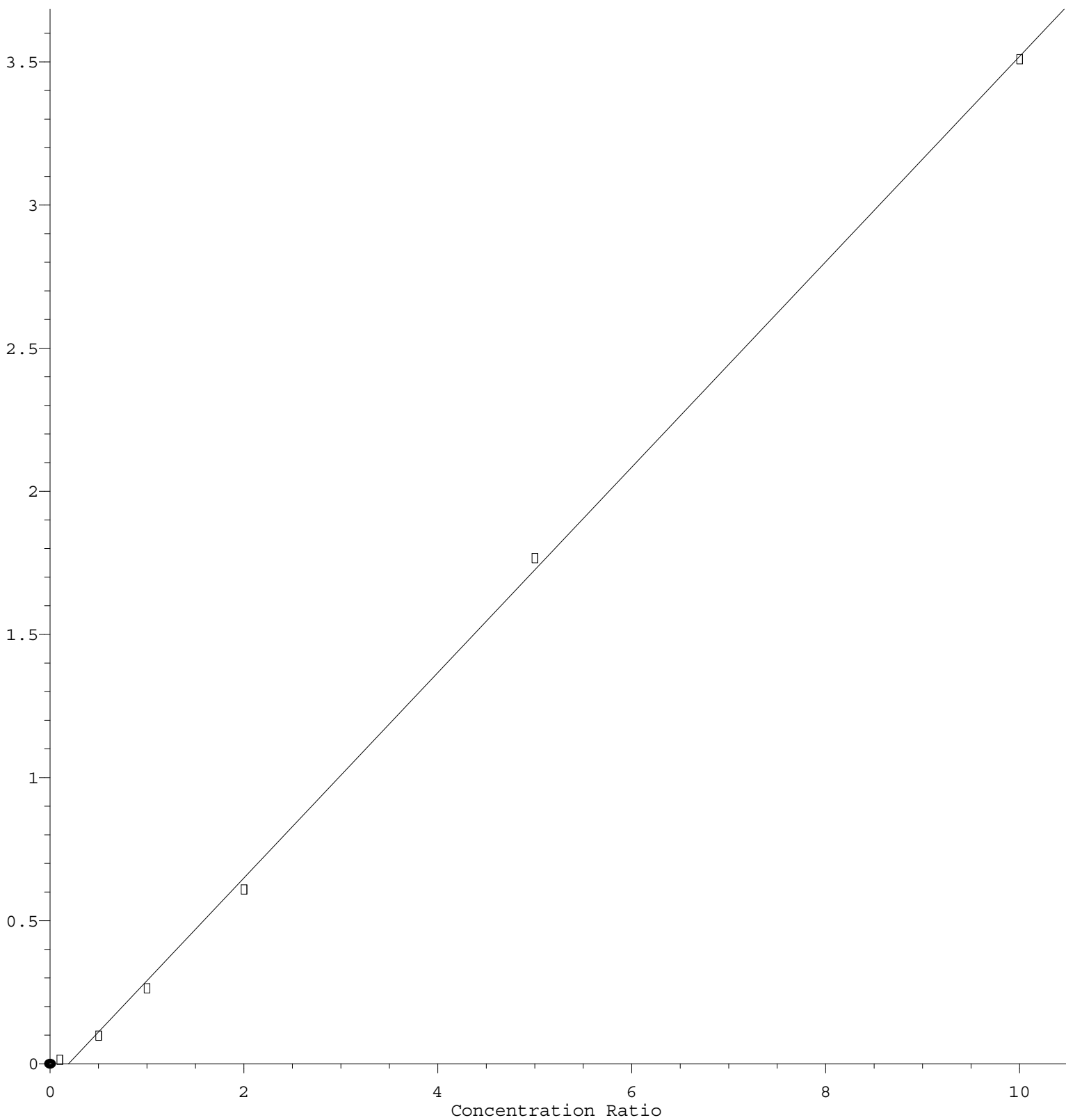
Coef of Det (r^2) = 0.999719 Curve Fit: Linear

Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M

Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

2-Chloroethyl Vinyl ether

Response Ratio



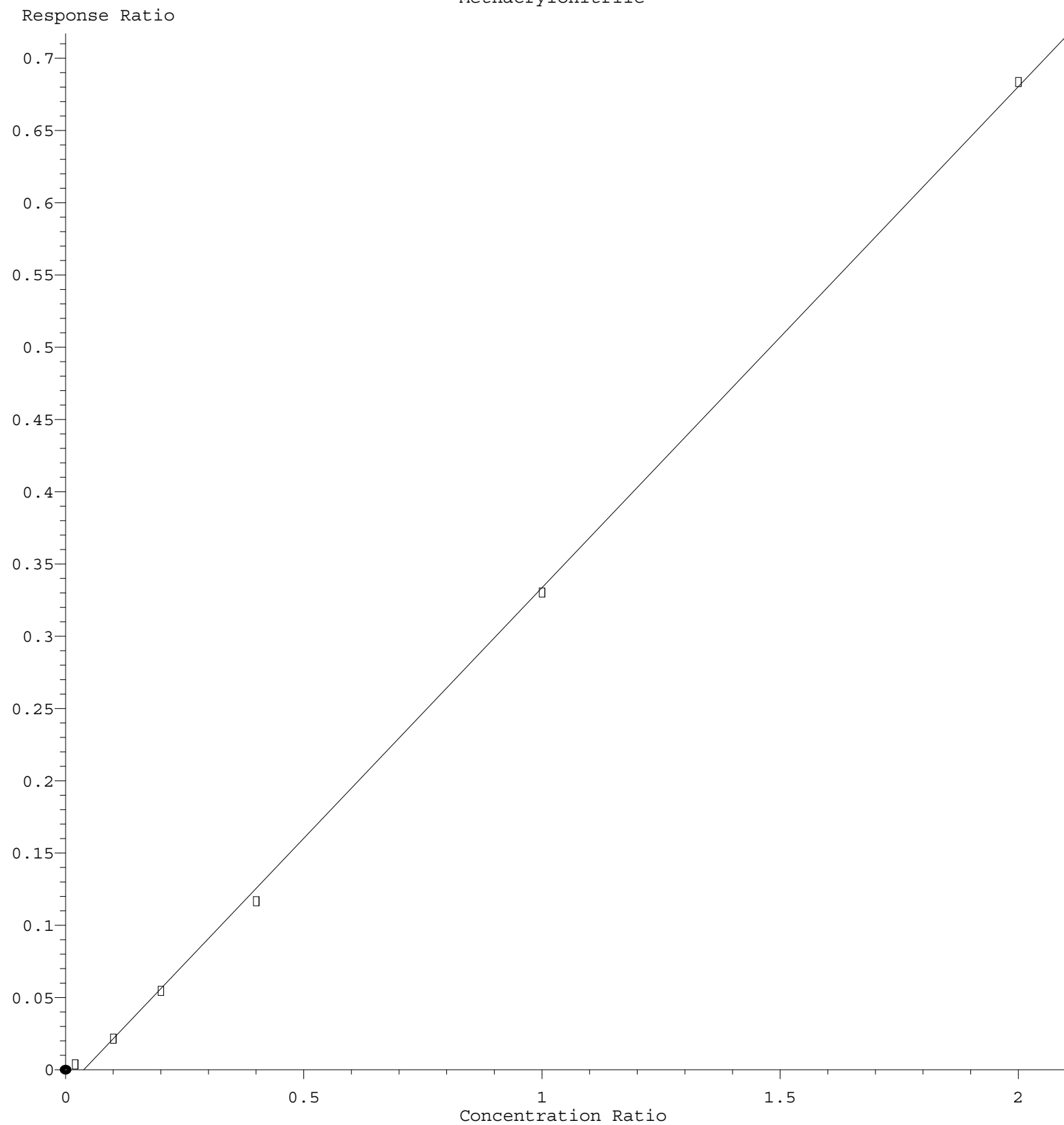
$$\text{Response} = 3.588\text{e-}001 * \text{Amt} - 6.905\text{e-}002$$

Coef of Det (r^2) = 0.999314 Curve Fit: Linear

Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M

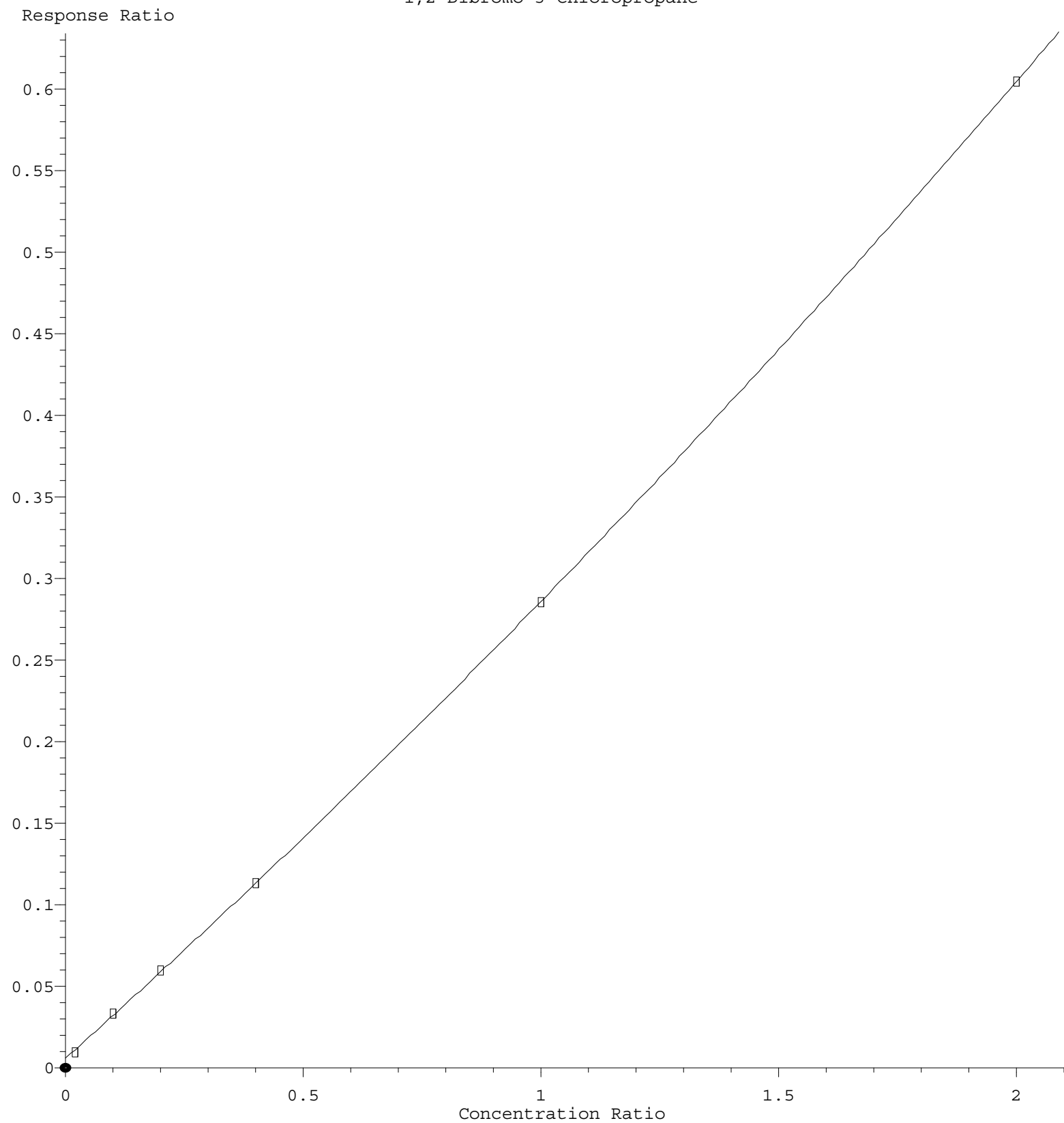
Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

Methacrylonitrile



Response = $3.467 \times 10^{-1} \times \text{Amt} - 1.325 \times 10^{-2}$
Coef of Det (r^2) = 0.999425 Curve Fit: Linear
Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M
Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

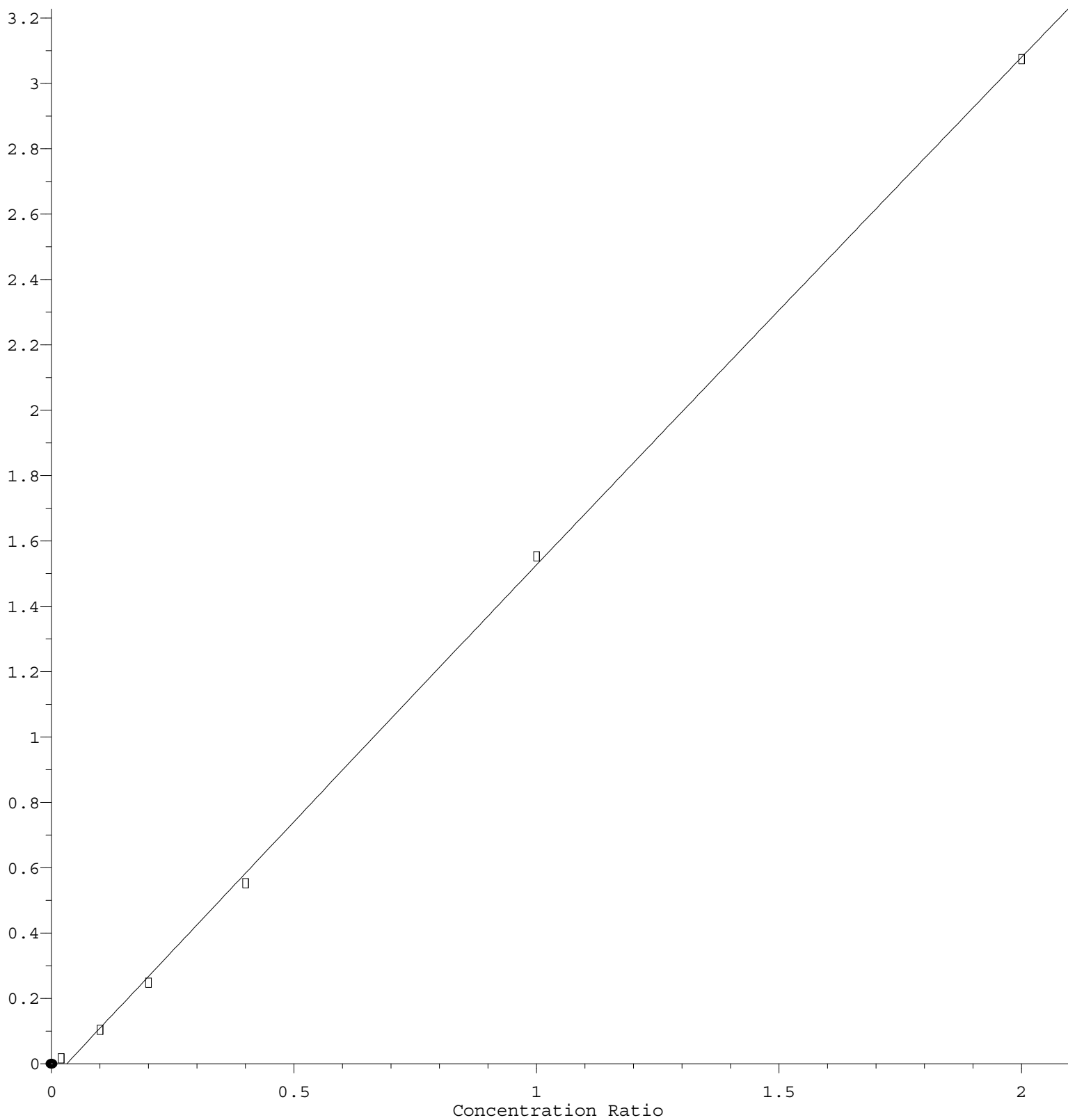
1,2-Dibromo-3-Chloropropane



$R = 1.931e-002 A^2 + 2.606e-001 A + 5.917e-003$
 Coef of Det (r^2) = 0.999981 Curve Fit: Quadratic
 Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M
 Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

Styrene

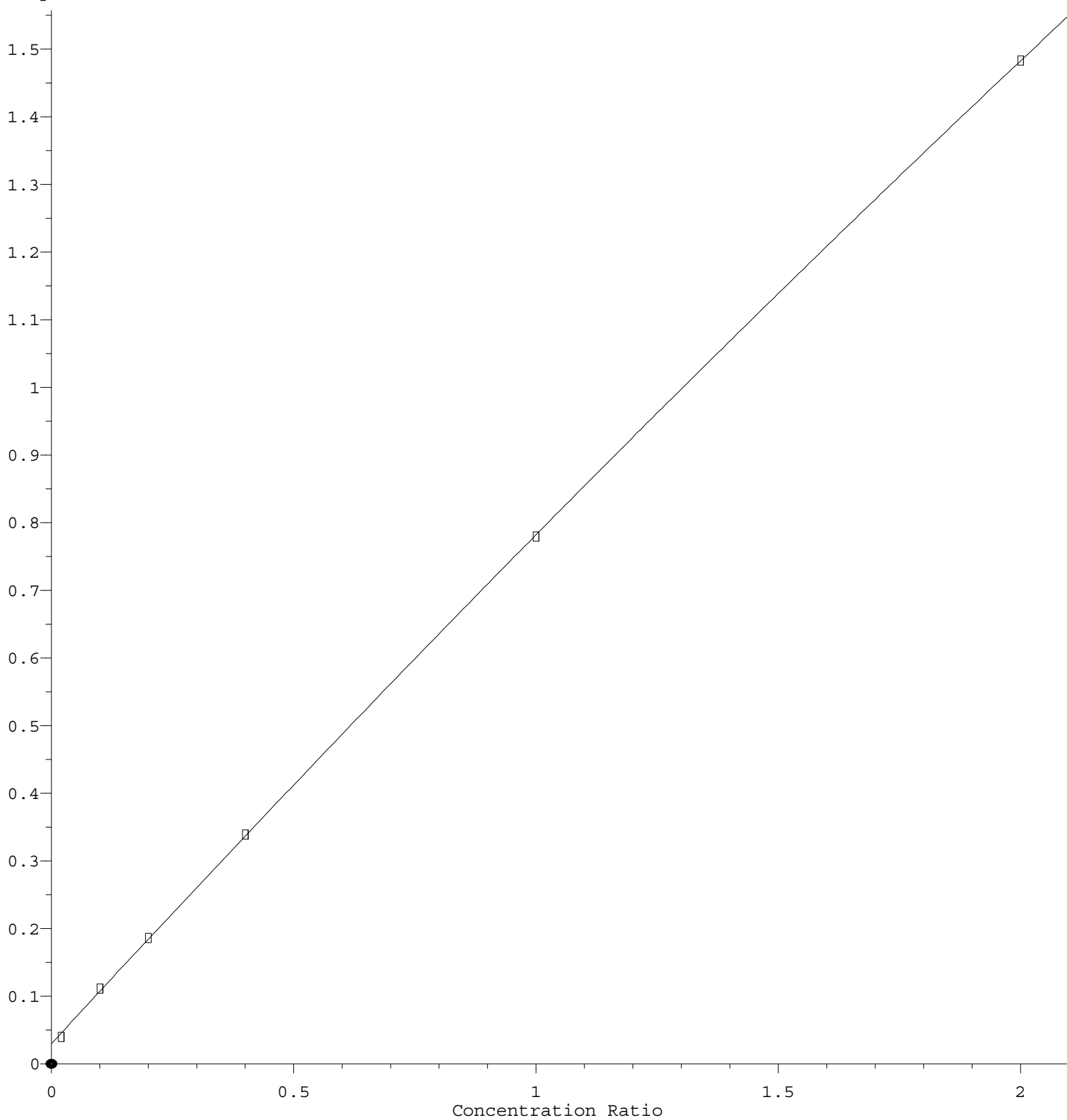
Response Ratio



$R = -1.124e-002 A^2 + 1.587e+000 A - 4.955e-002$
 Coef of Det (r^2) = 0.999540 Curve Fit: Quadratic
 Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M
 Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

Methylene Chloride

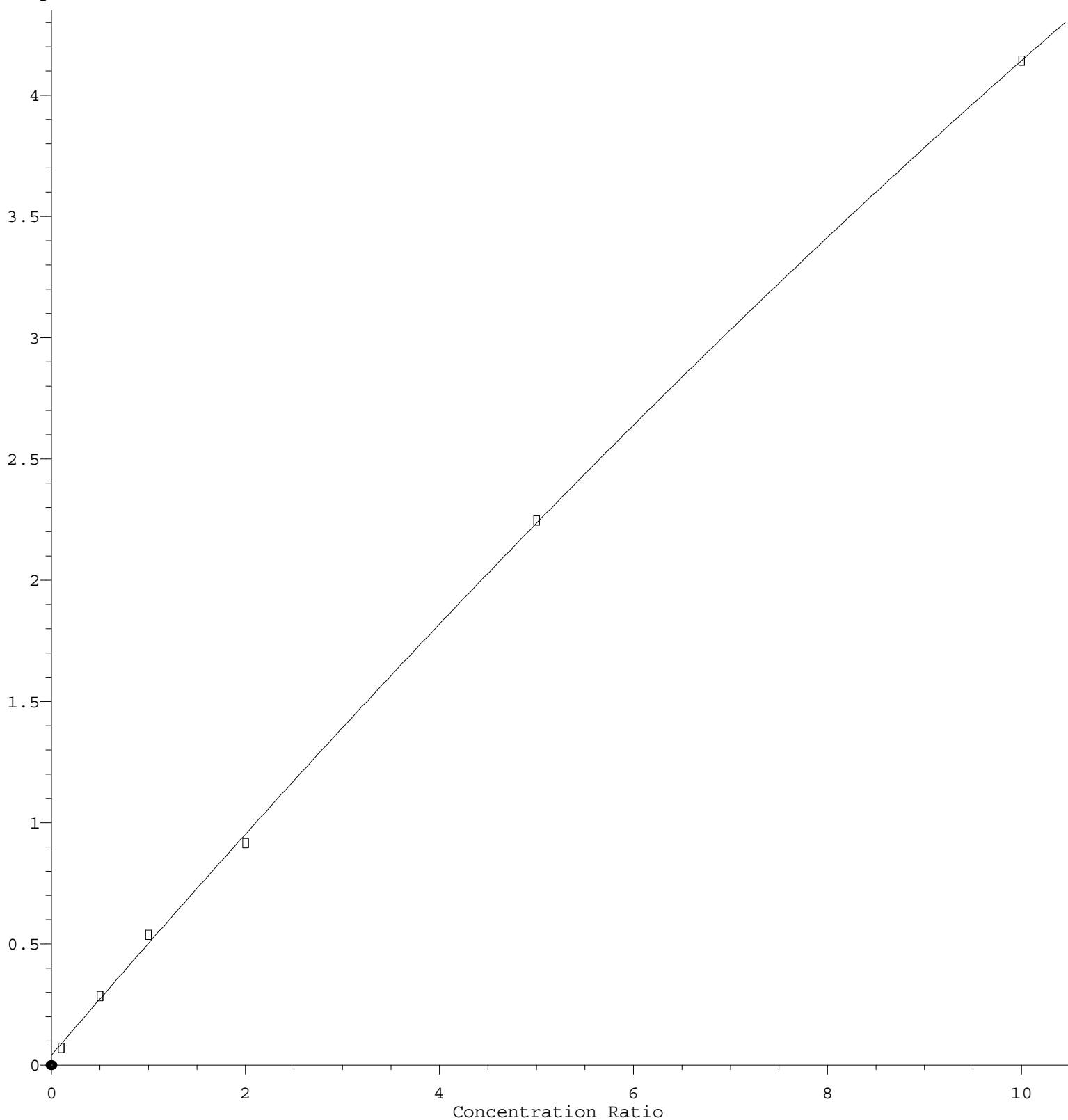
Response Ratio



$R = -2.570e-002 A^2 + 7.777e-001 A + 2.991e-002$
Coef of Det (r^2) = 0.999960 Curve Fit: Quadratic
Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M
Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

Acetone

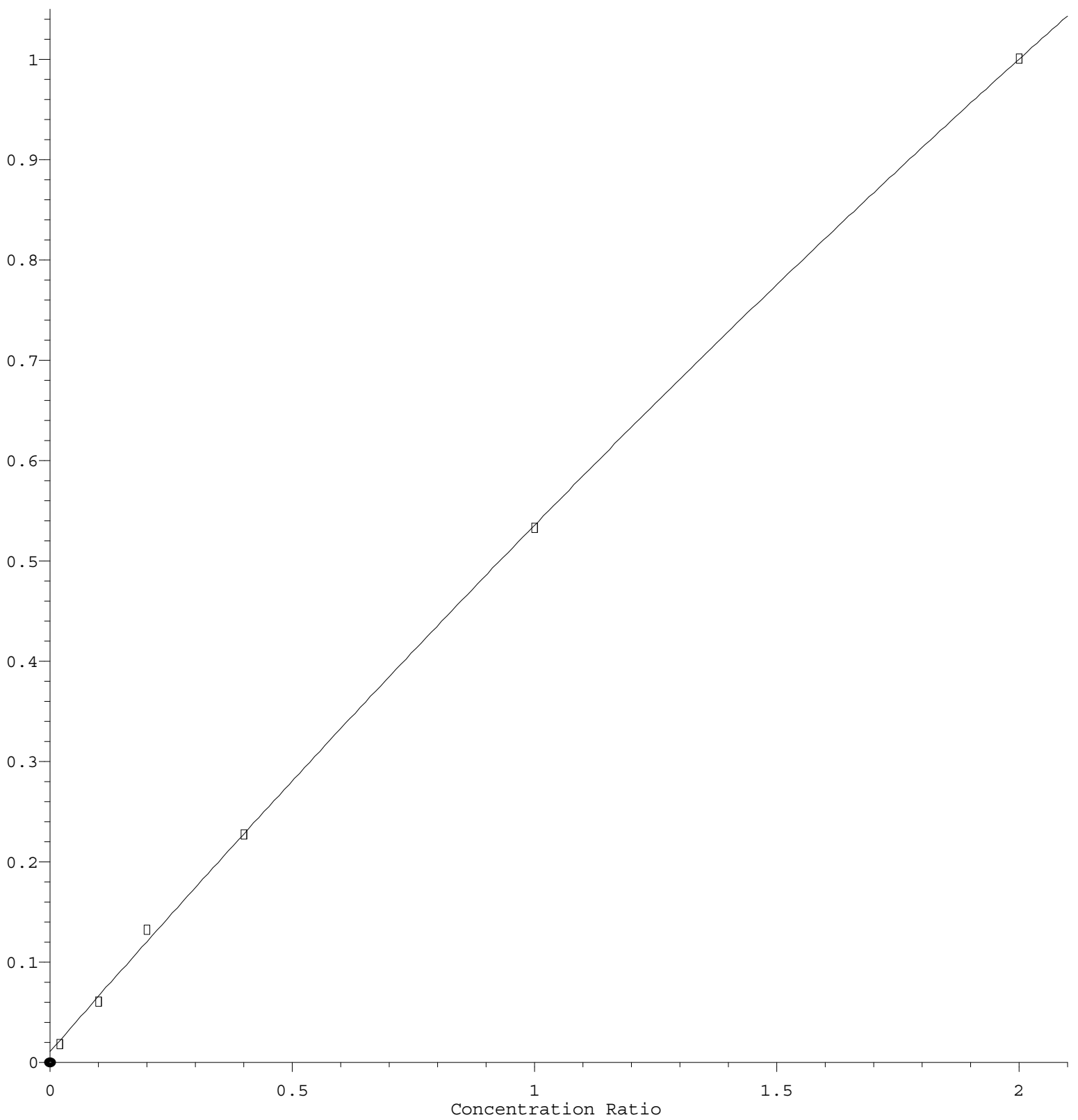
Response Ratio



$R = -5.738e-003 A^2 + 4.676e-001 A + 4.104e-002$
 Coef of Det (r^2) = 0.999745 Curve Fit: Quadratic
 Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M
 Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

Chloroethane

Response Ratio



$R = -3.006e-002 A^2 + 5.547e-001 A + 1.084e-002$
 Coef of Det (r^2) = 0.999734 Curve Fit: Quadratic
 Method Name: Z:\voasrv\HPCHEM1\MSVOA V\Method\82V092225W.M
 Calibration Table Last Updated: Wed Sep 24 08:08:08 2025

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039137.D
 Acq On : 22 Sep 2025 12:26
 Operator : SY/MD
 Sample : VSTDICC010
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 4 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICC010

Quant Time: Sep 24 07:21:49 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.597	168	501722	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.529	114	837272	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.773	117	824583	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	463438	50.000	ug/l	0.00

System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.944	65	77610	10.688	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	21.380%#
35) Dibromofluoromethane	4.503	113	72914	10.833	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	21.660%#
50) Toluene-d8	7.240	98	195300	9.879	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	19.760%#
62) 4-Bromofluorobenzene	10.005	95	80587	9.902	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	19.800%#

Target Compounds					Qvalue	
2) Dichlorodifluoromethane	1.121	85	85113	10.716	ug/l	99
3) Chloromethane	1.230	50	95215	11.167	ug/l	95
4) Vinyl Chloride	1.298	62	101023	11.068	ug/l	99
5) Bromomethane	1.500	94	63359	11.011	ug/l	98
6) Chloroethane	1.561	64	66412	11.087	ug/l	98
7) Trichlorofluoromethane	1.729	101	139927	10.588	ug/l	98
8) Diethyl Ether	1.918	74	50669	10.328	ug/l	99
9) 1,1,2-Trichlorotrifluo...	2.082	101	84221	10.711	ug/l	99
10) Methyl Iodide	2.198	142	77448	10.343	ug/l	98
11) Tert butyl alcohol	2.568	59	84609	55.130	ug/l	99
12) 1,1-Dichloroethene	2.082	96	80704	10.669	ug/l	97
13) Acrolein	2.015	56	13557	50.908	ug/l #	80
14) Allyl chloride	2.359	41	139249	10.305	ug/l	98
15) Acrylonitrile	2.674	53	254235	51.676	ug/l	99
16) Acetone	2.121	43	269573	53.773	ug/l	99
17) Carbon Disulfide	2.253	76	246125	10.639	ug/l	99
18) Methyl Acetate	2.381	43	115005	10.689	ug/l	97
19) Methyl tert-butyl Ether	2.712	73	255535	10.601	ug/l	99
20) Methylene Chloride	2.455	84	93350	10.106	ug/l	99
21) trans-1,2-Dichloroethene	2.703	96	84891	10.541	ug/l	95
22) Diisopropyl ether	3.214	45	276604	10.699	ug/l #	68
23) Vinyl Acetate	3.191	43	1200618	53.087	ug/l	100
24) 1,1-Dichloroethane	3.118	63	167496	10.594	ug/l	100
25) 2-Butanone	3.873	43	303016	53.015	ug/l	100
26) 2,2-Dichloropropane	3.809	77	136919	9.806	ug/l	100
27) cis-1,2-Dichloroethene	3.822	96	91069	10.254	ug/l	99
28) Bromochloromethane	4.146	49	83889	12.019	ug/l	98
29) Tetrahydrofuran	4.240	42	188818	51.879	ug/l	96
30) Chloroform	4.278	83	175847	10.652	ug/l	97
31) Cyclohexane	4.584	56	132996	10.269	ug/l	96
32) 1,1,1-Trichloroethane	4.513	97	156733	10.966	ug/l	96
36) 1,1-Dichloropropene	4.744	75	104420	10.187	ug/l	99
37) Ethyl Acetate	3.982	43	119710	9.520	ug/l #	91
38) Carbon Tetrachloride	4.732	117	124321	10.069	ug/l	98
39) Methylcyclohexane	6.043	83	117354	9.549	ug/l	94
40) Benzene	5.008	78	347728	10.493	ug/l	100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039137.D
 Acq On : 22 Sep 2025 12:26
 Operator : SY/MD
 Sample : VSTDICC010
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 4 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICC010

Quant Time: Sep 24 07:21:49 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.169	41	45674	9.764	ug/l	90
42) 1,2-Dichloroethane	5.043	62	121437	10.458	ug/l	100
43) Isopropyl Acetate	5.195	43	172880	9.980	ug/l	97
44) Trichloroethene	5.834	130	78943	10.324	ug/l	93
45) 1,2-Dichloropropane	6.092	63	85295	10.355	ug/l	99
46) Dibromomethane	6.230	93	65717	10.424	ug/l	98
47) Bromodichloromethane	6.426	83	139261	10.685	ug/l	97
48) Methyl methacrylate	6.304	41	68465	8.639	ug/l	97
49) 1,4-Dioxane	6.297	88	23699	190.260	ug/l	99
51) 4-Methyl-2-Pentanone	7.150	43	598702	54.051	ug/l	97
52) Toluene	7.310	92	202741	10.552	ug/l	100
53) t-1,3-Dichloropropene	7.580	75	117314	10.329	ug/l	97
54) cis-1,3-Dichloropropene	6.950	75	134529	10.729	ug/l	98
55) 1,1,2-Trichloroethane	7.764	97	89390	10.376	ug/l	98
56) Ethyl methacrylate	7.719	69	95279	9.313	ug/l	99
57) 1,3-Dichloropropane	7.934	76	144218	10.607	ug/l	98
58) 2-Chloroethyl Vinyl ether	6.815	63	220693	46.148	ug/l	99
59) 2-Hexanone	8.066	43	422703	49.020	ug/l	93
60) Dibromochloromethane	8.169	129	101236	10.313	ug/l	97
61) 1,2-Dibromoethane	8.275	107	93182	10.576	ug/l	100
64) Tetrachloroethene	7.899	164	71811	10.397	ug/l	98
65) Chlorobenzene	8.805	112	231329	10.072	ug/l	99
66) 1,1,1,2-Tetrachloroethane	8.899	131	85678	10.324	ug/l	99
67) Ethyl Benzene	8.937	91	334394	9.618	ug/l	99
68) m/p-Xylenes	9.063	106	270452	20.155	ug/l	99
69) o-Xylene	9.468	106	115922	9.756	ug/l	96
70) Styrene	9.487	104	204368	9.382	ug/l	100
71) Bromoform	9.657	173	65302	10.074	ug/l #	98
73) Isopropylbenzene	9.857	105	330305	9.827	ug/l	99
74) N-amyl acetate	9.725	43	132108	9.853	ug/l	96
75) 1,1,2,2-Tetrachloroethane	10.165	83	148267	10.067	ug/l	99
76) 1,2,3-Trichloropropane	10.198	75	105275	9.977	ug/l	99
77) Bromobenzene	10.143	156	87828	10.071	ug/l	96
78) n-propylbenzene	10.278	91	401316	9.804	ug/l	100
79) 2-Chlorotoluene	10.349	91	255053	10.413	ug/l	99
80) 1,3,5-Trimethylbenzene	10.464	105	270977	9.699	ug/l	98
81) trans-1,4-Dichloro-2-b...	9.931	75	44314	10.082	ug/l	97
82) 4-Chlorotoluene	10.464	91	292863	10.312	ug/l	100
83) tert-Butylbenzene	10.789	119	267528	9.603	ug/l	98
84) 1,2,4-Trimethylbenzene	10.841	105	256505	9.457	ug/l	99
85) sec-Butylbenzene	11.011	105	354312	9.588	ug/l	100
86) p-Isopropyltoluene	11.169	119	300032	9.755	ug/l	99
87) 1,3-Dichlorobenzene	11.108	146	178990	10.141	ug/l	98
88) 1,4-Dichlorobenzene	11.198	146	191843	9.835	ug/l	97
89) n-Butylbenzene	11.580	91	281258	9.575	ug/l	97
90) Hexachloroethane	11.821	117	74910	9.891	ug/l	96
91) 1,2-Dichlorobenzene	11.567	146	162271	9.841	ug/l	98
92) 1,2-Dibromo-3-Chloropr...	12.355	75	27674	10.167	ug/l	98
93) 1,2,4-Trichlorobenzene	13.188	180	85982	9.096	ug/l	98
94) Hexachlorobutadiene	13.368	225	47413	9.653	ug/l	97
95) Naphthalene	13.426	128	267645	8.803	ug/l	99
96) 1,2,3-Trichlorobenzene	13.667	180	90238	9.334	ug/l	100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039137.D
Acq On : 22 Sep 2025 12:26
Operator : SY/MD
Sample : VSTDICC010
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 4 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC010

Quant Time: Sep 24 07:21:49 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 07:14:23 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC010

[illegible]

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039138.D
 Acq On : 22 Sep 2025 12:48
 Operator : SY/MD
 Sample : VSTDIC020
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 5 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDIC020

Quant Time: Sep 24 07:22:45 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.600	168	545775	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	880963	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.773	117	860340	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	509460	50.000	ug/l	0.00

System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.947	65	144689	18.317	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	36.640%#
35) Dibromofluoromethane	4.503	113	134662	19.014	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	38.020%#
50) Toluene-d8	7.236	98	384827	18.501	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	37.000%#
62) 4-Bromofluorobenzene	10.002	95	163597	19.105	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	38.220%#

Target Compounds				Qvalue		
2) Dichlorodifluoromethane	1.121	85	165175	19.118	ug/l	96
3) Chloromethane	1.230	50	177694	19.159	ug/l	99
4) Vinyl Chloride	1.298	62	186887	18.822	ug/l	99
5) Bromomethane	1.500	94	119578	19.104	ug/l	98
6) Chloroethane	1.561	64	124021	19.935	ug/l	99
7) Trichlorofluoromethane	1.729	101	273532	19.027	ug/l	100
8) Diethyl Ether	1.921	74	100661	18.862	ug/l	99
9) 1,1,2-Trichlorotrifluo...	2.085	101	159719	18.673	ug/l	99
10) Methyl Iodide	2.198	142	161932	19.880	ug/l	99
11) Tert butyl alcohol	2.568	59	171137	102.509	ug/l	98
12) 1,1-Dichloroethene	2.085	96	157427	19.131	ug/l	97
13) Acrolein	2.011	56	29609	102.210	ug/l	97
14) Allyl chloride	2.359	41	280200	19.062	ug/l	99
15) Acrylonitrile	2.674	53	504022	94.179	ug/l	99
16) Acetone	2.121	43	499669	95.756	ug/l	99
17) Carbon Disulfide	2.253	76	476229	18.923	ug/l	100
18) Methyl Acetate	2.381	43	226208	19.328	ug/l	98
19) Methyl tert-butyl Ether	2.712	73	498918	19.027	ug/l	99
20) Methylene Chloride	2.458	84	185050	20.143	ug/l	97
21) trans-1,2-Dichloroethene	2.703	96	164898	18.823	ug/l	97
22) Diisopropyl ether	3.217	45	552331	19.640	ug/l	83
23) Vinyl Acetate	3.191	43	2445787	99.414	ug/l	100
24) 1,1-Dichloroethane	3.118	63	320146	18.615	ug/l	99
25) 2-Butanone	3.870	43	637413	102.519	ug/l	100
26) 2,2-Dichloropropane	3.812	77	278362	18.326	ug/l	98
27) cis-1,2-Dichloroethene	3.822	96	183867	19.031	ug/l	98
28) Bromochloromethane	4.146	49	164406	21.654	ug/l	99
29) Tetrahydrofuran	4.240	42	405777	102.491	ug/l	99
30) Chloroform	4.281	83	342700	19.083	ug/l	99
31) Cyclohexane	4.587	56	266753	18.934	ug/l	98
32) 1,1,1-Trichloroethane	4.516	97	288341	18.546	ug/l	99
36) 1,1-Dichloropropene	4.748	75	212835	19.734	ug/l	99
37) Ethyl Acetate	3.979	43	260591	19.696	ug/l	97
38) Carbon Tetrachloride	4.735	117	250103	19.252	ug/l	98
39) Methylcyclohexane	6.047	83	253693	19.619	ug/l	95
40) Benzene	5.011	78	697141	19.994	ug/l	99

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VW092225\
 Data File : VW039138.D
 Acq On : 22 Sep 2025 12:48
 Operator : SY/MD
 Sample : VSTDIC020
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 5 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDIC020

Quant Time: Sep 24 07:22:45 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.169	41	102755	18.721	ug/l	95
42) 1,2-Dichloroethane	5.043	62	244487	20.012	ug/l	99
43) Isopropyl Acetate	5.195	43	362512	19.889	ug/l	98
44) Trichloroethene	5.831	130	160692	19.972	ug/l	96
45) 1,2-Dichloropropane	6.092	63	182757	21.087	ug/l	96
46) Dibromomethane	6.227	93	129319	19.495	ug/l	98
47) Bromodichloromethane	6.429	83	269262	19.635	ug/l	97
48) Methyl methacrylate	6.301	41	158251	18.977	ug/l	99
49) 1,4-Dioxane	6.294	88	47860	365.172	ug/l #	85
51) 4-Methyl-2-Pentanone	7.150	43	1296796	111.268	ug/l	98
52) Toluene	7.310	92	422367	20.892	ug/l	100
53) t-1,3-Dichloropropene	7.577	75	245181	20.516	ug/l	98
54) cis-1,3-Dichloropropene	6.950	75	273906	20.761	ug/l	100
55) 1,1,2-Trichloroethane	7.760	97	176812	19.505	ug/l	98
56) Ethyl methacrylate	7.719	69	212320	19.723	ug/l	97
57) 1,3-Dichloropropane	7.934	76	291304	20.363	ug/l	100
58) 2-Chloroethyl Vinyl ether	6.815	63	536254	94.274	ug/l	99
59) 2-Hexanone	8.066	43	940265	101.225	ug/l	98
60) Dibromochloromethane	8.169	129	200793	19.440	ug/l	99
61) 1,2-Dibromoethane	8.275	107	189555	20.447	ug/l	99
64) Tetrachloroethene	7.899	164	141876	19.688	ug/l	99
65) Chlorobenzene	8.805	112	470841	19.648	ug/l	97
66) 1,1,1,2-Tetrachloroethane	8.899	131	167936	19.395	ug/l	99
67) Ethyl Benzene	8.937	91	737391	20.328	ug/l	99
68) m/p-Xylenes	9.063	106	607362	43.381	ug/l	99
69) o-Xylene	9.468	106	255212	20.587	ug/l	100
70) Styrene	9.487	104	475382	19.022	ug/l	99
71) Bromoform	9.654	173	132242	19.552	ug/l	98
73) Isopropylbenzene	9.857	105	741305	20.062	ug/l	99
74) N-amyl acetate	9.725	43	306963	20.826	ug/l	99
75) 1,1,2,2-Tetrachloroethane	10.165	83	297741	18.390	ug/l	100
76) 1,2,3-Trichloropropane	10.198	75	213532	18.409	ug/l	99
77) Bromobenzene	10.140	156	187188	19.525	ug/l	96
78) n-propylbenzene	10.278	91	905281	20.118	ug/l	99
79) 2-Chlorotoluene	10.349	91	547836	20.346	ug/l	99
80) 1,3,5-Trimethylbenzene	10.464	105	640175	20.843	ug/l	99
81) trans-1,4-Dichloro-2-b...	9.928	75	91848	19.009	ug/l	96
82) 4-Chlorotoluene	10.461	91	655427	20.993	ug/l	99
83) tert-Butylbenzene	10.789	119	587151	19.172	ug/l	99
84) 1,2,4-Trimethylbenzene	10.841	105	631679	21.185	ug/l	100
85) sec-Butylbenzene	11.011	105	808869	19.912	ug/l	99
86) p-Isopropyltoluene	11.165	119	682814	20.196	ug/l	100
87) 1,3-Dichlorobenzene	11.104	146	374208	19.287	ug/l	99
88) 1,4-Dichlorobenzene	11.198	146	397526	18.539	ug/l	100
89) n-Butylbenzene	11.580	91	640044	19.822	ug/l	99
90) Hexachloroethane	11.818	117	150843	18.118	ug/l	99
91) 1,2-Dichlorobenzene	11.567	146	336566	18.567	ug/l	98
92) 1,2-Dibromo-3-Chloropr...	12.352	75	57671	19.989	ug/l	98
93) 1,2,4-Trichlorobenzene	13.185	180	196334	18.894	ug/l	99
94) Hexachlorobutadiene	13.368	225	96610	17.893	ug/l	98
95) Naphthalene	13.426	128	634557	18.986	ug/l	99
96) 1,2,3-Trichlorobenzene	13.667	180	198308	18.659	ug/l	97

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039138.D
Acq On : 22 Sep 2025 12:48
Operator : SY/MD
Sample : VSTDICC020
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 5 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC020

Quant Time: Sep 24 07:22:45 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 07:14:23 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC020

[illegible]

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039139.D
 Acq On : 22 Sep 2025 13:26
 Operator : SY/MD
 Sample : VSTDICCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 6 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICCC050

Quant Time: Sep 24 07:23:42 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.597	168	563410	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.529	114	894173	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.773	117	863906	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	516167	50.000	ug/l	0.00

System Monitoring Compounds

33) 1,2-Dichloroethane-d4	4.937	65	363821	44.618	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	89.240%
35) Dibromofluoromethane	4.497	113	337386	46.935	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	93.860%
50) Toluene-d8	7.233	98	1016135	48.130	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	96.260%
62) 4-Bromofluorobenzene	9.998	95	453081	52.130	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	104.260%

Target Compounds

						Qvalue

2) Dichlorodifluoromethane	1.121	85	416798	46.733	ug/l	100
3) Chloromethane	1.230	50	433999	45.329	ug/l	100
4) Vinyl Chloride	1.298	62	470936	45.944	ug/l	100
5) Bromomethane	1.494	94	286332	44.312	ug/l	100
6) Chloroethane	1.558	64	300232	49.734	ug/l	100
7) Trichlorofluoromethane	1.725	101	672325	45.304	ug/l	100
8) Diethyl Ether	1.918	74	251813	45.707	ug/l	100
9) 1,1,2-Trichlorotrifluo...	2.082	101	395615	44.803	ug/l	100
10) Methyl Iodide	2.195	142	442822	52.662	ug/l	100
11) Tert butyl alcohol	2.564	59	385746	223.826	ug/l	100
12) 1,1-Dichloroethene	2.082	96	379710	44.700	ug/l	100
13) Acrolein	2.008	56	74486	249.076	ug/l	100
14) Allyl chloride	2.355	41	700342	46.154	ug/l	100
15) Acrylonitrile	2.671	53	1228907	222.439	ug/l	100
16) Acetone	2.118	43	1265424	251.263	ug/l	100
17) Carbon Disulfide	2.253	76	1172348	45.126	ug/l	100
18) Methyl Acetate	2.378	43	553676	45.827	ug/l	100
19) Methyl tert-butyl Ether	2.709	73	1258060	46.476	ug/l	100
20) Methylene Chloride	2.455	84	439142	49.828	ug/l	100
21) trans-1,2-Dichloroethene	2.699	96	407938	45.109	ug/l	100
22) Diisopropyl ether	3.214	45	1419527	48.895	ug/l	100
23) Vinyl Acetate	3.185	43	6209687	244.506	ug/l	100
24) 1,1-Dichloroethane	3.114	63	781832	44.037	ug/l	100
25) 2-Butanone	3.860	43	1650213	257.105	ug/l	100
26) 2,2-Dichloropropane	3.806	77	698283	44.534	ug/l	100
27) cis-1,2-Dichloroethene	3.818	96	488972	49.026	ug/l	100
28) Bromochloromethane	4.143	49	382640	48.820	ug/l	100
29) Tetrahydrofuran	4.230	42	1039169	254.258	ug/l	100
30) Chloroform	4.275	83	839217	45.268	ug/l	100
31) Cyclohexane	4.580	56	701482	48.232	ug/l	100
32) 1,1,1-Trichloroethane	4.510	97	731030	45.547	ug/l	100
36) 1,1-Dichloropropene	4.741	75	554798	50.680	ug/l	100
37) Ethyl Acetate	3.970	43	638131	47.519	ug/l	100
38) Carbon Tetrachloride	4.732	117	628516	47.666	ug/l	100
39) Methylcyclohexane	6.043	83	736231	56.095	ug/l	100
40) Benzene	5.005	78	1779301	50.276	ug/l	100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VW092225\
 Data File : VW039139.D
 Acq On : 22 Sep 2025 13:26
 Operator : SY/MD
 Sample : VSTDICCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 6 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICCC050

Quant Time: Sep 24 07:23:42 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.153	41	295316	49.541	ug/l	100
42) 1,2-Dichloroethane	5.037	62	595216	48.000	ug/l	100
43) Isopropyl Acetate	5.188	43	956288	51.691	ug/l	100
44) Trichloroethene	5.828	130	414171	50.715	ug/l	100
45) 1,2-Dichloropropane	6.085	63	448750	51.014	ug/l	100
46) Dibromomethane	6.220	93	329073	48.876	ug/l	100
47) Bromodichloromethane	6.423	83	673646	48.398	ug/l	100
48) Methyl methacrylate	6.291	41	450230	53.193	ug/l	100
49) 1,4-Dioxane	6.288	88	133834	1006.069	ug/l	100
51) 4-Methyl-2-Pentanone	7.146	43	3273840	276.753	ug/l	100
52) Toluene	7.304	92	1099325	53.574	ug/l	100
53) t-1,3-Dichloropropene	7.571	75	650976	53.666	ug/l	100
54) cis-1,3-Dichloropropene	6.944	75	725015	54.142	ug/l	100
55) 1,1,2-Trichloroethane	7.757	97	441822	48.020	ug/l	100
56) Ethyl methacrylate	7.712	69	640636	58.632	ug/l	100
57) 1,3-Dichloropropane	7.931	76	733341	50.505	ug/l	100
58) 2-Chloroethyl Vinyl ether	6.809	63	1579407	255.700	ug/l	100
59) 2-Hexanone	8.059	43	2441880	255.630	ug/l	100
60) Dibromochloromethane	8.165	129	507043	48.365	ug/l	100
61) 1,2-Dibromoethane	8.272	107	473037	50.273	ug/l	100
64) Tetrachloroethene	7.895	164	356209	49.227	ug/l	100
65) Chlorobenzene	8.802	112	1194769	49.651	ug/l	100
66) 1,1,1,2-Tetrachloroethane	8.895	131	421762	48.507	ug/l	100
67) Ethyl Benzene	8.934	91	2057558	56.487	ug/l	100
68) m/p-Xylenes	9.059	106	1639130	116.591	ug/l	100
69) o-Xylene	9.468	106	730558	58.688	ug/l	100
70) Styrene	9.484	104	1341298	50.845	ug/l	100
71) Bromoform	9.651	173	341871	50.337	ug/l	100
73) Isopropylbenzene	9.854	105	2033373	54.313	ug/l	100
74) N-amyl acetate	9.722	43	863632	57.833	ug/l	100
75) 1,1,2,2-Tetrachloroethane	10.165	83	735590	44.844	ug/l	100
76) 1,2,3-Trichloropropane	10.194	75	541552	46.080	ug/l	100
77) Bromobenzene	10.140	156	477608	49.171	ug/l	100
78) n-propylbenzene	10.275	91	2528458	55.461	ug/l	100
79) 2-Chlorotoluene	10.345	91	1462530	53.612	ug/l	100
80) 1,3,5-Trimethylbenzene	10.464	105	1748435	56.188	ug/l	100
81) trans-1,4-Dichloro-2-b...	9.928	75	255259	52.143	ug/l	100
82) 4-Chlorotoluene	10.461	91	1722899	54.466	ug/l	100
83) tert-Butylbenzene	10.789	119	1663245	53.602	ug/l	100
84) 1,2,4-Trimethylbenzene	10.837	105	1743425	57.709	ug/l	100
85) sec-Butylbenzene	11.011	105	2252781	54.737	ug/l	100
86) p-Isopropyltoluene	11.165	119	1900412	55.479	ug/l	100
87) 1,3-Dichlorobenzene	11.104	146	958701	48.769	ug/l	100
88) 1,4-Dichlorobenzene	11.194	146	998426	45.957	ug/l	100
89) n-Butylbenzene	11.577	91	1864844	57.003	ug/l	100
90) Hexachloroethane	11.818	117	375818	44.553	ug/l	100
91) 1,2-Dichlorobenzene	11.567	146	897135	48.848	ug/l	100
92) 1,2-Dibromo-3-Chloropr...	12.349	75	147329	49.928	ug/l	100
93) 1,2,4-Trichlorobenzene	13.185	180	555323	52.747	ug/l	100
94) Hexachlorobutadiene	13.368	225	247695	45.279	ug/l	100
95) Naphthalene	13.423	128	1968567	58.135	ug/l	100
96) 1,2,3-Trichlorobenzene	13.664	180	571563	53.081	ug/l	100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039139.D
Acq On : 22 Sep 2025 13:26
Operator : SY/MD
Sample : VSTDICCC050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 6 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VSTDICCC050

Quant Time: Sep 24 07:23:42 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 07:14:23 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VSTDICCC050

[illegible]

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039140.D
 Acq On : 22 Sep 2025 13:49
 Operator : SY/MD
 Sample : VSTDICC100
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 7 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICC100

Quant Time: Sep 24 07:24:39 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.600	168	587274	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	935392	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	904902	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	523452	50.000	ug/l	0.00

System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.941	65	725543	85.362	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	= 170.720%#	
35) Dibromofluoromethane	4.500	113	674408	89.684	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	= 179.360%#	
50) Toluene-d8	7.236	98	2052477	92.932	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	= 185.860%#	
62) 4-Bromofluorobenzene	9.998	95	937505	103.112	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	= 206.220%#	

Target Compounds					Qvalue	
2) Dichlorodifluoromethane	1.121	85	820676	88.278	ug/l	99
3) Chloromethane	1.230	50	856347	85.806	ug/l	98
4) Vinyl Chloride	1.298	62	940694	88.044	ug/l	99
5) Bromomethane	1.487	94	615924	91.446	ug/l	99
6) Chloroethane	1.555	64	587651	100.067	ug/l	99
7) Trichlorofluoromethane	1.722	101	1351410	87.363	ug/l	99
8) Diethyl Ether	1.918	74	511033	88.990	ug/l	100
9) 1,1,2-Trichlorotrifluo...	2.079	101	806216	87.593	ug/l	98
10) Methyl Iodide	2.195	142	931666	106.294	ug/l	100
11) Tert butyl alcohol	2.574	59	798702	444.608	ug/l	99
12) 1,1-Dichloroethene	2.079	96	778436	87.914	ug/l	98
13) Acrolein	2.008	56	160749	515.691	ug/l	99
14) Allyl chloride	2.355	41	1418889	89.708	ug/l	99
15) Acrylonitrile	2.674	53	2509475	435.771	ug/l	100
16) Acetone	2.118	43	2432444	499.795	ug/l	100
17) Carbon Disulfide	2.253	76	2332239	86.125	ug/l	99
18) Methyl Acetate	2.378	43	1142900	90.753	ug/l	100
19) Methyl tert-butyl Ether	2.712	73	2586428	91.667	ug/l	99
20) Methylene Chloride	2.455	84	870955	100.038	ug/l	99
21) trans-1,2-Dichloroethene	2.699	96	817275	86.701	ug/l	98
22) Diisopropyl ether	3.217	45	2857568	94.428	ug/l	94
23) Vinyl Acetate	3.188	43	12498986	472.148	ug/l	99
24) 1,1-Dichloroethane	3.117	63	1560774	84.338	ug/l	99
25) 2-Butanone	3.863	43	3404031	508.801	ug/l	99
26) 2,2-Dichloropropane	3.812	77	1414724	86.559	ug/l	100
27) cis-1,2-Dichloroethene	3.818	96	1010617	97.210	ug/l	99
28) Bromochloromethane	4.146	49	748549	91.624	ug/l	98
29) Tetrahydrofuran	4.233	42	2198745	516.116	ug/l	99
30) Chloroform	4.278	83	1682895	87.087	ug/l	98
31) Cyclohexane	4.580	56	1473260	97.182	ug/l	97
32) 1,1,1-Trichloroethane	4.513	97	1483938	88.700	ug/l	99
36) 1,1-Dichloropropene	4.741	75	1144520	99.944	ug/l	100
37) Ethyl Acetate	3.973	43	1328655	94.580	ug/l	98
38) Carbon Tetrachloride	4.735	117	1263028	91.566	ug/l	100
39) Methylcyclohexane	6.047	83	1603169	116.766	ug/l	98
40) Benzene	5.008	78	3611886	97.560	ug/l	98

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VW092225\
 Data File : VW039140.D
 Acq On : 22 Sep 2025 13:49
 Operator : SY/MD
 Sample : VSTDICC100
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 7 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICC100

Quant Time: Sep 24 07:24:39 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.156	41	639301	100.494	ug/l	99
42) 1,2-Dichloroethane	5.037	62	1202991	92.737	ug/l	100
43) Isopropyl Acetate	5.191	43	2058622	106.372	ug/l	100
44) Trichloroethene	5.828	130	849813	99.474	ug/l	97
45) 1,2-Dichloropropane	6.088	63	909288	98.813	ug/l	95
46) Dibromomethane	6.220	93	661336	93.898	ug/l	98
47) Bromodichloromethane	6.426	83	1355845	93.118	ug/l	99
48) Methyl methacrylate	6.294	41	984431	111.182	ug/l	99
49) 1,4-Dioxane	6.288	88	304943	2191.331	ug/l	95
51) 4-Methyl-2-Pentanone	7.149	43	6674594	539.372	ug/l	99
52) Toluene	7.307	92	2271340	105.813	ug/l	99
53) t-1,3-Dichloropropene	7.571	75	1408125	110.970	ug/l	97
54) cis-1,3-Dichloropropene	6.947	75	1501380	107.178	ug/l	100
55) 1,1,2-Trichloroethane	7.757	97	897804	93.280	ug/l	99
56) Ethyl methacrylate	7.712	69	1424432	124.621	ug/l	99
57) 1,3-Dichloropropane	7.931	76	1501225	98.832	ug/l	100
58) 2-Chloroethyl Vinyl ether	6.812	63	3282267	498.705	ug/l	100
59) 2-Hexanone	8.063	43	4988835	497.186	ug/l	99
60) Dibromochloromethane	8.165	129	1044126	95.206	ug/l	99
61) 1,2-Dibromoethane	8.272	107	960914	97.622	ug/l	98
64) Tetrachloroethene	7.895	164	736462	97.167	ug/l	97
65) Chlorobenzene	8.802	112	2457021	97.480	ug/l	97
66) 1,1,1,2-Tetrachloroethane	8.895	131	863523	94.816	ug/l	100
67) Ethyl Benzene	8.934	91	4376266	114.700	ug/l	100
68) m/p-Xylenes	9.059	106	3396653	230.657	ug/l	100
69) o-Xylene	9.468	106	1592963	122.170	ug/l	98
70) Styrene	9.484	104	2781887	99.832	ug/l	100
71) Bromoform	9.651	173	724467	101.838	ug/l	99
73) Isopropylbenzene	9.854	105	4288139	112.946	ug/l	100
74) N-amyl acetate	9.722	43	1846779	121.948	ug/l	98
75) 1,1,2,2-Tetrachloroethane	10.165	83	1453989	87.406	ug/l	100
76) 1,2,3-Trichloropropane	10.198	75	1070580	89.827	ug/l	95
77) Bromobenzene	10.140	156	993863	100.897	ug/l	99
78) n-propylbenzene	10.278	91	5220220	112.910	ug/l	99
79) 2-Chlorotoluene	10.345	91	3018051	109.093	ug/l	99
80) 1,3,5-Trimethylbenzene	10.464	105	3638030	115.284	ug/l	100
81) trans-1,4-Dichloro-2-b...	9.927	75	545452	109.871	ug/l	97
82) 4-Chlorotoluene	10.461	91	3573507	111.396	ug/l	100
83) tert-Butylbenzene	10.789	119	3572813	113.541	ug/l	99
84) 1,2,4-Trimethylbenzene	10.837	105	3601355	117.550	ug/l	100
85) sec-Butylbenzene	11.011	105	4763689	114.135	ug/l	100
86) p-Isopropyltoluene	11.165	119	3948556	113.666	ug/l	99
87) 1,3-Dichlorobenzene	11.104	146	1970594	98.849	ug/l	99
88) 1,4-Dichlorobenzene	11.194	146	2008676	91.172	ug/l	99
89) n-Butylbenzene	11.577	91	3939852	118.754	ug/l	99
90) Hexachloroethane	11.818	117	776688	90.794	ug/l	99
91) 1,2-Dichlorobenzene	11.564	146	1886394	101.282	ug/l	100
92) 1,2-Dibromo-3-Chloropr...	12.349	75	316446	100.015	ug/l	96
93) 1,2,4-Trichlorobenzene	13.185	180	1244510	116.564	ug/l	100
94) Hexachlorobutadiene	13.368	225	524789	94.597	ug/l	99
95) Naphthalene	13.423	128	4442612	129.371	ug/l	100
96) 1,2,3-Trichlorobenzene	13.667	180	1246406	114.143	ug/l	99

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039140.D
Acq On : 22 Sep 2025 13:49
Operator : SY/MD
Sample : VSTDICC100
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 7 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC100

Quant Time: Sep 24 07:24:39 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 07:14:23 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039142.D
 Acq On : 22 Sep 2025 15:37
 Operator : SY/MD
 Sample : VSTDICC005
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 10 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICC005

Manual Integrations
 APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
 Supervised By :Semsettin Yesilyurt 09/26/2025

Quant Time: Sep 24 07:25:48 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.596	168	492799	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.529	114	835334	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.773	117	807740	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	425480	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.947	65	45267	6.347	ug/l	0.00
Spiked Amount 50.000	Range 81 - 118		Recovery =	12.700%#		
35) Dibromofluoromethane	4.503	113	37959	5.653	ug/l	0.00
Spiked Amount 50.000	Range 80 - 119		Recovery =	11.300%#		
50) Toluene-d8	7.239	98	117860	5.976	ug/l	0.00
Spiked Amount 50.000	Range 89 - 112		Recovery =	11.960%#		
62) 4-Bromofluorobenzene	10.008	95	39819	4.904	ug/l	0.00
Spiked Amount 50.000	Range 85 - 114		Recovery =	9.800%#		
Target Compounds					Qvalue	
2) Dichlorodifluoromethane	1.121	85	38031	4.875	ug/l	93
3) Chloromethane	1.230	50	43830	5.234	ug/l	97
4) Vinyl Chloride	1.298	62	45941	5.124	ug/l	99
5) Bromomethane	1.494	94	32301	5.715	ug/l	100
6) Chloroethane	1.558	64	29857	4.506	ug/l	93
7) Trichlorofluoromethane	1.725	101	67248	5.181	ug/l	100
8) Diethyl Ether	1.918	74	25304	5.251	ug/l	95
9) 1,1,2-Trichlorotrifluo...	2.082	101	39608	5.128	ug/l	99
10) Methyl Iodide	2.195	142	31463	4.278	ug/l	95
11) Tert butyl alcohol	2.571	59	40994	27.195	ug/l	97
12) 1,1-Dichloroethene	2.082	96	38859	5.230	ug/l	96
13) Acrolein	2.015	56	6095	23.302	ug/l	98
14) Allyl chloride	2.355	41	70683	5.326	ug/l	96
15) Acrylonitrile	2.677	53	128729	26.639	ug/l	99
16) Acetone	2.121	43	140190	26.199	ug/l	98
17) Carbon Disulfide	2.253	76	121350	5.340	ug/l	99
18) Methyl Acetate	2.381	43	54053	5.115	ug/l	97
19) Methyl tert-butyl Ether	2.712	73	125944	5.319	ug/l	96
20) Methylene Chloride	2.455	84	54859	5.252	ug/l	96
21) trans-1,2-Dichloroethene	2.703	96	41151	5.202	ug/l	98
22) Diisopropyl ether	3.217	45	126181	4.969	ug/l #	64
23) Vinyl Acetate	3.191	43	570043	25.661	ug/l	99
24) 1,1-Dichloroethane	3.117	63	81764	5.265	ug/l	96
25) 2-Butanone	3.879	43	140360	25.002	ug/l	99
26) 2,2-Dichloropropane	3.812	77	68656	5.006	ug/l	99
27) cis-1,2-Dichloroethene	3.825	96	40984	4.698	ug/l	88
28) Bromochloromethane	4.153	49	25874	3.774	ug/l	97
29) Tetrahydrofuran	4.243	42	86190	24.110	ug/l	94
30) Chloroform	4.281	83	86066	5.308	ug/l	99
31) Cyclohexane	4.580	56	69326	5.450	ug/l #	94
32) 1,1,1-Trichloroethane	4.513	97	74539	5.310	ug/l	95
36) 1,1-Dichloropropene	4.744	75	44786	4.379	ug/l	99
37) Ethyl Acetate	3.995	43	60285	4.805	ug/l #	73
38) Carbon Tetrachloride	4.732	117	61498	4.992	ug/l	96
39) Methylcyclohexane	6.047	83	56805	4.633	ug/l	92
40) Benzene	5.014	78	163948	4.959	ug/l	100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039142.D
 Acq On : 22 Sep 2025 15:37
 Operator : SY/MD
 Sample : VSTDICC005
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 10 Sample Multiplier: 1

Instrument :

MSVOA_V

ClientSampleId :

VSTDICC005

Manual Integrations

APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025

Supervised By :Semsettin Yesilyurt 09/26/2025

Quant Time: Sep 24 07:25:48 2025

Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M

Quant Title : SW846 8260

QLast Update : Wed Sep 24 07:14:23 2025

Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.188	41	17959	4.995	ug/l #	86
42) 1,2-Dichloroethane	5.047	62	57857	4.994	ug/l	96
43) Isopropyl Acetate	5.204	43	77935	4.509	ug/l #	93
44) Trichloroethene	5.838	130	38395	5.033	ug/l	89
45) 1,2-Dichloropropane	6.098	63	39327	4.786	ug/l	100
46) Dibromomethane	6.233	93	31863	5.066	ug/l	98
47) Bromodichloromethane	6.429	83	64039	4.925	ug/l	95
48) Methyl methacrylate	6.313	41	29705	3.757	ug/l	93
49) 1,4-Dioxane	6.307	88	8999	72.413	ug/l #	79
51) 4-Methyl-2-Pentanone	7.153	43	255889	23.155	ug/l	99
52) Toluene	7.313	92	94414	4.925	ug/l	99
53) t-1,3-Dichloropropene	7.583	75	53878	4.755	ug/l	94
54) cis-1,3-Dichloropropene	6.957	75	58800	4.700	ug/l	98
55) 1,1,2-Trichloroethane	7.764	97	43788	5.094	ug/l	99
56) Ethyl methacrylate	7.722	69	42598	4.173	ug/l	97
57) 1,3-Dichloropropane	7.937	76	63818	4.705	ug/l	99
58) 2-Chloroethyl Vinyl ether	6.818	63	81984	23.077	ug/l	100
59) 2-Hexanone	8.069	43	179185	22.071	ug/l	88
60) Dibromochloromethane	8.169	129	49953	5.100	ug/l	96
61) 1,2-Dibromoethane	8.278	107	44729	5.088	ug/l	98
64) Tetrachloroethene	7.899	164	34795	5.143	ug/l	98
65) Chlorobenzene	8.805	112	114424	5.086	ug/l	98
66) 1,1,1,2-Tetrachloroethane	8.899	131	40690	5.005	ug/l	95
67) Ethyl Benzene	8.940	91	157790	4.633	ug/l	97
68) m/p-Xylenes	9.066	106	115177	8.762	ug/l	98
69) o-Xylene	9.471	106	51281	4.406	ug/l	100
70) Styrene	9.490	104	83873	4.836	ug/l	96
71) Bromoform	9.657	173	31133	4.903	ug/l #	97
73) Isopropylbenzene	9.857	105	140251	4.545	ug/l	99
74) N-amyl acetate	9.731	43	54436	4.422	ug/l	98
75) 1,1,2,2-Tetrachloroethane	10.165	83	71656	5.299	ug/l	99
76) 1,2,3-Trichloropropane	10.201	75	51079m	5.273	ug/l	
77) Bromobenzene	10.146	156	39428	4.924	ug/l	95
78) n-propylbenzene	10.281	91	168794	4.492	ug/l	99
79) 2-Chlorotoluene	10.349	91	110002	4.892	ug/l	99
80) 1,3,5-Trimethylbenzene	10.468	105	114749	4.474	ug/l	98
81) trans-1,4-Dichloro-2-b...	9.934	75	18154	4.499	ug/l	90
82) 4-Chlorotoluene	10.468	91	122429	4.695	ug/l	99
83) tert-Butylbenzene	10.789	119	117310	4.586	ug/l	99
84) 1,2,4-Trimethylbenzene	10.841	105	108157	4.343	ug/l	96
85) sec-Butylbenzene	11.014	105	153526	4.525	ug/l	99
86) p-Isopropyltoluene	11.169	119	127377	4.511	ug/l	98
87) 1,3-Dichlorobenzene	11.111	146	82673	5.102	ug/l	98
88) 1,4-Dichlorobenzene	11.198	146	91001	5.082	ug/l	94
89) n-Butylbenzene	11.583	91	119822	4.443	ug/l	96
90) Hexachloroethane	11.818	117	35762	5.143	ug/l	95
91) 1,2-Dichlorobenzene	11.570	146	76477	5.052	ug/l	98
92) 1,2-Dibromo-3-Chloropr...	12.358	75	14163	5.211	ug/l	92
93) 1,2,4-Trichlorobenzene	13.191	180	39775	4.583	ug/l	100
94) Hexachlorobutadiene	13.368	225	22549	5.001	ug/l	98
95) Naphthalene	13.429	128	113395	4.062	ug/l	99
96) 1,2,3-Trichlorobenzene	13.670	180	38315	4.317	ug/l	96

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039142.D
Acq On : 22 Sep 2025 15:37
Operator : SY/MD
Sample : VSTDICC005
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 10 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC005

Manual Integrations
APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
Supervised By :Semsettin Yesilyurt 09/26/2025

Quant Time: Sep 24 07:25:48 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 07:14:23 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC005

Manual Integrations APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
Supervised By :Semsettin Yesilyurt 09/26/2025



Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039143.D
 Acq On : 22 Sep 2025 16:35
 Operator : SY/MD
 Sample : VSTDICC001
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 11 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDICC001

Manual Integrations
 APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
 Supervised By :Semsettin Yesilyurt 09/26/2025

Quant Time: Sep 24 07:26:45 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 07:14:23 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.603	168	406498	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.535	114	735855	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	700418	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.175	152	322677	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	0.000	65	0d	0.000	ug/l	
Spiked Amount	50.000	Range	81 - 118	Recovery	=	0.000%#
35) Dibromofluoromethane	0.000	113	0d	0.000	ug/l	
Spiked Amount	50.000	Range	80 - 119	Recovery	=	0.000%#
50) Toluene-d8	0.000	98	0d	0.000	ug/l	
Spiked Amount	50.000	Range	89 - 112	Recovery	=	0.000%#
62) 4-Bromofluorobenzene	0.000	95	0d	0.000	ug/l	
Spiked Amount	50.000	Range	85 - 114	Recovery	=	0.000%#
Target Compounds					Qvalue	
2) Dichlorodifluoromethane	1.121	85	7593	1.180	ug/l	98
3) Chloromethane	1.230	50	7695	1.114	ug/l	99
4) Vinyl Chloride	1.297	62	8342	1.128	ug/l	98
6) Chloroethane	1.564	64	7440	0.673	ug/l	96
7) Trichlorofluoromethane	1.725	101	12570	1.174	ug/l	85
8) Diethyl Ether	1.921	74	4650	1.170	ug/l	92
9) 1,1,2-Trichlorotrifluo...	2.082	101	7630	1.198	ug/l	96
12) 1,1-Dichloroethene	2.085	96	7094	1.157	ug/l #	66
14) Allyl chloride	2.362	41	12383	1.131	ug/l	97
15) Acrylonitrile	2.683	53	23873	5.989	ug/l	96
16) Acetone	2.127	43	28927	3.224	ug/l	96
17) Carbon Disulfide	2.256	76	21708	1.158	ug/l	98
18) Methyl Acetate	2.391	43	9742	1.118	ug/l	98
19) Methyl tert-butyl Ether	2.715	73	21063	1.078	ug/l #	89
20) Methylene Chloride	2.458	84	16152	0.632	ug/l	90
21) trans-1,2-Dichloroethene	2.709	96	7797	1.195	ug/l	92
22) Diisopropyl ether	3.220	45	21620	1.032	ug/l #	34
23) Vinyl Acetate	3.204	43	91192	4.977	ug/l	99
24) 1,1-Dichloroethane	3.121	63	15790	1.233	ug/l #	94
25) 2-Butanone	3.921	43	22515m	4.862	ug/l	
26) 2,2-Dichloropropane	3.815	77	15492m	1.369	ug/l	
27) cis-1,2-Dichloroethene	3.834	96	8138	1.131	ug/l	91
28) Bromochloromethane	4.162	49	6039	1.068	ug/l #	53
29) Tetrahydrofuran	4.262	42	13621	4.619	ug/l #	65
30) Chloroform	4.288	83	15288	1.143	ug/l	96
32) 1,1,1-Trichloroethane	4.519	97	12926	1.116	ug/l	93
36) 1,1-Dichloropropene	4.764	75	9572m	1.063	ug/l	
37) Ethyl Acetate	4.030	43	12136m	1.098	ug/l	
38) Carbon Tetrachloride	4.735	117	13098m	1.207	ug/l	
39) Methylcyclohexane	6.050	83	9159	0.848	ug/l	91
40) Benzene	5.027	78	28487	0.978	ug/l	98
41) Methacrylonitrile	4.223	41	2717m	2.427	ug/l	
42) 1,2-Dichloroethane	5.059	62	10892	1.067	ug/l	95
43) Isopropyl Acetate	5.236	43	12786m	0.840	ug/l	
44) Trichloroethene	5.850	130	6408	0.953	ug/l	84
45) 1,2-Dichloropropane	6.104	63	6838	0.945	ug/l	93

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039143.D
 Acq On : 22 Sep 2025 16:35
 Operator : SY/MD
 Sample : VSTDICC001
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 11 Sample Multiplier: 1

Instrument :

MSVOA_V

ClientSampleId :

VSTDICC001

Manual Integrations

APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025

Supervised By :Semsettin Yesilyurt 09/26/2025

Quant Time: Sep 24 07:26:45 2025

Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M

Quant Title : SW846 8260

QLast Update : Wed Sep 24 07:14:23 2025

Response via : Initial Calibration

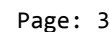
Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
46) Dibromomethane	6.249	93	5835	1.053	ug/l	89
47) Bromodichloromethane	6.439	83	12206	1.066	ug/l #	90
48) Methyl methacrylate	6.371	41	7232m	1.038	ug/l	
49) 1,4-Dioxane	6.333	88	2721m	24.855	ug/l	
51) 4-Methyl-2-Pentanone	7.165	43	33797	3.472	ug/l	96
52) Toluene	7.326	92	13265	0.786	ug/l	87
53) t-1,3-Dichloropropene	7.606	75	8060	0.807	ug/l	90
54) cis-1,3-Dichloropropene	6.973	75	8754	0.794	ug/l #	93
55) 1,1,2-Trichloroethane	7.776	97	8140	1.075	ug/l	91
56) Ethyl methacrylate	7.747	69	7455m	0.829	ug/l	
57) 1,3-Dichloropropane	7.956	76	11732	0.982	ug/l	94
58) 2-Chloroethyl Vinyl ether	6.841	63	10011	11.289	ug/l	98
59) 2-Hexanone	8.091	43	17015	4.307	ug/l	98
60) Dibromochloromethane	8.172	129	9122	1.057	ug/l	87
61) 1,2-Dibromoethane	8.288	107	7129	0.921	ug/l	90
64) Tetrachloroethene	7.899	164	5814	0.991	ug/l	90
65) Chlorobenzene	8.812	112	20007	1.025	ug/l #	88
66) 1,1,1,2-Tetrachloroethane	8.902	131	7603	1.079	ug/l #	61
67) Ethyl Benzene	8.947	91	24171	0.818	ug/l	94
68) m/p-Xylenes	9.072	106	16239	1.425	ug/l	98
69) o-Xylene	9.477	106	7250	0.718	ug/l	79
70) Styrene	9.503	104	11594	2.083	ug/l #	80
71) Bromoform	9.667	173	5558	1.009	ug/l #	94
73) Isopropylbenzene	9.860	105	20820	0.890	ug/l	98
74) N-amyl acetate	9.747	43	6654	0.713	ug/l #	91
75) 1,1,2,2-Tetrachloroethane	10.169	83	12746	1.243	ug/l	95
76) 1,2,3-Trichloropropane	10.207	75	8871	1.207	ug/l	100
77) Bromobenzene	10.159	156	6311	1.039	ug/l	93
78) n-propylbenzene	10.284	91	24996	0.877	ug/l	96
79) 2-Chlorotoluene	10.355	91	13640	0.800	ug/l	89
80) 1,3,5-Trimethylbenzene	10.471	105	15886	0.817	ug/l	96
82) 4-Chlorotoluene	10.474	91	15363	0.777	ug/l	91
83) tert-Butylbenzene	10.789	119	18552	0.956	ug/l	99
84) 1,2,4-Trimethylbenzene	10.847	105	15048	0.797	ug/l	95
85) sec-Butylbenzene	11.014	105	23268	0.904	ug/l	98
86) p-Isopropyltoluene	11.168	119	18549	0.866	ug/l	86
87) 1,3-Dichlorobenzene	11.117	146	12747	1.037	ug/l	97
88) 1,4-Dichlorobenzene	11.201	146	16279m	1.199	ug/l	
89) n-Butylbenzene	11.586	91	17079	0.835	ug/l	94
90) Hexachloroethane	11.818	117	6736	1.277	ug/l	94
91) 1,2-Dichlorobenzene	11.577	146	12486	1.088	ug/l	98
92) 1,2-Dibromo-3-Chloropr...	12.361	75	3071	0.690	ug/l	75
93) 1,2,4-Trichlorobenzene	13.197	180	6637	1.008	ug/l	99
94) Hexachlorobutadiene	13.368	225	4406	1.288	ug/l	95
95) Naphthalene	13.438	128	19706m	0.931	ug/l	
96) 1,2,3-Trichlorobenzene	13.676	180	7184	1.067	ug/l	97

(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VSTDICC001

Manual Integrations APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
Supervised By :Semsettin Yesilyurt 09/26/2025



Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039144.D
 Acq On : 22 Sep 2025 16:58
 Operator : SY/MD
 Sample : VSTDICV050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 12 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 ICVV092225

Quant Time: Sep 24 08:10:27 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Manual Integrations APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
 Supervised By :Semsettin Yesilyurt 09/26/2025

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.600	168	530687	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.535	114	821122	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	811351	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	500088	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.940	65	368982	48.041	ug/l	0.00
Spiked Amount 50.000	Range 81 - 118		Recovery =	96.080%		
35) Dibromofluoromethane	4.500	113	335549	50.832	ug/l	0.00
Spiked Amount 50.000	Range 80 - 119		Recovery =	101.660%		
50) Toluene-d8	7.236	98	1012009	52.199	ug/l	0.00
Spiked Amount 50.000	Range 89 - 112		Recovery =	104.400%		
62) 4-Bromofluorobenzene	10.001	95	453523	56.823	ug/l	0.00
Spiked Amount 50.000	Range 85 - 114		Recovery =	113.640%		
Target Compounds						
					Qvalue	
2) Dichlorodifluoromethane	1.121	85	380034	45.238	ug/l	98
3) Chloromethane	1.230	50	419560	46.523	ug/l	99
4) Vinyl Chloride	1.298	62	438971	45.467	ug/l	100
5) Bromomethane	1.494	94	274408	45.085	ug/l	100
6) Chloroethane	1.558	64	288768	50.873	ug/l	97
7) Trichlorofluoromethane	1.725	101	649327	46.452	ug/l	98
8) Diethyl Ether	1.918	74	239410	46.136	ug/l	100
9) 1,1,2-Trichlorotrifluo...	2.082	101	386460	46.465	ug/l	98
10) Methyl Iodide	2.198	142	394299	49.783	ug/l	100
11) Tert butyl alcohol	2.571	59	409358	252.173	ug/l	99
12) 1,1-Dichloroethene	2.082	96	367440	45.922	ug/l	99
13) Acrolein	2.011	56	76596	271.925	ug/l	95
14) Allyl chloride	2.359	41	673070	47.092	ug/l	100
15) Acrylonitrile	2.674	53	1247029	239.637	ug/l	100
16) Acetone	2.117	43	1131047	237.325	ug/l	99
17) Carbon Disulfide	2.252	76	1114937	45.563	ug/l	99
18) Methyl Acetate	2.381	43	564211	49.579	ug/l	99
19) Methyl tert-butyl Ether	2.712	73	1204021	47.223	ug/l	100
20) Methylene Chloride	2.458	84	426790	51.536	ug/l	99
21) trans-1,2-Dichloroethene	2.703	96	391103	45.915	ug/l	100
22) Diisopropyl ether	3.217	45	1355158	49.556	ug/l	95
23) Vinyl Acetate	3.188	43	5946893	248.597	ug/l	100
24) 1,1-Dichloroethane	3.117	63	761495	45.536	ug/l	100
25) 2-Butanone	3.867	43	1611713	262.050	ug/l	99
26) 2,2-Dichloropropane	3.812	77	648277	43.721	ug/l	99
27) cis-1,2-Dichloroethene	3.818	96	459958	48.960	ug/l	99
28) Bromochloromethane	4.146	49	384995	52.149	ug/l	100
29) Tetrahydrofuran	4.236	42	1064545	276.528	ug/l	99
30) Chloroform	4.278	83	820367	46.980	ug/l	99
31) Cyclohexane	4.584	56	654806	47.799	ug/l	99
32) 1,1,1-Trichloroethane	4.513	97	709351	46.922	ug/l	99
36) 1,1-Dichloropropene	4.744	75	537027	53.809	ug/l	99
37) Ethyl Acetate	3.976	43	630342	52.050	ug/l	99
38) Carbon Tetrachloride	4.735	117	603187	49.452	ug/l	99
39) Methylcyclohexane	6.047	83	692131	57.426	ug/l	98
40) Benzene	5.008	78	1699505	52.293	ug/l	98

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039144.D
 Acq On : 22 Sep 2025 16:58
 Operator : SY/MD
 Sample : VSTDICV050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 12 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 ICVV092225

Quant Time: Sep 24 08:10:27 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Manual Integrations APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
 Supervised By :Semsettin Yesilyurt 09/26/2025

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.159	41	282895	51.602	ug/l	99
42) 1,2-Dichloroethane	5.040	62	583550	51.245	ug/l	99
43) Isopropyl Acetate	5.194	43	924654	55.998	ug/l	99
44) Trichloroethene	5.831	130	384092	51.216	ug/l	95
45) 1,2-Dichloropropane	6.088	63	440211	54.495	ug/l	97
46) Dibromomethane	6.223	93	316032	51.116	ug/l	97
47) Bromodichloromethane	6.426	83	654836	51.232	ug/l	99
48) Methyl methacrylate	6.297	41	442395	59.104	ug/l	98
49) 1,4-Dioxane	6.291	88	136899	1133.346	ug/l	99
51) 4-Methyl-2-Pentanone	7.149	43	3292794	303.119	ug/l	100
52) Toluene	7.307	92	1064972	56.517	ug/l	99
53) t-1,3-Dichloropropene	7.574	75	621814	55.823	ug/l	99
54) cis-1,3-Dichloropropene	6.947	75	690532	56.154	ug/l	98
55) 1,1,2-Trichloroethane	7.760	97	432856	51.231	ug/l	99
56) Ethyl methacrylate	7.715	69	610159	60.810	ug/l	99
57) 1,3-Dichloropropane	7.934	76	716955	53.769	ug/l	100
58) 2-Chloroethyl Vinyl ether	6.812	63	1474938	259.936	ug/l	99
59) 2-Hexanone	8.063	43	2462493	280.545	ug/l	100
60) Dibromochloromethane	8.169	129	499908	51.926	ug/l	99
61) 1,2-Dibromoethane	8.272	107	460510	53.295	ug/l	100
64) Tetrachloroethene	7.895	164	346271	50.954	ug/l	99
65) Chlorobenzene	8.802	112	1153503	51.041	ug/l	99
66) 1,1,1,2-Tetrachloroethane	8.899	131	409401	50.136	ug/l	100
67) Ethyl Benzene	8.937	91	1964943	57.438	ug/l	100
68) m/p-Xylenes	9.062	106	1592118	120.582	ug/l	100
69) o-Xylene	9.468	106	706301	60.414	ug/l	98
70) Styrene	9.484	104	1306032	52.671	ug/l	99
71) Bromoform	9.654	173	335410	52.585	ug/l #	98
73) Isopropylbenzene	9.857	105	1966280	54.210	ug/l	100
74) N-amyl acetate	9.725	43	841099	58.135	ug/l	99
75) 1,1,2,2-Tetrachloroethane	10.165	83	724860	45.610	ug/l	100
76) 1,2,3-Trichloropropane	10.197	75	563785m	49.515	ug/l	
77) Bromobenzene	10.140	156	462931	49.193	ug/l	99
78) n-propylbenzene	10.278	91	2444658	55.347	ug/l	99
79) 2-Chlorotoluene	10.349	91	1412046	53.426	ug/l	99
80) 1,3,5-Trimethylbenzene	10.464	105	1709353	56.698	ug/l	99
81) trans-1,4-Dichloro-2-b...	9.927	75	247871	52.262	ug/l	99
82) 4-Chlorotoluene	10.461	91	1680201	54.824	ug/l	100
83) tert-Butylbenzene	10.789	119	1610870	53.584	ug/l	100
84) 1,2,4-Trimethylbenzene	10.837	105	1695994	57.944	ug/l	100
85) sec-Butylbenzene	11.011	105	2189540	54.911	ug/l	100
86) p-Isopropyltoluene	11.165	119	1841747	55.495	ug/l	100
87) 1,3-Dichlorobenzene	11.104	146	939483	49.328	ug/l	99
88) 1,4-Dichlorobenzene	11.194	146	971665	46.502	ug/l	99
89) n-Butylbenzene	11.580	91	1798412	56.740	ug/l	100
90) Hexachloroethane	11.818	117	370188	45.296	ug/l	98
91) 1,2-Dichlorobenzene	11.567	146	878347	49.362	ug/l	99
92) 1,2-Dibromo-3-Chloropr...	12.349	75	150399	52.479	ug/l	98
93) 1,2,4-Trichlorobenzene	13.185	180	533045	52.259	ug/l	99
94) Hexachlorobutadiene	13.368	225	243496	45.942	ug/l	99
95) Naphthalene	13.422	128	1919772	58.231	ug/l	100
96) 1,2,3-Trichlorobenzene	13.667	180	549079	52.633	ug/l	98

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039144.D
Acq On : 22 Sep 2025 16:58
Operator : SY/MD
Sample : VSTDICV050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 12 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
ICVVV092225

Manual Integrations
APPROVED

Reviewed By :Mahesh Dadoda 09/26/2025
Supervised By :Semsettin Yesilyurt 09/26/2025

Quant Time: Sep 24 08:10:27 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

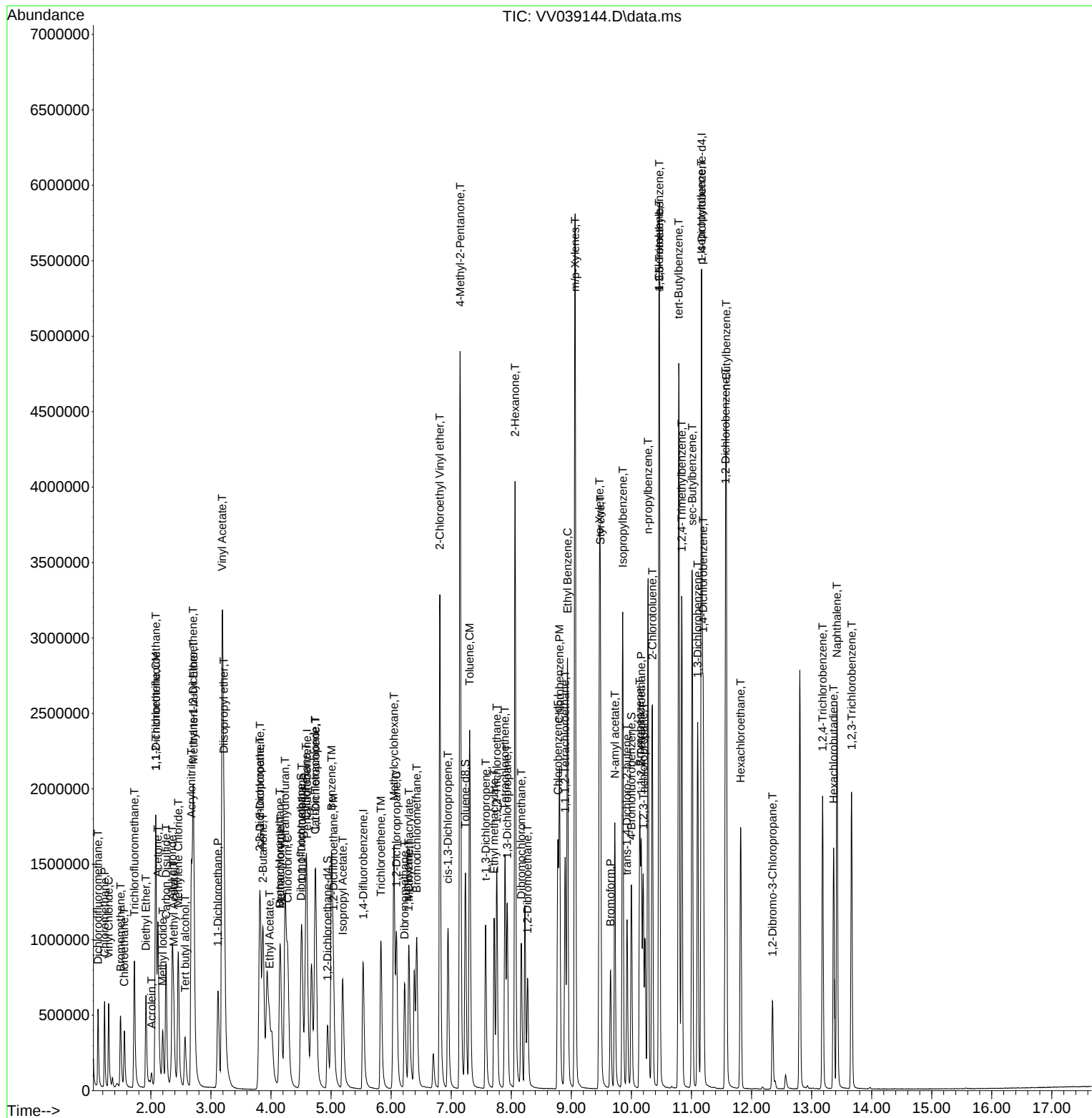
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Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VW092225\
Data File : VW039144.D
Acq On    : 22 Sep 2025  16:58
Operator  : SY/MD
Sample    : VSTDICV050
Misc      : 5.0mL/MSVOA_V/WATER
ALS Vial  : 12   Sample Multiplier: 1
```

Instrument :
MSVOA_V
ClientSampleId :
ICVV092225

Manual Integrations

Reviewed By :Mahesh Dadoda 09/26/2025
Supervised By :Semsettin Yesilyurt 09/26/2025

Quant Time: Sep 24 08:10:27 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration



Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039144.D
 Acq On : 22 Sep 2025 16:58
 Operator : SY/MD
 Sample : VSTDICV050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 12 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 ICVV092225

Quant Time: Sep 24 08:10:27 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 I	Pentafluorobenzene	1.000	1.000	0.0	94	0.00
2 T	Dichlorodifluoromethane	0.791	0.716	9.5	91	0.00
3 P	Chloromethane	0.850	0.791	6.9	97	0.00
4 C	Vinyl Chloride	0.910	0.827	9.1#	93	0.00
5 T	Bromomethane	0.573	0.517	9.8	96	0.00
6 T	Chloroethane	0.631	0.544	13.8	96	0.00
7 T	Trichlorofluoromethane	1.317	1.224	7.1	97	0.00
8 T	Diethyl Ether	0.489	0.451	7.8	95	0.00
9 T	1,1,2-Trichlorotrifluoroeth	0.784	0.728	7.1	98	0.00
10 T	Methyl Iodide	0.746	0.743	0.4	89	0.00
11 T	Tert butyl alcohol	0.153	0.154	-0.7	106	0.00
12 CM	1,1-Dichloroethene	0.754	0.692	8.2#	97	0.00
13 T	Acrolein	0.027	0.029	-7.4	103	0.00
14 T	Allyl chloride	1.347	1.268	5.9	96	0.00
15 T	Acrylonitrile	0.490	0.470	4.1	101	0.00
16 T	Acetone	0.523	0.426	18.5	89	0.00
17 T	Carbon Disulfide	2.306	2.101	8.9	95	0.00
18 T	Methyl Acetate	1.072	1.063	0.8	102	0.00
19 T	Methyl tert-butyl Ether	2.402	2.269	5.5	96	0.00
20 T	Methylene Chloride	1.066	0.804	24.6#	97	0.00
21 T	trans-1,2-Dichloroethene	0.803	0.737	8.2	96	0.00
22 T	Diisopropyl ether	2.576	2.554	0.9	95	0.00
23 T	Vinyl Acetate	2.254	2.241	0.6	96	0.00
24 P	1,1-Dichloroethane	1.576	1.435	8.9	97	0.00
25 T	2-Butanone	0.579	0.607	-4.8	98	0.00
26 T	2,2-Dichloropropane	1.397	1.222	12.5	93	0.00
27 T	cis-1,2-Dichloroethene	0.885	0.867	2.0	94	0.00
28 T	Bromochloromethane	0.696	0.725	-4.2	101	0.00
29 T	Tetrahydrofuran	0.363	0.401	-10.5	102	0.00
30 C	Chloroform	1.645	1.546	6.0#	98	0.00
31 T	Cyclohexane	1.291	1.234	4.4	93	0.00
32 T	1,1,1-Trichloroethane	1.424	1.337	6.1	97	0.00
33 S	1,2-Dichloroethane-d4	0.724	0.695	4.0	101	0.00
34 I	1,4-Difluorobenzene	1.000	1.000	0.0	92	0.00
35 S	Dibromofluoromethane	0.402	0.409	-1.7	99	0.00
36 T	1,1-Dichloropropene	0.608	0.654	-7.6	97	0.00
37 T	Ethyl Acetate	0.737	0.768	-4.2	99	0.00
38 T	Carbon Tetrachloride	0.743	0.735	1.1	96	0.00
39 T	Methylcyclohexane	0.734	0.843	-14.9	94	0.00
40 TM	Benzene	1.979	2.070	-4.6	96	0.00
41 T	Methacrylonitrile	0.273	0.345	-26.4#	96	0.00
42 TM	1,2-Dichloroethane	0.693	0.711	-2.6	98	0.00
43 T	Isopropyl Acetate	1.005	1.126	-12.0	97	0.00
44 TM	Trichloroethene	0.457	0.468	-2.4	93	0.00
45 C	1,2-Dichloropropane	0.492	0.536	-8.9#	98	0.00
46 T	Dibromomethane	0.376	0.385	-2.4	96	0.00
47 T	Bromodichloromethane	0.778	0.797	-2.4	97	0.00
48 T	Methyl methacrylate	0.456	0.539	-18.2	98	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039144.D
 Acq On : 22 Sep 2025 16:58
 Operator : SY/MD
 Sample : VSTDICV050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 12 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 ICVV092225

Quant Time: Sep 24 08:10:27 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
49 T	1,4-Dioxane	0.007	0.008	-14.3	102	0.00
50 S	Toluene-d8	1.181	1.232	-4.3	100	0.00
51 T	4-Methyl-2-Pentanone	0.661	0.802	-21.3#	101	0.00
52 CM	Toluene	1.147	1.297	-13.1#	97	0.00
53 T	t-1,3-Dichloropropene	0.678	0.757	-11.7	96	0.00
54 T	cis-1,3-Dichloropropene	0.749	0.841	-12.3	95	0.00
55 T	1,1,2-Trichloroethane	0.514	0.527	-2.5	98	0.00
56 T	Ethyl methacrylate	0.611	0.743	-21.6#	95	0.00
57 T	1,3-Dichloropropane	0.812	0.873	-7.5	98	0.00
58 T	2-Chloroethyl Vinyl ether	0.267	0.359	-34.5#	93	0.00
59 T	2-Hexanone	0.463	0.600	-29.6#	101	0.00
60 T	Dibromochloromethane	0.586	0.609	-3.9	99	0.00
61 T	1,2-Dibromoethane	0.526	0.561	-6.7	97	0.00
62 S	4-Bromofluorobenzene	0.486	0.552	-13.6	100	0.00
63 I	Chlorobenzene-d5	1.000	1.000	0.0	94	0.00
64 T	Tetrachloroethene	0.419	0.427	-1.9	97	0.00
65 PM	Chlorobenzene	1.393	1.422	-2.1	97	0.00
66 T	1,1,1,2-Tetrachloroethane	0.503	0.505	-0.4	97	0.00
67 C	Ethyl Benzene	2.108	2.422	-14.9#	95	0.00
68 T	m/p-Xylenes	0.814	0.981	-20.5#	97	0.00
69 T	o-Xylene	0.720	0.871	-21.0#	97	0.00
70 T	Styrene	1.263	1.610	-27.5#	97	0.00
71 P	Bromoform	0.393	0.413	-5.1	98	0.00
72 I	1,4-Dichlorobenzene-d4	1.000	1.000	0.0	97	0.00
73 T	Isopropylbenzene	3.627	3.932	-8.4	97	0.00
74 T	N-amyl acetate	1.447	1.682	-16.2	97	0.00
75 P	1,1,2,2-Tetrachloroethane	1.589	1.449	8.8	99	0.00
76 T	1,2,3-Trichloropropane	1.138	1.127	1.0	104	0.00
77 T	Bromobenzene	0.941	0.926	1.6	97	0.00
78 T	n-propylbenzene	4.416	4.888	-10.7	97	0.00
79 T	2-Chlorotoluene	2.643	2.824	-6.8	97	0.00
80 T	1,3,5-Trimethylbenzene	3.014	3.418	-13.4	98	0.00
81 T	trans-1,4-Dichloro-2-butene	0.474	0.496	-4.6	97	0.00
82 T	4-Chlorotoluene	3.064	3.360	-9.7	98	0.00
83 T	tert-Butylbenzene	3.006	3.221	-7.2	97	0.00
84 T	1,2,4-Trimethylbenzene	2.926	3.391	-15.9	97	0.00
85 T	sec-Butylbenzene	3.987	4.378	-9.8	97	0.00
86 T	p-Isopropyltoluene	3.318	3.683	-11.0	97	0.00
87 T	1,3-Dichlorobenzene	1.904	1.879	1.3	98	0.00
88 T	1,4-Dichlorobenzene	2.089	1.943	7.0	97	0.00
89 T	n-Butylbenzene	3.169	3.596	-13.5	96	0.00
90 T	Hexachloroethane	0.817	0.740	9.4	99	0.00
91 T	1,2-Dichlorobenzene	1.779	1.756	1.3	98	0.00
92 T	1,2-Dibromo-3-Chloropropane	0.330	0.301	8.8	102	0.00
93 T	1,2,4-Trichlorobenzene	1.020	1.066	-4.5	96	0.00
94 T	Hexachlorobutadiene	0.530	0.487	8.1	98	0.00
95 T	Naphthalene	3.296	3.839	-16.5	98	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039144.D
Acq On : 22 Sep 2025 16:58
Operator : SY/MD
Sample : VSTDICV050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 12 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
ICVVV092225

Quant Time: Sep 24 08:10:27 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
96 T	1,2,3-Trichlorobenzene	1.043	1.098	-5.3	96	0.00

(#) = Out of Range

SPCC's out = 0 CCC's out = 6

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039144.D
 Acq On : 22 Sep 2025 16:58
 Operator : SY/MD
 Sample : VSTDICV050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 12 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 ICSVV092225

Quant Time: Sep 24 08:10:27 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Pentafluorobenzene	50.000	50.000	0.0	94	0.00
2 T	Dichlorodifluoromethane	50.000	45.238	9.5	91	0.00
3 P	Chloromethane	50.000	46.523	7.0	97	0.00
4 C	Vinyl Chloride	50.000	45.467	9.1#	93	0.00
5 T	Bromomethane	50.000	45.085	9.8	96	0.00
6 T	Chloroethane	50.000	50.873	-1.7	96	0.00
7 T	Trichlorofluoromethane	50.000	46.452	7.1	97	0.00
8 T	Diethyl Ether	50.000	46.136	7.7	95	0.00
9 T	1,1,2-Trichlorotrifluoroeth	50.000	46.465	7.1	98	0.00
10 T	Methyl Iodide	50.000	49.783	0.4	89	0.00
11 T	Tert butyl alcohol	250.000	252.173	-0.9	106	0.00
12 CM	1,1-Dichloroethene	50.000	45.922	8.2#	97	0.00
13 T	Acrolein	250.000	271.925	-8.8	103	0.00
14 T	Allyl chloride	50.000	47.092	5.8	96	0.00
15 T	Acrylonitrile	250.000	239.637	4.1	101	0.00
16 T	Acetone	250.000	237.325	5.1	89	0.00
17 T	Carbon Disulfide	50.000	45.563	8.9	95	0.00
18 T	Methyl Acetate	50.000	49.579	0.8	102	0.00
19 T	Methyl tert-butyl Ether	50.000	47.223	5.6	96	0.00
20 T	Methylene Chloride	50.000	51.536	-3.1	97	0.00
21 T	trans-1,2-Dichloroethene	50.000	45.915	8.2	96	0.00
22 T	Diisopropyl ether	50.000	49.556	0.9	95	0.00
23 T	Vinyl Acetate	250.000	248.597	0.6	96	0.00
24 P	1,1-Dichloroethane	50.000	45.536	8.9	97	0.00
25 T	2-Butanone	250.000	262.050	-4.8	98	0.00
26 T	2,2-Dichloropropane	50.000	43.721	12.6	93	0.00
27 T	cis-1,2-Dichloroethene	50.000	48.960	2.1	94	0.00
28 T	Bromochloromethane	50.000	52.149	-4.3	101	0.00
29 T	Tetrahydrofuran	250.000	276.528	-10.6	102	0.00
30 C	Chloroform	50.000	46.980	6.0#	98	0.00
31 T	Cyclohexane	50.000	47.799	4.4	93	0.00
32 T	1,1,1-Trichloroethane	50.000	46.922	6.2	97	0.00
33 S	1,2-Dichloroethane-d4	50.000	48.041	3.9	101	0.00
34 I	1,4-Difluorobenzene	50.000	50.000	0.0	92	0.00
35 S	Dibromofluoromethane	50.000	50.832	-1.7	99	0.00
36 T	1,1-Dichloropropene	50.000	53.809	-7.6	97	0.00
37 T	Ethyl Acetate	50.000	52.050	-4.1	99	0.00
38 T	Carbon Tetrachloride	50.000	49.452	1.1	96	0.00
39 T	Methylcyclohexane	50.000	57.426	-14.9	94	0.00
40 TM	Benzene	50.000	52.293	-4.6	96	0.00
41 T	Methacrylonitrile	50.000	51.602	-3.2	96	0.00
42 TM	1,2-Dichloroethane	50.000	51.245	-2.5	98	0.00
43 T	Isopropyl Acetate	50.000	55.998	-12.0	97	0.00
44 TM	Trichloroethene	50.000	51.216	-2.4	93	0.00
45 C	1,2-Dichloropropane	50.000	54.495	-9.0#	98	0.00
46 T	Dibromomethane	50.000	51.116	-2.2	96	0.00
47 T	Bromodichloromethane	50.000	51.232	-2.5	97	0.00
48 T	Methyl methacrylate	50.000	59.104	-18.2	98	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
 Data File : VV039144.D
 Acq On : 22 Sep 2025 16:58
 Operator : SY/MD
 Sample : VSTDICV050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 12 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 ICSVV092225

Quant Time: Sep 24 08:10:27 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
49 T	1,4-Dioxane	1000.000	1133.346	-13.3	102	0.00
50 S	Toluene-d8	50.000	52.199	-4.4	100	0.00
51 T	4-Methyl-2-Pentanone	250.000	303.119	-21.2#	101	0.00
52 CM	Toluene	50.000	56.517	-13.0#	97	0.00
53 T	t-1,3-Dichloropropene	50.000	55.823	-11.6	96	0.00
54 T	cis-1,3-Dichloropropene	50.000	56.154	-12.3	95	0.00
55 T	1,1,2-Trichloroethane	50.000	51.231	-2.5	98	0.00
56 T	Ethyl methacrylate	50.000	60.810	-21.6#	95	0.00
57 T	1,3-Dichloropropane	50.000	53.769	-7.5	98	0.00
58 T	2-Chloroethyl Vinyl ether	250.000	259.936	-4.0	93	0.00
59 T	2-Hexanone	250.000	280.545	-12.2	101	0.00
60 T	Dibromochloromethane	50.000	51.926	-3.9	99	0.00
61 T	1,2-Dibromoethane	50.000	53.295	-6.6	97	0.00
62 S	4-Bromofluorobenzene	50.000	56.823	-13.6	100	0.00
63 I	Chlorobenzene-d5	50.000	50.000	0.0	94	0.00
64 T	Tetrachloroethene	50.000	50.954	-1.9	97	0.00
65 PM	Chlorobenzene	50.000	51.041	-2.1	97	0.00
66 T	1,1,1,2-Tetrachloroethane	50.000	50.136	-0.3	97	0.00
67 C	Ethyl Benzene	50.000	57.438	-14.9#	95	0.00
68 T	m/p-Xylenes	100.000	120.582	-20.6#	97	0.00
69 T	o-Xylene	50.000	60.414	-20.8#	97	0.00
70 T	Styrene	50.000	52.671	-5.3	97	0.00
71 P	Bromoform	50.000	52.585	-5.2	98	0.00
72 I	1,4-Dichlorobenzene-d4	50.000	50.000	0.0	97	0.00
73 T	Isopropylbenzene	50.000	54.210	-8.4	97	0.00
74 T	N-amyl acetate	50.000	58.135	-16.3	97	0.00
75 P	1,1,2,2-Tetrachloroethane	50.000	45.610	8.8	99	0.00
76 T	1,2,3-Trichloropropane	50.000	49.515	1.0	104	0.00
77 T	Bromobenzene	50.000	49.193	1.6	97	0.00
78 T	n-propylbenzene	50.000	55.347	-10.7	97	0.00
79 T	2-Chlorotoluene	50.000	53.426	-6.9	97	0.00
80 T	1,3,5-Trimethylbenzene	50.000	56.698	-13.4	98	0.00
81 T	trans-1,4-Dichloro-2-butene	50.000	52.262	-4.5	97	0.00
82 T	4-Chlorotoluene	50.000	54.824	-9.6	98	0.00
83 T	tert-Butylbenzene	50.000	53.584	-7.2	97	0.00
84 T	1,2,4-Trimethylbenzene	50.000	57.944	-15.9	97	0.00
85 T	sec-Butylbenzene	50.000	54.911	-9.8	97	0.00
86 T	p-Isopropyltoluene	50.000	55.495	-11.0	97	0.00
87 T	1,3-Dichlorobenzene	50.000	49.328	1.3	98	0.00
88 T	1,4-Dichlorobenzene	50.000	46.502	7.0	97	0.00
89 T	n-Butylbenzene	50.000	56.740	-13.5	96	0.00
90 T	Hexachloroethane	50.000	45.296	9.4	99	0.00
91 T	1,2-Dichlorobenzene	50.000	49.362	1.3	98	0.00
92 T	1,2-Dibromo-3-Chloropropane	50.000	52.479	-5.0	102	0.00
93 T	1,2,4-Trichlorobenzene	50.000	52.259	-4.5	96	0.00
94 T	Hexachlorobutadiene	50.000	45.942	8.1	98	0.00
95 T	Naphthalene	50.000	58.231	-16.5	98	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039144.D
Acq On : 22 Sep 2025 16:58
Operator : SY/MD
Sample : VSTDICV050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 12 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
ICVVV092225

Quant Time: Sep 24 08:10:27 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
96 T	1,2,3-Trichlorobenzene	50.000	52.633	-5.3	96	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 6



284 Sheffield Street, Mountainside, New Jersey 07092, Phone : 908 789 8900,
Fax : 908 789 8922

VOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	<u>Alliance</u>	Contract:	<u>ATGG01</u>
Lab Code:	<u>ACE</u>	SDG No.:	<u>Q3164</u>
Instrument ID:	<u>MSVOA_V</u>	Calibration Date/Time:	<u>09/24/2025</u> <u>10:49</u>
Lab File ID:	<u>VV039146.D</u>	Init. Calib. Date(s):	<u>09/22/2025</u> <u>09/22/2025</u>
Heated Purge: (Y/N)	<u>N</u>	Init. Calib. Time(s):	<u>12:26</u> <u>16:35</u>
GC Column:	<u>DB-624UI</u>	ID:	<u>0.18</u> (mm)

COMPOUND	RRF	RRF050	MIN RRF	%D	MAX%D
Benzene	1.979	1.817		-8.19	20
Toluene	1.147	1.120		-2.35	20
Ethyl Benzene	2.108	1.997		-5.27	20
m/p-Xylenes	0.814	0.814		0	20
o-Xylene	0.720	0.732		1.67	20
1,2-Dichloroethane-d4	0.724	0.661		-8.7	20
Dibromofluoromethane	0.402	0.366		-8.95	20
Toluene-d8	1.181	1.053		-10.84	20
4-Bromofluorobenzene	0.486	0.489		0.62	20

All other compounds must meet a minimum RRF of 0.010.
RRF of 1,4-Dioxane = Value should be divide by 1000.

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039146.D
 Acq On : 24 Sep 2025 10:49
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.600	168	558154	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	907128	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.777	117	923608	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	551620	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.941	65	369186	45.702	ug/l	0.00
Spiked Amount 50.000	Range 81 - 118		Recovery =	91.400%		
35) Dibromofluoromethane	4.500	113	332304	45.567	ug/l	0.00
Spiked Amount 50.000	Range 80 - 119		Recovery =	91.140%		
50) Toluene-d8	7.236	98	954978	44.587	ug/l	0.00
Spiked Amount 50.000	Range 89 - 112		Recovery =	89.180%		
62) 4-Bromofluorobenzene	10.002	95	443437	50.291	ug/l	0.00
Spiked Amount 50.000	Range 85 - 114		Recovery =	100.580%		
Target Compounds					Qvalue	
2) Dichlorodifluoromethane	1.121	85	349754	39.585	ug/l	96
3) Chloromethane	1.227	50	407305	42.941	ug/l	100
4) Vinyl Chloride	1.295	62	414391	40.808	ug/l	100
5) Bromomethane	1.494	94	281451	43.967	ug/l	100
6) Chloroethane	1.558	64	272410	45.230	ug/l	97
7) Trichlorofluoromethane	1.722	101	592362	40.292	ug/l	99
8) Diethyl Ether	1.918	74	252704	46.301	ug/l	99
9) 1,1,2-Trichlorotrifluo...	2.079	101	350274	40.042	ug/l	98
10) Methyl Iodide	2.195	142	408452	49.032	ug/l	100
11) Tert butyl alcohol	2.571	59	439522	257.431	ug/l	98
12) 1,1-Dichloroethene	2.079	96	343170	40.778	ug/l	97
13) Acrolein	2.008	56	82132	277.230	ug/l	99
14) Allyl chloride	2.356	41	678394	45.129	ug/l	97
15) Acrylonitrile	2.674	53	1317414	240.705	ug/l	100
16) Acetone	2.118	43	1249190	250.295	ug/l	99
17) Carbon Disulfide	2.249	76	1032299	40.110	ug/l	99
18) Methyl Acetate	2.378	43	608422	50.833	ug/l	100
19) Methyl tert-butyl Ether	2.712	73	1280541	47.752	ug/l	100
20) Methylene Chloride	2.455	84	433679	49.660	ug/l	100
21) trans-1,2-Dichloroethene	2.700	96	381303	42.561	ug/l	100
22) Diisopropyl ether	3.214	45	1373474	47.754	ug/l	92
23) Vinyl Acetate	3.188	43	6174734	245.419	ug/l	99
24) 1,1-Dichloroethane	3.114	63	738164	41.969	ug/l	100
25) 2-Butanone	3.864	43	1696999	262.339	ug/l	98
26) 2,2-Dichloropropane	3.809	77	608343	39.009	ug/l	100
27) cis-1,2-Dichloroethene	3.815	96	451121	45.657	ug/l	99
28) Bromochloromethane	4.143	49	381004	49.069	ug/l	97
29) Tetrahydrofuran	4.233	42	1116689	275.798	ug/l	100
30) Chloroform	4.275	83	808825	44.039	ug/l	100
31) Cyclohexane	4.581	56	584423	40.562	ug/l	99
32) 1,1,1-Trichloroethane	4.513	97	652596	41.043	ug/l	99
36) 1,1-Dichloropropene	4.741	75	487252	44.192	ug/l	98
37) Ethyl Acetate	3.973	43	663808	49.616	ug/l	100
38) Carbon Tetrachloride	4.732	117	552633	41.012	ug/l	99
39) Methylcyclohexane	6.047	83	605143	45.449	ug/l	99
40) Benzene	5.005	78	1647897	45.898	ug/l	100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039146.D
 Acq On : 24 Sep 2025 10:49
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.156	41	299616	49.549	ug/l	99
42) 1,2-Dichloroethane	5.037	62	594558	47.262	ug/l	100
43) Isopropyl Acetate	5.191	43	967617	53.044	ug/l	100
44) Trichloroethene	5.828	130	367209	44.323	ug/l	96
45) 1,2-Dichloropropane	6.085	63	431931	48.401	ug/l	98
46) Dibromomethane	6.224	93	328651	48.117	ug/l	99
47) Bromodichloromethane	6.426	83	659260	46.688	ug/l	99
48) Methyl methacrylate	6.294	41	457609	55.340	ug/l	99
49) 1,4-Dioxane	6.291	88	155709	1166.851	ug/l	97
51) 4-Methyl-2-Pentanone	7.146	43	3404796	283.713	ug/l	100
52) Toluene	7.307	92	1016283	48.820	ug/l	99
53) t-1,3-Dichloropropene	7.571	75	638267	51.867	ug/l	98
54) cis-1,3-Dichloropropene	6.947	75	697782	51.364	ug/l	100
55) 1,1,2-Trichloroethane	7.760	97	453142	48.547	ug/l	98
56) Ethyl methacrylate	7.715	69	630396	56.871	ug/l	100
57) 1,3-Dichloropropane	7.931	76	727051	49.356	ug/l	100
58) 2-Chloroethyl Vinyl ether	6.809	63	1515308	242.405	ug/l	100
59) 2-Hexanone	8.063	43	2528312	260.898	ug/l	99
60) Dibromochloromethane	8.166	129	506709	47.642	ug/l	99
61) 1,2-Dibromoethane	8.272	107	480605	50.348	ug/l	100
64) Tetrachloroethene	7.896	164	320555	41.437	ug/l	99
65) Chlorobenzene	8.802	112	1127747	43.836	ug/l	98
66) 1,1,1,2-Tetrachloroethane	8.895	131	407431	43.830	ug/l	100
67) Ethyl Benzene	8.934	91	1844871	47.374	ug/l	100
68) m/p-Xylenes	9.063	106	1502908	99.991	ug/l	100
69) o-Xylene	9.468	106	676401	50.825	ug/l	99
70) Styrene	9.484	104	1276348	45.393	ug/l	99
71) Bromoform	9.651	173	348806	48.038	ug/l #	100
73) Isopropylbenzene	9.857	105	1824633	45.605	ug/l	100
74) N-amyl acetate	9.725	43	854806	53.563	ug/l	98
75) 1,1,2,2-Tetrachloroethane	10.166	83	747109	42.619	ug/l	99
76) 1,2,3-Trichloropropane	10.198	75	575337	45.809	ug/l	98
77) Bromobenzene	10.140	156	460599	44.372	ug/l	99
78) n-propylbenzene	10.278	91	2250074	46.182	ug/l	99
79) 2-Chlorotoluene	10.349	91	1358304	46.591	ug/l	99
80) 1,3,5-Trimethylbenzene	10.465	105	1607013	48.324	ug/l	99
81) trans-1,4-Dichloro-2-b...	9.928	75	256388	49.007	ug/l	99
82) 4-Chlorotoluene	10.461	91	1605527	47.493	ug/l	99
83) tert-Butylbenzene	10.789	119	1497786	45.168	ug/l	99
84) 1,2,4-Trimethylbenzene	10.838	105	1608644	49.826	ug/l	100
85) sec-Butylbenzene	11.011	105	1967674	44.737	ug/l	99
86) p-Isopropyltoluene	11.165	119	1677532	45.825	ug/l	100
87) 1,3-Dichlorobenzene	11.104	146	909637	43.299	ug/l	99
88) 1,4-Dichlorobenzene	11.194	146	954565	41.416	ug/l	100
89) n-Butylbenzene	11.580	91	1645246	47.058	ug/l	100
90) Hexachloroethane	11.821	117	343097	38.060	ug/l	99
91) 1,2-Dichlorobenzene	11.567	146	862856	43.962	ug/l	99
92) 1,2-Dibromo-3-Chloropr...	12.349	75	158828	50.345	ug/l	97
93) 1,2,4-Trichlorobenzene	13.185	180	522821	46.468	ug/l	100
94) Hexachlorobutadiene	13.368	225	224298	38.367	ug/l	100
95) Naphthalene	13.426	128	1960806	53.919	ug/l	99
96) 1,2,3-Trichlorobenzene	13.667	180	551841	47.956	ug/l	99

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039146.D
Acq On : 24 Sep 2025 10:49
Operator : SY/MD
Sample : VSTDCCC050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 2 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

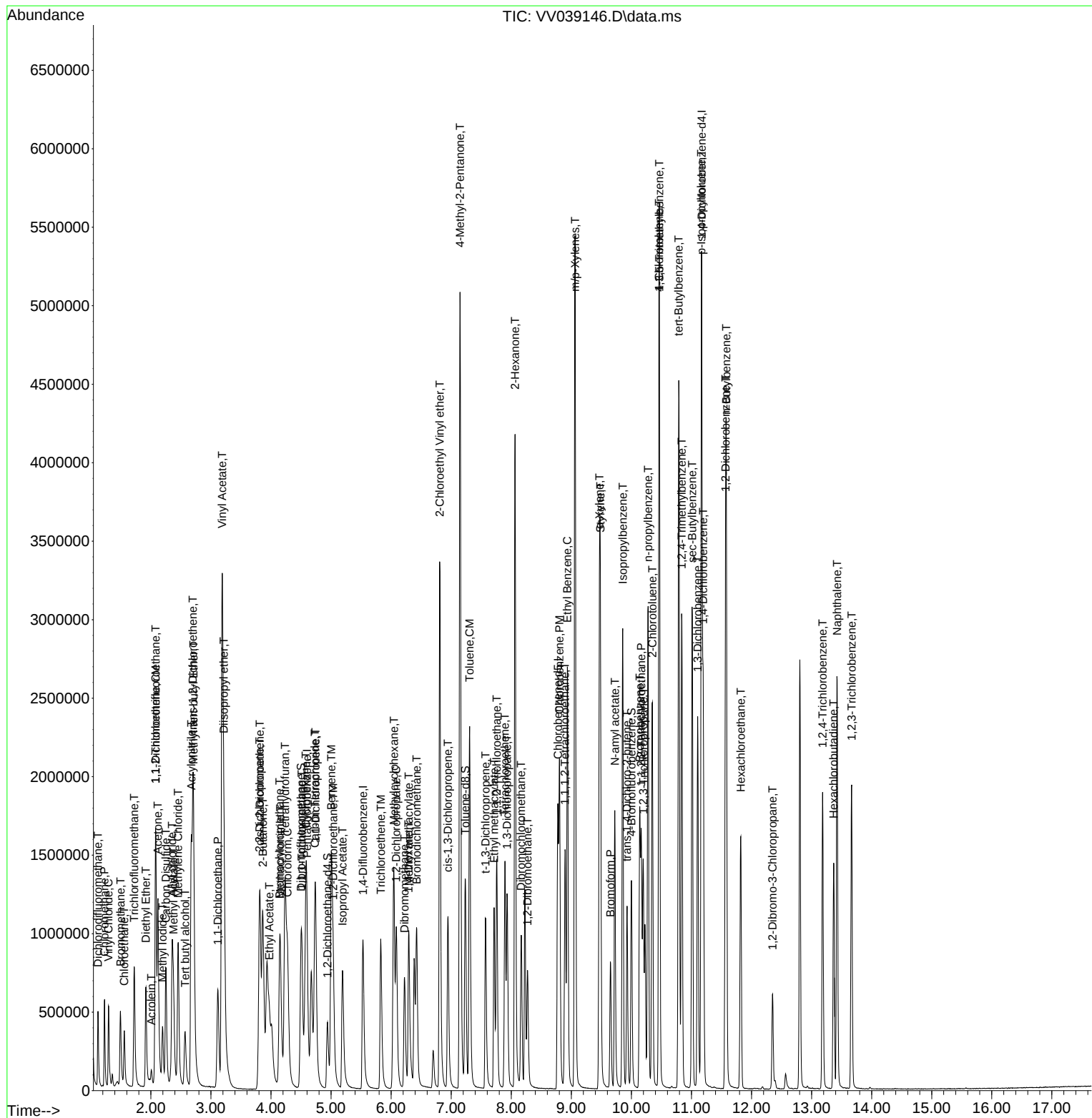
Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

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Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039146.D
Acq On    : 24 Sep 2025  10:49
Operator  : SY/MD
Sample    : VSTDCCC050
Misc      : 5.0mL/MSVOA_V/WATER
ALS Vial  : 2      Sample Multiplier: 1
```

Instrument :
MSVOA_V
ClientSampleId :
VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration



Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039146.D
 Acq On : 24 Sep 2025 10:49
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
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Instrument :
 MSVOA_V
 LabSampleId :
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Quant Time: Sep 25 04:42:18 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 I	Pentafluorobenzene	1.000	1.000	0.0	99	0.00
2 T	Dichlorodifluoromethane	0.791	0.627	20.7#	84	0.00
3 P	Chloromethane	0.850	0.730	14.1	94	0.00
4 C	Vinyl Chloride	0.910	0.742	18.5#	88	0.00
5 T	Bromomethane	0.573	0.504	12.0	98	0.00
6 T	Chloroethane	0.631	0.488	22.7#	91	0.00
7 T	Trichlorofluoromethane	1.317	1.061	19.4	88	0.00
8 T	Diethyl Ether	0.489	0.453	7.4	100	0.00
9 T	1,1,2-Trichlorotrifluoroeth	0.784	0.628	19.9	89	0.00
10 T	Methyl Iodide	0.746	0.732	1.9	92	0.00
11 T	Tert butyl alcohol	0.153	0.157	-2.6	114	0.00
12 CM	1,1-Dichloroethene	0.754	0.615	18.4#	90	0.00
13 T	Acrolein	0.027	0.029	-7.4	110	0.00
14 T	Allyl chloride	1.347	1.215	9.8	97	0.00
15 T	Acrylonitrile	0.490	0.472	3.7	107	0.00
16 T	Acetone	0.523	0.448	14.3	99	0.00
17 T	Carbon Disulfide	2.306	1.849	19.8	88	0.00
18 T	Methyl Acetate	1.072	1.090	-1.7	110	0.00
19 T	Methyl tert-butyl Ether	2.402	2.294	4.5	102	0.00
20 T	Methylene Chloride	1.066	0.777	27.1#	99	0.00
21 T	trans-1,2-Dichloroethene	0.803	0.683	14.9	93	0.00
22 T	Diisopropyl ether	2.576	2.461	4.5	97	0.00
23 T	Vinyl Acetate	2.254	2.213	1.8	99	0.00
24 P	1,1-Dichloroethane	1.576	1.323	16.1	94	0.00
25 T	2-Butanone	0.579	0.608	-5.0	103	0.00
26 T	2,2-Dichloropropane	1.397	1.090	22.0#	87	0.00
27 T	cis-1,2-Dichloroethene	0.885	0.808	8.7	92	0.00
28 T	Bromochloromethane	0.696	0.683	1.9	100	0.00
29 T	Tetrahydrofuran	0.363	0.400	-10.2	107	0.00
30 C	Chloroform	1.645	1.449	11.9#	96	0.00
31 T	Cyclohexane	1.291	1.047	18.9	83	0.00
32 T	1,1,1-Trichloroethane	1.424	1.169	17.9	89	0.00
33 S	1,2-Dichloroethane-d4	0.724	0.661	8.7	101	0.00
34 I	1,4-Difluorobenzene	1.000	1.000	0.0	101	0.00
35 S	Dibromofluoromethane	0.402	0.366	9.0	98	0.00
36 T	1,1-Dichloropropene	0.608	0.537	11.7	88	0.00
37 T	Ethyl Acetate	0.737	0.732	0.7	104	0.00
38 T	Carbon Tetrachloride	0.743	0.609	18.0	88	0.00
39 T	Methylcyclohexane	0.734	0.667	9.1	82	0.00
40 TM	Benzene	1.979	1.817	8.2	93	0.00
41 T	Methacrylonitrile	0.273	0.330	-20.9#	101	0.00
42 TM	1,2-Dichloroethane	0.693	0.655	5.5	100	0.00
43 T	Isopropyl Acetate	1.005	1.067	-6.2	101	0.00
44 TM	Trichloroethene	0.457	0.405	11.4	89	0.00
45 C	1,2-Dichloropropane	0.492	0.476	3.3#	96	0.00
46 T	Dibromomethane	0.376	0.362	3.7	100	0.00
47 T	Bromodichloromethane	0.778	0.727	6.6	98	0.00
48 T	Methyl methacrylate	0.456	0.504	-10.5	102	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039146.D
 Acq On : 24 Sep 2025 10:49
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 LabSampleId :
 VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
49 T	1,4-Dioxane	0.007	0.009	-28.6#	116	0.00
50 S	Toluene-d8	1.181	1.053	10.8	94	0.00
51 T	4-Methyl-2-Pentanone	0.661	0.751	-13.6	104	0.00
52 CM	Toluene	1.147	1.120	2.4#	92	0.00
53 T	t-1,3-Dichloropropene	0.678	0.704	-3.8	98	0.00
54 T	cis-1,3-Dichloropropene	0.749	0.769	-2.7	96	0.00
55 T	1,1,2-Trichloroethane	0.514	0.500	2.7	103	0.00
56 T	Ethyl methacrylate	0.611	0.695	-13.7	98	0.00
57 T	1,3-Dichloropropane	0.812	0.801	1.4	99	0.00
58 T	2-Chloroethyl Vinyl ether	0.267	0.334	-25.1#	96	0.00
59 T	2-Hexanone	0.463	0.557	-20.3#	104	0.00
60 T	Dibromochloromethane	0.586	0.559	4.6	100	0.00
61 T	1,2-Dibromoethane	0.526	0.530	-0.8	102	0.00
62 S	4-Bromofluorobenzene	0.486	0.489	-0.6	98	0.00
63 I	Chlorobenzene-d5	1.000	1.000	0.0	107	0.00
64 T	Tetrachloroethene	0.419	0.347	17.2	90	0.00
65 PM	Chlorobenzene	1.393	1.221	12.3	94	0.00
66 T	1,1,1,2-Tetrachloroethane	0.503	0.441	12.3	97	0.00
67 C	Ethyl Benzene	2.108	1.997	5.3#	90	0.00
68 T	m/p-Xylenes	0.814	0.814	0.0	92	0.00
69 T	o-Xylene	0.720	0.732	-1.7	93	0.00
70 T	Styrene	1.263	1.382	-9.4	95	0.00
71 P	Bromoform	0.393	0.378	3.8	102	0.00
72 I	1,4-Dichlorobenzene-d4	1.000	1.000	0.0	107	0.00
73 T	Isopropylbenzene	3.627	3.308	8.8	90	0.00
74 T	N-amyl acetate	1.447	1.550	-7.1	99	0.00
75 P	1,1,2,2-Tetrachloroethane	1.589	1.354	14.8	102	0.00
76 T	1,2,3-Trichloropropane	1.138	1.043	8.3	106	0.00
77 T	Bromobenzene	0.941	0.835	11.3	96	0.00
78 T	n-propylbenzene	4.416	4.079	7.6	89	0.00
79 T	2-Chlorotoluene	2.643	2.462	6.8	93	0.00
80 T	1,3,5-Trimethylbenzene	3.014	2.913	3.4	92	0.00
81 T	trans-1,4-Dichloro-2-butene	0.474	0.465	1.9	100	0.00
82 T	4-Chlorotoluene	3.064	2.911	5.0	93	0.00
83 T	tert-Butylbenzene	3.006	2.715	9.7	90	0.00
84 T	1,2,4-Trimethylbenzene	2.926	2.916	0.3	92	0.00
85 T	sec-Butylbenzene	3.987	3.567	10.5	87	0.00
86 T	p-Isopropyltoluene	3.318	3.041	8.3	88	0.00
87 T	1,3-Dichlorobenzene	1.904	1.649	13.4	95	0.00
88 T	1,4-Dichlorobenzene	2.089	1.730	17.2	96	0.00
89 T	n-Butylbenzene	3.169	2.983	5.9	88	0.00
90 T	Hexachloroethane	0.817	0.622	23.9#	91	0.00
91 T	1,2-Dichlorobenzene	1.779	1.564	12.1	96	0.00
92 T	1,2-Dibromo-3-Chloropropane	0.330	0.288	12.7	108	0.00
93 T	1,2,4-Trichlorobenzene	1.020	0.948	7.1	94	0.00
94 T	Hexachlorobutadiene	0.530	0.407	23.2#	91	0.00
95 T	Naphthalene	3.296	3.555	-7.9	100	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039146.D
Acq On : 24 Sep 2025 10:49
Operator : SY/MD
Sample : VSTDCCC050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 2 Sample Multiplier: 1

Instrument :
MSVOA_V
LabSampleId :
VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
96 T	1,2,3-Trichlorobenzene	1.043	1.000	4.1	97	0.00

(#) = Out of Range

SPCC's out = 0 CCC's out = 6

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039146.D
 Acq On : 24 Sep 2025 10:49
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 LabSampleId :
 VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Pentafluorobenzene	50.000	50.000	0.0	99	0.00
2 T	Dichlorodifluoromethane	50.000	39.585	20.8#	84	0.00
3 P	Chloromethane	50.000	42.941	14.1	94	0.00
4 C	Vinyl Chloride	50.000	40.808	18.4#	88	0.00
5 T	Bromomethane	50.000	43.967	12.1	98	0.00
6 T	Chloroethane	50.000	45.230	9.5	91	0.00
7 T	Trichlorofluoromethane	50.000	40.292	19.4	88	0.00
8 T	Diethyl Ether	50.000	46.301	7.4	100	0.00
9 T	1,1,2-Trichlorotrifluoroeth	50.000	40.042	19.9	89	0.00
10 T	Methyl Iodide	50.000	49.032	1.9	92	0.00
11 T	Tert butyl alcohol	250.000	257.431	-3.0	114	0.00
12 CM	1,1-Dichloroethene	50.000	40.778	18.4#	90	0.00
13 T	Acrolein	250.000	277.230	-10.9	110	0.00
14 T	Allyl chloride	50.000	45.129	9.7	97	0.00
15 T	Acrylonitrile	250.000	240.705	3.7	107	0.00
16 T	Acetone	250.000	250.295	-0.1	99	0.00
17 T	Carbon Disulfide	50.000	40.110	19.8	88	0.00
18 T	Methyl Acetate	50.000	50.833	-1.7	110	0.00
19 T	Methyl tert-butyl Ether	50.000	47.752	4.5	102	0.00
20 T	Methylene Chloride	50.000	49.660	0.7	99	0.00
21 T	trans-1,2-Dichloroethene	50.000	42.561	14.9	93	0.00
22 T	Diisopropyl ether	50.000	47.754	4.5	97	0.00
23 T	Vinyl Acetate	250.000	245.419	1.8	99	0.00
24 P	1,1-Dichloroethane	50.000	41.969	16.1	94	0.00
25 T	2-Butanone	250.000	262.339	-4.9	103	0.00
26 T	2,2-Dichloropropane	50.000	39.009	22.0#	87	0.00
27 T	cis-1,2-Dichloroethene	50.000	45.657	8.7	92	0.00
28 T	Bromochloromethane	50.000	49.069	1.9	100	0.00
29 T	Tetrahydrofuran	250.000	275.798	-10.3	107	0.00
30 C	Chloroform	50.000	44.039	11.9#	96	0.00
31 T	Cyclohexane	50.000	40.562	18.9	83	0.00
32 T	1,1,1-Trichloroethane	50.000	41.043	17.9	89	0.00
33 S	1,2-Dichloroethane-d4	50.000	45.702	8.6	101	0.00
34 I	1,4-Difluorobenzene	50.000	50.000	0.0	101	0.00
35 S	Dibromofluoromethane	50.000	45.567	8.9	98	0.00
36 T	1,1-Dichloropropene	50.000	44.192	11.6	88	0.00
37 T	Ethyl Acetate	50.000	49.616	0.8	104	0.00
38 T	Carbon Tetrachloride	50.000	41.012	18.0	88	0.00
39 T	Methylcyclohexane	50.000	45.449	9.1	82	0.00
40 TM	Benzene	50.000	45.898	8.2	93	0.00
41 T	Methacrylonitrile	50.000	49.549	0.9	101	0.00
42 TM	1,2-Dichloroethane	50.000	47.262	5.5	100	0.00
43 T	Isopropyl Acetate	50.000	53.044	-6.1	101	0.00
44 TM	Trichloroethene	50.000	44.323	11.4	89	0.00
45 C	1,2-Dichloropropane	50.000	48.401	3.2#	96	0.00
46 T	Dibromomethane	50.000	48.117	3.8	100	0.00
47 T	Bromodichloromethane	50.000	46.688	6.6	98	0.00
48 T	Methyl methacrylate	50.000	55.340	-10.7	102	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039146.D
 Acq On : 24 Sep 2025 10:49
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 LabSampleId :
 VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
49 T	1,4-Dioxane	1000.000	1166.851	-16.7	116	0.00
50 S	Toluene-d8	50.000	44.587	10.8	94	0.00
51 T	4-Methyl-2-Pentanone	250.000	283.713	-13.5	104	0.00
52 CM	Toluene	50.000	48.820	2.4#	92	0.00
53 T	t-1,3-Dichloropropene	50.000	51.867	-3.7	98	0.00
54 T	cis-1,3-Dichloropropene	50.000	51.364	-2.7	96	0.00
55 T	1,1,2-Trichloroethane	50.000	48.547	2.9	103	0.00
56 T	Ethyl methacrylate	50.000	56.871	-13.7	98	0.00
57 T	1,3-Dichloropropane	50.000	49.356	1.3	99	0.00
58 T	2-Chloroethyl Vinyl ether	250.000	242.405	3.0	96	0.00
59 T	2-Hexanone	250.000	260.898	-4.4	104	0.00
60 T	Dibromochloromethane	50.000	47.642	4.7	100	0.00
61 T	1,2-Dibromoethane	50.000	50.348	-0.7	102	0.00
62 S	4-Bromofluorobenzene	50.000	50.291	-0.6	98	0.00
63 I	Chlorobenzene-d5	50.000	50.000	0.0	107	0.00
64 T	Tetrachloroethene	50.000	41.437	17.1	90	0.00
65 PM	Chlorobenzene	50.000	43.836	12.3	94	0.00
66 T	1,1,1,2-Tetrachloroethane	50.000	43.830	12.3	97	0.00
67 C	Ethyl Benzene	50.000	47.374	5.3#	90	0.00
68 T	m/p-Xylenes	100.000	99.991	0.0	92	0.00
69 T	o-Xylene	50.000	50.825	-1.7	93	0.00
70 T	Styrene	50.000	45.393	9.2	95	0.00
71 P	Bromoform	50.000	48.038	3.9	102	0.00
72 I	1,4-Dichlorobenzene-d4	50.000	50.000	0.0	107	0.00
73 T	Isopropylbenzene	50.000	45.605	8.8	90	0.00
74 T	N-amyl acetate	50.000	53.563	-7.1	99	0.00
75 P	1,1,2,2-Tetrachloroethane	50.000	42.619	14.8	102	0.00
76 T	1,2,3-Trichloropropane	50.000	45.809	8.4	106	0.00
77 T	Bromobenzene	50.000	44.372	11.3	96	0.00
78 T	n-propylbenzene	50.000	46.182	7.6	89	0.00
79 T	2-Chlorotoluene	50.000	46.591	6.8	93	0.00
80 T	1,3,5-Trimethylbenzene	50.000	48.324	3.4	92	0.00
81 T	trans-1,4-Dichloro-2-butene	50.000	49.007	2.0	100	0.00
82 T	4-Chlorotoluene	50.000	47.493	5.0	93	0.00
83 T	tert-Butylbenzene	50.000	45.168	9.7	90	0.00
84 T	1,2,4-Trimethylbenzene	50.000	49.826	0.3	92	0.00
85 T	sec-Butylbenzene	50.000	44.737	10.5	87	0.00
86 T	p-Isopropyltoluene	50.000	45.825	8.3	88	0.00
87 T	1,3-Dichlorobenzene	50.000	43.299	13.4	95	0.00
88 T	1,4-Dichlorobenzene	50.000	41.416	17.2	96	0.00
89 T	n-Butylbenzene	50.000	47.058	5.9	88	0.00
90 T	Hexachloroethane	50.000	38.060	23.9#	91	0.00
91 T	1,2-Dichlorobenzene	50.000	43.962	12.1	96	0.00
92 T	1,2-Dibromo-3-Chloropropane	50.000	50.345	-0.7	108	0.00
93 T	1,2,4-Trichlorobenzene	50.000	46.468	7.1	94	0.00
94 T	Hexachlorobutadiene	50.000	38.367	23.3#	91	0.00
95 T	Naphthalene	50.000	53.919	-7.8	100	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039146.D
Acq On : 24 Sep 2025 10:49
Operator : SY/MD
Sample : VSTDCCC050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 2 Sample Multiplier: 1

Instrument :
MSVOA_V
LabSampleId :
VSTDCCC050

Quant Time: Sep 25 04:42:18 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
96 T	1,2,3-Trichlorobenzene	50.000	47.956	4.1	97	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 6



284 Sheffield Street, Mountainside, New Jersey 07092, Phone : 908 789 8900,
Fax : 908 789 8922

VOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	<u>Alliance</u>	Contract:	<u>ATGG01</u>
Lab Code:	<u>ACE</u>	SDG No.:	<u>Q3164</u>
Instrument ID:	<u>MSVOA_V</u>	Calibration Date/Time:	<u>09/25/2025</u> <u>09:59</u>
Lab File ID:	<u>VV039168.D</u>	Init. Calib. Date(s):	<u>09/22/2025</u> <u>09/22/2025</u>
Heated Purge: (Y/N)	<u>N</u>	Init. Calib. Time(s):	<u>12:26</u> <u>16:35</u>
GC Column:	<u>DB-624UI</u>	ID:	<u>0.18</u> (mm)

COMPOUND	RRF	RRF050	MIN RRF	%D	MAX%D
Benzene	1.979	1.926		-2.68	20
Toluene	1.147	1.204		4.97	20
Ethyl Benzene	2.108	2.320		10.06	20
m/p-Xylenes	0.814	0.936		14.99	20
o-Xylene	0.720	0.841		16.81	20
1,2-Dichloroethane-d4	0.724	0.609		-15.88	20
Dibromofluoromethane	0.402	0.384		-4.48	20
Toluene-d8	1.181	1.121		-5.08	20
4-Bromofluorobenzene	0.486	0.508		4.53	20

All other compounds must meet a minimum RRF of 0.010.
RRF of 1,4-Dioxane = Value should be divide by 1000.

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039168.D
 Acq On : 25 Sep 2025 09:59
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.596	168	605297	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.529	114	933338	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.773	117	901987	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	535710	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.937	65	368829	42.102	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	84.200%
35) Dibromofluoromethane	4.497	113	358851	47.826	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	95.660%
50) Toluene-d8	7.233	98	1046447	47.485	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	94.980%
62) 4-Bromofluorobenzene	9.998	95	473731	52.218	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	104.440%
Target Compounds						
					Qvalue	
2) Dichlorodifluoromethane	1.121	85	412859	43.088	ug/l	96
3) Chloromethane	1.227	50	441526	42.924	ug/l	99
4) Vinyl Chloride	1.294	62	495985	45.040	ug/l	99
5) Bromomethane	1.494	94	277073	39.912	ug/l	99
6) Chloroethane	1.558	64	307095	47.163	ug/l	96
7) Trichlorofluoromethane	1.722	101	725756	45.520	ug/l	99
8) Diethyl Ether	1.915	74	273800	46.259	ug/l	95
9) 1,1,2-Trichlorotrifluo...	2.079	101	429502	45.275	ug/l	98
10) Methyl Iodide	2.195	142	394240	43.640	ug/l	99
11) Tert butyl alcohol	2.571	59	462139	249.596	ug/l	98
12) 1,1-Dichloroethene	2.079	96	422387	46.283	ug/l	93
13) Acrolein	2.008	56	73826	229.786	ug/l	96
14) Allyl chloride	2.355	41	725818	44.523	ug/l	97
15) Acrylonitrile	2.670	53	1328524	223.829	ug/l	99
16) Acetone	2.117	43	1227421	224.844	ug/l	98
17) Carbon Disulfide	2.249	76	1213128	43.464	ug/l	99
18) Methyl Acetate	2.378	43	602375	46.408	ug/l	97
19) Methyl tert-butyl Ether	2.709	73	1360066	46.768	ug/l	99
20) Methylene Chloride	2.455	84	477153	50.438	ug/l	94
21) trans-1,2-Dichloroethene	2.699	96	438480	45.131	ug/l	98
22) Diisopropyl ether	3.214	45	1401924	44.947	ug/l	97
23) Vinyl Acetate	3.185	43	6223966	228.109	ug/l	98
24) 1,1-Dichloroethane	3.114	63	827604	43.389	ug/l	100
25) 2-Butanone	3.863	43	1647490	234.849	ug/l	96
26) 2,2-Dichloropropane	3.805	77	690029	40.800	ug/l	100
27) cis-1,2-Dichloroethene	3.815	96	499109	46.579	ug/l	98
28) Bromochloromethane	4.143	49	370658	44.019	ug/l	91
29) Tetrahydrofuran	4.233	42	1075144	244.856	ug/l	97
30) Chloroform	4.275	83	862902	43.324	ug/l	98
31) Cyclohexane	4.580	56	694256	44.432	ug/l	97
32) 1,1,1-Trichloroethane	4.510	97	759916	44.071	ug/l	98
36) 1,1-Dichloropropene	4.741	75	562517	49.586	ug/l	99
37) Ethyl Acetate	3.973	43	650966	47.290	ug/l	98
38) Carbon Tetrachloride	4.731	117	642615	46.350	ug/l	99
39) Methylcyclohexane	6.043	83	743575	54.277	ug/l	99
40) Benzene	5.005	78	1797939	48.671	ug/l	99

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039168.D
 Acq On : 25 Sep 2025 09:59
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.156	41	291884	47.017	ug/l	98
42) 1,2-Dichloroethane	5.037	62	596048	46.050	ug/l	98
43) Isopropyl Acetate	5.191	43	954350	50.848	ug/l	98
44) Trichloroethene	5.828	130	432014	50.680	ug/l	99
45) 1,2-Dichloropropane	6.085	63	459902	50.088	ug/l	99
46) Dibromomethane	6.220	93	337738	48.059	ug/l	97
47) Bromodichloromethane	6.423	83	680557	46.843	ug/l	98
48) Methyl methacrylate	6.294	41	449431	52.825	ug/l	98
49) 1,4-Dioxane	6.291	88	155938	1135.751	ug/l	97
51) 4-Methyl-2-Pentanone	7.146	43	3274291	265.177	ug/l	98
52) Toluene	7.307	92	1123906	52.474	ug/l	99
53) t-1,3-Dichloropropene	7.571	75	661472	52.243	ug/l	99
54) cis-1,3-Dichloropropene	6.944	75	731190	52.312	ug/l	97
55) 1,1,2-Trichloroethane	7.757	97	465236	48.443	ug/l	98
56) Ethyl methacrylate	7.712	69	658147	57.707	ug/l	95
57) 1,3-Dichloropropane	7.931	76	748312	49.373	ug/l	100
58) 2-Chloroethyl Vinyl ether	6.808	63	1657102	257.039	ug/l	99
59) 2-Hexanone	8.062	43	2465950	247.438	ug/l	97
60) Dibromochloromethane	8.165	129	535532	48.938	ug/l	99
61) 1,2-Dibromoethane	8.271	107	498142	50.719	ug/l	100
64) Tetrachloroethene	7.895	164	376666	49.857	ug/l	97
65) Chlorobenzene	8.802	112	1242390	49.450	ug/l	98
66) 1,1,1,2-Tetrachloroethane	8.895	131	435516	47.975	ug/l	98
67) Ethyl Benzene	8.934	91	2092639	55.024	ug/l	99
68) m/p-Xylenes	9.059	106	1688294	115.018	ug/l	100
69) o-Xylene	9.468	106	758328	58.347	ug/l	98
70) Styrene	9.484	104	1379790	50.114	ug/l	99
71) Bromoform	9.651	173	362481	51.118	ug/l #	99
73) Isopropylbenzene	9.853	105	2113988	54.407	ug/l	99
74) N-amyl acetate	9.725	43	853417	55.064	ug/l	98
75) 1,1,2,2-Tetrachloroethane	10.165	83	747856	43.928	ug/l	99
76) 1,2,3-Trichloropropane	10.197	75	573502	47.019	ug/l	98
77) Bromobenzene	10.140	156	497564	49.357	ug/l	98
78) n-propylbenzene	10.278	91	2554851	53.995	ug/l	99
79) 2-Chlorotoluene	10.345	91	1498849	52.939	ug/l	100
80) 1,3,5-Trimethylbenzene	10.464	105	1809221	56.020	ug/l	100
81) trans-1,4-Dichloro-2-b...	9.927	75	258743	50.926	ug/l	97
82) 4-Chlorotoluene	10.461	91	1769699	53.904	ug/l	100
83) tert-Butylbenzene	10.789	119	1725454	53.579	ug/l	100
84) 1,2,4-Trimethylbenzene	10.837	105	1777572	56.693	ug/l	98
85) sec-Butylbenzene	11.011	105	2294341	53.713	ug/l	99
86) p-Isopropyltoluene	11.165	119	1933306	54.380	ug/l	99
87) 1,3-Dichlorobenzene	11.104	146	995077	48.773	ug/l	100
88) 1,4-Dichlorobenzene	11.194	146	1019216	45.535	ug/l	99
89) n-Butylbenzene	11.580	91	1854102	54.607	ug/l	100
90) Hexachloroethane	11.821	117	387482	44.260	ug/l	99
91) 1,2-Dichlorobenzene	11.567	146	930166	48.799	ug/l	99
92) 1,2-Dibromo-3-Chloropr...	12.348	75	156226	50.961	ug/l	97
93) 1,2,4-Trichlorobenzene	13.184	180	568781	52.055	ug/l	99
94) Hexachlorobutadiene	13.368	225	261028	45.975	ug/l	99
95) Naphthalene	13.422	128	2039508	57.749	ug/l	100
96) 1,2,3-Trichlorobenzene	13.667	180	579422	51.848	ug/l	97

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
Data File : VV039168.D
Acq On : 25 Sep 2025 09:59
Operator : SY/MD
Sample : VSTDCCC050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 2 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

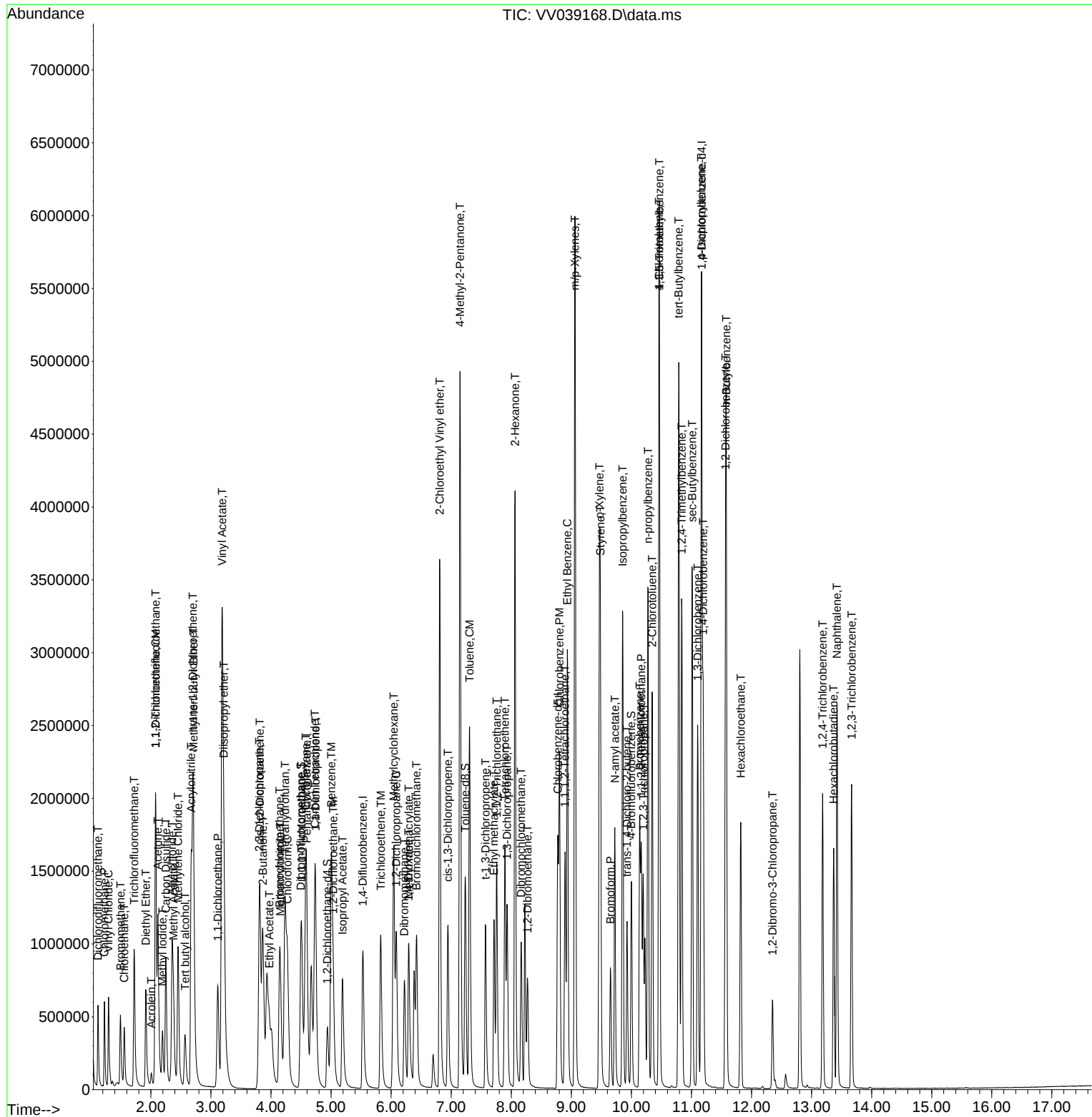
Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
----------	------	------	----------	------	-------	----------

(#) = qualifier out of range (m) = manual integration (+) = signals summed

```
Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VW092525\  
Data File : VW039168.D  
Acq On    : 25 Sep 2025   09:59  
Operator  : SY/MD  
Sample    : VSTDCCC050  
Misc      : 5.0mL/MSVOA_V/WATER  
ALS Vial  : 2      Sample Multiplier: 1
```

Instrument :
MSVOA_V
ClientSampleId :
VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration



Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039168.D
 Acq On : 25 Sep 2025 09:59
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 LabSampleId :
 VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
1 I	Pentafluorobenzene	1.000	1.000	0.0	107	0.00
2 T	Dichlorodifluoromethane	0.791	0.682	13.8	99	0.00
3 P	Chloromethane	0.850	0.729	14.2	102	0.00
4 C	Vinyl Chloride	0.910	0.819	10.0#	105	0.00
5 T	Bromomethane	0.573	0.458	20.1#	97	0.00
6 T	Chloroethane	0.631	0.507	19.7	102	0.00
7 T	Trichlorofluoromethane	1.317	1.199	9.0	108	0.00
8 T	Diethyl Ether	0.489	0.452	7.6	109	0.00
9 T	1,1,2-Trichlorotrifluoroeth	0.784	0.710	9.4	109	0.00
10 T	Methyl Iodide	0.746	0.651	12.7	89	0.00
11 T	Tert butyl alcohol	0.153	0.153	0.0	120	0.00
12 CM	1,1-Dichloroethene	0.754	0.698	7.4#	111	0.00
13 T	Acrolein	0.027	0.024	11.1	99	0.00
14 T	Allyl chloride	1.347	1.199	11.0	104	0.00
15 T	Acrylonitrile	0.490	0.439	10.4	108	0.00
16 T	Acetone	0.523	0.406	22.4#	97	0.00
17 T	Carbon Disulfide	2.306	2.004	13.1	103	0.00
18 T	Methyl Acetate	1.072	0.995	7.2	109	0.00
19 T	Methyl tert-butyl Ether	2.402	2.247	6.5	108	0.00
20 T	Methylene Chloride	1.066	0.788	26.1#	109	0.00
21 T	trans-1,2-Dichloroethene	0.803	0.724	9.8	107	0.00
22 T	Diisopropyl ether	2.576	2.316	10.1	99	0.00
23 T	Vinyl Acetate	2.254	2.056	8.8	100	0.00
24 P	1,1-Dichloroethane	1.576	1.367	13.3	106	0.00
25 T	2-Butanone	0.579	0.544	6.0	100	0.00
26 T	2,2-Dichloropropane	1.397	1.140	18.4	99	0.00
27 T	cis-1,2-Dichloroethene	0.885	0.825	6.8	102	0.00
28 T	Bromochloromethane	0.696	0.612	12.1	97	0.00
29 T	Tetrahydrofuran	0.363	0.355	2.2	103	0.00
30 C	Chloroform	1.645	1.426	13.3#	103	0.00
31 T	Cyclohexane	1.291	1.147	11.2	99	0.00
32 T	1,1,1-Trichloroethane	1.424	1.255	11.9	104	0.00
33 S	1,2-Dichloroethane-d4	0.724	0.609	15.9	101	0.00
34 I	1,4-Difluorobenzene	1.000	1.000	0.0	104	0.00
35 S	Dibromofluoromethane	0.402	0.384	4.5	106	0.00
36 T	1,1-Dichloropropene	0.608	0.603	0.8	101	0.00
37 T	Ethyl Acetate	0.737	0.697	5.4	102	0.00
38 T	Carbon Tetrachloride	0.743	0.689	7.3	102	0.00
39 T	Methylcyclohexane	0.734	0.797	-8.6	101	0.00
40 TM	Benzene	1.979	1.926	2.7	101	0.00
41 T	Methacrylonitrile	0.273	0.313	-14.7	99	0.00
42 TM	1,2-Dichloroethane	0.693	0.639	7.8	100	0.00
43 T	Isopropyl Acetate	1.005	1.023	-1.8	100	0.00
44 TM	Trichloroethene	0.457	0.463	-1.3	104	0.00
45 C	1,2-Dichloropropane	0.492	0.493	-0.2#	102	0.00
46 T	Dibromomethane	0.376	0.362	3.7	103	0.00
47 T	Bromodichloromethane	0.778	0.729	6.3	101	0.00
48 T	Methyl methacrylate	0.456	0.482	-5.7	100	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039168.D
 Acq On : 25 Sep 2025 09:59
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 LabSampleId :
 VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
49 T	1,4-Dioxane	0.007	0.008	-14.3	117	0.00
50 S	Toluene-d8	1.181	1.121	5.1	103	0.00
51 T	4-Methyl-2-Pentanone	0.661	0.702	-6.2	100	0.00
52 CM	Toluene	1.147	1.204	-5.0#	102	0.00
53 T	t-1,3-Dichloropropene	0.678	0.709	-4.6	102	0.00
54 T	cis-1,3-Dichloropropene	0.749	0.783	-4.5	101	0.00
55 T	1,1,2-Trichloroethane	0.514	0.498	3.1	105	0.00
56 T	Ethyl methacrylate	0.611	0.705	-15.4	103	0.00
57 T	1,3-Dichloropropane	0.812	0.802	1.2	102	0.00
58 T	2-Chloroethyl Vinyl ether	0.267	0.355	-33.0#	105	0.00
59 T	2-Hexanone	0.463	0.528	-14.0	101	0.00
60 T	Dibromochloromethane	0.586	0.574	2.0	106	0.00
61 T	1,2-Dibromoethane	0.526	0.534	-1.5	105	0.00
62 S	4-Bromofluorobenzene	0.486	0.508	-4.5	105	0.00
63 I	Chlorobenzene-d5	1.000	1.000	0.0	104	0.00
64 T	Tetrachloroethene	0.419	0.418	0.2	106	0.00
65 PM	Chlorobenzene	1.393	1.377	1.1	104	0.00
66 T	1,1,1,2-Tetrachloroethane	0.503	0.483	4.0	103	0.00
67 C	Ethyl Benzene	2.108	2.320	-10.1#	102	0.00
68 T	m/p-Xylenes	0.814	0.936	-15.0	103	0.00
69 T	o-Xylene	0.720	0.841	-16.8	104	0.00
70 T	Styrene	1.263	1.530	-21.1#	103	0.00
71 P	Bromoform	0.393	0.402	-2.3	106	0.00
72 I	1,4-Dichlorobenzene-d4	1.000	1.000	0.0	104	0.00
73 T	Isopropylbenzene	3.627	3.946	-8.8	104	0.00
74 T	N-amyl acetate	1.447	1.593	-10.1	99	0.00
75 P	1,1,2,2-Tetrachloroethane	1.589	1.396	12.1	102	0.00
76 T	1,2,3-Trichloropropane	1.138	1.071	5.9	106	0.00
77 T	Bromobenzene	0.941	0.929	1.3	104	0.00
78 T	n-propylbenzene	4.416	4.769	-8.0	101	0.00
79 T	2-Chlorotoluene	2.643	2.798	-5.9	102	0.00
80 T	1,3,5-Trimethylbenzene	3.014	3.377	-12.0	103	0.00
81 T	trans-1,4-Dichloro-2-butene	0.474	0.483	-1.9	101	0.00
82 T	4-Chlorotoluene	3.064	3.303	-7.8	103	0.00
83 T	tert-Butylbenzene	3.006	3.221	-7.2	104	0.00
84 T	1,2,4-Trimethylbenzene	2.926	3.318	-13.4	102	0.00
85 T	sec-Butylbenzene	3.987	4.283	-7.4	102	0.00
86 T	p-Isopropyltoluene	3.318	3.609	-8.8	102	0.00
87 T	1,3-Dichlorobenzene	1.904	1.857	2.5	104	0.00
88 T	1,4-Dichlorobenzene	2.089	1.903	8.9	102	0.00
89 T	n-Butylbenzene	3.169	3.461	-9.2	99	0.00
90 T	Hexachloroethane	0.817	0.723	11.5	103	0.00
91 T	1,2-Dichlorobenzene	1.779	1.736	2.4	104	0.00
92 T	1,2-Dibromo-3-Chloropropane	0.330	0.292	11.5	106	0.00
93 T	1,2,4-Trichlorobenzene	1.020	1.062	-4.1	102	0.00
94 T	Hexachlorobutadiene	0.530	0.487	8.1	105	0.00
95 T	Naphthalene	3.296	3.807	-15.5	104	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
Data File : VV039168.D
Acq On : 25 Sep 2025 09:59
Operator : SY/MD
Sample : VSTDCCC050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 2 Sample Multiplier: 1

Instrument :
MSVOA_V
LabSampleId :
VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	AvgRF	CCRF	%Dev	Area%	Dev(min)
96 T	1,2,3-Trichlorobenzene	1.043	1.082	-3.7	101	0.00

(#) = Out of Range

SPCC's out = 0 CCC's out = 6

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039168.D
 Acq On : 25 Sep 2025 09:59
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 LabSampleId :
 VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
1 I	Pentafluorobenzene	50.000	50.000	0.0	107	0.00
2 T	Dichlorodifluoromethane	50.000	43.088	13.8	99	0.00
3 P	Chloromethane	50.000	42.924	14.2	102	0.00
4 C	Vinyl Chloride	50.000	45.040	9.9#	105	0.00
5 T	Bromomethane	50.000	39.912	20.2#	97	0.00
6 T	Chloroethane	50.000	47.163	5.7	102	0.00
7 T	Trichlorofluoromethane	50.000	45.520	9.0	108	0.00
8 T	Diethyl Ether	50.000	46.259	7.5	109	0.00
9 T	1,1,2-Trichlorotrifluoroeth	50.000	45.275	9.5	109	0.00
10 T	Methyl Iodide	50.000	43.640	12.7	89	0.00
11 T	Tert butyl alcohol	250.000	249.596	0.2	120	0.00
12 CM	1,1-Dichloroethene	50.000	46.283	7.4#	111	0.00
13 T	Acrolein	250.000	229.786	8.1	99	0.00
14 T	Allyl chloride	50.000	44.523	11.0	104	0.00
15 T	Acrylonitrile	250.000	223.829	10.5	108	0.00
16 T	Acetone	250.000	224.844	10.1	97	0.00
17 T	Carbon Disulfide	50.000	43.464	13.1	103	0.00
18 T	Methyl Acetate	50.000	46.408	7.2	109	0.00
19 T	Methyl tert-butyl Ether	50.000	46.768	6.5	108	0.00
20 T	Methylene Chloride	50.000	50.438	-0.9	109	0.00
21 T	trans-1,2-Dichloroethene	50.000	45.131	9.7	107	0.00
22 T	Diisopropyl ether	50.000	44.947	10.1	99	0.00
23 T	Vinyl Acetate	250.000	228.109	8.8	100	0.00
24 P	1,1-Dichloroethane	50.000	43.389	13.2	106	0.00
25 T	2-Butanone	250.000	234.849	6.1	100	0.00
26 T	2,2-Dichloropropane	50.000	40.800	18.4	99	0.00
27 T	cis-1,2-Dichloroethene	50.000	46.579	6.8	102	0.00
28 T	Bromochloromethane	50.000	44.019	12.0	97	0.00
29 T	Tetrahydrofuran	250.000	244.856	2.1	103	0.00
30 C	Chloroform	50.000	43.324	13.4#	103	0.00
31 T	Cyclohexane	50.000	44.432	11.1	99	0.00
32 T	1,1,1-Trichloroethane	50.000	44.071	11.9	104	0.00
33 S	1,2-Dichloroethane-d4	50.000	42.102	15.8	101	0.00
34 I	1,4-Difluorobenzene	50.000	50.000	0.0	104	0.00
35 S	Dibromofluoromethane	50.000	47.826	4.3	106	0.00
36 T	1,1-Dichloropropene	50.000	49.586	0.8	101	0.00
37 T	Ethyl Acetate	50.000	47.290	5.4	102	0.00
38 T	Carbon Tetrachloride	50.000	46.350	7.3	102	0.00
39 T	Methylcyclohexane	50.000	54.277	-8.6	101	0.00
40 TM	Benzene	50.000	48.671	2.7	101	0.00
41 T	Methacrylonitrile	50.000	47.017	6.0	99	0.00
42 TM	1,2-Dichloroethane	50.000	46.050	7.9	100	0.00
43 T	Isopropyl Acetate	50.000	50.848	-1.7	100	0.00
44 TM	Trichloroethene	50.000	50.680	-1.4	104	0.00
45 C	1,2-Dichloropropane	50.000	50.088	-0.2#	102	0.00
46 T	Dibromomethane	50.000	48.059	3.9	103	0.00
47 T	Bromodichloromethane	50.000	46.843	6.3	101	0.00
48 T	Methyl methacrylate	50.000	52.825	-5.7	100	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039168.D
 Acq On : 25 Sep 2025 09:59
 Operator : SY/MD
 Sample : VSTDCCC050
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 2 Sample Multiplier: 1

Instrument :
 MSVOA_V
 LabSampleId :
 VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
 Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
49 T	1,4-Dioxane	1000.000	1135.751	-13.6	117	0.00
50 S	Toluene-d8	50.000	47.485	5.0	103	0.00
51 T	4-Methyl-2-Pentanone	250.000	265.177	-6.1	100	0.00
52 CM	Toluene	50.000	52.474	-4.9#	102	0.00
53 T	t-1,3-Dichloropropene	50.000	52.243	-4.5	102	0.00
54 T	cis-1,3-Dichloropropene	50.000	52.312	-4.6	101	0.00
55 T	1,1,2-Trichloroethane	50.000	48.443	3.1	105	0.00
56 T	Ethyl methacrylate	50.000	57.707	-15.4	103	0.00
57 T	1,3-Dichloropropane	50.000	49.373	1.3	102	0.00
58 T	2-Chloroethyl Vinyl ether	250.000	257.039	-2.8	105	0.00
59 T	2-Hexanone	250.000	247.438	1.0	101	0.00
60 T	Dibromochloromethane	50.000	48.938	2.1	106	0.00
61 T	1,2-Dibromoethane	50.000	50.719	-1.4	105	0.00
62 S	4-Bromofluorobenzene	50.000	52.218	-4.4	105	0.00
63 I	Chlorobenzene-d5	50.000	50.000	0.0	104	0.00
64 T	Tetrachloroethene	50.000	49.857	0.3	106	0.00
65 PM	Chlorobenzene	50.000	49.450	1.1	104	0.00
66 T	1,1,1,2-Tetrachloroethane	50.000	47.975	4.0	103	0.00
67 C	Ethyl Benzene	50.000	55.024	-10.0#	102	0.00
68 T	m/p-Xylenes	100.000	115.018	-15.0	103	0.00
69 T	o-Xylene	50.000	58.347	-16.7	104	0.00
70 T	Styrene	50.000	50.114	-0.2	103	0.00
71 P	Bromoform	50.000	51.118	-2.2	106	0.00
72 I	1,4-Dichlorobenzene-d4	50.000	50.000	0.0	104	0.00
73 T	Isopropylbenzene	50.000	54.407	-8.8	104	0.00
74 T	N-amyl acetate	50.000	55.064	-10.1	99	0.00
75 P	1,1,2,2-Tetrachloroethane	50.000	43.928	12.1	102	0.00
76 T	1,2,3-Trichloropropane	50.000	47.019	6.0	106	0.00
77 T	Bromobenzene	50.000	49.357	1.3	104	0.00
78 T	n-propylbenzene	50.000	53.995	-8.0	101	0.00
79 T	2-Chlorotoluene	50.000	52.939	-5.9	102	0.00
80 T	1,3,5-Trimethylbenzene	50.000	56.020	-12.0	103	0.00
81 T	trans-1,4-Dichloro-2-butene	50.000	50.926	-1.9	101	0.00
82 T	4-Chlorotoluene	50.000	53.904	-7.8	103	0.00
83 T	tert-Butylbenzene	50.000	53.579	-7.2	104	0.00
84 T	1,2,4-Trimethylbenzene	50.000	56.693	-13.4	102	0.00
85 T	sec-Butylbenzene	50.000	53.713	-7.4	102	0.00
86 T	p-Isopropyltoluene	50.000	54.380	-8.8	102	0.00
87 T	1,3-Dichlorobenzene	50.000	48.773	2.5	104	0.00
88 T	1,4-Dichlorobenzene	50.000	45.535	8.9	102	0.00
89 T	n-Butylbenzene	50.000	54.607	-9.2	99	0.00
90 T	Hexachloroethane	50.000	44.260	11.5	103	0.00
91 T	1,2-Dichlorobenzene	50.000	48.799	2.4	104	0.00
92 T	1,2-Dibromo-3-Chloropropane	50.000	50.961	-1.9	106	0.00
93 T	1,2,4-Trichlorobenzene	50.000	52.055	-4.1	102	0.00
94 T	Hexachlorobutadiene	50.000	45.975	8.0	105	0.00
95 T	Naphthalene	50.000	57.749	-15.5	104	0.00

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
Data File : VV039168.D
Acq On : 25 Sep 2025 09:59
Operator : SY/MD
Sample : VSTDCCC050
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 2 Sample Multiplier: 1

Instrument :
MSVOA_V
LabSampleId :
VSTDCCC050

Quant Time: Sep 26 01:18:56 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Min. RRF : 0.000 Min. Rel. Area : 50% Max. R.T. Dev 0.50min
Max. RRF Dev : 20% Max. Rel. Area : 150%

	Compound	Amount	Calc.	%Dev	Area%	Dev(min)
96 T	1,2,3-Trichlorobenzene	50.000	51.848	-3.7	101	0.00

(#) = Out of Range SPCC's out = 0 CCC's out = 6



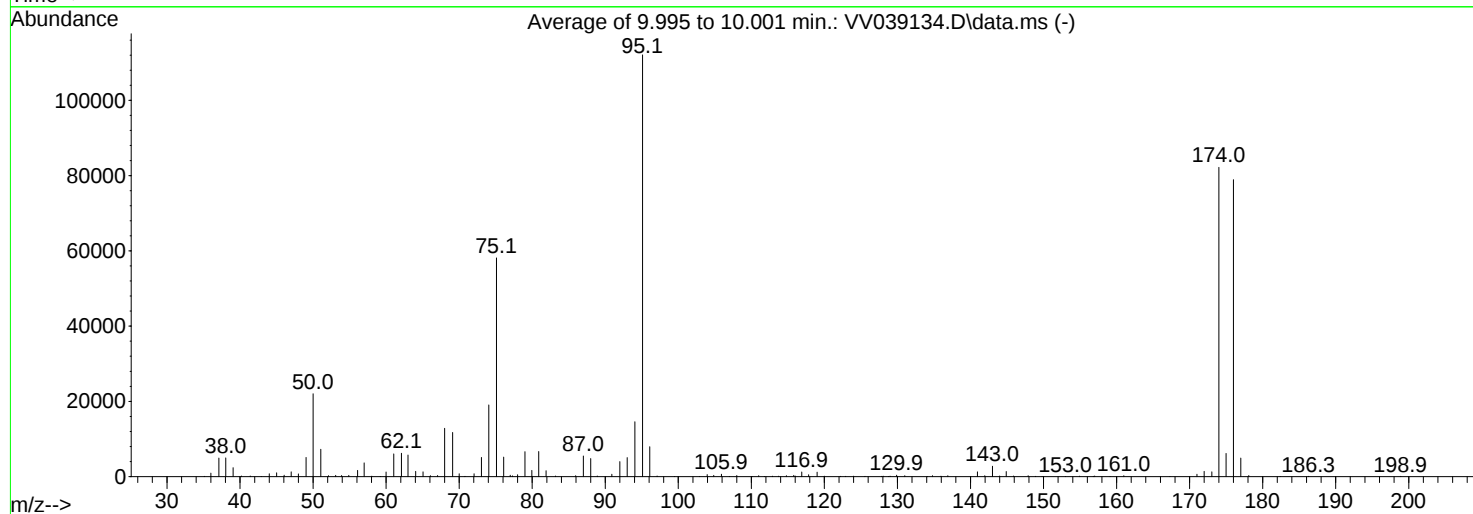
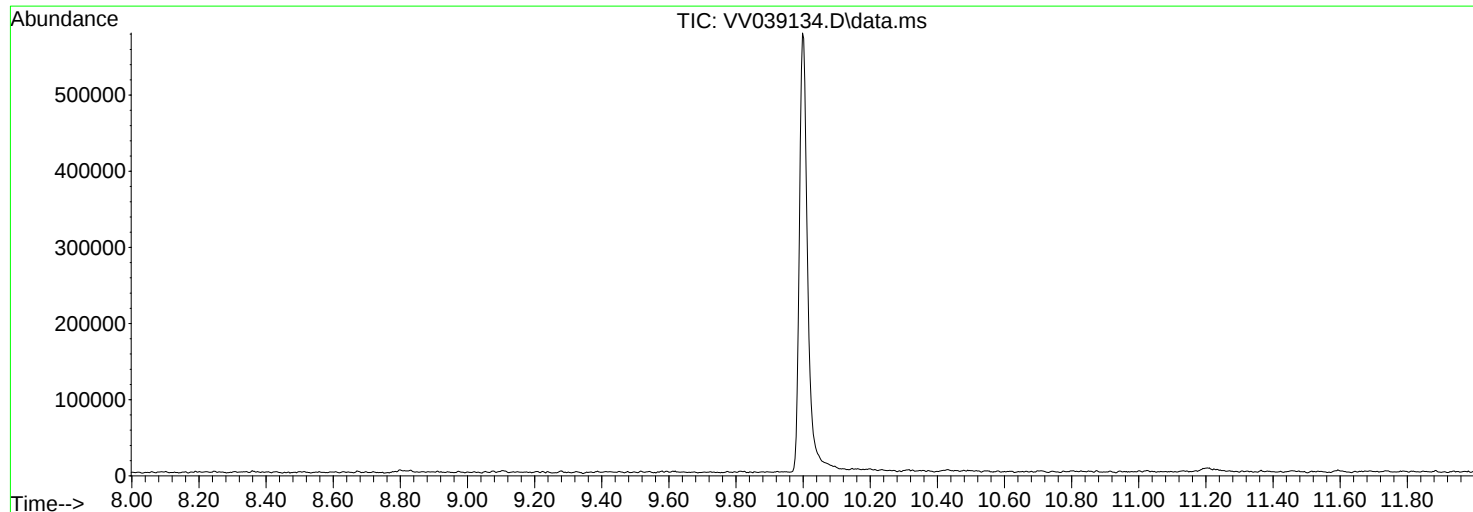
QC SAMPLE DATA

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092225\
Data File : VV039134.D
Acq On : 22 Sep 2025 10:15
Operator : SY/MD
Sample : BFB
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 1 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
BFB

Integration File: RTEINT.P

Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Title : SW846 8260
Last Update : Wed Sep 24 08:08:08 2025



AutoFind: Scans 2785, 2786, 2787; Background Corrected with Scan 2774

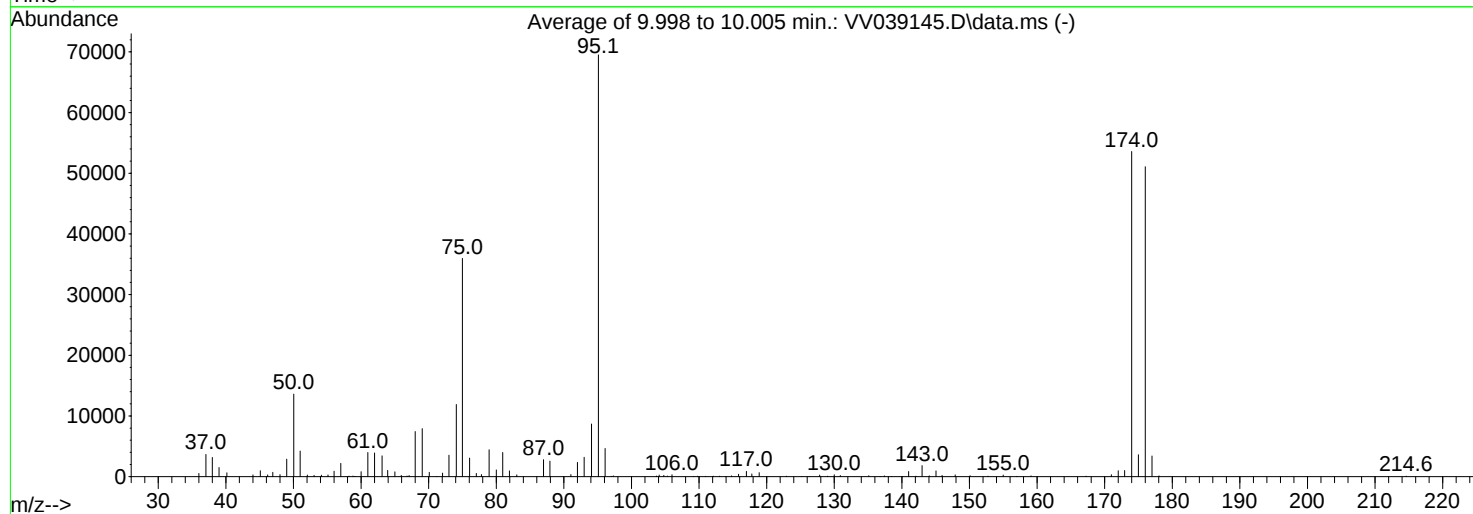
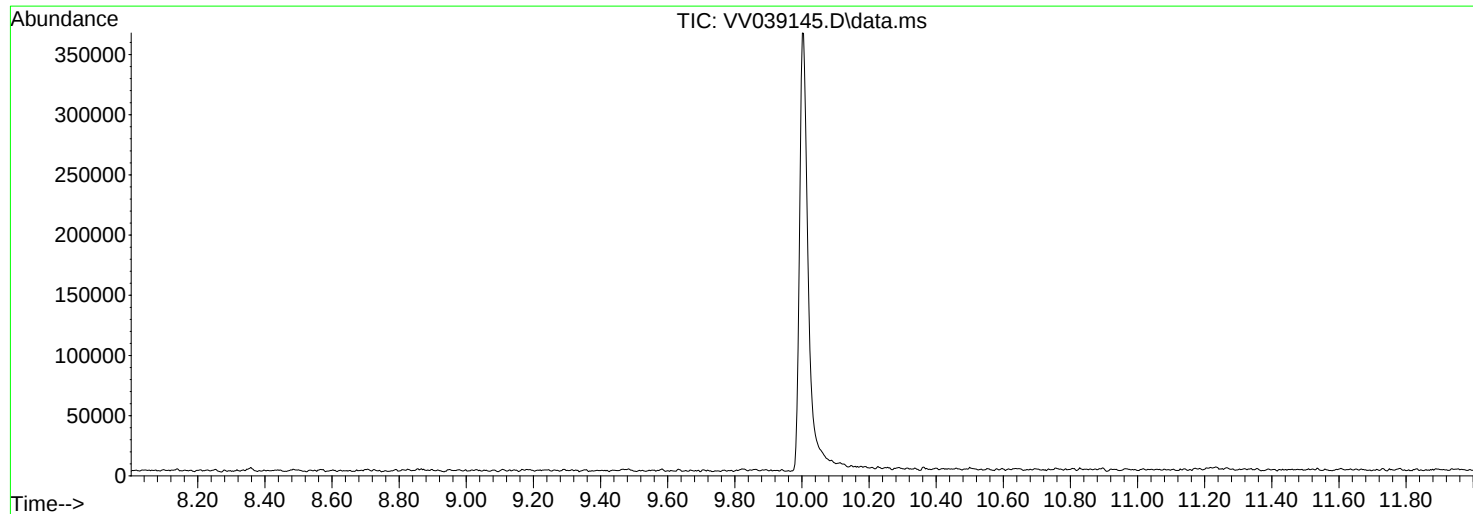
Target	Rel. to	Lower	Upper	Rel.	Raw	Result
Mass	Mass	Limit%	Limit%	Abn%	Abn	Pass/Fail
50	95	15	40	19.6	22014	PASS
75	95	30	60	51.9	58147	PASS
95	95	100	100	100.0	112096	PASS
96	95	5	9	7.1	7955	PASS
173	174	0.00	2	1.6	1275	PASS
174	95	50	100	73.3	82200	PASS
175	174	5	9	7.5	6152	PASS
176	174	95	101	96.0	78917	PASS
177	176	5	9	6.3	4933	PASS

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039145.D
Acq On : 24 Sep 2025 09:52
Operator : SY/MD
Sample : BFB
Misc : 25.0mL/MSVOA_V/WATER
ALS Vial : 1 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
BFB

Integration File: RTEINT.P

Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Title : SW846 8260
Last Update : Wed Sep 24 08:08:08 2025



AutoFind: Scans 2786, 2787, 2788; Background Corrected with Scan 2776

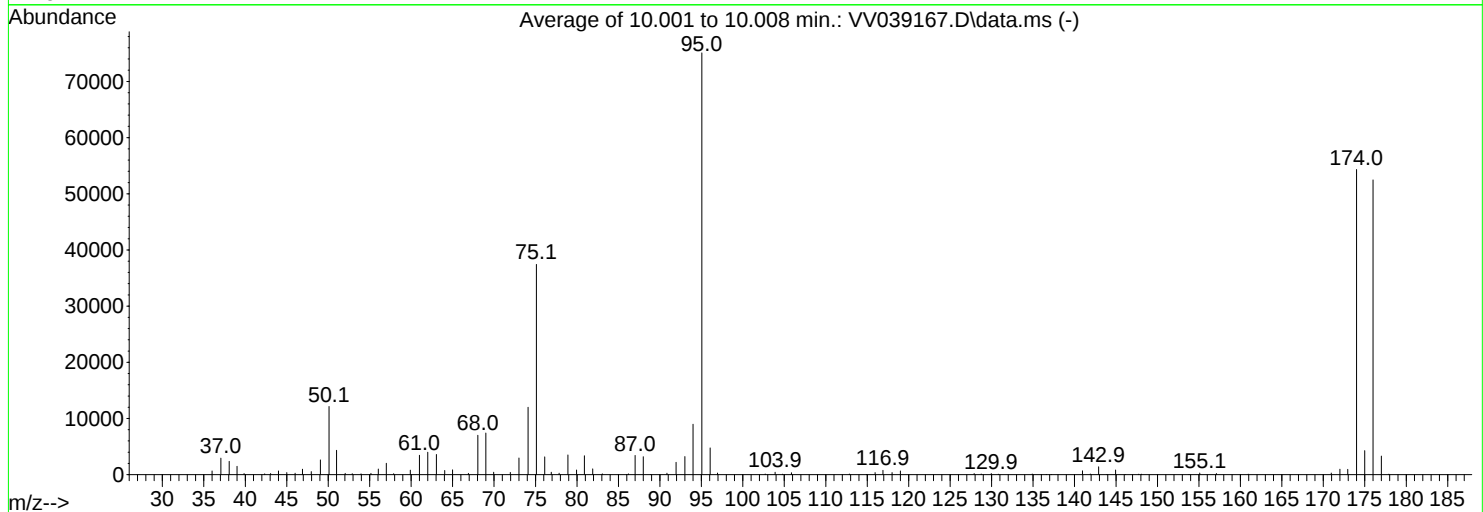
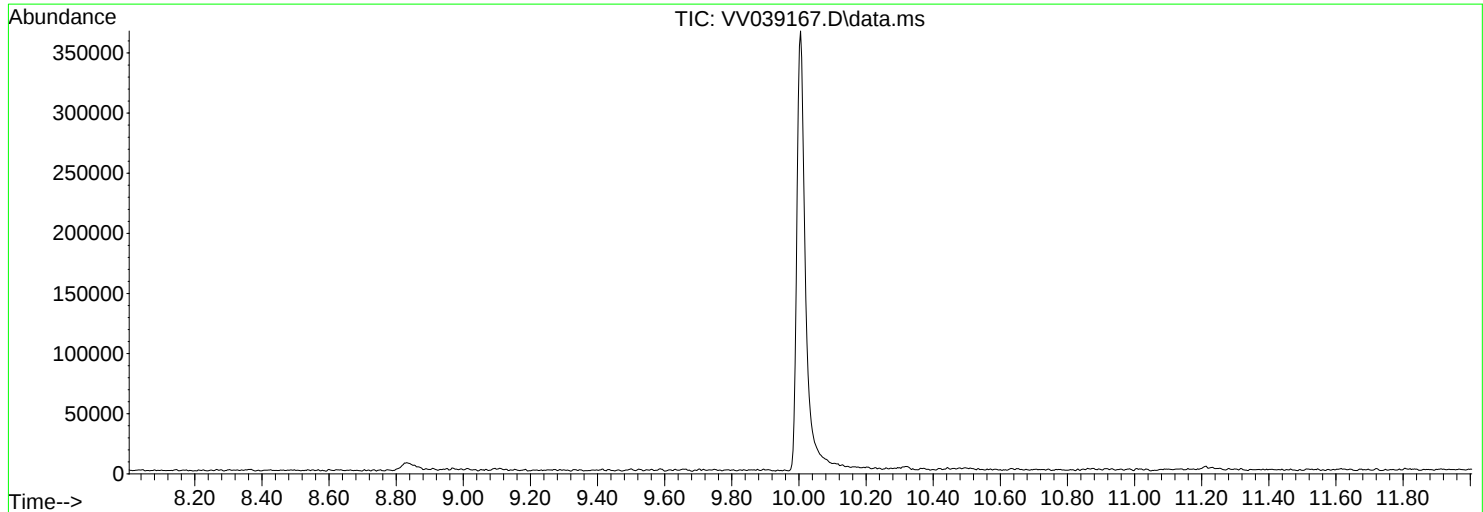
Target Mass	Rel. to Mass	Lower Limit%	Upper Limit%	Rel. Abn%	Raw Abn	Result Pass/Fail
50	95	15	40	19.6	13613	PASS
75	95	30	60	51.7	35965	PASS
95	95	100	100	100.0	69512	PASS
96	95	5	9	6.7	4635	PASS
173	174	0.00	2	1.9	995	PASS
174	95	50	100	77.1	53579	PASS
175	174	5	9	6.7	3610	PASS
176	174	95	101	95.3	51064	PASS
177	176	5	9	6.6	3395	PASS

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
Data File : VV039167.D
Acq On : 25 Sep 2025 09:14
Operator : SY/MD
Sample : BFB
Misc : 25.0mL/MSVOA_V/WATER
ALS Vial : 1 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
BFB

Integration File: RTEINT.P

Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Title : SW846 8260
Last Update : Wed Sep 24 08:08:08 2025



AutoFind: Scans 2787, 2788, 2789; Background Corrected with Scan 2776

Target	Rel. to	Lower	Upper	Rel.	Raw	Result
Mass	Mass	Limit%	Limit%	Abn%	Abn	Pass/Fail
50	95	15	40	16.2	12139	PASS
75	95	30	60	49.9	37445	PASS
95	95	100	100	100.0	75093	PASS
96	95	5	9	6.3	4756	PASS
173	174	0.00	2	1.7	944	PASS
174	95	50	100	72.3	54320	PASS
175	174	5	9	7.9	4279	PASS
176	174	95	101	96.6	52461	PASS
177	176	5	9	6.3	3317	PASS

Report of Analysis

Client:	ATG-GREENVILLE AEC	Date Collected:	
Project:	AEC-2025-0013 CSC 4151 Coram	Date Received:	
Client Sample ID:	VV0924WBL01	SDG No.:	Q3164
Lab Sample ID:	VV0924WBL01	Matrix:	Water
Analytical Method:	8260D	% Solid:	0
Sample Wt/Vol:	5 Units: mL	Final Vol:	5000 uL
Soil Aliquot Vol:	uL	Test:	VOC-BTEX
GC Column:	DB-624UI ID : 0.18	Level :	LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039148.D	1	09/24/25 13:09	VV092425

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	0.15	U	0.15	1.00	ug/L
108-88-3	Toluene	0.14	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	0.13	U	0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	0.24	U	0.24	2.00	ug/L
95-47-6	o-Xylene	0.12	U	0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	62.2		74 - 125	124%	SPK: 50
1868-53-7	Dibromofluoromethane	54.6		75 - 124	109%	SPK: 50
2037-26-5	Toluene-d8	44.1		86 - 113	88%	SPK: 50
460-00-4	4-Bromofluorobenzene	48.0		77 - 121	96%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	464000	4.596			
540-36-3	1,4-Difluorobenzene	859000	5.532			
3114-55-4	Chlorobenzene-d5	832000	8.773			
3855-82-1	1,4-Dichlorobenzene-d4	379000	11.175			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039148.D
 Acq On : 24 Sep 2025 13:09
 Operator : SY/MD
 Sample : VV0924WBL01
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 4 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0924WBL01

Quant Time: Sep 25 04:44:19 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.596	168	463770	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	858892	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.773	117	831675	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.175	152	379013	50.000	ug/l	0.00

System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.937	65	417696	62.230	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	124.460%#
35) Dibromofluoromethane	4.497	113	376781	54.568	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	109.140%
50) Toluene-d8	7.236	98	894968	44.132	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	88.260%#
62) 4-Bromofluorobenzene	10.001	95	400947	48.026	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	96.060%

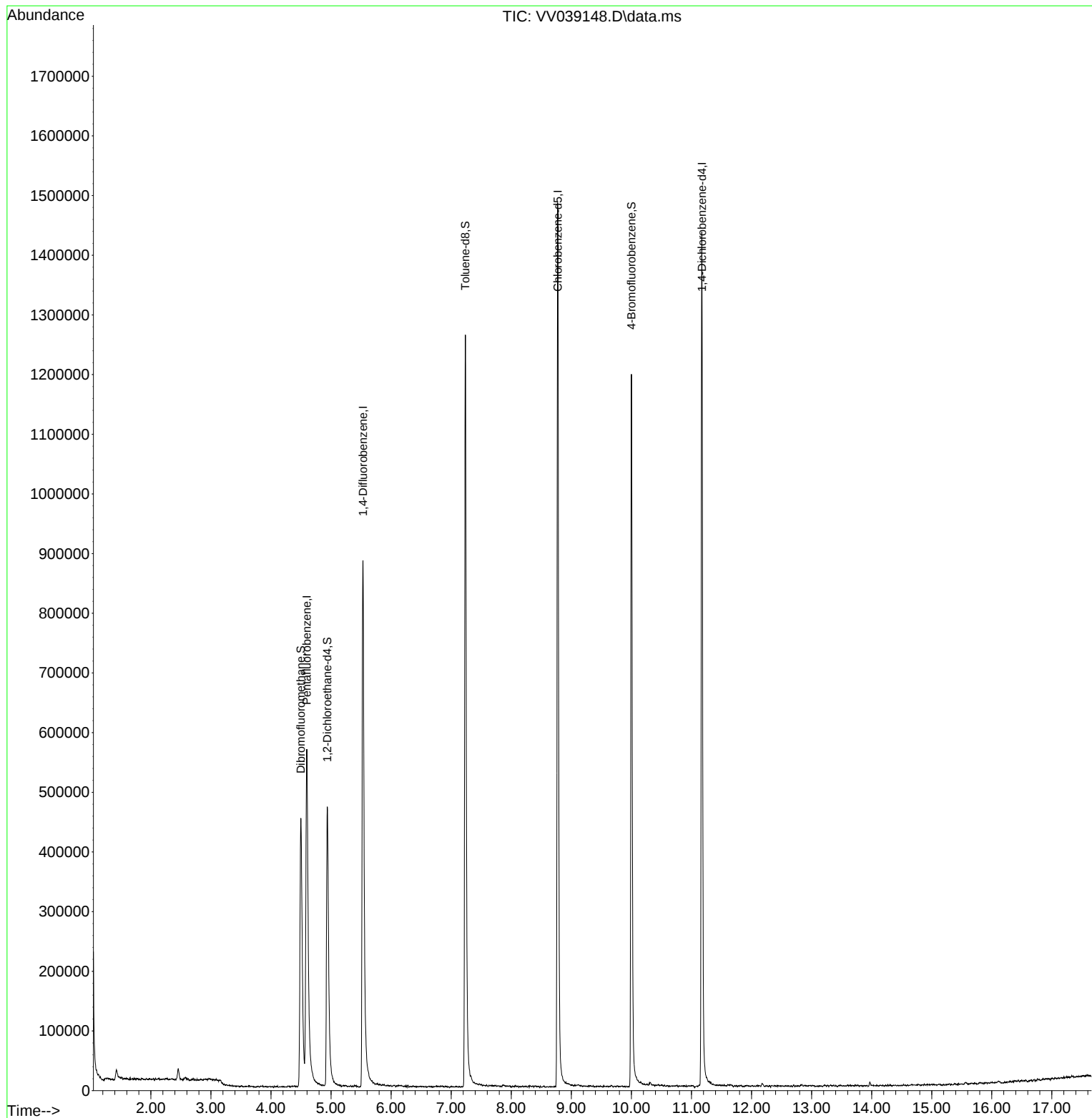
Target Compounds	Qvalue

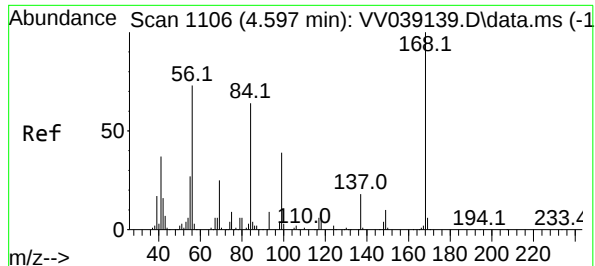
(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039148.D
Acq On : 24 Sep 2025 13:09
Operator : SY/MD
Sample : VV0924WBL01
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 4 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VV0924WBL01

Quant Time: Sep 25 04:44:19 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

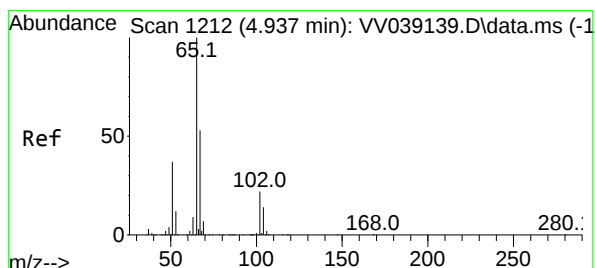
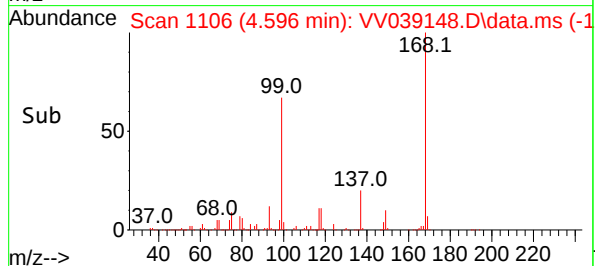
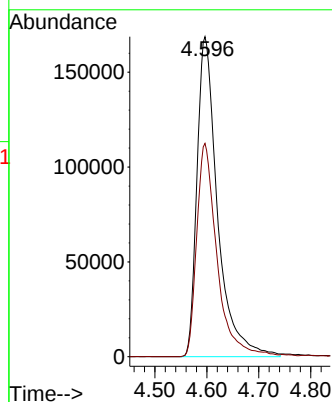
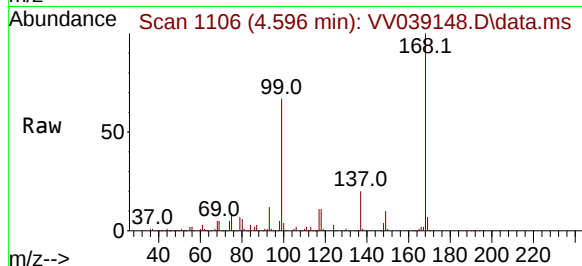




#1
Pentafluorobenzene
Concen: 50.000 ug/l
RT: 4.596 min Scan# 1106
Delta R.T. -0.001 min
Lab File: VV039148.D
Acq: 24 Sep 2025 13:09

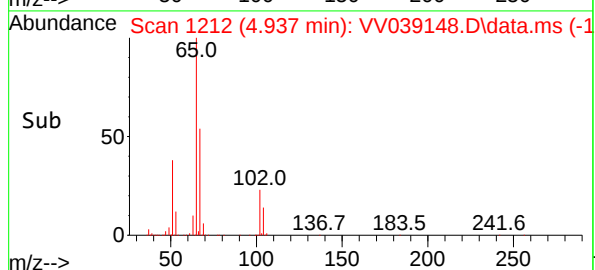
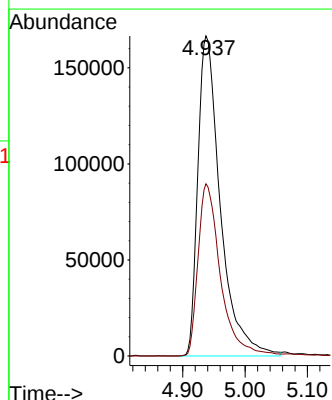
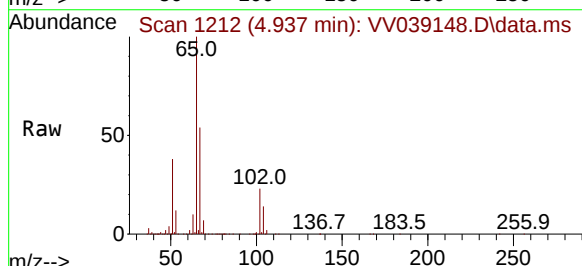
Instrument :
MSVOA_V
ClientSampleId :
VV0924WBL01

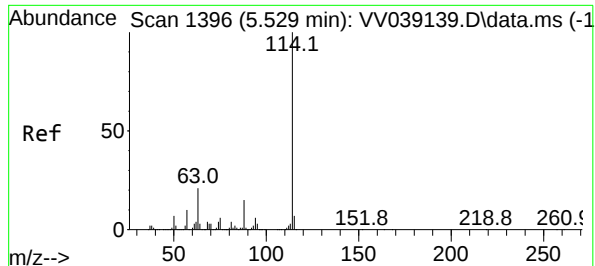
Tgt Ion:168 Resp: 463770
Ion Ratio Lower Upper
168 100
99 66.7 49.6 74.4



#33
1,2-Dichloroethane-d4
Concen: 62.230 ug/l
RT: 4.937 min Scan# 1212
Delta R.T. -0.000 min
Lab File: VV039148.D
Acq: 24 Sep 2025 13:09

Tgt Ion: 65 Resp: 417696
Ion Ratio Lower Upper
65 100
67 53.6 0.0 107.0





#34

1,4-Difluorobenzene

Concen: 50.000 ug/l

RT: 5.532 min Scan# 1396

Delta R.T. 0.003 min

Lab File: VV039148.D

Acq: 24 Sep 2025 13:09

Instrument :

MSVOA_V

ClientSampleId :

VV0924WBL01

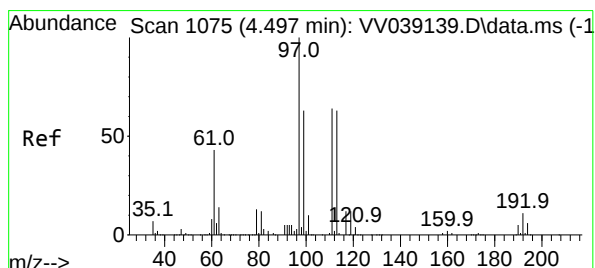
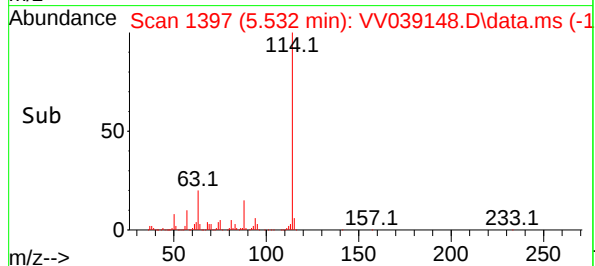
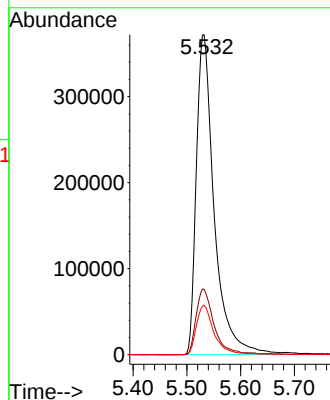
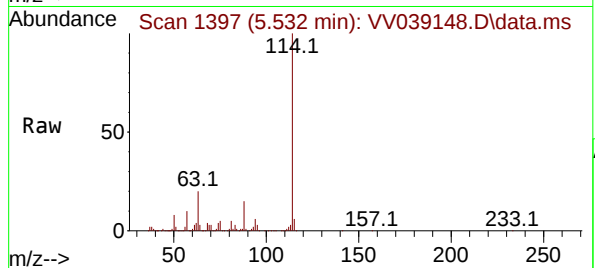
Tgt Ion:114 Resp: 858892

Ion Ratio Lower Upper

114 100

63 20.4 0.0 42.6

88 15.4 0.0 30.8



#35

Dibromofluoromethane

Concen: 54.568 ug/l

RT: 4.497 min Scan# 1075

Delta R.T. -0.000 min

Lab File: VV039148.D

Acq: 24 Sep 2025 13:09

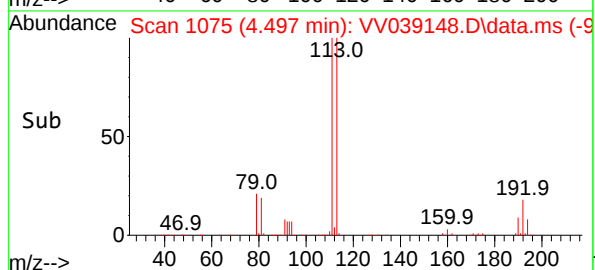
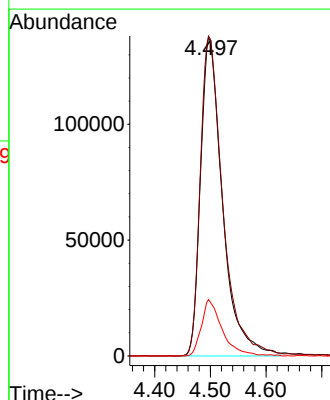
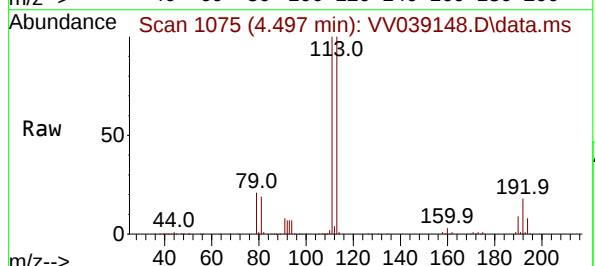
Tgt Ion:113 Resp: 376781

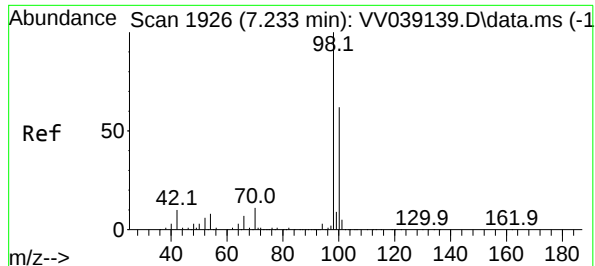
Ion Ratio Lower Upper

113 100

111 102.7 82.4 123.6

192 16.6 13.8 20.8





#50

Toluene-d8

Concen: 44.132 ug/l

RT: 7.236 min Scan# 1926

Delta R.T. 0.003 min

Lab File: VV039148.D

Acq: 24 Sep 2025 13:09

Instrument :

MSVOA_V

ClientSampleId :

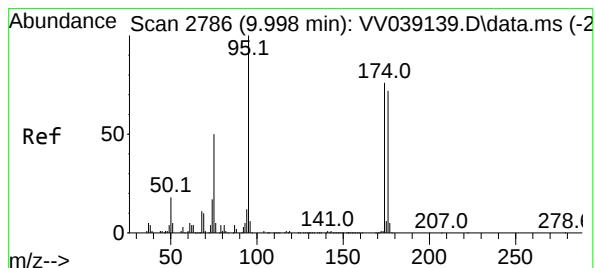
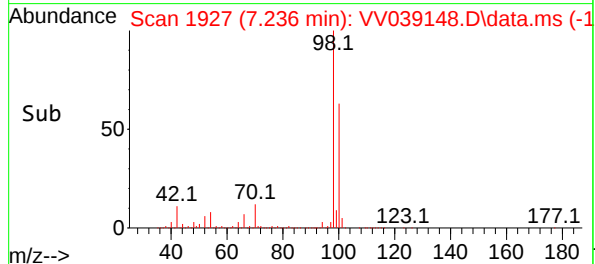
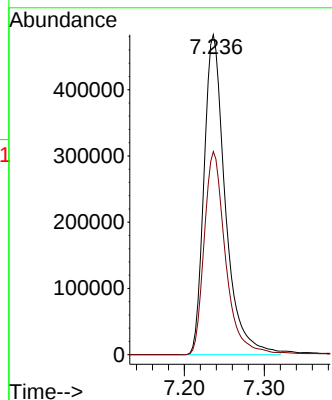
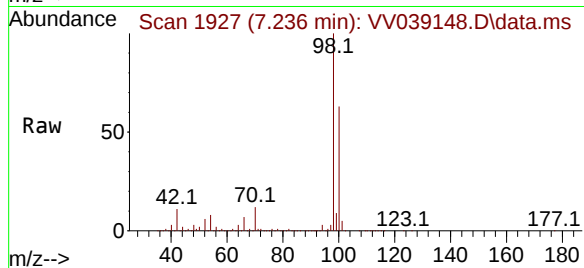
VV0924WBL01

Tgt Ion: 98 Resp: 894968

Ion Ratio Lower Upper

98 100

100 63.4 52.2 78.2



#62

4-Bromofluorobenzene

Concen: 48.026 ug/l

RT: 10.001 min Scan# 2787

Delta R.T. 0.003 min

Lab File: VV039148.D

Acq: 24 Sep 2025 13:09

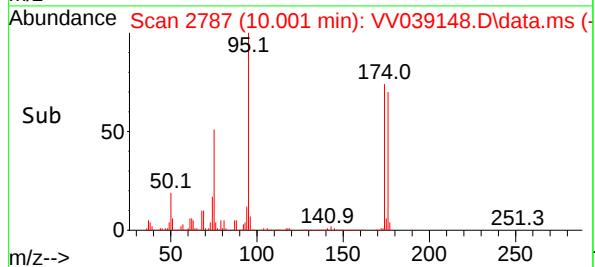
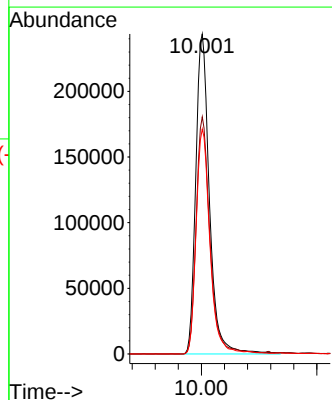
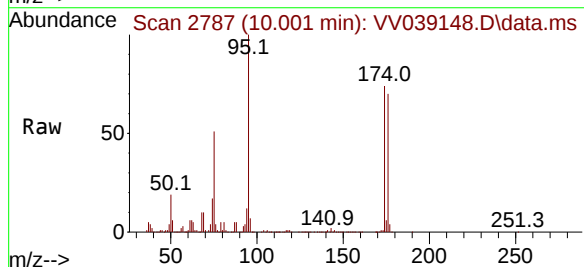
Tgt Ion: 95 Resp: 400947

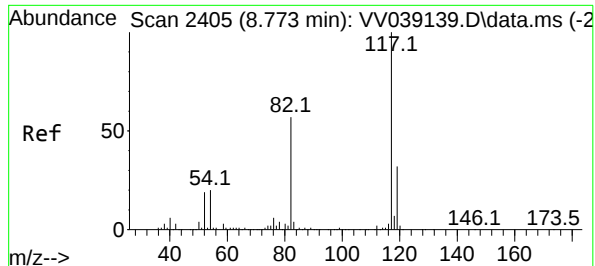
Ion Ratio Lower Upper

95 100

174 74.8 0.0 150.4

176 70.4 0.0 144.2





#63

Chlorobenzene-d5

Concen: 50.000 ug/l

RT: 8.773 min Scan# 2405

Delta R.T. 0.000 min

Lab File: VV039148.D

Acq: 24 Sep 2025 13:09

Instrument :

MSVOA_V

ClientSampleId :

VV0924WBL01

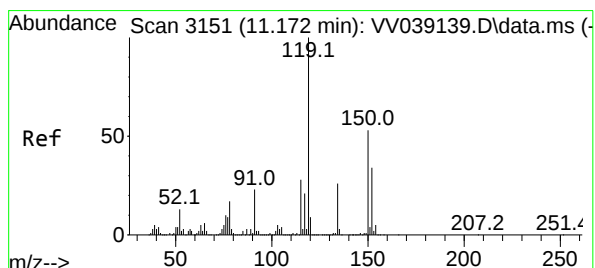
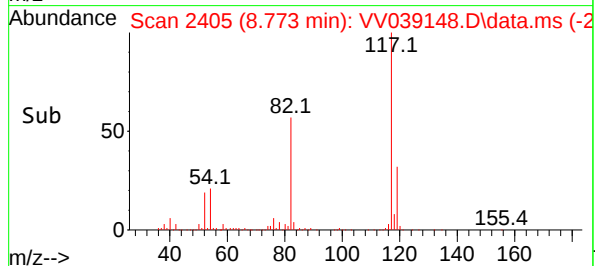
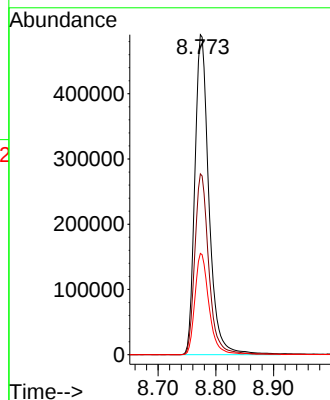
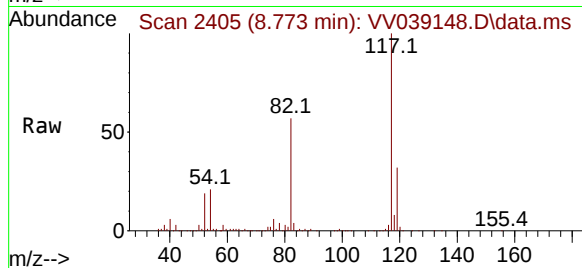
Tgt Ion:117 Resp: 831675

Ion Ratio Lower Upper

117 100

82 56.5 45.4 68.2

119 31.7 25.6 38.4



#72

1,4-Dichlorobenzene-d4

Concen: 50.000 ug/l

RT: 11.175 min Scan# 3152

Delta R.T. 0.003 min

Lab File: VV039148.D

Acq: 24 Sep 2025 13:09

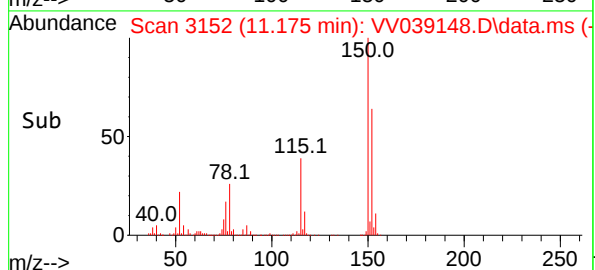
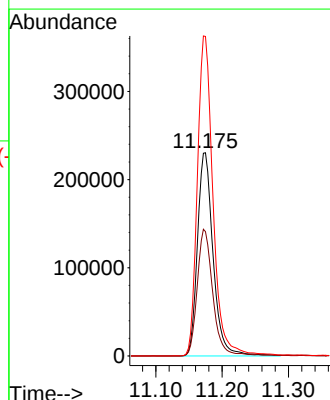
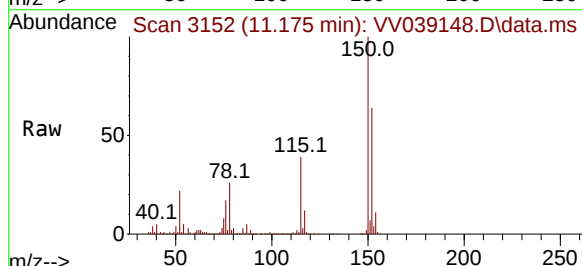
Tgt Ion:152 Resp: 379013

Ion Ratio Lower Upper

152 100

115 61.8 44.8 134.4

150 157.4 0.0 353.8



Report of Analysis

Client:	ATG-GREENVILLE AEC			Date Collected:			
Project:	AEC-2025-0013 CSC 4151 Coram			Date Received:			
Client Sample ID:	VV0925WBL01			SDG No.:	Q3164		
Lab Sample ID:	VV0925WBL01			Matrix:	Water		
Analytical Method:	8260D			% Solid:	0		
Sample Wt/Vol:	5	Units:	mL	Final Vol:	5000	uL	
Soil Aliquot Vol:			uL	Test:	VOC-BTEX		
GC Column:	DB-624UI	ID :	0.18	Level :	LOW		
Prep Method :							

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039170.D	1	09/25/25 11:01	VV092525

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	0.15	U	0.15	1.00	ug/L
108-88-3	Toluene	0.14	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	0.13	U	0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	0.24	U	0.24	2.00	ug/L
95-47-6	o-Xylene	0.12	U	0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	60.1		74 - 125	120%	SPK: 50
1868-53-7	Dibromofluoromethane	53.3		75 - 124	107%	SPK: 50
2037-26-5	Toluene-d8	47.3		86 - 113	95%	SPK: 50
460-00-4	4-Bromofluorobenzene	44.1		77 - 121	88%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	480000	4.597			
540-36-3	1,4-Difluorobenzene	865000	5.532			
3114-55-4	Chlorobenzene-d5	871000	8.776			
3855-82-1	1,4-Dichlorobenzene-d4	414000	11.175			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039170.D
 Acq On : 25 Sep 2025 11:01
 Operator : SY/MD
 Sample : VV0925WBL01
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 4 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0925WBL01

Quant Time: Sep 26 01:20:45 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.597	168	480047	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	864912	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	870930	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.175	152	413592	50.000	ug/l	0.00

System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.941	65	417535	60.097	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	120.200%#
35) Dibromofluoromethane	4.500	113	370419	53.273	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	106.540%
50) Toluene-d8	7.236	98	965988	47.302	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	94.600%
62) 4-Bromofluorobenzene	10.001	95	371030	44.133	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	88.260%

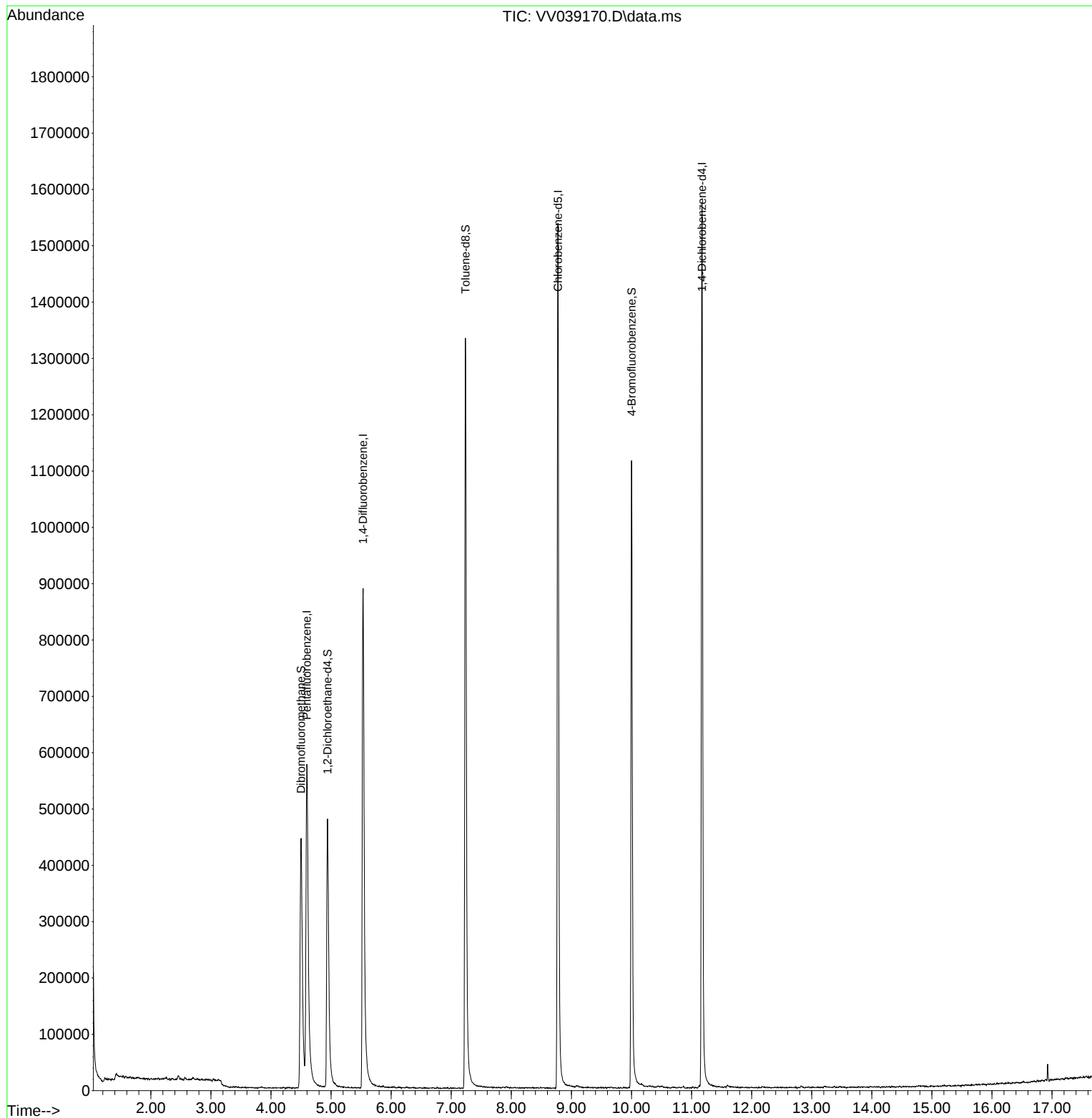
Target Compounds	Qvalue

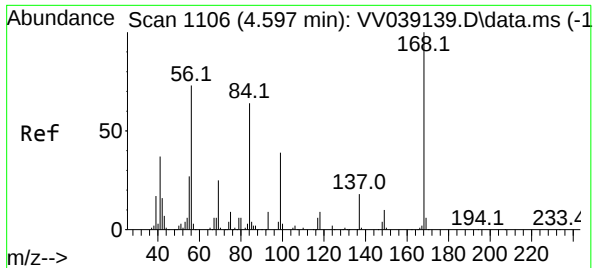
(#) = qualifier out of range (m) = manual integration (+) = signals summed

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
Data File : VV039170.D
Acq On : 25 Sep 2025 11:01
Operator : SY/MD
Sample : VV0925WBL01
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 4 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VV0925WBL01

Quant Time: Sep 26 01:20:45 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

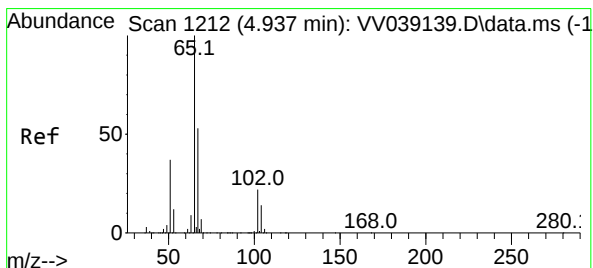
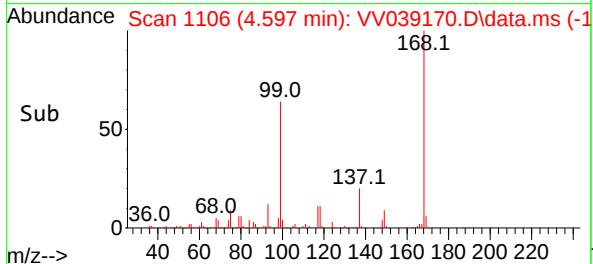
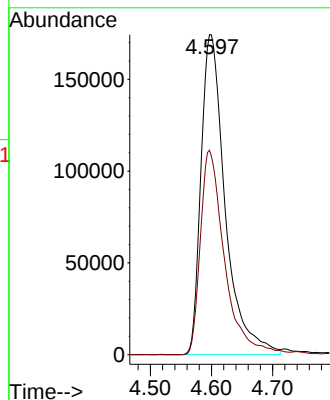
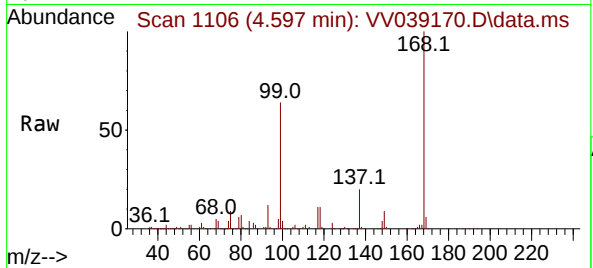




#1
Pentafluorobenzene
Concen: 50.000 ug/l
RT: 4.597 min Scan# 1106
Delta R.T. -0.000 min
Lab File: VV039170.D
Acq: 25 Sep 2025 11:01

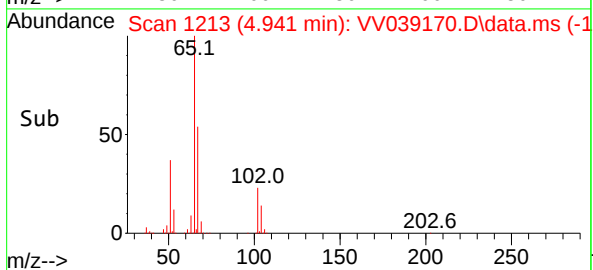
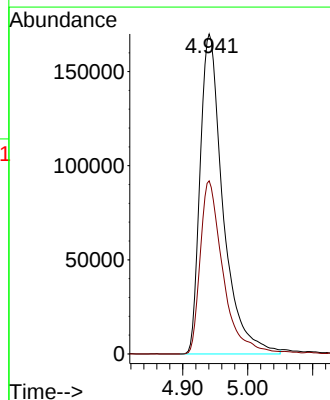
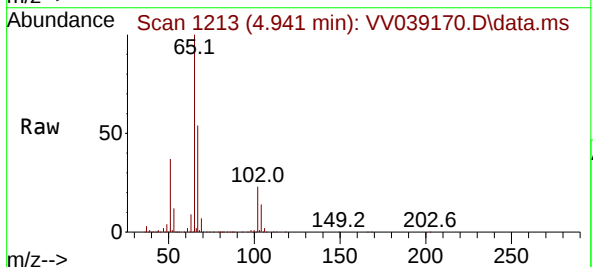
Instrument :
MSVOA_V
ClientSampleId :
VV0925WBL01

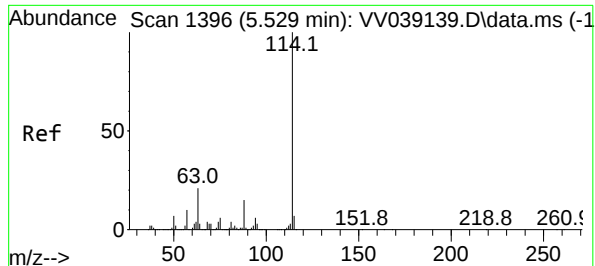
Tgt Ion:168 Resp: 480047
Ion Ratio Lower Upper
168 100
99 63.9 49.6 74.4



#33
1,2-Dichloroethane-d4
Concen: 60.097 ug/l
RT: 4.941 min Scan# 1213
Delta R.T. 0.003 min
Lab File: VV039170.D
Acq: 25 Sep 2025 11:01

Tgt Ion: 65 Resp: 417535
Ion Ratio Lower Upper
65 100
67 54.2 0.0 107.0





#34

1,4-Difluorobenzene

Concen: 50.000 ug/l

RT: 5.532 min Scan# 11

Delta R.T. 0.003 min

Lab File: VV039170.D

Acq: 25 Sep 2025 11:01

Instrument :

MSVOA_V

ClientSampleId :

VV0925WBL01

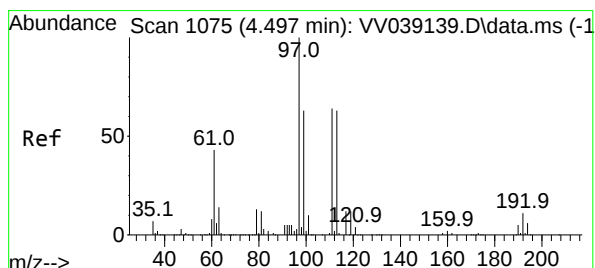
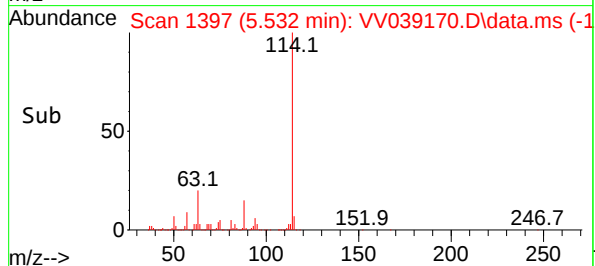
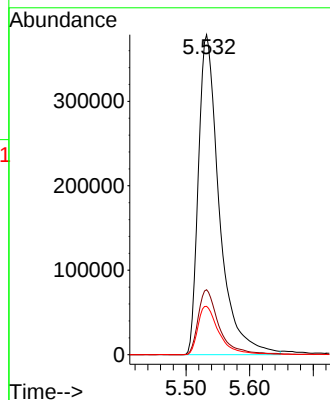
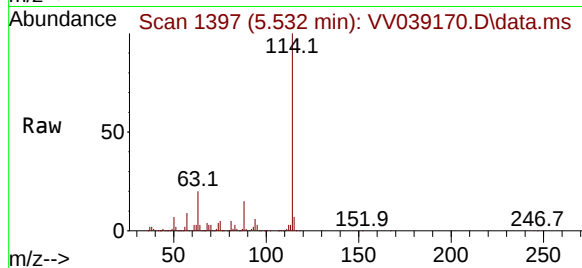
Tgt Ion:114 Resp: 864912

Ion Ratio Lower Upper

114 100

63 20.3 0.0 42.6

88 15.1 0.0 30.8



#35

Dibromofluoromethane

Concen: 53.273 ug/l

RT: 4.500 min Scan# 1076

Delta R.T. 0.003 min

Lab File: VV039170.D

Acq: 25 Sep 2025 11:01

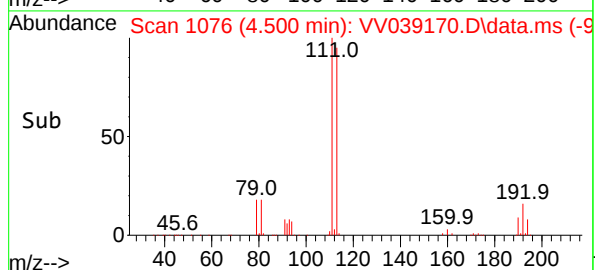
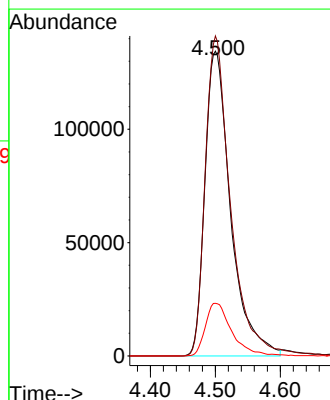
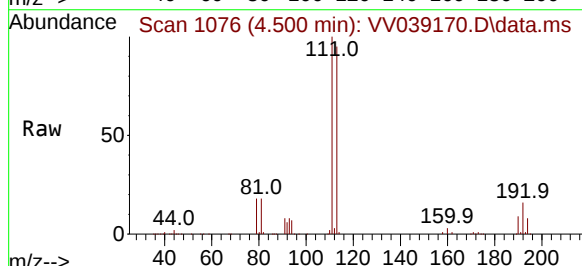
Tgt Ion:113 Resp: 370419

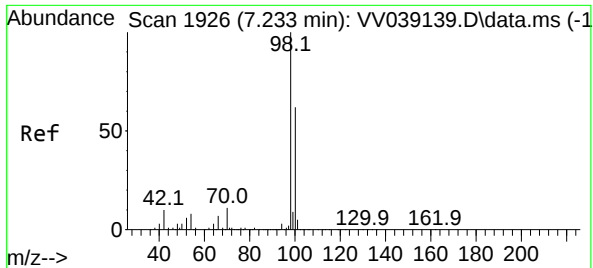
Ion Ratio Lower Upper

113 100

111 104.0 82.4 123.6

192 17.8 13.8 20.8

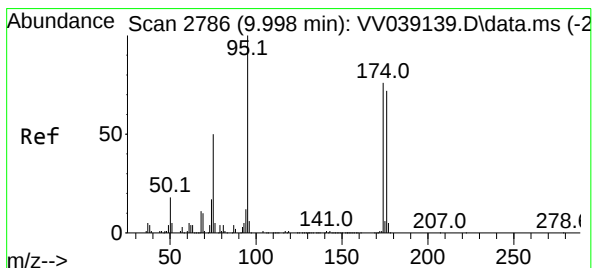
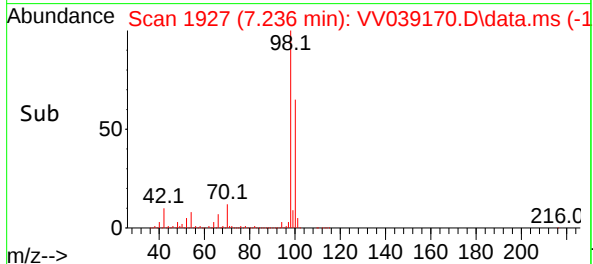
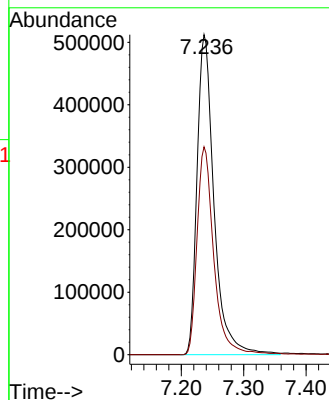
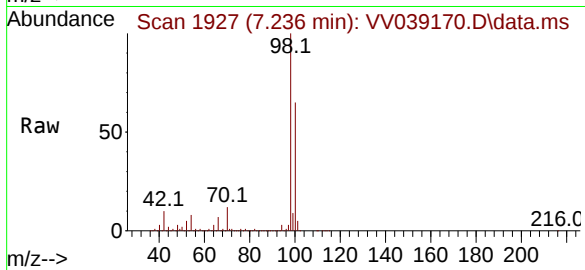




#50
Toluene-d8
Concen: 47.302 ug/l
RT: 7.236 min Scan# 1926
Delta R.T. 0.003 min
Lab File: VV039170.D
Acq: 25 Sep 2025 11:01

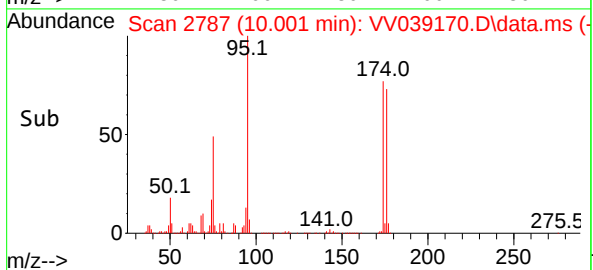
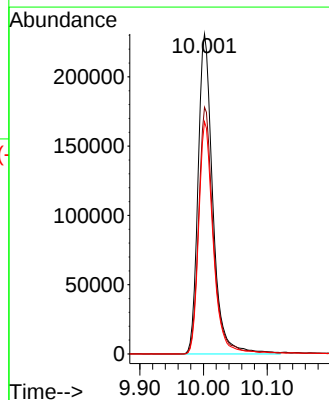
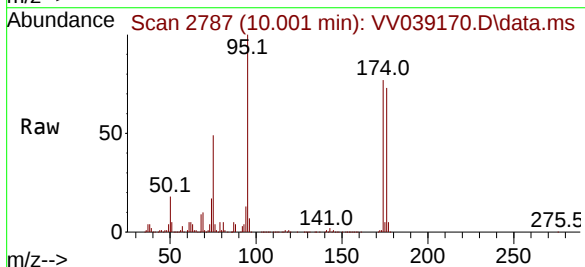
Instrument :
MSVOA_V
ClientSampleId :
VV0925WBL01

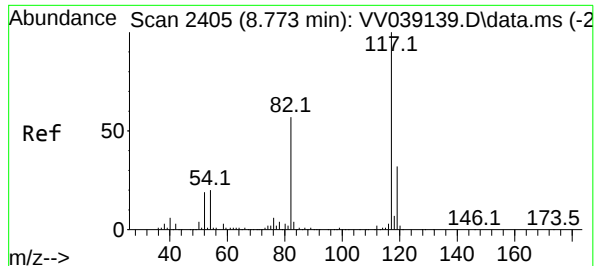
Tgt Ion: 98 Resp: 965988
Ion Ratio Lower Upper
98 100
100 64.0 52.2 78.2



#62
4-Bromofluorobenzene
Concen: 44.133 ug/l
RT: 10.001 min Scan# 2787
Delta R.T. 0.003 min
Lab File: VV039170.D
Acq: 25 Sep 2025 11:01

Tgt Ion: 95 Resp: 371030
Ion Ratio Lower Upper
95 100
174 77.2 0.0 150.4
176 71.4 0.0 144.2





#63

Chlorobenzene-d5

Concen: 50.000 ug/l

RT: 8.776 min Scan# 2405

Delta R.T. 0.003 min

Lab File: VV039170.D

Acq: 25 Sep 2025 11:01

Instrument :

MSVOA_V

ClientSampleId :

VV0925WBL01

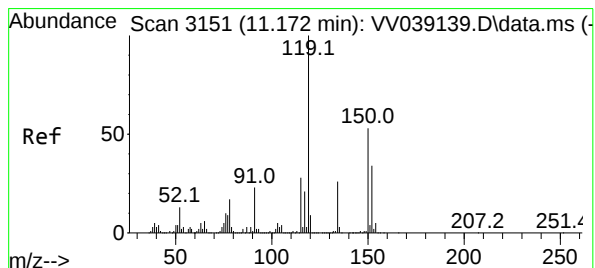
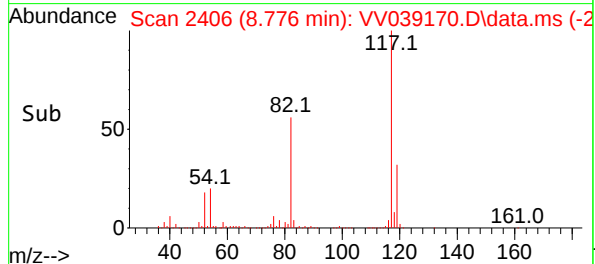
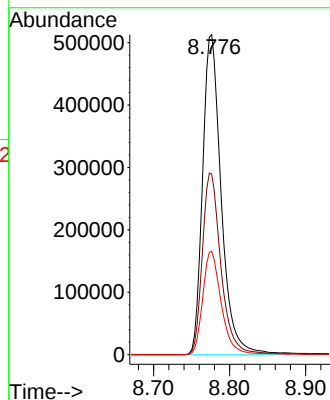
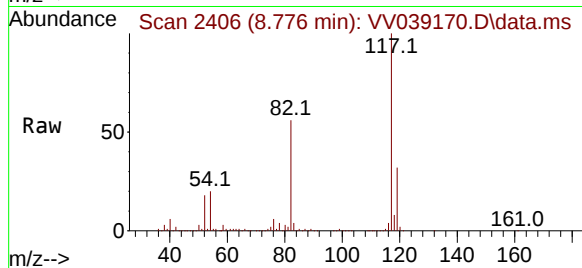
Tgt Ion:117 Resp: 870930

Ion Ratio Lower Upper

117 100

82 56.4 45.4 68.2

119 32.3 25.6 38.4



#72

1,4-Dichlorobenzene-d4

Concen: 50.000 ug/l

RT: 11.175 min Scan# 3152

Delta R.T. 0.003 min

Lab File: VV039170.D

Acq: 25 Sep 2025 11:01

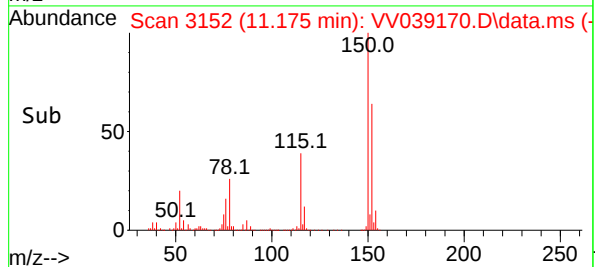
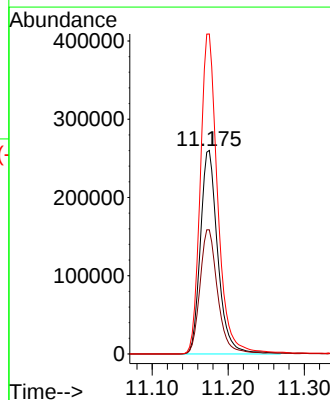
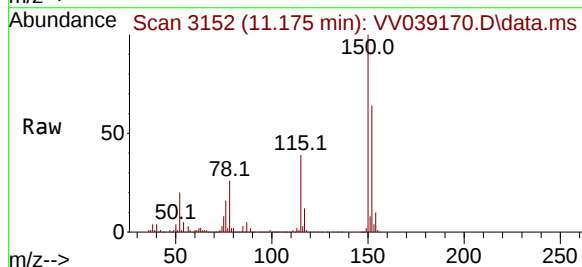
Tgt Ion:152 Resp: 413592

Ion Ratio Lower Upper

152 100

115 62.1 44.8 134.4

150 158.0 0.0 353.8



Report of Analysis

Client:	ATG-GREENVILLE AEC			Date Collected:	
Project:	AEC-2025-0013 CSC 4151 Coram			Date Received:	
Client Sample ID:	VV0924WBS02			SDG No.:	Q3164
Lab Sample ID:	VV0924WBS02			Matrix:	Water
Analytical Method:	8260D			% Solid:	0
Sample Wt/Vol:	5	Units:	mL	Final Vol:	5000 uL
Soil Aliquot Vol:			uL	Test:	VOC-BTEX
GC Column:	DB-624UI	ID :	0.18	Level :	LOW
Prep Method :					

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039150.D	1	09/24/25 14:01	VV092425

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	20.1		0.15	1.00	ug/L
108-88-3	Toluene	21.8		0.14	1.00	ug/L
100-41-4	Ethyl Benzene	19.9		0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	43.2		0.24	2.00	ug/L
95-47-6	o-Xylene	20.4		0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	49.1		74 - 125	98%	SPK: 50
1868-53-7	Dibromofluoromethane	50.0		75 - 124	100%	SPK: 50
2037-26-5	Toluene-d8	49.5		86 - 113	99%	SPK: 50
460-00-4	4-Bromofluorobenzene	52.0		77 - 121	104%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	492000	4.6			
540-36-3	1,4-Difluorobenzene	796000	5.532			
3114-55-4	Chlorobenzene-d5	785000	8.776			
3855-82-1	1,4-Dichlorobenzene-d4	469000	11.172			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039150.D
 Acq On : 24 Sep 2025 14:01
 Operator : SY/MD
 Sample : VV0924WBS02
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 6 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0924WBS02

Quant Time: Sep 25 04:45:44 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.600	168	492318	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	795602	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	784865	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	468522	50.000	ug/l	0.00
System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.941	65	349518	49.053	ug/l	0.00
Spiked Amount 50.000	Range 81 - 118		Recovery =	98.100%		
35) Dibromofluoromethane	4.500	113	319944	50.023	ug/l	0.00
Spiked Amount 50.000	Range 80 - 119		Recovery =	100.040%		
50) Toluene-d8	7.236	98	929285	49.469	ug/l	0.00
Spiked Amount 50.000	Range 89 - 112		Recovery =	98.940%		
62) 4-Bromofluorobenzene	10.002	95	402187	52.007	ug/l	0.00
Spiked Amount 50.000	Range 85 - 114		Recovery =	104.020%		
Target Compounds						Qvalue
2) Dichlorodifluoromethane	1.121	85	148756	19.087	ug/l	99
3) Chloromethane	1.230	50	159926	19.115	ug/l	98
4) Vinyl Chloride	1.298	62	173519	19.373	ug/l	100
5) Bromomethane	1.497	94	117993	20.897	ug/l	100
6) Chloroethane	1.561	64	117810	21.073	ug/l	99
7) Trichlorofluoromethane	1.725	101	259123	19.982	ug/l	99
8) Diethyl Ether	1.918	74	101938	21.175	ug/l	99
9) 1,1,2-Trichlorotrifluo...	2.082	101	151673	19.657	ug/l	99
10) Methyl Iodide	2.198	142	155103	21.109	ug/l	98
11) Tert butyl alcohol	2.568	59	166515	110.571	ug/l	99
12) 1,1-Dichloroethene	2.082	96	148258	19.973	ug/l	96
13) Acrolein	2.011	56	30535	116.851	ug/l	97
14) Allyl chloride	2.359	41	270301	20.386	ug/l	98
15) Acrylonitrile	2.674	53	510399	105.726	ug/l	100
16) Acetone	2.118	43	500511	107.135	ug/l	98
17) Carbon Disulfide	2.253	76	445608	19.629	ug/l	99
18) Methyl Acetate	2.381	43	236287	22.381	ug/l	97
19) Methyl tert-butyl Ether	2.712	73	494520	20.907	ug/l	96
20) Methylene Chloride	2.458	84	185787	22.678	ug/l	97
21) trans-1,2-Dichloroethene	2.703	96	155320	19.655	ug/l	98
22) Diisopropyl ether	3.217	45	514292	20.273	ug/l	85
23) Vinyl Acetate	3.192	43	2297382	103.522	ug/l	100
24) 1,1-Dichloroethane	3.118	63	304665	19.638	ug/l	99
25) 2-Butanone	3.870	43	603602	105.789	ug/l	98
26) 2,2-Dichloropropane	3.812	77	246178	17.897	ug/l	100
27) cis-1,2-Dichloroethene	3.822	96	170071	19.514	ug/l	100
28) Bromochloromethane	4.146	49	138833	20.271	ug/l	96
29) Tetrahydrofuran	4.240	42	389964	109.192	ug/l	100
30) Chloroform	4.278	83	322270	19.894	ug/l	99
31) Cyclohexane	4.584	56	238107	18.736	ug/l	99
32) 1,1,1-Trichloroethane	4.513	97	279806	19.951	ug/l	97
36) 1,1-Dichloropropene	4.745	75	187909	19.432	ug/l	98
37) Ethyl Acetate	3.982	43	243790	20.776	ug/l #	92
38) Carbon Tetrachloride	4.735	117	232933	19.709	ug/l	100
39) Methylcyclohexane	6.047	83	228816	19.594	ug/l	96
40) Benzene	5.011	78	633991	20.134	ug/l	97

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039150.D
 Acq On : 24 Sep 2025 14:01
 Operator : SY/MD
 Sample : VV0924WBS02
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 6 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0924WBS02

Quant Time: Sep 25 04:45:44 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.166	41	96391	19.385	ug/l	93
42) 1,2-Dichloroethane	5.040	62	228142	20.677	ug/l	99
43) Isopropyl Acetate	5.195	43	337530	21.097	ug/l	99
44) Trichloroethene	5.831	130	147483	20.297	ug/l	99
45) 1,2-Dichloropropane	6.092	63	163901	20.941	ug/l	96
46) Dibromomethane	6.227	93	126157	21.059	ug/l	99
47) Bromodichloromethane	6.426	83	254692	20.565	ug/l	99
48) Methyl methacrylate	6.301	41	145228	20.025	ug/l	98
49) 1,4-Dioxane	6.294	88	46282	395.445	ug/l	91
51) 4-Methyl-2-Pentanone	7.150	43	1236058	117.436	ug/l	98
52) Toluene	7.310	92	398034	21.801	ug/l	99
53) t-1,3-Dichloropropene	7.577	75	233157	21.603	ug/l	96
54) cis-1,3-Dichloropropene	6.950	75	256176	21.501	ug/l	96
55) 1,1,2-Trichloroethane	7.764	97	171861	20.993	ug/l	97
56) Ethyl methacrylate	7.719	69	198833	20.452	ug/l	100
57) 1,3-Dichloropropane	7.934	76	277597	21.486	ug/l	97
58) 2-Chloroethyl Vinyl ether	6.815	63	517964	100.347	ug/l	99
59) 2-Hexanone	8.066	43	894828	106.669	ug/l	98
60) Dibromochloromethane	8.166	129	191297	20.508	ug/l	99
61) 1,2-Dibromoethane	8.275	107	179580	21.450	ug/l	99
64) Tetrachloroethene	7.899	164	132876	20.213	ug/l	98
65) Chlorobenzene	8.805	112	432358	19.777	ug/l	97
66) 1,1,1,2-Tetrachloroethane	8.899	131	156609	19.826	ug/l	100
67) Ethyl Benzene	8.937	91	658467	19.898	ug/l	100
68) m/p-Xylenes	9.063	106	551462	43.176	ug/l	99
69) o-Xylene	9.468	106	230630	20.393	ug/l	98
70) Styrene	9.487	104	431651	18.940	ug/l	98
71) Bromoform	9.654	173	131945	21.384	ug/l #	98
73) Isopropylbenzene	9.857	105	671517	19.761	ug/l	100
74) N-amyl acetate	9.725	43	283249	20.897	ug/l	99
75) 1,1,2,2-Tetrachloroethane	10.166	83	286264	19.226	ug/l	100
76) 1,2,3-Trichloropropane	10.198	75	206903	19.396	ug/l	100
77) Bromobenzene	10.140	156	173461	19.674	ug/l	99
78) n-propylbenzene	10.278	91	836225	20.207	ug/l	100
79) 2-Chlorotoluene	10.349	91	511892	20.673	ug/l	98
80) 1,3,5-Trimethylbenzene	10.465	105	583511	20.659	ug/l	99
81) trans-1,4-Dichloro-2-b...	9.931	75	86872	19.550	ug/l	99
82) 4-Chlorotoluene	10.465	91	594451	20.703	ug/l	100
83) tert-Butylbenzene	10.789	119	525337	18.652	ug/l	100
84) 1,2,4-Trimethylbenzene	10.841	105	564955	20.602	ug/l	100
85) sec-Butylbenzene	11.014	105	728021	19.488	ug/l	100
86) p-Isopropyltoluene	11.169	119	612071	19.685	ug/l	100
87) 1,3-Dichlorobenzene	11.108	146	349017	19.560	ug/l	100
88) 1,4-Dichlorobenzene	11.198	146	375846	19.199	ug/l	99
89) n-Butylbenzene	11.580	91	577604	19.451	ug/l	98
90) Hexachloroethane	11.821	117	135944	17.755	ug/l	99
91) 1,2-Dichlorobenzene	11.567	146	316105	18.962	ug/l	100
92) 1,2-Dibromo-3-Chloropr...	12.352	75	56358	21.271	ug/l	98
93) 1,2,4-Trichlorobenzene	13.185	180	178967	18.728	ug/l	99
94) Hexachlorobutadiene	13.368	225	89231	17.970	ug/l	99
95) Naphthalene	13.426	128	578144	18.718	ug/l	100
96) 1,2,3-Trichlorobenzene	13.667	180	187044	19.137	ug/l	100

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039150.D
Acq On : 24 Sep 2025 14:01
Operator : SY/MD
Sample : VV0924WBS02
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 6 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VV0924WBS02

Quant Time: Sep 25 04:45:44 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VV0924WBS02

[illegible]

Report of Analysis

Client:	ATG-GREENVILLE AEC	Date Collected:	
Project:	AEC-2025-0013 CSC 4151 Coram	Date Received:	
Client Sample ID:	VV0925WBS01	SDG No.:	Q3164
Lab Sample ID:	VV0925WBS01	Matrix:	Water
Analytical Method:	8260D	% Solid:	0
Sample Wt/Vol:	5 Units: mL	Final Vol:	5000 uL
Soil Aliquot Vol:	uL	Test:	VOC-BTEX
GC Column:	DB-624UI ID : 0.18	Level :	LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039171.D	1	09/25/25 11:24	VV092525

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	19.9		0.15	1.00	ug/L
108-88-3	Toluene	21.5		0.14	1.00	ug/L
100-41-4	Ethyl Benzene	20.0		0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	43.1		0.24	2.00	ug/L
95-47-6	o-Xylene	21.0		0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	45.1		74 - 125	90%	SPK: 50
1868-53-7	Dibromofluoromethane	47.8		75 - 124	96%	SPK: 50
2037-26-5	Toluene-d8	47.9		86 - 113	96%	SPK: 50
460-00-4	4-Bromofluorobenzene	51.2		77 - 121	102%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	557000	4.6			
540-36-3	1,4-Difluorobenzene	899000	5.535			
3114-55-4	Chlorobenzene-d5	872000	8.776			
3855-82-1	1,4-Dichlorobenzene-d4	509000	11.172			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039171.D
 Acq On : 25 Sep 2025 11:24
 Operator : SY/MD
 Sample : VV0925WBS01
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 5 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0925WBS01

Quant Time: Sep 26 01:21:22 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.600	168	557017	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.535	114	898902	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	871912	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	508947	50.000	ug/l	0.00

System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.944	65	363796	45.127	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	90.260%
35) Dibromofluoromethane	4.500	113	345732	47.843	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	95.680%
50) Toluene-d8	7.236	98	1017460	47.939	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	95.880%
62) 4-Bromofluorobenzene	10.001	95	447073	51.168	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	102.340%

Target Compounds				Qvalue		
2) Dichlorodifluoromethane	1.121	85	160404	18.191	ug/l	99
3) Chloromethane	1.230	50	180629	19.082	ug/l	99
4) Vinyl Chloride	1.298	62	196152	19.356	ug/l	99
5) Bromomethane	1.497	94	119487	18.704	ug/l	98
6) Chloroethane	1.561	64	122694	19.279	ug/l	97
7) Trichlorofluoromethane	1.725	101	284580	19.396	ug/l	98
8) Diethyl Ether	1.918	74	106026	19.466	ug/l	99
9) 1,1,2-Trichlorotrifluo...	2.082	101	174658	20.007	ug/l	97
10) Methyl Iodide	2.198	142	135972	16.356	ug/l	99
11) Tert butyl alcohol	2.571	59	187721	110.174	ug/l	99
12) 1,1-Dichloroethene	2.082	96	166976	19.882	ug/l	98
13) Acrolein	2.011	56	34025	115.083	ug/l	98
14) Allyl chloride	2.359	41	286242	19.080	ug/l	98
15) Acrylonitrile	2.677	53	538540	98.597	ug/l	100
16) Acetone	2.121	43	476447	89.017	ug/l	97
17) Carbon Disulfide	2.253	76	487416	18.977	ug/l	99
18) Methyl Acetate	2.381	43	252836	21.167	ug/l	98
19) Methyl tert-butyl Ether	2.716	73	535106	19.995	ug/l	99
20) Methylene Chloride	2.458	84	192332	20.555	ug/l	95
21) trans-1,2-Dichloroethene	2.703	96	176159	19.703	ug/l	96
22) Diisopropyl ether	3.220	45	549959	19.161	ug/l	99
23) Vinyl Acetate	3.191	43	2415006	96.182	ug/l	98
24) 1,1-Dichloroethane	3.117	63	339160	19.322	ug/l	98
25) 2-Butanone	3.873	43	626977	97.122	ug/l	99
26) 2,2-Dichloropropane	3.812	77	266277	17.109	ug/l	98
27) cis-1,2-Dichloroethene	3.822	96	191963	19.468	ug/l	98
28) Bromochloromethane	4.150	49	158654	20.475	ug/l	97
29) Tetrahydrofuran	4.240	42	407358	100.814	ug/l	98
30) Chloroform	4.281	83	352541	19.234	ug/l	97
31) Cyclohexane	4.587	56	262018	18.223	ug/l	94
32) 1,1,1-Trichloroethane	4.513	97	301922	19.027	ug/l	98
36) 1,1-Dichloropropene	4.744	75	210259	19.244	ug/l	98
37) Ethyl Acetate	3.986	43	249892	18.849	ug/l	99
38) Carbon Tetrachloride	4.735	117	258780	19.380	ug/l	96
39) Methylcyclohexane	6.047	83	253234	19.193	ug/l	98
40) Benzene	5.011	78	706824	19.867	ug/l	98

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
 Data File : VV039171.D
 Acq On : 25 Sep 2025 11:24
 Operator : SY/MD
 Sample : VV0925WBS01
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 5 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0925WBS01

Quant Time: Sep 26 01:21:22 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.166	41	101017	18.119	ug/l	92
42) 1,2-Dichloroethane	5.043	62	243861	19.562	ug/l	99
43) Isopropyl Acetate	5.198	43	354785	19.627	ug/l	99
44) Trichloroethene	5.834	130	167082	20.352	ug/l	99
45) 1,2-Dichloropropane	6.092	63	181382	20.511	ug/l	96
46) Dibromomethane	6.227	93	135579	20.031	ug/l	98
47) Bromodichloromethane	6.429	83	269985	19.295	ug/l	100
48) Methyl methacrylate	6.301	41	154615	18.869	ug/l	97
49) 1,4-Dioxane	6.294	88	55817	422.109	ug/l	98
51) 4-Methyl-2-Pentanone	7.149	43	1295291	108.921	ug/l	99
52) Toluene	7.310	92	442966	21.474	ug/l	98
53) t-1,3-Dichloropropene	7.577	75	244520	20.052	ug/l	99
54) cis-1,3-Dichloropropene	6.950	75	272587	20.249	ug/l	98
55) 1,1,2-Trichloroethane	7.764	97	184964	19.997	ug/l	95
56) Ethyl methacrylate	7.719	69	220741	20.096	ug/l	97
57) 1,3-Dichloropropane	7.937	76	293432	20.102	ug/l	99
58) 2-Chloroethyl Vinyl ether	6.815	63	541689	93.599	ug/l	99
59) 2-Hexanone	8.066	43	942254	99.572	ug/l	99
60) Dibromochloromethane	8.169	129	212792	20.190	ug/l	100
61) 1,2-Dibromoethane	8.275	107	195149	20.631	ug/l	99
64) Tetrachloroethene	7.899	164	147205	20.157	ug/l	95
65) Chlorobenzene	8.805	112	488359	20.108	ug/l	98
66) 1,1,1,2-Tetrachloroethane	8.899	131	174648	19.902	ug/l	100
67) Ethyl Benzene	8.937	91	736424	20.032	ug/l	99
68) m/p-Xylenes	9.063	106	611502	43.097	ug/l	100
69) o-Xylene	9.468	106	263271	20.955	ug/l	99
70) Styrene	9.487	104	489732	19.311	ug/l	99
71) Bromoform	9.654	173	140287	20.466	ug/l #	98
73) Isopropylbenzene	9.857	105	759505	20.575	ug/l	100
74) N-amyl acetate	9.725	43	297254	20.188	ug/l	99
75) 1,1,2,2-Tetrachloroethane	10.165	83	313809	19.402	ug/l	100
76) 1,2,3-Trichloropropane	10.198	75	227880	19.665	ug/l	98
77) Bromobenzene	10.143	156	194148	20.272	ug/l	97
78) n-propylbenzene	10.278	91	917928	20.420	ug/l	99
79) 2-Chlorotoluene	10.349	91	569887	21.187	ug/l	98
80) 1,3,5-Trimethylbenzene	10.464	105	649406	21.165	ug/l	100
81) trans-1,4-Dichloro-2-b...	9.931	75	93631	19.398	ug/l	98
82) 4-Chlorotoluene	10.464	91	658860	21.124	ug/l	100
83) tert-Butylbenzene	10.789	119	599275	19.587	ug/l	99
84) 1,2,4-Trimethylbenzene	10.841	105	634517	21.301	ug/l	100
85) sec-Butylbenzene	11.014	105	814764	20.078	ug/l	99
86) p-Isopropyltoluene	11.169	119	692579	20.505	ug/l	99
87) 1,3-Dichlorobenzene	11.108	146	380476	19.629	ug/l	100
88) 1,4-Dichlorobenzene	11.198	146	400938	18.854	ug/l	99
89) n-Butylbenzene	11.580	91	629906	19.528	ug/l	98
90) Hexachloroethane	11.821	117	153004	18.396	ug/l	99
91) 1,2-Dichlorobenzene	11.567	146	359391	19.846	ug/l	99
92) 1,2-Dibromo-3-Chloropr...	12.352	75	63577	22.105	ug/l	98
93) 1,2,4-Trichlorobenzene	13.185	180	198844	19.155	ug/l	98
94) Hexachlorobutadiene	13.368	225	100713	18.672	ug/l	99
95) Naphthalene	13.426	128	654241	19.499	ug/l	99
96) 1,2,3-Trichlorobenzene	13.667	180	210809	19.856	ug/l	99

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092525\
Data File : VV039171.D
Acq On : 25 Sep 2025 11:24
Operator : SY/MD
Sample : VV0925WBS01
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 5 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VV0925WBS01

Quant Time: Sep 26 01:21:22 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

Instrument :
MSVOA_V
ClientSampleId :
VV0925WBS01

[illegible]

Report of Analysis

Client:	ATG-GREENVILLE AEC			Date Collected:	
Project:	AEC-2025-0013 CSC 4151 Coram			Date Received:	
Client Sample ID:	VV0924WBSD02			SDG No.:	Q3164
Lab Sample ID:	VV0924WBSD02			Matrix:	Water
Analytical Method:	8260D			% Solid:	0
Sample Wt/Vol:	5	Units:	mL	Final Vol:	5000 uL
Soil Aliquot Vol:			uL	Test:	VOC-BTEX
GC Column:	DB-624UI	ID :	0.18	Level :	LOW
Prep Method :					

File ID/Qc Batch:	Dilution:	Date Analyzed	Prep Batch ID
VV039165.D	1	09/24/25 19:59	VV092425

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	18.8		0.15	1.00	ug/L
108-88-3	Toluene	20.0		0.14	1.00	ug/L
100-41-4	Ethyl Benzene	19.2		0.13	1.00	ug/L
179601-23-1	m/p-Xylenes	40.4		0.24	2.00	ug/L
95-47-6	o-Xylene	19.8		0.12	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	49.5		74 - 125	99%	SPK: 50
1868-53-7	Dibromofluoromethane	50.1		75 - 124	100%	SPK: 50
2037-26-5	Toluene-d8	50.7		86 - 113	101%	SPK: 50
460-00-4	4-Bromofluorobenzene	55.0		77 - 121	110%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	545000	4.6			
540-36-3	1,4-Difluorobenzene	904000	5.532			
3114-55-4	Chlorobenzene-d5	885000	8.776			
3855-82-1	1,4-Dichlorobenzene-d4	511000	11.172			

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

E = Value Exceeds Calibration Range

Q = indicates LCS control criteria did not meet requirements

M = MS/MSD acceptance criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039165.D
 Acq On : 24 Sep 2025 19:59
 Operator : SY/MD
 Sample : VV0924WBSD02
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 22 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0924WBSD02

Quant Time: Sep 25 08:24:33 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)

Internal Standards						
1) Pentafluorobenzene	4.600	168	545364	50.000	ug/l	0.00
34) 1,4-Difluorobenzene	5.532	114	903558	50.000	ug/l	0.00
63) Chlorobenzene-d5	8.776	117	885424	50.000	ug/l	0.00
72) 1,4-Dichlorobenzene-d4	11.172	152	510716	50.000	ug/l	0.00

System Monitoring Compounds						
33) 1,2-Dichloroethane-d4	4.944	65	390603	49.487	ug/l	0.00
Spiked Amount	50.000	Range	81 - 118	Recovery	=	98.980%
35) Dibromofluoromethane	4.503	113	363843	50.089	ug/l	0.00
Spiked Amount	50.000	Range	80 - 119	Recovery	=	100.180%
50) Toluene-d8	7.236	98	1080761	50.659	ug/l	0.00
Spiked Amount	50.000	Range	89 - 112	Recovery	=	101.320%
62) 4-Bromofluorobenzene	10.001	95	483167	55.014	ug/l	0.00
Spiked Amount	50.000	Range	85 - 114	Recovery	=	110.020%

Target Compounds				Qvalue		
2) Dichlorodifluoromethane	1.121	85	156320	18.107	ug/l	98
3) Chloromethane	1.230	50	168337	18.164	ug/l	100
4) Vinyl Chloride	1.298	62	180006	18.142	ug/l	99
5) Bromomethane	1.497	94	91963	14.703	ug/l	99
6) Chloroethane	1.561	64	124416	20.020	ug/l	100
7) Trichlorofluoromethane	1.725	101	271403	18.893	ug/l	99
8) Diethyl Ether	1.918	74	98678	18.504	ug/l	98
9) 1,1,2-Trichlorotrifluo...	2.082	101	162784	19.045	ug/l	98
10) Methyl Iodide	2.198	142	97066	11.925	ug/l	99
11) Tert butyl alcohol	2.568	59	177476	106.387	ug/l	97
12) 1,1-Dichloroethene	2.082	96	159305	19.374	ug/l	92
13) Acrolein	2.011	56	32659	112.823	ug/l	95
14) Allyl chloride	2.359	41	262106	17.845	ug/l	98
15) Acrylonitrile	2.677	53	500749	93.637	ug/l	99
16) Acetone	2.121	43	420916	79.697	ug/l	98
17) Carbon Disulfide	2.253	76	461436	18.349	ug/l	99
18) Methyl Acetate	2.381	43	235110	20.104	ug/l	99
19) Methyl tert-butyl Ether	2.712	73	506464	19.329	ug/l	99
20) Methylene Chloride	2.458	84	191133	20.897	ug/l	98
21) trans-1,2-Dichloroethene	2.703	96	163017	18.623	ug/l	99
22) Diisopropyl ether	3.217	45	524404	18.661	ug/l	84
23) Vinyl Acetate	3.191	43	2227046	90.591	ug/l	99
24) 1,1-Dichloroethane	3.117	63	311443	18.122	ug/l	99
25) 2-Butanone	3.873	43	581279	91.967	ug/l	99
26) 2,2-Dichloropropane	3.812	77	223757	14.684	ug/l	99
27) cis-1,2-Dichloroethene	3.822	96	181034	18.752	ug/l	99
28) Bromochloromethane	4.150	49	145773	19.214	ug/l	95
29) Tetrahydrofuran	4.243	42	385403	97.419	ug/l	99
30) Chloroform	4.278	83	330400	18.412	ug/l	97
31) Cyclohexane	4.584	56	258917	18.392	ug/l	96
32) 1,1,1-Trichloroethane	4.513	97	285397	18.370	ug/l	98
36) 1,1-Dichloropropene	4.748	75	205485	18.711	ug/l	98
37) Ethyl Acetate	3.982	43	238455	17.894	ug/l	98
38) Carbon Tetrachloride	4.735	117	243831	18.166	ug/l	98
39) Methylcyclohexane	6.050	83	241644	18.220	ug/l	97
40) Benzene	5.011	78	673840	18.842	ug/l	99

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
 Data File : VV039165.D
 Acq On : 24 Sep 2025 19:59
 Operator : SY/MD
 Sample : VV0924WBSD02
 Misc : 5.0mL/MSVOA_V/WATER
 ALS Vial : 22 Sample Multiplier: 1

Instrument :
 MSVOA_V
 ClientSampleId :
 VV0924WBSD02

Quant Time: Sep 25 08:24:33 2025
 Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
 Quant Title : SW846 8260
 QLast Update : Wed Sep 24 08:08:08 2025
 Response via : Initial Calibration

Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
41) Methacrylonitrile	4.166	41	96133	17.256	ug/l	93
42) 1,2-Dichloroethane	5.043	62	232378	18.545	ug/l	100
43) Isopropyl Acetate	5.198	43	339665	18.694	ug/l	99
44) Trichloroethene	5.831	130	158174	19.167	ug/l	100
45) 1,2-Dichloropropane	6.092	63	172229	19.376	ug/l	97
46) Dibromomethane	6.230	93	128940	18.952	ug/l	97
47) Bromodichloromethane	6.429	83	257271	18.292	ug/l	97
48) Methyl methacrylate	6.301	41	148719	18.056	ug/l	99
49) 1,4-Dioxane	6.297	88	52205	392.759	ug/l	94
51) 4-Methyl-2-Pentanone	7.149	43	1220207	102.079	ug/l	99
52) Toluene	7.310	92	415722	20.049	ug/l	100
53) t-1,3-Dichloropropene	7.577	75	226898	18.511	ug/l	99
54) cis-1,3-Dichloropropene	6.950	75	259523	19.179	ug/l	98
55) 1,1,2-Trichloroethane	7.764	97	171629	18.460	ug/l	97
56) Ethyl methacrylate	7.719	69	204966	18.564	ug/l	98
57) 1,3-Dichloropropane	7.934	76	277934	18.942	ug/l	100
58) 2-Chloroethyl Vinyl ether	6.815	63	486078	84.589	ug/l	99
59) 2-Hexanone	8.066	43	867936	91.441	ug/l	99
60) Dibromochloromethane	8.169	129	195919	18.494	ug/l	100
61) 1,2-Dibromoethane	8.275	107	183626	19.312	ug/l	99
64) Tetrachloroethene	7.899	164	146324	19.730	ug/l	98
65) Chlorobenzene	8.805	112	491329	19.922	ug/l	99
66) 1,1,1,2-Tetrachloroethane	8.899	131	162674	18.255	ug/l	100
67) Ethyl Benzene	8.937	91	716157	19.183	ug/l	99
68) m/p-Xylenes	9.063	106	582336	40.415	ug/l	99
69) o-Xylene	9.468	106	252664	19.804	ug/l	98
70) Styrene	9.487	104	457807	17.897	ug/l	99
71) Bromoform	9.654	173	131160	18.843	ug/l #	99
73) Isopropylbenzene	9.857	105	724475	19.558	ug/l	99
74) N-amyl acetate	9.725	43	279661	18.927	ug/l	99
75) 1,1,2,2-Tetrachloroethane	10.165	83	290148	17.877	ug/l	100
76) 1,2,3-Trichloropropane	10.198	75	219335	18.862	ug/l	100
77) Bromobenzene	10.140	156	183095	19.051	ug/l	97
78) n-propylbenzene	10.278	91	881210	19.535	ug/l	100
79) 2-Chlorotoluene	10.349	91	539084	19.972	ug/l	99
80) 1,3,5-Trimethylbenzene	10.464	105	620336	20.148	ug/l	99
81) trans-1,4-Dichloro-2-b...	9.931	75	82955	17.126	ug/l	98
82) 4-Chlorotoluene	10.464	91	626744	20.025	ug/l	100
83) tert-Butylbenzene	10.789	119	563652	18.359	ug/l	100
84) 1,2,4-Trimethylbenzene	10.841	105	604069	20.209	ug/l	99
85) sec-Butylbenzene	11.014	105	783211	19.233	ug/l	100
86) p-Isopropyltoluene	11.165	119	661055	19.504	ug/l	100
87) 1,3-Dichlorobenzene	11.104	146	362792	18.652	ug/l	99
88) 1,4-Dichlorobenzene	11.198	146	381649	17.885	ug/l	98
89) n-Butylbenzene	11.580	91	608918	18.812	ug/l	98
90) Hexachloroethane	11.818	117	138165	16.554	ug/l	99
91) 1,2-Dichlorobenzene	11.567	146	335832	18.481	ug/l	100
92) 1,2-Dibromo-3-Chloropr...	12.352	75	57082	19.730	ug/l	99
93) 1,2,4-Trichlorobenzene	13.185	180	191728	18.406	ug/l	100
94) Hexachlorobutadiene	13.368	225	91410	16.888	ug/l	99
95) Naphthalene	13.426	128	623030	18.505	ug/l	100
96) 1,2,3-Trichlorobenzene	13.667	180	199614	18.736	ug/l	98

Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\
Data File : VV039165.D
Acq On : 24 Sep 2025 19:59
Operator : SY/MD
Sample : VV0924WBSD02
Misc : 5.0mL/MSVOA_V/WATER
ALS Vial : 22 Sample Multiplier: 1

Instrument :
MSVOA_V
ClientSampleId :
VV0924WBSD02

Quant Time: Sep 25 08:24:33 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration

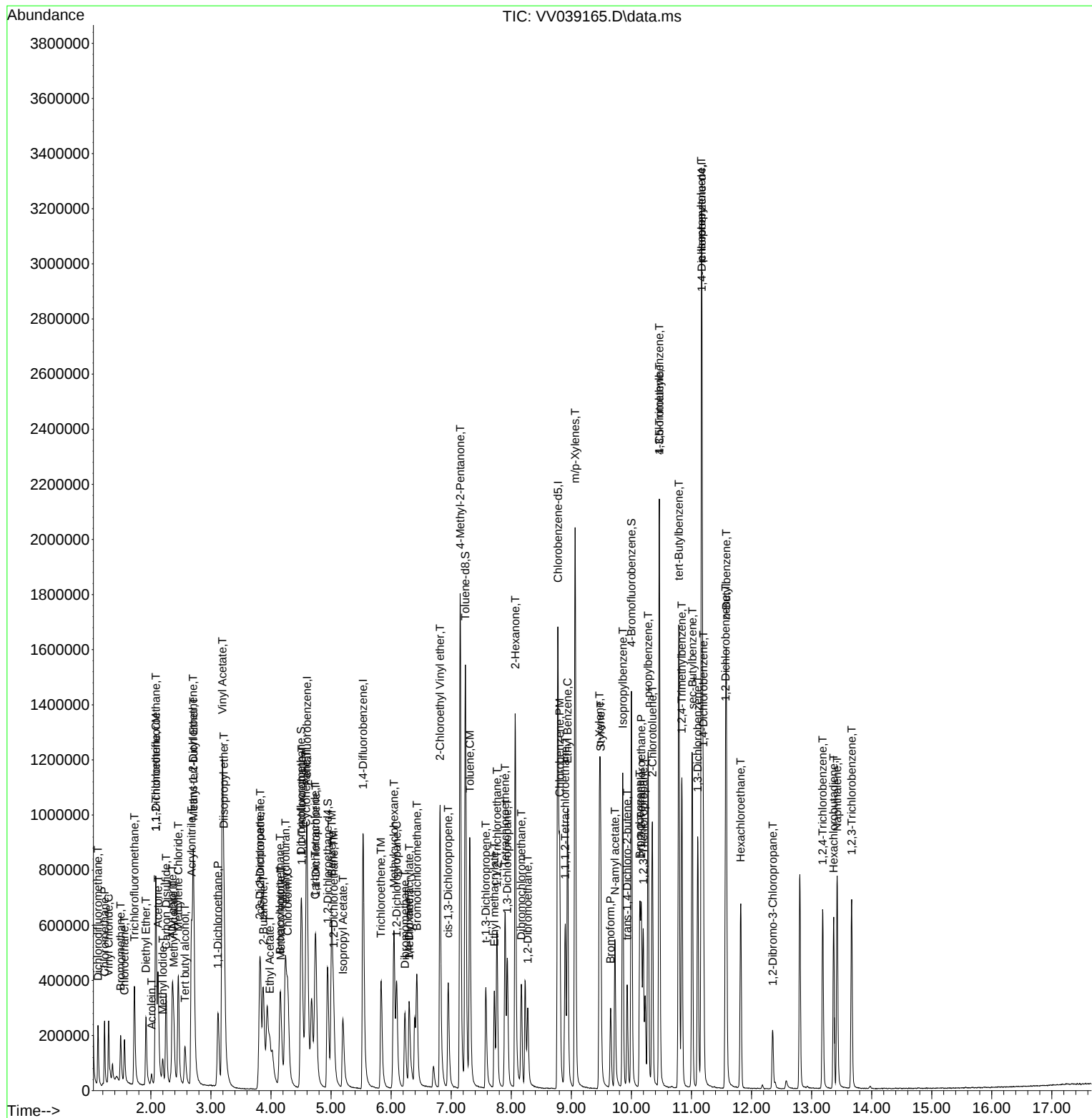
Compound	R.T.	QIon	Response	Conc	Units	Dev(Min)
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(#) = qualifier out of range (m) = manual integration (+) = signals summed

```
Data Path : Z:\voasrv\HPCHEM1\MSVOA_V\DATA\VV092425\  
Data File : VV039165.D  
Acq On    : 24 Sep 2025   19:59  
Operator  : SY/MD  
Sample    : VV0924WBSD02  
Misc      : 5.0mL/MSVOA_V/WATER  
ALS Vial  : 22   Sample Multiplier: 1
```

Instrument :
MSVOA_V
ClientSampleId :
VV0924WBSD02

Quant Time: Sep 25 08:24:33 2025
Quant Method : Z:\voasrv\HPCHEM1\MSVOA_V\Method\82V092225W.M
Quant Title : SW846 8260
QLast Update : Wed Sep 24 08:08:08 2025
Response via : Initial Calibration



Manual Integration Report

Sequence:	VV092225	Instrument	MSVOA_v
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Sample ID	File ID	Parameter	Review By	Review On	Supervised By	Supervised On	Reason
VSTDICC005	VV039142.D	1,2,3-Trichloropropane	MMDadod a	9/26/2025 1:38:50 AM	SAM	9/26/2025 1:57:11 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	1,1-Dichloropropene	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	1,4-Dichlorobenzene	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	1,4-Dioxane	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	2,2-Dichloropropane	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	2-Butanone	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	Carbon Tetrachloride	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	Ethyl Acetate	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	Ethyl methacrylate	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	Isopropyl Acetate	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	Methacrylonitrile	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	Methyl methacrylate	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software
VSTDICC001	VV039143.D	Naphthalene	MMDadod a	9/26/2025 1:38:52 AM	SAM	9/26/2025 1:57:13 AM	Peak Integrated by Software



284 Sheffield Street, Mountainside, New Jersey 07092, Phone : 908 789 8900, Fax : 908 789 8922

Manual Integration Report

Sequence:	VV092225	Instrument	MSVOA_v
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Sample ID	File ID	Parameter	Review By	Review On	Supervised By	Supervised On	Reason
VSTDICV050	VV039144.D	1,2,3-Trichloropropane	MMDadoda	9/26/2025 1:38:55 AM	SAM	9/26/2025 1:57:15 AM	Peak Integrated by Software



284 Sheffield Street, Mountainside, New Jersey 07092, Phone : 908 789 8900, Fax : 908 789 8922

Manual Integration Report

Sequence:

VV092425

Instrument

MSVOA_v

Sample ID

File ID

Parameter

Review By

Review On

Supervised
By

Supervised On

Reason



284 Sheffield Street, Mountainside, New Jersey 07092, Phone : 908 789 8900, Fax : 908 789 8922

Manual Integration Report

Sequence:

VV092525

Instrument

MSVOA_v

Sample ID

File ID

Parameter

Review By

Review On

Supervised
By

Supervised On

Reason

Instrument ID: **MSVOA_V**

Daily Analysis Runlog For Sequence/QCBatch ID # VV092225

Review By	Mahesh Dadoda	Review On	9/26/2025 1:39:32 AM		
Supervise By	Semsettin Yesilyurt	Supervise On	9/26/2025 1:56:56 AM		
SubDirectory	VV092225	HP Acquire Method	MSVOA_V	HP Processing Method	82v092225w.m
STD. NAME		STD REF.#			
Tune/Reschk		VP135580			
Initial Calibration Stds		VP135581,VP135582,VP135583,VP135584,VP135585,VP135586			
CCC					
Internal Standard/PEM		VP133935			
ICV/I.BLK		VP135587			
Surrogate Standard					
MS/MSD Standard					
LCS Standard					

Sr#	SampleId	Data File Name	Date-Time	Operator	Status
1	BFB	VV039134.D	22 Sep 2025 10:15	SY/MD	Ok
2	VSTDICC001	VV039135.D	22 Sep 2025 10:59	SY/MD	Not Ok
3	VSTDICC005	VV039136.D	22 Sep 2025 11:30	SY/MD	Not Ok
4	VSTDICC010	VV039137.D	22 Sep 2025 12:26	SY/MD	Ok
5	VSTDICC020	VV039138.D	22 Sep 2025 12:48	SY/MD	Ok
6	VSTDICCC050	VV039139.D	22 Sep 2025 13:26	SY/MD	Ok
7	VSTDICC100	VV039140.D	22 Sep 2025 13:49	SY/MD	Ok
8	VIBLK	VV039141.D	22 Sep 2025 14:21	SY/MD	Ok
9	VSTDICC005	VV039142.D	22 Sep 2025 15:37	SY/MD	Ok,M
10	VSTDICC001	VV039143.D	22 Sep 2025 16:35	SY/MD	Ok,M
11	VSTDICV050	VV039144.D	22 Sep 2025 16:58	SY/MD	Ok,M

M : Manual Integration

Instrument ID: **MSVOA_V**

Daily Analysis Runlog For Sequence/QC Batch ID # VV092425

Review By	Mahesh Dadoda	Review On	9/26/2025 1:45:06 AM
Supervise By	Semsettin Yesilyurt	Supervise On	9/26/2025 2:03:33 AM
SubDirectory	VV092425	HP Acquire Method	MSVOA_V
		HP Processing Method	
STD. NAME	STD REF.#		
Tune/Reschk Initial Calibration Stds	VP135622		
CCC Internal Standard/PEM ICV/I.BLK Surrogate Standard MS/MSD Standard LCS Standard	VP135623,VP135624 VP133935		

Sr#	SampleId	Data File Name	Date-Time	Operator	Status
1	BFB	VV039145.D	24 Sep 2025 09:52	SY/MD	Ok
2	VSTDCCC050	VV039146.D	24 Sep 2025 10:49	SY/MD	Ok
3	VV0924MBL01	VV039147.D	24 Sep 2025 12:08	SY/MD	Ok
4	VV0924WBL01	VV039148.D	24 Sep 2025 13:09	SY/MD	Ok
5	VV0924WBS01	VV039149.D	24 Sep 2025 13:37	SY/MD	Not Ok
6	VV0924WBS02	VV039150.D	24 Sep 2025 14:01	SY/MD	Ok
7	Q3173-02	VV039151.D	24 Sep 2025 14:46	SY/MD	Ok
8	Q3173-01	VV039152.D	24 Sep 2025 15:08	SY/MD	Ok
9	Q3175-01	VV039153.D	24 Sep 2025 15:31	SY/MD	Dilution
10	Q3175-02	VV039154.D	24 Sep 2025 15:53	SY/MD	Dilution
11	VIBLK	VV039155.D	24 Sep 2025 16:16	SY/MD	Ok
12	VIBLK	VV039156.D	24 Sep 2025 16:38	SY/MD	Ok
13	Q3172-01	VV039157.D	24 Sep 2025 17:00	SY/MD	Dilution
14	Q3172-02	VV039158.D	24 Sep 2025 17:22	SY/MD	Dilution
15	Q3172-03	VV039159.D	24 Sep 2025 17:45	SY/MD	ReRun
16	Q3164-01	VV039160.D	24 Sep 2025 18:07	SY/MD	Ok
17	Q3164-02	VV039161.D	24 Sep 2025 18:30	SY/MD	Not Ok
18	Q3164-03	VV039162.D	24 Sep 2025 18:52	SY/MD	Ok
19	VIBLK	VV039163.D	24 Sep 2025 19:15	SY/MD	Ok
20	VIBLK	VV039164.D	24 Sep 2025 19:37	SY/MD	Ok,M
21	VV0924WBSD02	VV039165.D	24 Sep 2025 19:59	SY/MD	Ok

Instrument ID: **MSVOA_V**

Daily Analysis Runlog For Sequence/QC Batch ID # VV092425

Review By	Mahesh Dadoda	Review On	9/26/2025 1:45:06 AM	
Supervise By	Semsettin Yesilyurt	Supervise On	9/26/2025 2:03:33 AM	
SubDirectory	VV092425	HP Acquire Method	MSVOA_V	HP Processing Method
STD. NAME	STD REF.#			
Tune/Reschk Initial Calibration Stds	VP135622			
CCC	VP135623,VP135624			
Internal Standard/PEM	VP133935			
ICV/I.BLK				
Surrogate Standard				
MS/MSD Standard				
LCS Standard				

22	VSTDCCC050	VV039166.D	24 Sep 2025 20:22	SY/MD	Ok
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M : Manual Integration

Instrument ID: **MSVOA_V**

Daily Analysis Runlog For Sequence/QC Batch ID # VV092525

Review By	Mahesh Dadoda	Review On	9/29/2025 12:56:22 AM
Supervise By	Semsettin Yesilyurt	Supervise On	9/29/2025 1:08:36 AM
SubDirectory	VV092525	HP Acquire Method	8260_V
		HP Processing Method	82v092225w.m
STD. NAME	STD REF.#		
Tune/Reschk Initial Calibration Stds	VP135649		
CCC Internal Standard/PEM ICV/I.BLK Surrogate Standard MS/MSD Standard LCS Standard	VP135650,VP135651 VP133935		

Sr#	SampleId	Data File Name	Date-Time	Operator	Status
1	BFB	VV039167.D	25 Sep 2025 09:14	SY/MD	Ok
2	VSTDCCC050	VV039168.D	25 Sep 2025 09:59	SY/MD	Ok
3	VV0925MBL01	VV039169.D	25 Sep 2025 10:31	SY/MD	Ok
4	VV0925WBL01	VV039170.D	25 Sep 2025 11:01	SY/MD	Ok
5	VV0925WBS01	VV039171.D	25 Sep 2025 11:24	SY/MD	Ok
6	VV0925WBSD01	VV039172.D	25 Sep 2025 11:48	SY/MD	Ok
7	Q3164-02	VV039173.D	25 Sep 2025 12:10	SY/MD	Ok
8	Q3175-01DL	VV039174.D	25 Sep 2025 12:33	SY/MD	Dilution
9	Q3175-02DL	VV039175.D	25 Sep 2025 12:55	SY/MD	Ok,M
10	VIBLK	VV039176.D	25 Sep 2025 13:18	SY/MD	Ok,M
11	Q3168-05	VV039177.D	25 Sep 2025 13:40	SY/MD	ReRun
12	Q3168-06	VV039178.D	25 Sep 2025 14:03	SY/MD	ReRun
13	Q3175-01DL2	VV039179.D	25 Sep 2025 14:25	SY/MD	Ok,M
14	Q3172-02DL	VV039180.D	25 Sep 2025 14:48	SY/MD	Ok
15	VIBLK	VV039181.D	25 Sep 2025 15:10	SY/MD	Ok
16	Q3172-01DL	VV039182.D	25 Sep 2025 15:33	SY/MD	Ok
17	Q3168-05RE	VV039183.D	25 Sep 2025 15:55	SY/MD	Confirms
18	Q3168-06RE	VV039184.D	25 Sep 2025 16:18	SY/MD	Confirms
19	Q3172-03	VV039185.D	25 Sep 2025 16:40	SY/MD	Ok
20	Q3195-22	VV039186.D	25 Sep 2025 17:03	SY/MD	Ok
21	Q3195-23	VV039187.D	25 Sep 2025 17:25	SY/MD	Ok

Instrument ID: **MSVOA_V**

Daily Analysis Runlog For Sequence/QC Batch ID # VV092525

Review By	Maresh Dadoda	Review On	9/29/2025 12:56:22 AM		
Supervise By	Semsettin Yesilyurt	Supervise On	9/29/2025 1:08:36 AM		
SubDirectory	VV092525	HP Acquire Method	8260_V	HP Processing Method	82v092225w.m
STD. NAME		STD REF.#			
Tune/Reschk Initial Calibration Stds		VP135649			
CCC		VP135650,VP135651			
Internal Standard/PEM		VP133935			
ICV/I.BLK					
Surrogate Standard					
MS/MSD Standard					
LCS Standard					

22	Q3195-24	VV039188.D	25 Sep 2025 17:47	SY/MD	ReRun
23	Q3195-25	VV039189.D	25 Sep 2025 18:10	SY/MD	Ok
24	Q3195-26	VV039190.D	25 Sep 2025 18:32	SY/MD	Ok
25	Q3195-27	VV039191.D	25 Sep 2025 18:55	SY/MD	Ok
26	Q3195-02	VV039192.D	25 Sep 2025 19:17	SY/MD	Ok
27	Q3195-09	VV039193.D	25 Sep 2025 19:40	SY/MD	Ok
28	Q3195-10	VV039194.D	25 Sep 2025 20:02	SY/MD	Ok
29	Q3195-11	VV039195.D	25 Sep 2025 20:25	SY/MD	Ok
30	VSTDCCC050	VV039196.D	25 Sep 2025 20:47	SY/MD	Ok

M : Manual Integration

Instrument ID: MSVOA_V

Daily Analysis Runlog For Sequence/QC Batch ID # VV092225

Review By	Mahesh Dadoda	Review On	9/26/2025 1:39:32 AM		
Supervise By	Semsettin Yesilyurt	Supervise On	9/26/2025 1:56:56 AM		
SubDirectory	VV092225	HP Acquire Method	MSVOA_V	HP Processing Method	82v092225w.m

STD. NAME	STD REF.#
Tune/Reschk	VP135580
Initial Calibration Stds	VP135581,VP135582,VP135583,VP135584,VP135585,VP135586
CCC	
Internal Standard/PEM	VP133935
ICV/I.BLK	VP135587
Surrogate Standard	
MS/MSD Standard	
LCS Standard	

Sr#	Sampleld	ClientID	Data File Name	Date-Time	Comment	Operator	Status
1	BFB	BFB	VV039134.D	22 Sep 2025 10:15		SY/MD	Ok
2	VSTDICC001	VSTDICC001	VV039135.D	22 Sep 2025 10:59	Not use	SY/MD	Not Ok
3	VSTDICC005	VSTDICC005	VV039136.D	22 Sep 2025 11:30	Not use	SY/MD	Not Ok
4	VSTDICC010	VSTDICC010	VV039137.D	22 Sep 2025 12:26		SY/MD	Ok
5	VSTDICC020	VSTDICC020	VV039138.D	22 Sep 2025 12:48	QR- 06,16,20,70,92	SY/MD	Ok
6	VSTDICCC050	VSTDICCC050	VV039139.D	22 Sep 2025 13:26	LR- 41,58,59	SY/MD	Ok
7	VSTDICC100	VSTDICC100	VV039140.D	22 Sep 2025 13:49		SY/MD	Ok
8	VIBLK	VIBLK	VV039141.D	22 Sep 2025 14:21		SY/MD	Ok
9	VSTDICC005	VSTDICC005	VV039142.D	22 Sep 2025 15:37		SY/MD	Ok,M
10	VSTDICC001	VSTDICC001	VV039143.D	22 Sep 2025 16:35		SY/MD	Ok,M
11	VSTDICV050	ICVVV092225	VV039144.D	22 Sep 2025 16:58	Goof for Non-DOD	SY/MD	Ok,M

M : Manual Integration

Instrument ID: MSVOA_V

Daily Analysis Runlog For Sequence/QC Batch ID # VV092425

Review By	Maresh Dadoda	Review On	9/26/2025 1:45:06 AM
Supervise By	Semsettin Yesilyurt	Supervise On	9/26/2025 2:03:33 AM
SubDirectory	VV092425	HP Acquire Method	MSVOA_V HP Processing Method

STD. NAME	STD REF.#
Tune/Reschk Initial Calibration Stds	VP135622
CCC Internal Standard/PEM ICV/I.BLK Surrogate Standard MS/MSD Standard LCS Standard	VP135623,VP135624 VP133935

Sr#	SampleID	ClientID	Data File Name	Date-Time	Comment	Operator	Status
1	BFB	BFB	VV039145.D	24 Sep 2025 09:52		SY/MD	Ok
2	VSTDCCC050	VSTDCCC050	VV039146.D	24 Sep 2025 10:49	pH#lot#v14222	SY/MD	Ok
3	VV0924MBL01	VV0924MBL01	VV039147.D	24 Sep 2025 12:08		SY/MD	Ok
4	VV0924WBL01	VV0924WBL01	VV039148.D	24 Sep 2025 13:09		SY/MD	Ok
5	VV0924WBS01	VV0924WBS01	VV039149.D	24 Sep 2025 13:37	Recovery fail	SY/MD	Not Ok
6	VV0924WBS02	VV0924WBS02	VV039150.D	24 Sep 2025 14:01		SY/MD	Ok
7	Q3173-02	Field Blank	VV039151.D	24 Sep 2025 14:46	pH<2.0 A,FB	SY/MD	Ok
8	Q3173-01	OBS1	VV039152.D	24 Sep 2025 15:08	pH<2.0 A	SY/MD	Ok
9	Q3175-01	MW2	VV039153.D	24 Sep 2025 15:31	pH<2.0 A ,Need 20X	SY/MD	Dilution
10	Q3175-02	MW4	VV039154.D	24 Sep 2025 15:53	pH<2.0 A ,Need 20X	SY/MD	Dilution
11	VIBLK	VIBLK	VV039155.D	24 Sep 2025 16:16		SY/MD	Ok
12	VIBLK	VIBLK	VV039156.D	24 Sep 2025 16:38		SY/MD	Ok
13	Q3172-01	FW7D	VV039157.D	24 Sep 2025 17:00	pH<2.0 A ,Surrogate Fail;Need 100X	SY/MD	Dilution
14	Q3172-02	FW6D	VV039158.D	24 Sep 2025 17:22	pH<2.0 A ,Need 4X	SY/MD	Dilution
15	Q3172-03	GB3W	VV039159.D	24 Sep 2025 17:45	pH<2.0 A ,E flag of previous sample	SY/MD	ReRun
16	Q3164-01	DA-1	VV039160.D	24 Sep 2025 18:07	pH<2.0 A	SY/MD	Ok
17	Q3164-02	DA-2	VV039161.D	24 Sep 2025 18:30	pH<2.0 A ,Confirm Concentration	SY/MD	Not Ok

Instrument ID: MSVOA_V

Daily Analysis Runlog For Sequence/QC Batch ID # VV092425

Review By	Mahesh Dadoda	Review On	9/26/2025 1:45:06 AM
Supervise By	Semsettin Yesilyurt	Supervise On	9/26/2025 2:03:33 AM
SubDirectory	VV092425	HP Acquire Method	MSVOA_V HP Processing Method
STD. NAME	STD REF.#		
Tune/Reschk Initial Calibration Stds	VP135622		
CCC	VP135623,VP135624		
Internal Standard/PEM	VP133935		
ICV/I.BLK			
Surrogate Standard			
MS/MSD Standard			
LCS Standard			

18	Q3164-03	DA-3	VV039162.D	24 Sep 2025 18:52	pH<2.0 A	SY/MD	Ok
19	VIBLK	VIBLK	VV039163.D	24 Sep 2025 19:15		SY/MD	Ok
20	VIBLK	VIBLK	VV039164.D	24 Sep 2025 19:37		SY/MD	Ok,M
21	VV0924WBSD02	VV0924WBSD02	VV039165.D	24 Sep 2025 19:59		SY/MD	Ok
22	VSTDCCC050	VSTDCCC050EC	VV039166.D	24 Sep 2025 20:22		SY/MD	Ok

M : Manual Integration

Instrument ID: MSVOA_V

Daily Analysis Runlog For Sequence/QC Batch ID # VV092525

Review By	Mahesh Dadoda	Review On	9/29/2025 12:56:22 AM		
Supervise By	Semsettin Yesilyurt	Supervise On	9/29/2025 1:08:36 AM		
SubDirectory	VV092525	HP Acquire Method	8260_V	HP Processing Method	82v092225w.m

STD. NAME	STD REF.#
Tune/Reschk Initial Calibration Stds	VP135649
CCC Internal Standard/PEM ICV/I.BLK Surrogate Standard MS/MSD Standard LCS Standard	VP135650,VP135651 VP133935

Sr#	SampleID	ClientID	Data File Name	Date-Time	Comment	Operator	Status
1	BFB	BFB	VV039167.D	25 Sep 2025 09:14		SY/MD	Ok
2	VSTDCCC050	VSTDCCC050	VV039168.D	25 Sep 2025 09:59	pH#lot#v14222	SY/MD	Ok
3	VV0925MBL01	VV0925MBL01	VV039169.D	25 Sep 2025 10:31		SY/MD	Ok
4	VV0925WBL01	VV0925WBL01	VV039170.D	25 Sep 2025 11:01		SY/MD	Ok
5	VV0925WBS01	VV0925WBS01	VV039171.D	25 Sep 2025 11:24		SY/MD	Ok
6	VV0925WBSD01	VV0925WBSD01	VV039172.D	25 Sep 2025 11:48		SY/MD	Ok
7	Q3164-02	DA-2	VV039173.D	25 Sep 2025 12:10	pH<2.0 B	SY/MD	Ok
8	Q3175-01DL	MW2DL	VV039174.D	25 Sep 2025 12:33	Need 100X	SY/MD	Dilution
9	Q3175-02DL	MW4DL	VV039175.D	25 Sep 2025 12:55		SY/MD	Ok,M
10	VIBLK	VIBLK	VV039176.D	25 Sep 2025 13:18		SY/MD	Ok,M
11	Q3168-05	TOTES COMP#1	VV039177.D	25 Sep 2025 13:40	pH<2.0 A, Surrogate fail	SY/MD	ReRun
12	Q3168-06	TOTES COMP#2	VV039178.D	25 Sep 2025 14:03	pH<2.0 A, Surrogate fail	SY/MD	ReRun
13	Q3175-01DL2	MW2DL2	VV039179.D	25 Sep 2025 14:25		SY/MD	Ok,M
14	Q3172-02DL	FW6DDL	VV039180.D	25 Sep 2025 14:48		SY/MD	Ok
15	VIBLK	VIBLK	VV039181.D	25 Sep 2025 15:10		SY/MD	Ok
16	Q3172-01DL	FW7DDL	VV039182.D	25 Sep 2025 15:33	Surrogate fail	SY/MD	Ok
17	Q3168-05RE	TOTES COMP#1RE	VV039183.D	25 Sep 2025 15:55	pH<2.0 B, Surrogate fail	SY/MD	Confirms
18	Q3168-06RE	TOTES COMP#2RE	VV039184.D	25 Sep 2025 16:18	pH<2.0 B, Surrogate fail	SY/MD	Confirms

Instrument ID: MSVOA_V

Daily Analysis Runlog For Sequence/QC Batch ID # VV092525

Review By	Mahesh Dadoda	Review On	9/29/2025 12:56:22 AM		
Supervise By	Semsettin Yesilyurt	Supervise On	9/29/2025 1:08:36 AM		
SubDirectory	VV092525	HP Acquire Method	8260_V	HP Processing Method	82v092225w.m
STD. NAME		STD REF.#			
Tune/Reschk Initial Calibration Stds		VP135649			
CCC		VP135650,VP135651			
Internal Standard/PEM		VP133935			
ICV/I.BLK					
Surrogate Standard					
MS/MSD Standard					
LCS Standard					

19	Q3172-03	GB3W	VV039185.D	25 Sep 2025 16:40	pH<2.0 A	SY/MD	Ok
20	Q3195-22	RW-1-092425	VV039186.D	25 Sep 2025 17:03	pH<2.0 A	SY/MD	Ok
21	Q3195-23	EB-01-091925	VV039187.D	25 Sep 2025 17:25	pH<2.0 A ,EB	SY/MD	Ok
22	Q3195-24	EB-02-092225	VV039188.D	25 Sep 2025 17:47	pH<2.0 A ,Internal Standard Fail	SY/MD	ReRun
23	Q3195-25	FB-01-091925	VV039189.D	25 Sep 2025 18:10	pH<2.0 A ,FB	SY/MD	Ok
24	Q3195-26	FB-02-092225	VV039190.D	25 Sep 2025 18:32	pH<2.0 A FB	SY/MD	Ok
25	Q3195-27	TB-01-091525	VV039191.D	25 Sep 2025 18:55	pH<2.0 A ,TB	SY/MD	Ok
26	Q3195-02	MW-1-091725	VV039192.D	25 Sep 2025 19:17	pH<2.0 A	SY/MD	Ok
27	Q3195-09	MW-22-091825	VV039193.D	25 Sep 2025 19:40	pH<2.0 A	SY/MD	Ok
28	Q3195-10	MW-23-092325	VV039194.D	25 Sep 2025 20:02	pH<2.0 A	SY/MD	Ok
29	Q3195-11	MW-24-091925	VV039195.D	25 Sep 2025 20:25	pH<2.0 A	SY/MD	Ok
30	VSTDCCC050	VSTDCCC050EC	VV039196.D	25 Sep 2025 20:47		SY/MD	Ok

M : Manual Integration



SHIPPING DOCUMENTS



284 Sheffield Street, Mountainside, NJ 07092

(908) 789-8900 Fax: (908) 788-9222

www.chemtech.net

CHAIN OF CUSTODY RECORD

Alliance Project Number: AEC-2025-0013

23164

COC Number:

CLIENT INFORMATION

COMPANY: Durham School Services

ADDRESS: 30 W Yaphank Road

CITY: Coram STATE: NY ZIP:

ATTENTION: staci.bruce@alliancetg.com

PHONE: FAX:

PROJECT INFORMATION

PROJECT NAME: CSC 4151 Coram

PROJECT #: AEC-2025-0013 LOCATION: Coram

PROJECT MANAGER: Staci Bruce

E-MAIL: staci.bruce@alliancetg.com

PHONE: 423-902-9441 FAX:

BILLING INFORMATION

BILL TO: staci.bruce@alliancetg.com

PO#

ADDRESS:

CITY: STATE: ZIP:

ATTENTION: PHONE:

ANALYSIS

COD	Total OG	BTEX							
1	2	3	4	5	6	7	8	9	

DATA TURNAROUND INFORMATION

FAX: DAYS*
HARD COPY: DAYS*
EDD DAYS*

* TO BE APPROVED BY ALLIANCE

STANDARD TURNAROUND TIME IS 10 BUSINESS DAYS

DATA DELIVERABLE INFORMATION

- ☐ RESULTS ONLY ☐ USEPA CLP
☐ RESULTS + QC ☐ New York State ASP "B"
☐ New Jersey REDUCED ☐ New York State ASP "A"
☐ New Jersey CLP ☐ Other _____
☐ EDD Format _____

PRESERVATIVES

COMMENTS

CHEMTECH SAMPLE ID	PROJECT SAMPLE IDENTIFICATION	SAMPLE MATRIX	SAMPLE TYPE		SAMPLE COLLECTION		# of Bottles	PRESERVATIVES									COMMENTS
			COMP	GRAB	DATE	TIME		A	C	HCL							
1.	DA-1	SW		X	9/18	09:18	4	1	1	2							i-voc-RCVD Becker
2.	DA-2	SW		X	↓	09:30	4	1	1	2							
3.	DA-3	SW		X	↓	09:52	4	1	1	2							i-voc-RCVD Becker
4.																	
5.																	
6.																	
7.																	
8.																	
9.																	
10.																	

SAMPLE CUSTODY MUST BE DOCUMENTED BELOW EACH TIME SAMPLES CHANGE POSSESSION INCLUDING COURIER DELIVERY

RELINQUISHED BY SAMPLER	DATE/TIME	RECEIVED BY	Conditions of bottles or coolers at receipt: ☐ Compliant ☐ Non Compliant ☐ Cooler Temp 24.9°C
1. Kristen Parrella	09/18/25	1. [Signature]	MeOH extraction requires an additional 4oz. Jar for percent solid
RELINQUISHED BY	DATE/TIME	RECEIVED BY	Comments: Sample Received without Ice
2. [Signature]	9/22/25	2. [Signature]	Ice in Cooler?: no
RELINQUISHED BY	DATE/TIME	RECEIVED FOR LAB BY	SHIPPED VIA: CLIENT: ☐ Hand Delivered ☐ Overnight
3. [Signature]		3. [Signature]	ALLIANCE: ☐ Picked Up ☐ Overnight
			Shipment Complete ☐ YES ☐ NO

WHITE - ALLIANCE COPY FOR RETURN TO CLIENT

YELLOW - ALLIANCE COPY

PINK - SAMPLER COPY

Laboratory Certification

Certified By	License No.
Connecticut	PH-0830
DOD ELAP (ANAB)	L2219
Maine	2024021
Maryland	296
New Hampshire	255425
New Jersey	20012
New York	11376
Pennsylvania	68-00548
Soil Permit	525-24-234-08441
Texas	TX-C25-00189
Virginia	460312

LOGIN REPORT/SAMPLE TRANSFER

Order ID : Q3164	ATGG01	Order Date : 9/22/2025 2:37:00 PM	Project Mgr :
Client Name : ATG-GREENVILLE AEC		Project Name : AEC-2025-0013 CSC 4151	Report Type : Level 1
Client Contact : Staci Bruce		Receive DateTime : 9/22/2025 2:29:00 PM	EDD Type : Excel NY
Invoice Name : ATG-GREENVILLE AEC		Purchase Order :	Hard Copy Date :
Invoice Contact : Staci Bruce			Date Signoff :

LAB ID	CLIENT ID	MATRIX	SAMPLE DATE	SAMPLE TIME	TEST	TEST GROUP	METHOD	FAX DATE	DUE DATES
Q3164-01	DA-1	Water	09/18/2025	09:18	VOC-BTEX		8260-Low	10 Bus. Days	
Q3164-02	DA-2	Water	09/18/2025	09:30	VOC-BTEX		8260-Low	10 Bus. Days	
Q3164-03	DA-3	Water	09/18/2025	09:52	VOC-BTEX		8260-Low	10 Bus. Days	

Relinquished By : SP

Date / Time : 9/22/25 15:20

Received By : Sam

Date / Time : 09/22/25 15:20

Storage Area : VOA Refridgerator Room