

ANALYTICAL RESULTS SUMMARY

SEMI-VOLATILE ORGANICS
VOLATILE ORGANICS

PROJECT NAME : CON ED UTEN MOUNT VERNON, NY

CDM SMITH
110 Fieldcrest Ave
Raritan Center
Edison, NJ - 08837
Phone No: 732-225-7000

ORDER ID : Q2075
ATTENTION : Marcie Ann Encinas



Laboratory Certification ID # 20012



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NEW YORK STATE DEPARTMENT OF ENVIRONMENTAL CONSERVATION
 FORM S-I

SAMPLE IDENTIFICATION AND ANALYTICAL REQUIREMENT SUMMARY

NYSDEC Sample ID/Code	Laboratory Sample ID/Code	VOA GC/MS (Method #)	BNA GC/MS (Method #)	VOA GC (Method #)	Pest PCBs (Method #)	Metals (Method #)	Other (Method #)
SS-10	Q2075-01	8260D	8270E				Chemtech -SOP
SS-910	Q2075-02	8260D	8270E				Chemtech -SOP
SS-11	Q2075-03	8260D	8270E				Chemtech -SOP
SS-MW1-11.5	Q2075-06	8260D	8270E				Chemtech -SOP
FB-05152025	Q2075-07	8260D, 8260-Low	8270E				Chemtech -SOP
FB-05162025	Q2075-08	8260D, 8260-Low	8270E				Chemtech -SOP

NEW YORK STATE DEPARTMENT OF ENVIRONMENTAL CONSERVATION

FORM S-IIa

SAMPLE PREPARATION AND ANALYSIS SUMMARY SEMIVOLATILE (BNA) ANALYSES

Laboratory Sample ID	Matrix	Date Collected	Date Rec'd at Lab	Date Extracted	Date Analyzed
Q2075-01	SOIL	05/15/25	05/16/25	05/20/25	05/22/25
Q2075-02	SOIL	05/15/25	05/16/25	05/20/25	05/22/25
Q2075-03	SOIL	05/15/25	05/16/25	05/20/25	05/21/25
Q2075-06	SOIL	05/16/25	05/16/25	05/20/25	05/21/25
Q2075-07	Water	05/15/25	05/16/25	05/21/25	05/22/25
Q2075-08	Water	05/16/25	05/16/25	05/21/25	05/21/25

*** Details For Test :** SVOCMS Group3

NEW YORK STATE DEPARTMENT OF ENVIRONMENTAL CONSERVATION

FORM S-IIb

SAMPLE PREPARATION AND ANALYSIS SUMMARY VOLATILE (VOA) ANALYSES

Laboratory Sample ID	Matrix	Date Collected	Date Rec'd at Lab	Date Extracted	Date Analyzed
Q2075-01	SOIL	05/15/25	05/16/25		05/20/25
Q2075-02	SOIL	05/15/25	05/16/25		05/20/25
Q2075-03	SOIL	05/15/25	05/16/25		05/20/25
Q2075-06	SOIL	05/16/25	05/16/25		05/22/25
Q2075-07	Water	05/15/25	05/16/25		05/21/25
Q2075-08	Water	05/16/25	05/16/25		05/21/25

*** Details For Test :** VOCMS Group3

NEW YORK STATE DEPARTMENT OF ENVIRONMENTAL CONSERVATION

FORM S-III

SAMPLE PREPARATION AND ANALYSIS SUMMARY MISCELLANEOUS ORGANIC ANALYSES

Laboratory Sample ID	Matrix	Analytical Protocol	Extraction Method	Auxiliary Cleanup	Dil/Conc Factor
Q2075-01	Solid	8260D	5035		
Q2075-02	Solid	8260D	5035		
Q2075-03	Solid	8260D	5035		
Q2075-04	Solid	8260D	5035		
Q2075-05	Solid	8260D	5035		
Q2075-06	Solid	8260D	5035		
Q2075-07	Water	8260-Low	5030		
Q2075-08	Water	8260-Low	5030		

NEW YORK STATE DEPARTMENT OF ENVIRONMENTAL CONSERVATION

FORM S-III

SAMPLE PREPARATION AND ANALYSIS SUMMARY MISCELLANEOUS ORGANIC ANALYSES

Laboratory Sample ID	Matrix	Analytical Protocol	Extraction Method	Auxiliary Cleanup	Dil/Conc Factor
Q2075-01	Solid	8270E	NA		
Q2075-02	Solid	8270E	NA		
Q2075-03	Solid	8270E	NA		
Q2075-04	Solid	8270E	NA		
Q2075-05	Solid	8270E	NA		
Q2075-06	Solid	8270E	NA		
Q2075-07	Water	8270E	NA		
Q2075-08	Water	8270E	NA		

Cover Page

Order ID : Q2075

Project ID : Con Ed UTEN Mount Vernon, NY

Client : CDM Smith

Lab Sample Number

Q2075-01	SS-10
Q2075-02	SS-910
Q2075-03	SS-11
Q2075-04	SS-11MS
Q2075-05	SS-11MSD
Q2075-06	SS-MW1-11.5
Q2075-07	FB-05152025
Q2075-08	FB-05162025

Client Sample Number

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.

APPROVED

Signature :

By Nimisha Pandya, QA/QC Supervisor at 11:50 am, May 28, 2025

Date: 5/27/2025

NYDOH CERTIFICATION NO - 11376

NJDEP CERTIFICATION NO - 20012

CASE NARRATIVE

CDM Smith

Project Name: Con Ed UTEN Mount Vernon, NY

Project # N/A

Order ID # Q2075

Test Name: VOCMS Group3

A. Number of Samples and Date of Receipt:

6 Solid samples were received on 05/16/2025.

2 Water samples were received on 05/16/2025.

B. Parameters

According to the Chain of Custody document, the following analyses were requested:
SVOCMS Group3 and VOCMS Group3. This data package contains results for VOCMS Group3.

C. Analytical Techniques:

The analysis performed on instrument MSVOA_X were done using GC column DB-624UI 20m 0.18mm 1.0 um. Cat#121-1324UIThe analysis performed on instrument MSVOA_Y were done using GC column Rxi-624SIL MS 30m, 0.25mm, 1.4 um, Cat. #13868.The analysis of VOCMS Group3 was based on method 8260D.

D. QA/ QC Samples:

The Holding Times were met for all analysis.

The Surrogate recoveries met the acceptable criteria.

The Internal Standards Areas met the acceptable requirements.

The Retention Times were acceptable for all samples.

The MS {Q2075-04MS} with File ID: VY022350.D recoveries met the requirements for all compounds except for m/p-Xylenes[148%], o-Xylene[162%] and 1,2,4-Trimethylbenzene[235%] due to matrix interference.

The MSD {Q2075-05MSD} with File ID: VY022351.D recoveries met the acceptable requirements except for Ethyl Benzene[136%], Toluene[138%] m/p-Xylenes[161%] and o-Xylene [180%] due to matrix interference.

The RPD for {Q2075-05MSD} with File ID: VY022351.D met criteria except for 1,2,4-Trimethylbenzene[36%], Benzene[30%], p-Isopropyltoluene[21%], tert-Butylbenzene[26%] and Toluene[29%] due to difference in results of MS and MSD.

The Blank Spike met requirements for all samples.

The Blank Spike Duplicate met requirements for all samples.

The Blank analysis did not indicate the presence of lab contamination.



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Phone: 908 789 8900 Fax: 908 789 8922

The Initial Calibration met the requirements.
The Continuous Calibration met the requirements.
The Tuning criteria met requirements.

Sample SS-MW1-11.5 was diluted due to high concentration.

E. Additional Comments:

Trip Blank was not provided with this set of samples.

The soil samples results are based on a dry weight basis.

The Sample #SS-MW1-11.5ME have the concentration of target compound below Method detection limits, therefore it is not reported as Hit in Form1.

Please use %D calculated based on Avg RF and CCRF for all compounds using Average Response Factor when the %RSD value for a compound is <20% for the Initial Calibration curve and use %D calculated based on Amount added and Calculated amount for all compounds using Linear Regression when the %RSD value for a compound is > 20% for the Initial Calibration curve for SW-846 analysis.

F. Manual Integration Comments:

Please refer to the Manual integration Report included with the Run Logs for information on the manual integrations performed.

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. The laboratory manager or his designee, as verified by the following signature has authorized release of the data contained in this hard copy data package.

APPROVED

By Nimisha Pandya, QA/QC Supervisor at 11:50 am, May 28, 2025

Signature _____



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Phone: 908 789 8900 Fax: 908 789 8922

CASE NARRATIVE

CDM Smith

Project Name: Con Ed UTEN Mount Vernon, NY

Project # N/A

Order ID # Q2075

Test Name: SVOCMS Group3

A. Number of Samples and Date of Receipt:

6 Solid samples were received on 05/16/2025.

2 Water samples were received on 05/16/2025.

B. Parameters

According to the Chain of Custody document, the following analyses were requested:

SVOCMS Group3 and VOCMS Group3. This data package contains results for

SVOCMS Group3.

C. Analytical Techniques:

The samples were analyzed on instrument BNA_F using GC Column DB-UI 8270D which is 20 meters, 0.18 mm ID, 0.36 um dfThe samples were analyzed on instrument BNA_P using GC Column ZB-SemiVolatiles Guardian which is 30 meters, 0.25 mm ID, 0.5 um df, Catalog # 7HG-G027-17-GGAThe analysis of SVOCMS Group3 was based on method 8270E and extraction was done based on method 3510.

D. QA/ QC Samples:

The Holding Times were met for all analysis.

The Surrogate recoveries met the acceptable criteria.

The Internal Standards Areas met the acceptable requirements except for SS-11MS and SS-11MSD. Due to matrix interference, hence no corrective action is required.

and

SS-MW1-11.5 due to the presence of non-targeted hydrocarbons which can be observed by the abnormal chromatogram. Hence no corrective action is required.

The Retention Times were acceptable for all samples.

The MS recoveries met the requirements for all compounds.

The MSD recoveries met the acceptable requirements.

The RPD met criteria.

The Blank Spike met requirements for all samples.

The Blank Spike Duplicate met requirements for all samples.

The Blank analysis did not indicate the presence of lab contamination.

The Initial Calibration met the requirements.



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The Continuous Calibration met the requirements.
The Tuning criteria met requirements.

E. Additional Comments:

The Form 6 is not included in the data package because the Initial Calibration was performed using 7 points.

The soil samples results are based on a dry weight basis.

Please use %D calculated based on Avg RF and CCRF for all compounds using Average Response Factor when the %RSD value for a compound is <20% for the Initial Calibration curve and use %D calculated based on Amount added and Calculated amount for all compounds using Linear Regression when the %RSD value for a compound is > 20% for the Initial Calibration curve for SW-846 analysis.

F. Manual Integration Comments:

Please refer to the Manual integration Report included with the Run Logs for information on the manual integrations performed.

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. The laboratory manager or his designee, as verified by the following signature has authorized release of the data contained in this hard copy data package.

Signature _____

APPROVED

By Nimisha Pandya, QA/QC Supervisor at 11:51 am, May 28, 2025

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 #: Q2075

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 Castody

✓

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: SOHIL JODHANI

Date: 05/27/2025

LAB CHRONICLE

OrderID:	Q2075	OrderDate:	5/16/2025 4:10:00 PM
Client:	CDM Smith	Project:	Con Ed UTEN Mount Vernon, NY
Contact:	Marcie Ann Encinas	Location:	L41, VOA Ref. #2 Soil, VOA Ref. #3 Water

LabID	ClientID	Matrix	Test	Method	Sample Date	Prep Date	Anal Date	Received
Q2075-01	SS-10	SOIL	VOCMS Group3	8260D	05/15/25		05/20/25	05/16/25
Q2075-02	SS-910	SOIL	VOCMS Group3	8260D	05/15/25		05/20/25	05/16/25
Q2075-03	SS-11	SOIL	VOCMS Group3	8260D	05/15/25		05/20/25	05/16/25
Q2075-06	SS-MW1-11.5	SOIL	VOCMS Group3	8260D	05/16/25		05/22/25	05/16/25
Q2075-06ME	SS-MW1-11.5ME	SOIL	VOCMS Group3	8260D	05/16/25		05/23/25	05/16/25
Q2075-07	FB-05152025	Water	VOCMS Group3	8260-Low	05/15/25		05/21/25	05/16/25
Q2075-08	FB-05162025	Water	VOCMS Group3	8260-Low	05/16/25		05/21/25	05/16/25

Hit Summary Sheet
SW-846

SDG No.: Q2075
Client: CDM Smith

Sample ID	Client ID	Matrix	Parameter	Concentration	C	MDL	RDL	Units
Client ID:	SS-10							
Q2075-01	SS-10	SOIL	Ethyl Benzene	21.1		0.54	4.00	ug/Kg
Q2075-01	SS-10	SOIL	Total Xylenes	93.8		1.66	12.1	ug/Kg
Q2075-01	SS-10	SOIL	Isopropylbenzene	5.70		0.63	4.00	ug/Kg
Q2075-01	SS-10	SOIL	n-propylbenzene	13.1		0.59	4.00	ug/Kg
Q2075-01	SS-10	SOIL	1,3,5-Trimethylbenzene	23.4		0.66	4.00	ug/Kg
Q2075-01	SS-10	SOIL	1,2,4-Trimethylbenzene	90.2		0.52	4.00	ug/Kg
Q2075-01	SS-10	SOIL	sec-Butylbenzene	3.50	J	0.53	4.00	ug/Kg
Q2075-01	SS-10	SOIL	p-Isopropyltoluene	2.10	J	0.50	4.00	ug/Kg
Q2075-01	SS-10	SOIL	n-Butylbenzene	3.70	J	1.20	4.00	ug/Kg
Total Voc :				257				
Total Concentration:				257				
Client ID:	SS-910							
Q2075-02	SS-910	SOIL	Ethyl Benzene	5.40		0.51	3.80	ug/Kg
Q2075-02	SS-910	SOIL	Total Xylenes	26.5		1.58	11.5	ug/Kg
Q2075-02	SS-910	SOIL	Isopropylbenzene	2.10	J	0.60	3.80	ug/Kg
Q2075-02	SS-910	SOIL	n-propylbenzene	5.00		0.56	3.80	ug/Kg
Q2075-02	SS-910	SOIL	1,3,5-Trimethylbenzene	9.10		0.63	3.80	ug/Kg
Q2075-02	SS-910	SOIL	1,2,4-Trimethylbenzene	32.9		0.49	3.80	ug/Kg
Q2075-02	SS-910	SOIL	sec-Butylbenzene	1.50	J	0.51	3.80	ug/Kg
Q2075-02	SS-910	SOIL	p-Isopropyltoluene	0.99	J	0.48	3.80	ug/Kg
Q2075-02	SS-910	SOIL	n-Butylbenzene	1.70	J	1.10	3.80	ug/Kg
Total Voc :				85.2				
Total Concentration:				85.2				
Client ID:	SS-11							
Q2075-03	SS-11	SOIL	Toluene	5.20		0.78	5.00	ug/Kg
Q2075-03	SS-11	SOIL	Ethyl Benzene	7.80		0.67	5.00	ug/Kg
Q2075-03	SS-11	SOIL	Total Xylenes	46.3		2.02	14.9	ug/Kg
Q2075-03	SS-11	SOIL	Isopropylbenzene	2.10	J	0.78	5.00	ug/Kg
Q2075-03	SS-11	SOIL	n-propylbenzene	5.40		0.73	5.00	ug/Kg
Q2075-03	SS-11	SOIL	1,3,5-Trimethylbenzene	9.80		0.82	5.00	ug/Kg
Q2075-03	SS-11	SOIL	1,2,4-Trimethylbenzene	40.2		0.64	5.00	ug/Kg
Q2075-03	SS-11	SOIL	sec-Butylbenzene	1.40	J	0.66	5.00	ug/Kg
Q2075-03	SS-11	SOIL	n-Butylbenzene	1.40	J	1.40	5.00	ug/Kg
Total Voc :				120				
Total Concentration:				120				
Client ID:	SS-MW1-11.5							
Q2075-06	SS-MW1-11.5	SOIL	Benzene	9.40		0.62	3.90	ug/Kg

Hit Summary Sheet
SW-846

SDG No.: Q2075
Client: CDM Smith

Sample ID	Client ID	Matrix	Parameter	Concentration	C	MDL	RDL	Units
Q2075-06	SS-MW1-11.5	SOIL	Toluene	44.7		0.61	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	Ethyl Benzene	18.9		0.52	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	Total Xylenes	1510	E	1.61	11.7	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	Isopropylbenzene	40.5		0.61	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	n-propylbenzene	56.3		0.57	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	1,3,5-Trimethylbenzene	770	E	0.64	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	1,2,4-Trimethylbenzene	740	E	0.50	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	sec-Butylbenzene	66.3		0.52	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	p-Isopropyltoluene	52.3		0.48	3.90	ug/Kg
Q2075-06	SS-MW1-11.5	SOIL	n-Butylbenzene	71.9		1.10	3.90	ug/Kg
Total Voc :				3380				
Total Concentration:				3380				
Client ID:	SS-MW1-11.5ME							
Q2075-06ME	SS-MW1-11.5ME	SOIL	Toluene	83.3	JD	25.6	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	Ethyl Benzene	110	JD	22.0	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	Total Xylenes	630	D	67.6	490	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	Isopropylbenzene	42.8	JD	25.6	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	n-propylbenzene	130	JD	24.0	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	1,3,5-Trimethylbenzene	220	D	26.9	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	1,2,4-Trimethylbenzene	810	D	21.0	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	sec-Butylbenzene	63.9	JD	21.7	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	p-Isopropyltoluene	43.7	JD	20.4	160	ug/Kg
Q2075-06ME	SS-MW1-11.5ME	SOIL	n-Butylbenzene	180	D	47.6	160	ug/Kg
Total Voc :				2310				
Total Concentration:				2310				



A
B
C
D
E
F
G

SAMPLE DATA

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-10	SDG No.:	Q2075
Lab Sample ID:	Q2075-01	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	79.8
Sample Wt/Vol:	7.75	Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022343.D	1		05/20/25 15:22	VY052025

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	4.00	U	0.64	4.00	ug/Kg
108-88-3	Toluene	4.00	U	0.63	4.00	ug/Kg
100-41-4	Ethyl Benzene	21.1		0.54	4.00	ug/Kg
1330-20-7	Total Xylenes	93.8		1.66	12.1	ug/Kg
98-82-8	Isopropylbenzene	5.70		0.63	4.00	ug/Kg
103-65-1	n-propylbenzene	13.1		0.59	4.00	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	23.4		0.66	4.00	ug/Kg
98-06-6	tert-Butylbenzene	4.00	U	0.54	4.00	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	90.2		0.52	4.00	ug/Kg
135-98-8	sec-Butylbenzene	3.50	J	0.53	4.00	ug/Kg
99-87-6	p-Isopropyltoluene	2.10	J	0.50	4.00	ug/Kg
104-51-8	n-Butylbenzene	3.70	J	1.20	4.00	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	53.3		63 - 155	107%	SPK: 50
1868-53-7	Dibromofluoromethane	51.9		70 - 134	104%	SPK: 50
2037-26-5	Toluene-d8	50.7		74 - 123	101%	SPK: 50
460-00-4	4-Bromofluorobenzene	42.9		38 - 136	86%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	237000	7.707			
540-36-3	1,4-Difluorobenzene	426000	8.616			
3114-55-4	Chlorobenzene-d5	353000	11.414			
3855-82-1	1,4-Dichlorobenzene-d4	139000	13.346			

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

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-910	SDG No.:	Q2075
Lab Sample ID:	Q2075-02	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	81.3
Sample Wt/Vol:	8.02	Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022344.D	1		05/20/25 15:45	VY052025

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	3.80	U	0.61	3.80	ug/Kg
108-88-3	Toluene	3.80	U	0.60	3.80	ug/Kg
100-41-4	Ethyl Benzene	5.40		0.51	3.80	ug/Kg
1330-20-7	Total Xylenes	26.5		1.58	11.5	ug/Kg
98-82-8	Isopropylbenzene	2.10	J	0.60	3.80	ug/Kg
103-65-1	n-propylbenzene	5.00		0.56	3.80	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	9.10		0.63	3.80	ug/Kg
98-06-6	tert-Butylbenzene	3.80	U	0.51	3.80	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	32.9		0.49	3.80	ug/Kg
135-98-8	sec-Butylbenzene	1.50	J	0.51	3.80	ug/Kg
99-87-6	p-Isopropyltoluene	0.99	J	0.48	3.80	ug/Kg
104-51-8	n-Butylbenzene	1.70	J	1.10	3.80	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	48.7		63 - 155	97%	SPK: 50
1868-53-7	Dibromofluoromethane	49.6		70 - 134	99%	SPK: 50
2037-26-5	Toluene-d8	48.7		74 - 123	97%	SPK: 50
460-00-4	4-Bromofluorobenzene	39.8		38 - 136	80%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	242000	7.707			
540-36-3	1,4-Difluorobenzene	431000	8.616			
3114-55-4	Chlorobenzene-d5	344000	11.414			
3855-82-1	1,4-Dichlorobenzene-d4	127000	13.347			

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

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Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-11	SDG No.:	Q2075
Lab Sample ID:	Q2075-03	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	87.4
Sample Wt/Vol:	5.75	Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022345.D	1		05/20/25 16:09	VY052025

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	5.00	U	0.79	5.00	ug/Kg
108-88-3	Toluene	5.20		0.78	5.00	ug/Kg
100-41-4	Ethyl Benzene	7.80		0.67	5.00	ug/Kg
1330-20-7	Total Xylenes	46.3		2.02	14.9	ug/Kg
98-82-8	Isopropylbenzene	2.10	J	0.78	5.00	ug/Kg
103-65-1	n-propylbenzene	5.40		0.73	5.00	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	9.80		0.82	5.00	ug/Kg
98-06-6	tert-Butylbenzene	5.00	U	0.67	5.00	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	40.2		0.64	5.00	ug/Kg
135-98-8	sec-Butylbenzene	1.40	J	0.66	5.00	ug/Kg
99-87-6	p-Isopropyltoluene	5.00	U	0.62	5.00	ug/Kg
104-51-8	n-Butylbenzene	1.40	J	1.40	5.00	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	46.7		63 - 155	93%	SPK: 50
1868-53-7	Dibromofluoromethane	48.8		70 - 134	98%	SPK: 50
2037-26-5	Toluene-d8	48.7		74 - 123	97%	SPK: 50
460-00-4	4-Bromofluorobenzene	39.5		38 - 136	79%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	240000	7.707			
540-36-3	1,4-Difluorobenzene	433000	8.615			
3114-55-4	Chlorobenzene-d5	346000	11.414			
3855-82-1	1,4-Dichlorobenzene-d4	131000	13.346			

U = Not Detected

LOQ = Limit of Quantitation

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J = Estimated Value

B = Analyte Found in Associated Method Blank

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Report of Analysis

Client:	CDM Smith	Date Collected:	05/16/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-MW1-11.5	SDG No.:	Q2075
Lab Sample ID:	Q2075-06	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	91
Sample Wt/Vol:	7.04	Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022403.D	1		05/22/25 18:03	VY052225

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	9.40		0.62	3.90	ug/Kg
108-88-3	Toluene	44.7		0.61	3.90	ug/Kg
100-41-4	Ethyl Benzene	18.9		0.52	3.90	ug/Kg
1330-20-7	Total Xylenes	1510	E	1.61	11.7	ug/Kg
98-82-8	Isopropylbenzene	40.5		0.61	3.90	ug/Kg
103-65-1	n-propylbenzene	56.3		0.57	3.90	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	770	E	0.64	3.90	ug/Kg
98-06-6	tert-Butylbenzene	3.90	U	0.52	3.90	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	740	E	0.50	3.90	ug/Kg
135-98-8	sec-Butylbenzene	66.3		0.52	3.90	ug/Kg
99-87-6	p-Isopropyltoluene	52.3		0.48	3.90	ug/Kg
104-51-8	n-Butylbenzene	71.9		1.10	3.90	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	44.5		63 - 155	89%	SPK: 50
1868-53-7	Dibromofluoromethane	50.4		70 - 134	101%	SPK: 50
2037-26-5	Toluene-d8	55.4		74 - 123	111%	SPK: 50
460-00-4	4-Bromofluorobenzene	49.9		38 - 136	100%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	235000	7.719			
540-36-3	1,4-Difluorobenzene	408000	8.622			
3114-55-4	Chlorobenzene-d5	331000	11.42			
3855-82-1	1,4-Dichlorobenzene-d4	156000	13.353			

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Report of Analysis

Client:	CDM Smith	Date Collected:	05/16/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-MW1-11.5ME	SDG No.:	Q2075
Lab Sample ID:	Q2075-06ME	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	91
Sample Wt/Vol:	8.36	Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:	100	uL	Test: VOCMS Group3
GC Column:	DB-624UI	ID : 0.18	Level : MED
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046336.D	1		05/23/25 11:46	VX052325

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	160	UD	26.0	160	ug/Kg
108-88-3	Toluene	83.3	JD	25.6	160	ug/Kg
100-41-4	Ethyl Benzene	110	JD	22.0	160	ug/Kg
1330-20-7	Total Xylenes	630	D	67.6	490	ug/Kg
98-82-8	Isopropylbenzene	42.8	JD	25.6	160	ug/Kg
103-65-1	n-propylbenzene	130	JD	24.0	160	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	220	D	26.9	160	ug/Kg
98-06-6	tert-Butylbenzene	160	UD	22.0	160	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	810	D	21.0	160	ug/Kg
135-98-8	sec-Butylbenzene	63.9	JD	21.7	160	ug/Kg
99-87-6	p-Isopropyltoluene	43.7	JD	20.4	160	ug/Kg
104-51-8	n-Butylbenzene	180	D	47.6	160	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	51.0		63 - 155	102%	SPK: 50
1868-53-7	Dibromofluoromethane	48.4		70 - 134	97%	SPK: 50
2037-26-5	Toluene-d8	50.7		74 - 123	101%	SPK: 50
460-00-4	4-Bromofluorobenzene	51.5		38 - 136	103%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	61400	5.544			
540-36-3	1,4-Difluorobenzene	119000	6.757			
3114-55-4	Chlorobenzene-d5	110000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	48700	12.024			

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Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	FB-05152025	SDG No.:	Q2075
Lab Sample ID:	Q2075-07	Matrix:	Water
Analytical Method:	8260D	% Solid:	0
Sample Wt/Vol:	5	Units: mL	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	DB-624UI	ID : 0.18	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046292.D	1		05/21/25 13:09	VX052125

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	1.00	U	0.15	1.00	ug/L
108-88-3	Toluene	1.00	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	1.00	U	0.13	1.00	ug/L
1330-20-7	Total Xylenes	3.00	U	0.36	3.00	ug/L
98-82-8	Isopropylbenzene	1.00	U	0.12	1.00	ug/L
103-65-1	n-propylbenzene	1.00	U	0.13	1.00	ug/L
108-67-8	1,3,5-Trimethylbenzene	1.00	U	0.15	1.00	ug/L
98-06-6	tert-Butylbenzene	1.00	U	0.14	1.00	ug/L
95-63-6	1,2,4-Trimethylbenzene	1.00	U	0.14	1.00	ug/L
135-98-8	sec-Butylbenzene	1.00	U	0.13	1.00	ug/L
99-87-6	p-Isopropyltoluene	1.00	U	0.13	1.00	ug/L
104-51-8	n-Butylbenzene	1.00	U	0.15	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	53.2		74 - 125	106%	SPK: 50
1868-53-7	Dibromofluoromethane	49.3		75 - 124	99%	SPK: 50
2037-26-5	Toluene-d8	50.7		86 - 113	101%	SPK: 50
460-00-4	4-Bromofluorobenzene	52.0		77 - 121	104%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	63600	5.55			
540-36-3	1,4-Difluorobenzene	127000	6.757			
3114-55-4	Chlorobenzene-d5	121000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	52600	12.018			

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A = Aldol-Condensation Reaction Products

Report of Analysis

Client:	CDM Smith	Date Collected:	05/16/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	FB-05162025	SDG No.:	Q2075
Lab Sample ID:	Q2075-08	Matrix:	Water
Analytical Method:	8260D	% Solid:	0
Sample Wt/Vol:	5	Units: mL	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	DB-624UI	ID : 0.18	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046293.D	1		05/21/25 13:32	VX052125

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	1.00	U	0.15	1.00	ug/L
108-88-3	Toluene	1.00	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	1.00	U	0.13	1.00	ug/L
1330-20-7	Total Xylenes	3.00	U	0.36	3.00	ug/L
98-82-8	Isopropylbenzene	1.00	U	0.12	1.00	ug/L
103-65-1	n-propylbenzene	1.00	U	0.13	1.00	ug/L
108-67-8	1,3,5-Trimethylbenzene	1.00	U	0.15	1.00	ug/L
98-06-6	tert-Butylbenzene	1.00	U	0.14	1.00	ug/L
95-63-6	1,2,4-Trimethylbenzene	1.00	U	0.14	1.00	ug/L
135-98-8	sec-Butylbenzene	1.00	U	0.13	1.00	ug/L
99-87-6	p-Isopropyltoluene	1.00	U	0.13	1.00	ug/L
104-51-8	n-Butylbenzene	1.00	U	0.15	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	52.2		74 - 125	104%	SPK: 50
1868-53-7	Dibromofluoromethane	50.1		75 - 124	100%	SPK: 50
2037-26-5	Toluene-d8	50.2		86 - 113	100%	SPK: 50
460-00-4	4-Bromofluorobenzene	51.7		77 - 121	103%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	68000	5.55			
540-36-3	1,4-Difluorobenzene	135000	6.757			
3114-55-4	Chlorobenzene-d5	128000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	56000	12.018			

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A
B
C
D
E
F
G

QC SUMMARY

Surrogate Summary

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260D

Lab Sample ID	Client ID	Parameter	Spike	Result	Recovery	Limits	
						Qual	Low
Q2075-01	SS-10	1,2-Dichloroethane-d4	50	53.3	107	63	155
		Dibromofluoromethane	50	51.9	104	70	134
		Toluene-d8	50	50.7	101	74	123
Q2075-02	SS-910	4-Bromofluorobenzene	50	42.9	86	38	136
		1,2-Dichloroethane-d4	50	48.7	97	63	155
		Dibromofluoromethane	50	49.5	99	70	134
Q2075-03	SS-11	Toluene-d8	50	48.7	97	74	123
		4-Bromofluorobenzene	50	39.8	80	38	136
		1,2-Dichloroethane-d4	50	46.7	93	63	155
Q2075-04MS	SS-11MS	Dibromofluoromethane	50	48.8	98	70	134
		Toluene-d8	50	48.7	97	74	123
		4-Bromofluorobenzene	50	39.5	79	38	136
Q2075-05MSD	SS-11MSD	1,2-Dichloroethane-d4	50	49.2	98	63	155
		Dibromofluoromethane	50	42.1	84	70	134
		Toluene-d8	50	39.0	78	74	123
Q2075-06	SS-MW1-11.5	4-Bromofluorobenzene	50	36.5	73	38	136
		1,2-Dichloroethane-d4	50	62.7	125	63	155
		Dibromofluoromethane	50	51.3	103	70	134
Q2075-06ME	SS-MW1-11.5ME	Toluene-d8	50	45.1	90	74	123
		4-Bromofluorobenzene	50	45.4	91	38	136
		1,2-Dichloroethane-d4	50	44.5	89	63	155
VX0523MBL01	VX0523MBL01	Dibromofluoromethane	50	50.4	101	70	134
		Toluene-d8	50	55.4	111	74	123
		4-Bromofluorobenzene	50	49.9	100	38	136
VX0523MBS01	VX0523MBS01	1,2-Dichloroethane-d4	50	51.0	102	63	155
		Dibromofluoromethane	50	48.4	97	70	134
		Toluene-d8	50	50.7	101	74	123
VY0520SBL01	VY0520SBL01	4-Bromofluorobenzene	50	51.5	103	38	136
		1,2-Dichloroethane-d4	50	53.1	106	63	155
		Dibromofluoromethane	50	50.8	102	70	134
VY0520SBS01	VY0520SBS01	Toluene-d8	50	50.0	100	74	123
		4-Bromofluorobenzene	50	51.6	103	38	136
		1,2-Dichloroethane-d4	50	52.0	104	63	155
VY0522SBL01	VY0522SBL01	Dibromofluoromethane	50	52.4	105	70	134
		Toluene-d8	50	50.3	101	74	123
		4-Bromofluorobenzene	50	51.8	104	38	136
VY0520SBS01	VY0520SBS01	1,2-Dichloroethane-d4	50	45.7	91	63	155
		Dibromofluoromethane	50	48.4	97	70	134
		Toluene-d8	50	48.6	97	74	123
VY0522SBL01	VY0522SBL01	4-Bromofluorobenzene	50	52.5	105	38	136
		1,2-Dichloroethane-d4	50	52.8	106	63	155
		Dibromofluoromethane	50	52.0	104	70	134
VY0522SBS01	VY0522SBS01	Toluene-d8	50	51.5	103	74	123
		4-Bromofluorobenzene	50	47.7	95	38	136
		1,2-Dichloroethane-d4	50	48.9	98	63	155
VY0522SBS01	VY0522SBS01	Dibromofluoromethane	50	49.8	100	70	134
		Toluene-d8	50	48.7	97	74	123
		4-Bromofluorobenzene	50	39.5	79	38	136
VY0522SBS01	VY0522SBS01	1,2-Dichloroethane-d4	50	48.4	97	63	155
		Dibromofluoromethane	50	50.3	101	70	134

Surrogate SummarySDG No.: Q2075Client: CDM SmithAnalytical Method: SW8260D

Lab Sample ID	Client ID	Parameter	Spike	Result	Recovery	Qual	Limits	
							Low	High
VY0522SBS01	VY0522SBS01	Toluene-d8	50	50.3	101	74	123	
		4-Bromofluorobenzene	50	47.7	95	38	136	

Surrogate Summary

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260-Low

Lab Sample ID	Client ID	Parameter	Spike	Result	RecoveryQual	Limits	
						Low	High
Q2075-07	FB-05152025	1,2-Dichloroethane-d4	50	53.2	106	74	125
		Dibromofluoromethane	50	49.3	99	75	124
		Toluene-d8	50	50.7	101	86	113
Q2075-08	FB-05162025	4-Bromofluorobenzene	50	52.0	104	77	121
		1,2-Dichloroethane-d4	50	52.2	104	74	125
		Dibromofluoromethane	50	50.1	100	75	124
VX0521WBL01	VX0521WBL01	Toluene-d8	50	50.2	100	86	113
		4-Bromofluorobenzene	50	51.7	103	77	121
		1,2-Dichloroethane-d4	50	53.5	107	74	125
VX0521WBS01	VX0521WBS01	Dibromofluoromethane	50	50.9	102	75	124
		Toluene-d8	50	50.8	102	86	113
		4-Bromofluorobenzene	50	50.7	101	77	121
VX0521WBSD0	VX0521WBSD01	1,2-Dichloroethane-d4	50	53.6	107	74	125
		Dibromofluoromethane	50	54.3	109	75	124
		Toluene-d8	50	52.4	105	86	113
VX0521WBSD0	VX0521WBSD01	4-Bromofluorobenzene	50	55.0	110	77	121
		1,2-Dichloroethane-d4	50	54.2	108	74	125
		Dibromofluoromethane	50	55.1	110	75	124
VX0521WBSD0	VX0521WBSD01	Toluene-d8	50	54.1	108	86	113
		4-Bromofluorobenzene	50	55.8	112	77	121

Matrix Spike/Matrix Spike Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260D

Parameter	Spike	Sample Result	Result	Units	Rec		RPD Qual	Limits			RPD
					Rec	Qual		Low	High		
Lab Sample ID :	Q2075-04MS	Client Sample ID :	SS-11MS					Datafile :			VY022350.D
Benzene	34	0	31.6	ug/Kg	93			59	140		
Toluene	34	5.20	40.1	ug/Kg	103			61	134		
Ethyl Benzene	34	7.80	46.5	ug/Kg	114			54	134		
m/p-Xylenes	67.9	29.5	130	ug/Kg	148	*		51	137		
o-Xylene	34	16.8	71.9	ug/Kg	162	*		57	139		
Isopropylbenzene	34	2.10	34.0	ug/Kg	94			44	160		
N-propylbenzene	34	5.40	39.5	ug/Kg	100			10	173		
1,3,5-Trimethylbenzene	34	9.80	47.6	ug/Kg	111			10	175		
tert-Butylbenzene	34	0	28.7	ug/Kg	84			10	173		
1,2,4-Trimethylbenzene	34	40.2	120	ug/Kg	235	*		10	175		
Sec-butylbenzene	34	1.40	30.9	ug/Kg	87			10	172		
p-Isopropyltoluene	34	0	29.0	ug/Kg	85			26	153		
n-Butylbenzene	34	1.40	30.5	ug/Kg	86			10	175		

Matrix Spike/Matrix Spike Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260D

Parameter	Spike	Sample		Result	Units	Rec		RPD	Limits		RPD	
		Result	Units			Rec	Qual		Low	High		
Lab Sample ID :	Q2075-05MSD	Client Sample ID :		SS-11MSD				Datafile :	VY022351.D			
Benzene	34.3	0	ug/Kg	43.1	ug/Kg	126		30	*	59	140	20
Toluene	34.3	5.20	ug/Kg	52.7	ug/Kg	138	*	29	*	61	134	20
Ethyl Benzene	34.3	7.80	ug/Kg	54.6	ug/Kg	136	*	18		54	134	20
m/p-Xylenes	68.7	29.5	ug/Kg	140	ug/Kg	161	*	8		51	137	20
o-Xylene	34.3	16.8	ug/Kg	78.5	ug/Kg	180	*	11		57	139	20
Isopropylbenzene	34.3	2.10	ug/Kg	40.5	ug/Kg	112		17		44	160	20
N-propylbenzene	34.3	5.40	ug/Kg	43.6	ug/Kg	111		10		10	173	20
1,3,5-Trimethylbenzene	34.3	9.80	ug/Kg	49.0	ug/Kg	114		3		10	175	20
tert-Butylbenzene	34.3	0	ug/Kg	37.7	ug/Kg	110		26	*	10	173	20
1,2,4-Trimethylbenzene	34.3	40.2	ug/Kg	96.4	ug/Kg	164		36	*	10	175	20
Sec-butylbenzene	34.3	1.40	ug/Kg	37.6	ug/Kg	106		20		10	172	20
p-Isopropyltoluene	34.3	0	ug/Kg	36.2	ug/Kg	106		21	*	26	153	20
n-Butylbenzene	34.3	1.40	ug/Kg	36.4	ug/Kg	102		17		10	175	20

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260-Low

Datafile : VX046287.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Limits		RPD
								Low	High	
VX0521WBS01	Benzene	20	20.3	ug/L	102			82	109	
	Toluene	20	20.5	ug/L	103			82	110	
	Ethyl Benzene	20	20.1	ug/L	101			83	109	
	m/p-Xylenes	40	40.3	ug/L	101			82	110	
	o-Xylene	20	20.9	ug/L	104			83	109	
	Isopropylbenzene	20	20.4	ug/L	102			83	112	
	N-propylbenzene	20	20.4	ug/L	102			83	112	
	1,3,5-Trimethylbenzene	20	21.1	ug/L	106			85	112	
	tert-Butylbenzene	20	20.2	ug/L	101			83	112	
	1,2,4-Trimethylbenzene	20	20.7	ug/L	104			85	111	
	Sec-butylbenzene	20	20.3	ug/L	102			81	114	
	p-Isopropyltoluene	20	20.5	ug/L	103			78	116	
	n-Butylbenzene	20	19.7	ug/L	99			75	115	

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260-Low

Datafile : VX046288.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Limits		
								Low	High	RPD
VX0521WBSD01	Benzene	20	20.4	ug/L	102	0		82	109	20
	Toluene	20	21.1	ug/L	106	3		82	110	20
	Ethyl Benzene	20	20.4	ug/L	102	1		83	109	20
	m/p-Xylenes	40	41.1	ug/L	103	2		82	110	20
	o-Xylene	20	21.0	ug/L	105	1		83	109	20
	Isopropylbenzene	20	20.8	ug/L	104	2		83	112	20
	N-propylbenzene	20	20.7	ug/L	104	2		83	112	20
	1,3,5-Trimethylbenzene	20	21.0	ug/L	105	1		85	112	20
	tert-Butylbenzene	20	20.5	ug/L	103	2		83	112	20
	1,2,4-Trimethylbenzene	20	20.9	ug/L	104	0		85	111	20
	Sec-butylbenzene	20	21.1	ug/L	106	4		81	114	20
	p-Isopropyltoluene	20	20.7	ug/L	104	1		78	116	20
	n-Butylbenzene	20	20.2	ug/L	101	2		75	115	20

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260D

Datafile : VX046335.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Limits		RPD
								Low	High	
VX0523MBS01	Benzene	2000	2000	ug/Kg	100			84	121	
	Toluene	2000	2000	ug/Kg	100			83	122	
	Ethyl Benzene	2000	2000	ug/Kg	100			82	124	
	m/p-Xylenes	4000	4000	ug/Kg	100			83	124	
	o-Xylene	2000	2100	ug/Kg	105			83	123	
	Isopropylbenzene	2000	2000	ug/Kg	100			82	124	
	N-propylbenzene	2000	1900	ug/Kg	95			81	123	
	1,3,5-Trimethylbenzene	2000	2100	ug/Kg	105			82	124	
	tert-Butylbenzene	2000	2000	ug/Kg	100			81	125	
	1,2,4-Trimethylbenzene	2000	2000	ug/Kg	100			81	125	
	Sec-butylbenzene	2000	2000	ug/Kg	100			80	124	
	p-Isopropyltoluene	2000	2000	ug/Kg	100			81	125	
	n-Butylbenzene	2000	1900	ug/Kg	95			78	126	

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260D

Datafile : VY022331.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Limits		RPD
								Low	High	
VY0520SBS01	Benzene	20	20.8	ug/Kg	104			84	121	
	Toluene	20	20.0	ug/Kg	100			83	122	
	Ethyl Benzene	20	19.8	ug/Kg	99			82	124	
	m/p-Xylenes	40	39.6	ug/Kg	99			83	124	
	o-Xylene	20	20.2	ug/Kg	101			83	123	
	Isopropylbenzene	20	20.1	ug/Kg	101			82	124	
	N-propylbenzene	20	20.3	ug/Kg	102			81	123	
	1,3,5-Trimethylbenzene	20	19.5	ug/Kg	98			82	124	
	tert-Butylbenzene	20	19.8	ug/Kg	99			81	125	
	1,2,4-Trimethylbenzene	20	19.3	ug/Kg	97			81	125	
	Sec-butylbenzene	20	20.0	ug/Kg	100			80	124	
	p-Isopropyltoluene	20	19.3	ug/Kg	97			81	125	
	n-Butylbenzene	20	19.7	ug/Kg	99			78	126	

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8260D

Datafile : VY022382.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Limits		RPD
								Low	High	
VY0522SBS01	Benzene	20	19.5	ug/Kg	98			84	121	
	Toluene	20	19.4	ug/Kg	97			83	122	
	Ethyl Benzene	20	18.7	ug/Kg	94			82	124	
	m/p-Xylenes	40	38.0	ug/Kg	95			83	124	
	o-Xylene	20	18.8	ug/Kg	94			83	123	
	Isopropylbenzene	20	19.3	ug/Kg	97			82	124	
	N-propylbenzene	20	19.4	ug/Kg	97			81	123	
	1,3,5-Trimethylbenzene	20	18.9	ug/Kg	95			82	124	
	tert-Butylbenzene	20	19.2	ug/Kg	96			81	125	
	1,2,4-Trimethylbenzene	20	18.8	ug/Kg	94			81	125	
	Sec-butylbenzene	20	19.4	ug/Kg	97			80	124	
	p-Isopropyltoluene	20	19.0	ug/Kg	95			81	125	
	n-Butylbenzene	20	19.3	ug/Kg	97			78	126	

VOLATILE METHOD BLANK SUMMARY

EPA SAMPLE NO.

VX0521WBL01

Lab Name: CHEMTECHContract: CAMP02Lab Code: CHEM Case No.: Q2075SAS No.: Q2075 SDG NO.: Q2075Lab File ID: VX046286.DLab Sample ID: VX0521WBL01Date Analyzed: 05/21/2025Time Analyzed: 10:46GC Column: DB-624UI ID: 0.18 (mm)Heated Purge: (Y/N) NInstrument ID: MSVOA_X

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
VX0521WBS01	VX0521WBS01	VX046287.D	05/21/2025
VX0521WBSD01	VX0521WBSD01	VX046288.D	05/21/2025
FB-05152025	Q2075-07	VX046292.D	05/21/2025
FB-05162025	Q2075-08	VX046293.D	05/21/2025

COMMENTS:

VOLATILE METHOD BLANK SUMMARY

EPA SAMPLE NO.

VX0523MBL01

Lab Name: CHEMTECHContract: CAMP02Lab Code: CHEM Case No.: Q2075SAS No.: Q2075 SDG NO.: Q2075Lab File ID: VX046332.DLab Sample ID: VX0523MBL01Date Analyzed: 05/23/2025Time Analyzed: 09:53GC Column: DB-624UI ID: 0.18 (mm)Heated Purge: (Y/N) NInstrument ID: MSVOA_X

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
VX0523MBS01	VX0523MBS01	VX046335.D	05/23/2025
SS-MW1-11.5ME	Q2075-06ME	VX046336.D	05/23/2025

COMMENTS:

VOLATILE METHOD BLANK SUMMARY

EPA SAMPLE NO.

VY0520SBL01

Lab Name: CHEMTECHContract: CAMP02Lab Code: CHEM Case No.: Q2075SAS No.: Q2075 SDG NO.: Q2075Lab File ID: VY022330.DLab Sample ID: VY0520SBL01Date Analyzed: 05/20/2025Time Analyzed: 09:48GC Column: RXI-624 ID: 0.25 (mm)Heated Purge: (Y/N) YInstrument ID: MSVOA_Y

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
VY0520SBS01	VY0520SBS01	VY022331.D	05/20/2025
SS-10	Q2075-01	VY022343.D	05/20/2025
SS-910	Q2075-02	VY022344.D	05/20/2025
SS-11	Q2075-03	VY022345.D	05/20/2025
SS-11MS	Q2075-04MS	VY022350.D	05/20/2025
SS-11MSD	Q2075-05MSD	VY022351.D	05/20/2025

COMMENTS:

VOLATILE METHOD BLANK SUMMARY

EPA SAMPLE NO.

VY0522SBL01

Lab Name: CHEMTECHContract: CAMP02Lab Code: CHEM Case No.: Q2075SAS No.: Q2075 SDG NO.: Q2075Lab File ID: VY022381.DLab Sample ID: VY0522SBL01Date Analyzed: 05/22/2025Time Analyzed: 09:07GC Column: RXI-624 ID: 0.25 (mm)Heated Purge: (Y/N) YInstrument ID: MSVOA_Y

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
VY0522SBS01	VY0522SBS01	VY022382.D	05/22/2025
SS-MW1-11.5	Q2075-06	VY022403.D	05/22/2025

COMMENTS:

VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VX046038.D	SAS No.:	Q2075
Instrument ID:	MSVOA_X	BFB Injection Date:	05/05/2025
GC Column:	DB-624UI ID: 0.18 (mm)	BFB Injection Time:	09:37
		Heated Purge:	Y/N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	22.1
75	30.0 - 60.0% of mass 95	56.2
95	Base Peak, 100% relative abundance	100
96	5.0 - 9.0% of mass 95	6.4
173	Less than 2.0% of mass 174	0.5 (0.7) 1
174	50.0 - 100.0% of mass 95	68.8
175	5.0 - 9.0% of mass 174	5 (7.3) 1
176	95.0 - 101.0% of mass 174	66.7 (97) 1
177	5.0 - 9.0% of mass 176	4.6 (6.9) 2

1-Value is % mass 174

2-Value is % mass 176

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDICC020	VSTDICC020	VX046041.D	05/05/2025	11:35
VSTDICCC050	VSTDICCC050	VX046042.D	05/05/2025	11:58
VSTDICC100	VSTDICC100	VX046043.D	05/05/2025	12:21
VSTDICC150	VSTDICC150	VX046044.D	05/05/2025	12:45
VSTDICC005	VSTDICC005	VX046046.D	05/05/2025	16:04
VSTDICC001	VSTDICC001	VX046047.D	05/05/2025	16:27

VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VX046283.D	SAS No.:	Q2075
Instrument ID:	MSVOA_X	BFB Injection Date:	05/21/2025
GC Column:	DB-624UI ID: 0.18 (mm)	BFB Injection Time:	09:13
		Heated Purge:	Y/N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	21.4
75	30.0 - 60.0% of mass 95	56.9
95	Base Peak, 100% relative abundance	100
96	5.0 - 9.0% of mass 95	7
173	Less than 2.0% of mass 174	0.6 (0.9) 1
174	50.0 - 100.0% of mass 95	70.1
175	5.0 - 9.0% of mass 174	5.4 (7.7) 1
176	95.0 - 101.0% of mass 174	67.1 (95.7) 1
177	5.0 - 9.0% of mass 176	4.3 (6.5) 2

1-Value is % mass 174

2-Value is % mass 176

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDCCC050	VSTDCCC050	VX046284.D	05/21/2025	09:55
VX0521WBL01	VX0521WBL01	VX046286.D	05/21/2025	10:46
VX0521WBS01	VX0521WBS01	VX046287.D	05/21/2025	11:09
VX0521WBSD01	VX0521WBSD01	VX046288.D	05/21/2025	11:36
FB-05152025	Q2075-07	VX046292.D	05/21/2025	13:09
FB-05162025	Q2075-08	VX046293.D	05/21/2025	13:32

VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VX046330.D	SAS No.:	Q2075
Instrument ID:	MSVOA_X	BFB Injection Date:	05/23/2025
GC Column:	DB-624UI ID: 0.18 (mm)	BFB Injection Time:	08:25
		Heated Purge:	Y/N

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	22
75	30.0 - 60.0% of mass 95	56.7
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	0.8 (1.2) 1
174	50.0 - 100.0% of mass 95	66.8
175	5.0 - 9.0% of mass 174	4.6 (6.9) 1
176	95.0 - 101.0% of mass 174	64 (95.9) 1
177	5.0 - 9.0% of mass 176	4.1 (6.4) 2

1-Value is % mass 174

2-Value is % mass 176

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDCCC050	VSTDCCC050	VX046331.D	05/23/2025	09:24
VX0523MBL01	VX0523MBL01	VX046332.D	05/23/2025	09:53
VX0523MBS01	VX0523MBS01	VX046335.D	05/23/2025	11:23
SS-MW1-11.5ME	Q2075-06ME	VX046336.D	05/23/2025	11:46

VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VY022252.D	SAS No.:	Q2075
Instrument ID:	MSVOA_Y	BFB Injection Date:	05/15/2025
GC Column:	RXI-624 ID: 0.25 (mm)	BFB Injection Time:	07:52
		Heated Purge: Y/N	Y

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	24
75	30.0 - 60.0% of mass 95	55.9
95	Base Peak, 100% relative abundance	100
96	5.0 - 9.0% of mass 95	6.8
173	Less than 2.0% of mass 174	0.7 (0.9) 1
174	50.0 - 100.0% of mass 95	80.5
175	5.0 - 9.0% of mass 174	5.9 (7.3) 1
176	95.0 - 101.0% of mass 174	77.5 (96.3) 1
177	5.0 - 9.0% of mass 176	5.3 (6.8) 2

1-Value is % mass 174

2-Value is % mass 176

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDICC005	VSTDICC005	VY022253.D	05/15/2025	09:46
VSTDICC010	VSTDICC010	VY022254.D	05/15/2025	10:16
VSTDICC020	VSTDICC020	VY022255.D	05/15/2025	10:39
VSTDICCC050	VSTDICCC050	VY022256.D	05/15/2025	11:02
VSTDICC100	VSTDICC100	VY022257.D	05/15/2025	11:24
VSTDICC150	VSTDICC150	VY022258.D	05/15/2025	11:47

VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VY022328.D	SAS No.:	Q2075
Instrument ID:	MSVOA_Y	SDG NO.:	Q2075
GC Column:	RXI-624 ID: 0.25 (mm)	BFB Injection Date:	05/20/2025
		BFB Injection Time:	08:09
		Heated Purge: Y/N	Y

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	24.5
75	30.0 - 60.0% of mass 95	58.5
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 (1.2) 1
174	50.0 - 100.0% of mass 95	82.9
175	5.0 - 9.0% of mass 174	6.4 (7.7) 1
176	95.0 - 101.0% of mass 174	80.3 (96.8) 1
177	5.0 - 9.0% of mass 176	5.4 (6.7) 2

1-Value is % mass 174

2-Value is % mass 176

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDCCC050	VSTDCCC050	VY022329.D	05/20/2025	08:40
VY0520SBL01	VY0520SBL01	VY022330.D	05/20/2025	09:48
VY0520SBS01	VY0520SBS01	VY022331.D	05/20/2025	10:21
SS-10	Q2075-01	VY022343.D	05/20/2025	15:22
SS-910	Q2075-02	VY022344.D	05/20/2025	15:45
SS-11	Q2075-03	VY022345.D	05/20/2025	16:09
SS-11MS	Q2075-04MS	VY022350.D	05/20/2025	18:04
SS-11MSD	Q2075-05MSD	VY022351.D	05/20/2025	18:26

VOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
BROMOFLUOROBENZENE (BFB)

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VY022379.D	SAS No.:	Q2075
Instrument ID:	MSVOA_Y	BFB Injection Date:	05/22/2025
GC Column:	RXI-624 ID: 0.25 (mm)	BFB Injection Time:	08:05
		Heated Purge: Y/N	Y

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
50	15.0 - 40.0% of mass 95	24.1
75	30.0 - 60.0% of mass 95	57.8
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	1.1 (1.3) 1
174	50.0 - 100.0% of mass 95	86.1
175	5.0 - 9.0% of mass 174	6.4 (7.5) 1
176	95.0 - 101.0% of mass 174	83.8 (97.4) 1
177	5.0 - 9.0% of mass 176	5.5 (6.5) 2

1-Value is % mass 174

2-Value is % mass 176

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
VSTDCCC050	VSTDCCC050	VY022380.D	05/22/2025	08:35
VY0522SBL01	VY0522SBL01	VY022381.D	05/22/2025	09:07
VY0522SBS01	VY0522SBS01	VY022382.D	05/22/2025	09:36
SS-MW1-11.5	Q2075-06	VY022403.D	05/22/2025	18:03

VOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: CHEMTECH Contract: CAMP02
 Lab Code: CHEM Case No.: Q2075 SAS No.: Q2075 SDG No.: Q2075
 Lab File ID: VX046284.D Date Analyzed: 05/21/2025
 Instrument ID: MSVOA_X Time Analyzed: 09:55
 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	95080	5.54	161730	6.75	140445	10.05
UPPER LIMIT	190160	6.037	323460	7.251	280890	10.549
LOWER LIMIT	47540	5.037	80865	6.251	70222.5	9.549
EPA SAMPLE NO.						
FB-05152025	63566	5.55	126684	6.76	120601	10.05
FB-05162025	67975	5.55	135035	6.76	128288	10.05
VX0521WBL01	66690	5.54	132814	6.76	124685	10.05
VX0521WBS01	83102	5.54	146697	6.76	130479	10.05
VX0521WBSD01	82656	5.54	142441	6.76	127622	10.05

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:	CHEMTECH	Contract:	CAMP02		
Lab Code:	<u>CHEM</u>	SAS No.:	<u>Q2075</u>	SDG NO.:	<u>Q2075</u>
Lab File ID:	<u>VX046284.D</u>	Date Analyzed:	<u>05/21/2025</u>		
Instrument ID:	<u>MSVOA_X</u>	Time Analyzed:	<u>09:55</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	67683	12.018				
UPPER LIMIT	135366	12.518				
LOWER LIMIT	33841.5	11.518				
EPA SAMPLE NO.						
FB-05152025	52644	12.02				
FB-05162025	55957	12.02				
VX0521WBL01	53099	12.02				
VX0521WBS01	62651	12.02				
VX0521WBSD01	60955	12.02				

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:	<u>CHEMTECH</u>	Contract:	<u>CAMP02</u>
Lab Code:	<u>CHEM</u>	Case No.:	<u>Q2075</u>
Lab File ID:	<u>VX046331.D</u>	Date Analyzed:	<u>05/23/2025</u>
Instrument ID:	<u>MSVOA_X</u>	Time Analyzed:	<u>09:24</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	89943	5.54	154938	6.76	137123	10.05
UPPER LIMIT	179886	6.044	309876	7.257	274246	10.549
LOWER LIMIT	44971.5	5.044	77469	6.257	68561.5	9.549
EPA SAMPLE NO.						
SS-MW1-11.5ME	61392	5.54	119098	6.76	110366	10.05
VX0523MBL01	59074	5.54	117281	6.76	111918	10.05
VX0523MBS01	83340	5.54	146324	6.76	129426	10.05

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>CHEMTECH</u>		Contract:	<u>CAMP02</u>			
Lab Code:	<u>CHEM</u>	Case No.:	<u>Q2075</u>	SAS No.:	<u>Q2075</u>	SDG NO.:	<u>Q2075</u>
Lab File ID:	<u>VX046331.D</u>		Date Analyzed:	<u>05/23/2025</u>			
Instrument ID:	<u>MSVOA_X</u>		Time Analyzed:	<u>09:24</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	66629	12.018				
	133258	12.518				
	33314.5	11.518				
EPA SAMPLE NO.						
SS-MW1-11.5ME	48680	12.02				
VX0523MBL01	48375	12.02				
VX0523MBS01	63342	12.02				

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:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VY022329.D	Date Analyzed:	05/20/2025
Instrument ID:	MSVOA_Y	Time Analyzed:	08:40
GC Column:	RXI-624	ID: 0.25 (mm)	Heated Purge: (Y/N) Y

	IS1 AREA #	RT #	IS2 AREA #	RT #	IS3 AREA #	RT #
12 HOUR STD	180374	7.71	298369	8.62	252391	11.41
UPPER LIMIT	360748	8.207	596738	9.116	504782	11.914
LOWER LIMIT	90187	7.207	149185	8.116	126196	10.914
EPA SAMPLE NO.						
SS-10	237248	7.71	425729	8.62	352723	11.41
SS-910	241551	7.71	430984	8.62	344445	11.41
SS-11	239791	7.71	432569	8.62	345893	11.41
SS-11MS	134739	7.71	248613	8.62	210937	11.41
SS-11MSD	117744	7.71	220285	8.62	197296	11.41
VY0520SBL01	253729	7.71	453295	8.62	355808	11.42
VY0520SBS01	162404	7.71	280923	8.62	236555	11.42

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:	CHEMTECH		Contract:	CAMP02			
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075	SDG NO.:	Q2075
Lab File ID:	VY022329.D		Date Analyzed:	05/20/2025			
Instrument ID:	MSVOA_Y		Time Analyzed:	08:40			
GC Column:	RXI-624	ID: 0.25 (mm)	Heated Purge: (Y/N)	Y			

	IS4 AREA #	RT #				
12 HOUR STD	120747	13.346				
	241494	13.846				
	60373.5	12.846				
EPA SAMPLE NO.						
SS-10	138928	13.35				
SS-910	126954	13.35				
SS-11	130644	13.35				
SS-11MS	91777	13.35				
SS-11MSD	91227	13.35				
VY0520SBL01	124571	13.35				
VY0520SBS01	111469	13.35				

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:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	Case No.:	Q2075
Lab File ID:	VY022380.D	Date Analyzed:	05/22/2025
Instrument ID:	MSVOA_Y	Time Analyzed:	08:35
GC Column:	RXI-624	ID: 0.25 (mm)	Heated Purge: (Y/N) Y

	IS1 AREA #	RT #	IS2 AREA #	RT #	IS3 AREA #	RT #
12 HOUR STD	161495	7.72	274168	8.62	235238	11.42
UPPER LIMIT	322990	8.22	548336	9.122	470476	11.92
LOWER LIMIT	80747.5	7.22	137084	8.122	117619	10.92
EPA SAMPLE NO.						
SS-MW1-11.5	234597	7.72	407953	8.62	330547	11.42
VY0522SBL01	263712	7.72	465826	8.62	376365	11.42
VY0522SBS01	174604	7.72	293942	8.62	250842	11.42

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:	CHEMTECH	Contract:	CAMP02		
Lab Code:	<u>CHEM</u>	SAS No.:	<u>Q2075</u>	SDG NO.:	<u>Q2075</u>
Lab File ID:	<u>VY022380.D</u>	Date Analyzed:	<u>05/22/2025</u>		
Instrument ID:	<u>MSVOA_Y</u>	Time Analyzed:	<u>08:35</u>		
GC Column:	<u>RXI-624</u>	ID: 0.25 (mm)	Heated Purge: (Y/N)	<u>Y</u>	

	IS4 AREA #	RT #				
12 HOUR STD	110586	13.353				
UPPER LIMIT	221172	13.853				
LOWER LIMIT	55293	12.853				
EPA SAMPLE NO.						
SS-MW1-11.5	155760	13.35				
VY0522SBL01	137350	13.35				
VY0522SBS01	118480	13.35				

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.



A
B
C
D
E
F
G

QC SAMPLE

DATA

Report of Analysis

Client:	CDM Smith	Date Collected:
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:
Client Sample ID:	VX0521WBL01	SDG No.: Q2075
Lab Sample ID:	VX0521WBL01	Matrix: Water
Analytical Method:	8260D	% Solid: 0
Sample Wt/Vol:	5 mL	Final Vol: 5000 uL
Soil Aliquot Vol:	uL	Test: VOCMS Group3
GC Column:	DB-624UI ID : 0.18	Level : LOW
Prep Method :		

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046286.D	1		05/21/25 10:46	VX052125

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	1.00	U	0.15	1.00	ug/L
108-88-3	Toluene	1.00	U	0.14	1.00	ug/L
100-41-4	Ethyl Benzene	1.00	U	0.13	1.00	ug/L
1330-20-7	Total Xylenes	3.00	U	0.36	3.00	ug/L
98-82-8	Isopropylbenzene	1.00	U	0.12	1.00	ug/L
103-65-1	n-propylbenzene	1.00	U	0.13	1.00	ug/L
108-67-8	1,3,5-Trimethylbenzene	1.00	U	0.15	1.00	ug/L
98-06-6	tert-Butylbenzene	1.00	U	0.14	1.00	ug/L
95-63-6	1,2,4-Trimethylbenzene	1.00	U	0.14	1.00	ug/L
135-98-8	sec-Butylbenzene	1.00	U	0.13	1.00	ug/L
99-87-6	p-Isopropyltoluene	1.00	U	0.13	1.00	ug/L
104-51-8	n-Butylbenzene	1.00	U	0.15	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	53.5		74 - 125	107%	SPK: 50
1868-53-7	Dibromofluoromethane	50.9		75 - 124	102%	SPK: 50
2037-26-5	Toluene-d8	50.8		86 - 113	102%	SPK: 50
460-00-4	4-Bromofluorobenzene	50.7		77 - 121	101%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	66700	5.544			
540-36-3	1,4-Difluorobenzene	133000	6.757			
3114-55-4	Chlorobenzene-d5	125000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	53100	12.018			

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

Report of Analysis

Client:	CDM Smith	Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	
Client Sample ID:	VX0523MBL01	SDG No.:	Q2075
Lab Sample ID:	VX0523MBL01	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	100
Sample Wt/Vol:	5	Units: g	Final Vol: 10000 uL
Soil Aliquot Vol:	100	uL	Test: VOCMS Group3
GC Column:	DB-624UI	ID : 0.18	Level : MED
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046332.D	1		05/23/25 09:53	VX052325

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	500	U	79.0	500	ug/Kg
108-88-3	Toluene	500	U	78.0	500	ug/Kg
100-41-4	Ethyl Benzene	500	U	67.0	500	ug/Kg
1330-20-7	Total Xylenes	1500	U	202	1500	ug/Kg
98-82-8	Isopropylbenzene	500	U	78.0	500	ug/Kg
103-65-1	n-propylbenzene	500	U	73.0	500	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	500	U	82.0	500	ug/Kg
98-06-6	tert-Butylbenzene	500	U	67.0	500	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	500	U	64.0	500	ug/Kg
135-98-8	sec-Butylbenzene	500	U	66.0	500	ug/Kg
99-87-6	p-Isopropyltoluene	500	U	62.0	500	ug/Kg
104-51-8	n-Butylbenzene	500	U	150	500	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	53.1		63 - 155	106%	SPK: 50
1868-53-7	Dibromofluoromethane	50.8		70 - 134	102%	SPK: 50
2037-26-5	Toluene-d8	50.0		74 - 123	100%	SPK: 50
460-00-4	4-Bromofluorobenzene	51.6		38 - 136	103%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	59100	5.544			
540-36-3	1,4-Difluorobenzene	117000	6.757			
3114-55-4	Chlorobenzene-d5	112000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	48400	12.018			

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

Report of Analysis

Client:	CDM Smith	Date Collected:
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:
Client Sample ID:	VY0520SBL01	SDG No.: Q2075
Lab Sample ID:	VY0520SBL01	Matrix: SOIL
Analytical Method:	8260D	% Solid: 100
Sample Wt/Vol:	5 Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:	uL	Test: VOCMS Group3
GC Column:	RXI-624 ID : 0.25	Level : LOW
Prep Method :		

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022330.D	1		05/20/25 09:48	VY052025

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	5.00	U	0.79	5.00	ug/Kg
108-88-3	Toluene	5.00	U	0.78	5.00	ug/Kg
100-41-4	Ethyl Benzene	5.00	U	0.67	5.00	ug/Kg
1330-20-7	Total Xylenes	15.0	U	2.02	15.0	ug/Kg
98-82-8	Isopropylbenzene	5.00	U	0.78	5.00	ug/Kg
103-65-1	n-propylbenzene	5.00	U	0.73	5.00	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	5.00	U	0.82	5.00	ug/Kg
98-06-6	tert-Butylbenzene	5.00	U	0.67	5.00	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	5.00	U	0.64	5.00	ug/Kg
135-98-8	sec-Butylbenzene	5.00	U	0.66	5.00	ug/Kg
99-87-6	p-Isopropyltoluene	5.00	U	0.62	5.00	ug/Kg
104-51-8	n-Butylbenzene	5.00	U	1.50	5.00	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	45.7		63 - 155	91%	SPK: 50
1868-53-7	Dibromofluoromethane	48.3		70 - 134	97%	SPK: 50
2037-26-5	Toluene-d8	48.6		74 - 123	97%	SPK: 50
460-00-4	4-Bromofluorobenzene	52.5		38 - 136	105%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	254000	7.707			
540-36-3	1,4-Difluorobenzene	453000	8.615			
3114-55-4	Chlorobenzene-d5	356000	11.42			
3855-82-1	1,4-Dichlorobenzene-d4	125000	13.346			

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

Report of Analysis

Client:	CDM Smith	Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	
Client Sample ID:	VY0522SBL01	SDG No.:	Q2075
Lab Sample ID:	VY0522SBL01	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	100
Sample Wt/Vol:	5 Units: g	Final Vol:	5000 uL
Soil Aliquot Vol:	uL	Test:	VOCMS Group3
GC Column:	RXI-624 ID : 0.25	Level :	LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022381.D	1		05/22/25 09:07	VY052225

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	5.00	U	0.79	5.00	ug/Kg
108-88-3	Toluene	5.00	U	0.78	5.00	ug/Kg
100-41-4	Ethyl Benzene	5.00	U	0.67	5.00	ug/Kg
1330-20-7	Total Xylenes	15.0	U	2.02	15.0	ug/Kg
98-82-8	Isopropylbenzene	5.00	U	0.78	5.00	ug/Kg
103-65-1	n-propylbenzene	5.00	U	0.73	5.00	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	5.00	U	0.82	5.00	ug/Kg
98-06-6	tert-Butylbenzene	5.00	U	0.67	5.00	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	5.00	U	0.64	5.00	ug/Kg
135-98-8	sec-Butylbenzene	5.00	U	0.66	5.00	ug/Kg
99-87-6	p-Isopropyltoluene	5.00	U	0.62	5.00	ug/Kg
104-51-8	n-Butylbenzene	5.00	U	1.50	5.00	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	48.9		63 - 155	98%	SPK: 50
1868-53-7	Dibromofluoromethane	49.8		70 - 134	100%	SPK: 50
2037-26-5	Toluene-d8	48.7		74 - 123	97%	SPK: 50
460-00-4	4-Bromofluorobenzene	39.5		38 - 136	79%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	264000	7.719			
540-36-3	1,4-Difluorobenzene	466000	8.622			
3114-55-4	Chlorobenzene-d5	376000	11.42			
3855-82-1	1,4-Dichlorobenzene-d4	137000	13.353			

U = Not Detected

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J = Estimated Value

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Report of Analysis

Client:	CDM Smith	Date Collected:
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:
Client Sample ID:	VX0521WBS01	SDG No.: Q2075
Lab Sample ID:	VX0521WBS01	Matrix: Water
Analytical Method:	8260D	% Solid: 0
Sample Wt/Vol:	5 mL	Final Vol: 5000 uL
Soil Aliquot Vol:	uL	Test: VOCMS Group3
GC Column:	DB-624UI ID : 0.18	Level : LOW
Prep Method :		

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046287.D	1		05/21/25 11:09	VX052125

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	20.3		0.15	1.00	ug/L
108-88-3	Toluene	20.5		0.14	1.00	ug/L
100-41-4	Ethyl Benzene	20.1		0.13	1.00	ug/L
1330-20-7	Total Xylenes	61.2		0.36	3.00	ug/L
98-82-8	Isopropylbenzene	20.4		0.12	1.00	ug/L
103-65-1	n-propylbenzene	20.4		0.13	1.00	ug/L
108-67-8	1,3,5-Trimethylbenzene	21.1		0.15	1.00	ug/L
98-06-6	tert-Butylbenzene	20.2		0.14	1.00	ug/L
95-63-6	1,2,4-Trimethylbenzene	20.7		0.14	1.00	ug/L
135-98-8	sec-Butylbenzene	20.3		0.13	1.00	ug/L
99-87-6	p-Isopropyltoluene	20.5		0.13	1.00	ug/L
104-51-8	n-Butylbenzene	19.7		0.15	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	53.7		74 - 125	107%	SPK: 50
1868-53-7	Dibromofluoromethane	54.3		75 - 124	109%	SPK: 50
2037-26-5	Toluene-d8	52.4		86 - 113	105%	SPK: 50
460-00-4	4-Bromofluorobenzene	55.0		77 - 121	110%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	83100	5.544			
540-36-3	1,4-Difluorobenzene	147000	6.757			
3114-55-4	Chlorobenzene-d5	130000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	62700	12.018			

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A = Aldol-Condensation Reaction Products

Report of Analysis

Client:	CDM Smith	Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	
Client Sample ID:	VX0523MBS01	SDG No.:	Q2075
Lab Sample ID:	VX0523MBS01	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	100
Sample Wt/Vol:	5	Units:	g
Soil Aliquot Vol:	100	uL	
GC Column:	DB-624UI	ID :	0.18
Prep Method :		Level :	MED

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046335.D	1		05/23/25 11:23	VX052325

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	2000		79.0	500	ug/Kg
108-88-3	Toluene	2000		78.0	500	ug/Kg
100-41-4	Ethyl Benzene	2000		67.0	500	ug/Kg
1330-20-7	Total Xylenes	6100		202	1500	ug/Kg
98-82-8	Isopropylbenzene	2000		78.0	500	ug/Kg
103-65-1	n-propylbenzene	1900		73.0	500	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	2100		82.0	500	ug/Kg
98-06-6	tert-Butylbenzene	2000		67.0	500	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	2000		64.0	500	ug/Kg
135-98-8	sec-Butylbenzene	2000		66.0	500	ug/Kg
99-87-6	p-Isopropyltoluene	2000		62.0	500	ug/Kg
104-51-8	n-Butylbenzene	1900		150	500	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	52.0		63 - 155	104%	SPK: 50
1868-53-7	Dibromofluoromethane	52.4		70 - 134	105%	SPK: 50
2037-26-5	Toluene-d8	50.3		74 - 123	101%	SPK: 50
460-00-4	4-Bromofluorobenzene	51.8		38 - 136	104%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	83300	5.544			
540-36-3	1,4-Difluorobenzene	146000	6.757			
3114-55-4	Chlorobenzene-d5	129000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	63300	12.018			

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Report of Analysis

Client:	CDM Smith			Date Collected:
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:
Client Sample ID:	VY0520SBS01		SDG No.:	Q2075
Lab Sample ID:	VY0520SBS01		Matrix:	SOIL
Analytical Method:	8260D		% Solid:	100
Sample Wt/Vol:	5	Units: g	Final Vol:	5000 uL
Soil Aliquot Vol:	uL		Test:	VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level :	LOW
Prep Method :				

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022331.D	1		05/20/25 10:21	VY052025

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	20.8	0.79		5.00	ug/Kg
108-88-3	Toluene	20.0	0.78		5.00	ug/Kg
100-41-4	Ethyl Benzene	19.8	0.67		5.00	ug/Kg
1330-20-7	Total Xylenes	59.8	2.02		15.0	ug/Kg
98-82-8	Isopropylbenzene	20.1	0.78		5.00	ug/Kg
103-65-1	n-propylbenzene	20.3	0.73		5.00	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	19.5	0.82		5.00	ug/Kg
98-06-6	tert-Butylbenzene	19.8	0.67		5.00	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	19.3	0.64		5.00	ug/Kg
135-98-8	sec-Butylbenzene	20.0	0.66		5.00	ug/Kg
99-87-6	p-Isopropyltoluene	19.3	0.62		5.00	ug/Kg
104-51-8	n-Butylbenzene	19.7	1.50		5.00	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	52.8	63 - 155		106%	SPK: 50
1868-53-7	Dibromofluoromethane	52.0	70 - 134		104%	SPK: 50
2037-26-5	Toluene-d8	51.5	74 - 123		103%	SPK: 50
460-00-4	4-Bromofluorobenzene	47.7	38 - 136		95%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	162000	7.707			
540-36-3	1,4-Difluorobenzene	281000	8.616			
3114-55-4	Chlorobenzene-d5	237000	11.42			
3855-82-1	1,4-Dichlorobenzene-d4	111000	13.347			

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Report of Analysis

Client:	CDM Smith			Date Collected:
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:
Client Sample ID:	VY0522SBS01		SDG No.:	Q2075
Lab Sample ID:	VY0522SBS01		Matrix:	SOIL
Analytical Method:	8260D		% Solid:	100
Sample Wt/Vol:	5	Units: g	Final Vol:	5000 uL
Soil Aliquot Vol:	uL		Test:	VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level :	LOW
Prep Method :				

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022382.D	1		05/22/25 09:36	VY052225

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	19.5	0.79		5.00	ug/Kg
108-88-3	Toluene	19.4	0.78		5.00	ug/Kg
100-41-4	Ethyl Benzene	18.7	0.67		5.00	ug/Kg
1330-20-7	Total Xylenes	56.8	2.02		15.0	ug/Kg
98-82-8	Isopropylbenzene	19.3	0.78		5.00	ug/Kg
103-65-1	n-propylbenzene	19.4	0.73		5.00	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	18.9	0.82		5.00	ug/Kg
98-06-6	tert-Butylbenzene	19.2	0.67		5.00	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	18.8	0.64		5.00	ug/Kg
135-98-8	sec-Butylbenzene	19.4	0.66		5.00	ug/Kg
99-87-6	p-Isopropyltoluene	19.0	0.62		5.00	ug/Kg
104-51-8	n-Butylbenzene	19.3	1.50		5.00	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	48.4	63 - 155		97%	SPK: 50
1868-53-7	Dibromofluoromethane	50.3	70 - 134		101%	SPK: 50
2037-26-5	Toluene-d8	50.3	74 - 123		101%	SPK: 50
460-00-4	4-Bromofluorobenzene	47.7	38 - 136		95%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	175000	7.719			
540-36-3	1,4-Difluorobenzene	294000	8.622			
3114-55-4	Chlorobenzene-d5	251000	11.42			
3855-82-1	1,4-Dichlorobenzene-d4	118000	13.353			

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Report of Analysis

Client:	CDM Smith	Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	
Client Sample ID:	VX0521WBSD01	SDG No.:	Q2075
Lab Sample ID:	VX0521WBSD01	Matrix:	Water
Analytical Method:	8260D	% Solid:	0
Sample Wt/Vol:	5	Units:	mL
Soil Aliquot Vol:		uL	
GC Column:	DB-624UI	ID :	0.18
Prep Method :		Level :	LOW

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VX046288.D	1		05/21/25 11:36	VX052125

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
71-43-2	Benzene	20.4		0.15	1.00	ug/L
108-88-3	Toluene	21.1		0.14	1.00	ug/L
100-41-4	Ethyl Benzene	20.4		0.13	1.00	ug/L
1330-20-7	Total Xylenes	62.1		0.36	3.00	ug/L
98-82-8	Isopropylbenzene	20.8		0.12	1.00	ug/L
103-65-1	n-propylbenzene	20.7		0.13	1.00	ug/L
108-67-8	1,3,5-Trimethylbenzene	21.0		0.15	1.00	ug/L
98-06-6	tert-Butylbenzene	20.5		0.14	1.00	ug/L
95-63-6	1,2,4-Trimethylbenzene	20.9		0.14	1.00	ug/L
135-98-8	sec-Butylbenzene	21.1		0.13	1.00	ug/L
99-87-6	p-Isopropyltoluene	20.7		0.13	1.00	ug/L
104-51-8	n-Butylbenzene	20.2		0.15	1.00	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	54.2		74 - 125	108%	SPK: 50
1868-53-7	Dibromofluoromethane	55.1		75 - 124	110%	SPK: 50
2037-26-5	Toluene-d8	54.1		86 - 113	108%	SPK: 50
460-00-4	4-Bromofluorobenzene	55.8		77 - 121	112%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	82700	5.544			
540-36-3	1,4-Difluorobenzene	142000	6.757			
3114-55-4	Chlorobenzene-d5	128000	10.049			
3855-82-1	1,4-Dichlorobenzene-d4	61000	12.018			

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Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-11MS	SDG No.:	Q2075
Lab Sample ID:	Q2075-04MS	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	87.4
Sample Wt/Vol:	8.42	Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022350.D	1		05/20/25 18:04	VY052025

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	31.6		0.54	3.40	ug/Kg
108-88-3	Toluene	40.1		0.53	3.40	ug/Kg
100-41-4	Ethyl Benzene	46.5		0.46	3.40	ug/Kg
1330-20-7	Total Xylenes	202		1.40	10.2	ug/Kg
98-82-8	Isopropylbenzene	34.0		0.53	3.40	ug/Kg
103-65-1	n-propylbenzene	39.5		0.50	3.40	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	47.6		0.56	3.40	ug/Kg
98-06-6	tert-Butylbenzene	28.7		0.46	3.40	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	120	E	0.43	3.40	ug/Kg
135-98-8	sec-Butylbenzene	30.9		0.45	3.40	ug/Kg
99-87-6	p-Isopropyltoluene	29.0		0.42	3.40	ug/Kg
104-51-8	n-Butylbenzene	30.5		0.99	3.40	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	49.2		63 - 155	98%	SPK: 50
1868-53-7	Dibromofluoromethane	42.1		70 - 134	84%	SPK: 50
2037-26-5	Toluene-d8	39.0		74 - 123	78%	SPK: 50
460-00-4	4-Bromofluorobenzene	36.5		38 - 136	73%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	135000		7.707		
540-36-3	1,4-Difluorobenzene	249000		8.616		
3114-55-4	Chlorobenzene-d5	211000		11.414		
3855-82-1	1,4-Dichlorobenzene-d4	91800		13.346		

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Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-11MSD	SDG No.:	Q2075
Lab Sample ID:	Q2075-05MSD	Matrix:	SOIL
Analytical Method:	8260D	% Solid:	87.4
Sample Wt/Vol:	8.33	Units: g	Final Vol: 5000 uL
Soil Aliquot Vol:		uL	Test: VOCMS Group3
GC Column:	RXI-624	ID : 0.25	Level : LOW
Prep Method :			

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
VY022351.D	1		05/20/25 18:26	VY052025

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
71-43-2	Benzene	43.1	0.54		3.40	ug/Kg
108-88-3	Toluene	52.7	0.54		3.40	ug/Kg
100-41-4	Ethyl Benzene	54.6	0.46		3.40	ug/Kg
1330-20-7	Total Xylenes	219	1.41		10.3	ug/Kg
98-82-8	Isopropylbenzene	40.5	0.54		3.40	ug/Kg
103-65-1	n-propylbenzene	43.6	0.50		3.40	ug/Kg
108-67-8	1,3,5-Trimethylbenzene	49.0	0.56		3.40	ug/Kg
98-06-6	tert-Butylbenzene	37.7	0.46		3.40	ug/Kg
95-63-6	1,2,4-Trimethylbenzene	96.4	0.44		3.40	ug/Kg
135-98-8	sec-Butylbenzene	37.6	0.45		3.40	ug/Kg
99-87-6	p-Isopropyltoluene	36.2	0.43		3.40	ug/Kg
104-51-8	n-Butylbenzene	36.4	1.00		3.40	ug/Kg
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	62.7		63 - 155	125%	SPK: 50
1868-53-7	Dibromofluoromethane	51.3		70 - 134	103%	SPK: 50
2037-26-5	Toluene-d8	45.1		74 - 123	90%	SPK: 50
460-00-4	4-Bromofluorobenzene	45.4		38 - 136	91%	SPK: 50
INTERNAL STANDARDS						
363-72-4	Pentafluorobenzene	118000	7.707			
540-36-3	1,4-Difluorobenzene	220000	8.616			
3114-55-4	Chlorobenzene-d5	197000	11.414			
3855-82-1	1,4-Dichlorobenzene-d4	91200	13.346			

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A
B
C
D
E
F
G

CALIBRATION

SUMMARY

VOLATILE ORGANICS INITIAL CALIBRATION DATA

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	SAS No.:	Q2075
Instrument ID:	MSVOA_X	SDG No.:	Q2075
Heated Purge:	(Y/N) N	Calibration Date(s):	05/05/2025
GC Column:	DB-624UI	Calibration Time(s):	11:35 16:27
	ID: 0.18 (mm)		

LAB FILE ID:	RRF020 = VX046041.D	RRF050 = VX046042.D	RRF100 = VX046043.D					
COMPOUND	RRF020	RRF050	RRF100	RRF150	RRF005	RRF001	RRF	% RSD
Benzene	1.426	1.474	1.441	1.477	1.337	1.348	1.417	4.3
Toluene	0.884	0.898	0.885	0.904	0.838	0.803	0.869	4.5
Ethyl Benzene	1.919	2.022	1.979	2.036	1.816	1.803	1.929	5.2
m/p-Xylenes	0.706	0.740	0.721	0.740	0.678	0.648	0.706	5.2
o-Xylene	0.688	0.727	0.706	0.726	0.639	0.642	0.688	5.7
Isopropylbenzene	3.843	4.130	3.876	4.156	3.562	3.789	3.893	5.7
n-propylbenzene	4.394	4.854	4.583	4.868	4.186	4.272	4.526	6.4
1,3,5-Trimethylbenzene	3.275	3.488	3.255	3.405	3.053	3.036	3.252	5.6
tert-Butylbenzene	3.115	3.435	3.255	3.411	3.098	3.341	3.276	4.4
1,2,4-Trimethylbenzene	3.274	3.522	3.335	3.444	3.034	3.150	3.293	5.5
sec-Butylbenzene	3.937	4.282	4.095	4.343	3.767	3.708	4.022	6.6
p-Isopropyltoluene	3.206	3.555	3.450	3.599	3.084	3.025	3.320	7.4
n-Butylbenzene	2.748	3.147	3.139	3.346	2.650	2.443	2.912	12
1,2-Dichloroethane-d4	0.953	0.910	0.930	0.932	0.935		0.932	1.6
Dibromofluoromethane	0.359	0.355	0.364	0.368	0.354		0.360	1.7
Toluene-d8	1.246	1.223	1.266	1.275	1.221		1.246	2
4-Bromofluorobenzene	0.455	0.470	0.500	0.500	0.464		0.478	4.4

* 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.

VOLATILE ORGANICS INITIAL CALIBRATION DATA

Lab Name:	CHEMTECH	Contract:	CAMP02
Lab Code:	CHEM	SAS No.:	Q2075
Instrument ID:	MSVOA_Y	SDG No.:	Q2075
Heated Purge:	(Y/N) Y	Calibration Date(s):	05/15/2025
GC Column:	RXI-624	Calibration Time(s):	09:46 11:47
	ID: 0.25 (mm)		

LAB FILE ID:	RRF005 = VY022253.D	RRF010 = VY022254.D	RRF020 = VY022255.D	RRF050 = VY022256.D	RRF100 = VY022257.D	RRF150 = VY022258.D	RRF	% RSD
COMPOUND	RRF005	RRF010	RRF020	RRF050	RRF100	RRF150		
Benzene	1.369	1.462	1.435	1.499	1.482	1.496	1.457	3.4
Toluene	0.871	0.924	0.890	0.944	0.947	0.961	0.923	3.8
Ethyl Benzene	2.014	2.043	2.049	2.175	2.173	2.222	2.113	4.1
m/p-Xylenes	0.749	0.770	0.763	0.818	0.825	0.852	0.796	5.2
o-Xylene	0.704	0.741	0.723	0.768	0.782	0.802	0.753	4.9
Isopropylbenzene	4.157	4.192	4.100	4.197	4.028	4.181	4.142	1.6
n-propylbenzene	5.031	5.047	4.904	5.127	4.865	5.006	4.996	1.9
1,3,5-Trimethylbenzene	3.393	3.386	3.275	3.410	3.251	3.349	3.344	2
tert-Butylbenzene	3.037	2.974	2.900	3.029	2.879	3.014	2.972	2.3
1,2,4-Trimethylbenzene	3.189	3.349	3.206	3.413	3.329	3.431	3.319	3.1
sec-Butylbenzene	4.390	4.461	4.318	4.573	4.307	4.421	4.412	2.2
p-Isopropyltoluene	3.598	3.610	3.573	3.818	3.754	3.911	3.710	3.7
n-Butylbenzene	3.419	3.500	3.491	3.668	3.452	3.533	3.510	2.5
1,2-Dichloroethane-d4	0.547	0.559	0.527	0.541	0.545	0.560	0.546	2.2
Dibromofluoromethane	0.294	0.294	0.294	0.301	0.297	0.310	0.298	2.2
Toluene-d8	1.224	1.200	1.195	1.230	1.214	1.272	1.222	2.3
4-Bromofluorobenzene	0.390	0.386	0.368	0.387	0.387	0.407	0.387	3.3

- * 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.

VOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH	Contract:	CAMP02				
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075	SDG No.:	Q2075
Instrument ID:	MSVOA_X	Calibration Date/Time:				05/21/2025	09:55
Lab File ID:	VX046284.D	Init. Calib. Date(s):				05/05/2025	05/05/2025
Heated Purge:	(Y/N) N	Init. Calib. Time(s):				11:35	16:27
GC Column:	DB-624UI	ID:	0.18	(mm)			

COMPOUND	RRF	RRF050	MIN RRF	%D	MAX%D
Benzene	1.417	1.412		-0.35	20
Toluene	0.869	0.857		-1.38	20
Ethyl Benzene	1.929	1.968		2.02	20
m/p-Xylenes	0.706	0.721		2.13	20
o-Xylene	0.688	0.712		3.49	20
Isopropylbenzene	3.893	4.007		2.93	20
n-propylbenzene	4.526	4.671		3.2	20
1,3,5-Trimethylbenzene	3.252	3.344		2.83	20
tert-Butylbenzene	3.276	3.302		0.79	20
1,2,4-Trimethylbenzene	3.293	3.417		3.77	20
sec-Butylbenzene	4.022	4.193		4.25	20
p-Isopropyltoluene	3.320	3.467		4.43	20
n-Butylbenzene	2.912	3.134		7.62	20
1,2-Dichloroethane-d4	0.932	0.913		-2.04	20
Dibromofluoromethane	0.360	0.371		3.06	20
Toluene-d8	1.246	1.223		-1.85	20
4-Bromofluorobenzene	0.478	0.481		0.63	20

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

VOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH		Contract:	CAMP02	
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075
Instrument ID:	MSVOA_X		Calibration Date/Time:	05/23/2025	09:24
Lab File ID:	VX046331.D		Init. Calib. Date(s):	05/05/2025	05/05/2025
Heated Purge:	(Y/N)	N	Init. Calib. Time(s):	11:35	16:27
GC Column:	DB-624UI	ID: 0.18 (mm)			

COMPOUND	RRF	RRF050	MIN RRF	%D	MAX%D
Benzene	1.417	1.421		0.28	20
Toluene	0.869	0.878		1.04	20
Ethyl Benzene	1.929	1.985		2.9	20
m/p-Xylenes	0.706	0.719		1.84	20
o-Xylene	0.688	0.714		3.78	20
Isopropylbenzene	3.893	3.986		2.39	20
n-propylbenzene	4.526	4.678		3.36	20
1,3,5-Trimethylbenzene	3.252	3.374		3.75	20
tert-Butylbenzene	3.276	3.334		1.77	20
1,2,4-Trimethylbenzene	3.293	3.388		2.88	20
sec-Butylbenzene	4.022	4.204		4.53	20
p-Isopropyltoluene	3.320	3.484		4.94	20
n-Butylbenzene	2.912	3.156		8.38	20
1,2-Dichloroethane-d4	0.932	0.966		3.65	20
Dibromofluoromethane	0.360	0.384		6.67	20
Toluene-d8	1.246	1.307		4.9	20
4-Bromofluorobenzene	0.478	0.514		7.53	20

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

VOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH	Contract:	CAMP02				
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075	SDG No.:	Q2075
Instrument ID:	MSVOA_Y	Calibration Date/Time:				05/20/2025	08:40
Lab File ID:	VY022329.D	Init. Calib. Date(s):				05/15/2025	05/15/2025
Heated Purge: (Y/N)	Y	Init. Calib. Time(s):				09:46	11:47
GC Column:	RXI-624	ID:	0.25	(mm)			

COMPOUND	RRF	RRF050	MIN RRF	%D	MAX%D
Benzene	1.457	1.473		1.1	20
Toluene	0.923	0.912		-1.19	20
Ethyl Benzene	2.113	2.120		0.33	20
m/p-Xylenes	0.796	0.803		0.88	20
o-Xylene	0.753	0.738		-1.99	20
Isopropylbenzene	4.142	4.147		0.12	20
n-propylbenzene	4.996	5.062		1.32	20
1,3,5-Trimethylbenzene	3.344	3.297		-1.41	20
tert-Butylbenzene	2.972	2.924		-1.62	20
1,2,4-Trimethylbenzene	3.319	3.232		-2.62	20
sec-Butylbenzene	4.412	4.438		0.59	20
p-Isopropyltoluene	3.710	3.717		0.19	20
n-Butylbenzene	3.510	3.562		1.48	20
1,2-Dichloroethane-d4	0.546	0.533		-2.38	20
Dibromofluoromethane	0.298	0.305		2.35	20
Toluene-d8	1.222	1.259		3.03	20
4-Bromofluorobenzene	0.387	0.375		-3.1	20

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

VOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH	Contract:	CAMP02				
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075	SDG No.:	Q2075
Instrument ID:	MSVOA_Y	Calibration Date/Time:			05/22/2025	08:35	
Lab File ID:	VY022380.D	Init. Calib. Date(s):			05/15/2025	05/15/2025	
Heated Purge: (Y/N)	Y	Init. Calib. Time(s):			09:46	11:47	
GC Column:	RXI-624	ID:	0.25	(mm)			

COMPOUND	RRF	RRF050	MIN RRF	%D	MAX%D
Benzene	1.457	1.548		6.25	20
Toluene	0.923	0.978		5.96	20
Ethyl Benzene	2.113	2.234		5.73	20
m/p-Xylenes	0.796	0.849		6.66	20
o-Xylene	0.753	0.791		5.05	20
Isopropylbenzene	4.142	4.432		7	20
n-propylbenzene	4.996	5.385		7.79	20
1,3,5-Trimethylbenzene	3.344	3.533		5.65	20
tert-Butylbenzene	2.972	3.200		7.67	20
1,2,4-Trimethylbenzene	3.319	3.526		6.24	20
sec-Butylbenzene	4.412	4.835		9.59	20
p-Isopropyltoluene	3.710	4.077		9.89	20
n-Butylbenzene	3.510	3.887		10.74	20
1,2-Dichloroethane-d4	0.546	0.561		2.75	20
Dibromofluoromethane	0.298	0.318		6.71	20
Toluene-d8	1.222	1.298		6.22	20
4-Bromofluorobenzene	0.387	0.387		0	20

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

LAB CHRONICLE

OrderID:	Q2075	OrderDate:	5/16/2025 4:10:00 PM
Client:	CDM Smith	Project:	Con Ed UTEN Mount Vernon, NY
Contact:	Marcie Ann Encinas	Location:	L41, VOA Ref. #2 Soil, VOA Ref. #3 Water

LabID	ClientID	Matrix	Test	Method	Sample Date	Prep Date	Anal Date	Received
Q2075-01	SS-10	SOIL	SVOCMS Group3	8270E	05/15/25	05/20/25	05/22/25	05/16/25
Q2075-02	SS-910	SOIL	SVOCMS Group3	8270E	05/15/25	05/20/25	05/22/25	05/16/25
Q2075-03	SS-11	SOIL	SVOCMS Group3	8270E	05/15/25	05/20/25	05/21/25	05/16/25
Q2075-06	SS-MW1-11.5	SOIL	SVOCMS Group3	8270E	05/16/25	05/20/25	05/21/25	05/16/25
Q2075-07	FB-05152025	Water	SVOCMS Group3	8270E	05/15/25	05/21/25	05/22/25	05/16/25
Q2075-08	FB-05162025	Water	SVOCMS Group3	8270E	05/16/25	05/21/25	05/21/25	05/16/25



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

Hit Summary Sheet SW-846

SDG No.: Q2075

Client: CDM Smith

Sample ID	Client ID	Matrix	Parameter	Concentration	C	MDL	RDL	Units
Client ID :	SS-10							
Q2075-01	SS-10	SOIL	Phenanthrene	830.000	26.2	210	ug/Kg	
Q2075-01	SS-10	SOIL	Anthracene	110.000 J	41.7	210	ug/Kg	
Q2075-01	SS-10	SOIL	Fluoranthene	1,200.000	37.6	210	ug/Kg	
Q2075-01	SS-10	SOIL	Pyrene	660.000	45.1	210	ug/Kg	
Q2075-01	SS-10	SOIL	Benzo(a)anthracene	420.000	28.8	210	ug/Kg	
Q2075-01	SS-10	SOIL	Chrysene	490.000	24.9	210	ug/Kg	
Q2075-01	SS-10	SOIL	Benzo(b)fluoranthene	640.000	23.8	210	ug/Kg	
Q2075-01	SS-10	SOIL	Benzo(k)fluoranthene	220.000	28.1	210	ug/Kg	
Q2075-01	SS-10	SOIL	Benzo(a)pyrene	350.000	37	210	ug/Kg	
Q2075-01	SS-10	SOIL	Indeno(1,2,3-cd)pyrene	160.000 J	36.5	210	ug/Kg	
Q2075-01	SS-10	SOIL	Benzo(g,h,i)perylene	170.000 J	32.2	210	ug/Kg	
Total Svoc :				5,250.00				
Total Concentration:				5,250.00				
Client ID :	SS-910							
Q2075-02	SS-910	SOIL	Phenanthrene	440.000	25.7	210	ug/Kg	
Q2075-02	SS-910	SOIL	Fluoranthene	940.000	36.8	210	ug/Kg	
Q2075-02	SS-910	SOIL	Pyrene	490.000	44.2	210	ug/Kg	
Q2075-02	SS-910	SOIL	Benzo(a)anthracene	290.000	28.2	210	ug/Kg	
Q2075-02	SS-910	SOIL	Chrysene	320.000	24.4	210	ug/Kg	
Q2075-02	SS-910	SOIL	Benzo(b)fluoranthene	520.000	23.3	210	ug/Kg	
Q2075-02	SS-910	SOIL	Benzo(k)fluoranthene	180.000 J	27.5	210	ug/Kg	
Q2075-02	SS-910	SOIL	Benzo(a)pyrene	310.000	36.2	210	ug/Kg	
Q2075-02	SS-910	SOIL	Indeno(1,2,3-cd)pyrene	150.000 J	35.7	210	ug/Kg	
Q2075-02	SS-910	SOIL	Benzo(g,h,i)perylene	170.000 J	31.5	210	ug/Kg	
Total Svoc :				3,810.00				
Total Concentration:				3,810.00				
Client ID :	SS-11							
Q2075-03	SS-11	SOIL	Fluoranthene	120.000 J	34.3	190	ug/Kg	
Total Svoc :				120.00				
Total Concentration:				120.00				



A
B
C
D
E
F
G

SAMPLE DATA

Report of Analysis

Client:	CDM Smith			Date Collected:	05/15/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	SS-10			SDG No.:	Q2075	
Lab Sample ID:	Q2075-01			Matrix:	SOIL	
Analytical Method:	8270E			% Solid:	79.8	
Sample Wt/Vol:	30.01	Units:	g	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3541					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142510.D	1	05/20/25 09:45	05/22/25 15:07	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	210	U	28.4	210	ug/Kg
208-96-8	Acenaphthylene	210	U	36.2	210	ug/Kg
83-32-9	Acenaphthene	210	U	26.7	210	ug/Kg
86-73-7	Fluorene	210	U	31.7	210	ug/Kg
85-01-8	Phenanthrene	830		26.2	210	ug/Kg
120-12-7	Anthracene	110	J	41.7	210	ug/Kg
206-44-0	Fluoranthene	1200		37.6	210	ug/Kg
129-00-0	Pyrene	660		45.1	210	ug/Kg
56-55-3	Benzo(a)anthracene	420		28.8	210	ug/Kg
218-01-9	Chrysene	490		24.9	210	ug/Kg
205-99-2	Benzo(b)fluoranthene	640		23.8	210	ug/Kg
207-08-9	Benzo(k)fluoranthene	220		28.1	210	ug/Kg
50-32-8	Benzo(a)pyrene	350		37.0	210	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	160	J	36.5	210	ug/Kg
53-70-3	Dibenz(a,h)anthracene	210	U	34.3	210	ug/Kg
191-24-2	Benzo(g,h,i)perylene	170	J	32.2	210	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	43.6		18 - 107	44%	SPK: 100
321-60-8	2-Fluorobiphenyl	37.5		20 - 109	38%	SPK: 100
1718-51-0	Terphenyl-d14	29.7		10 - 105	30%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	120000	6.898			
1146-65-2	Naphthalene-d8	453000	8.18			
15067-26-2	Acenaphthene-d10	223000	9.933			
1517-22-2	Phenanthrene-d10	308000	11.421			
1719-03-5	Chrysene-d12	229000	14.062			
1520-96-3	Perylene-d12	198000	15.556			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-10	SDG No.:	Q2075
Lab Sample ID:	Q2075-01	Matrix:	SOIL
Analytical Method:	8270E	% Solid:	79.8
Sample Wt/Vol:	30.01	Units:	g
Soil Aliquot Vol:		uL	
Extraction Type :		Decanted :	N
Injection Volume :		GPC Factor :	1.0
Prep Method :	SW3541	GPC Cleanup :	N
		Level :	LOW
		PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142510.D	1	05/20/25 09:45	05/22/25 15:07	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	05/15/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	SS-910			SDG No.:	Q2075	
Lab Sample ID:	Q2075-02			Matrix:	SOIL	
Analytical Method:	8270E			% Solid:	81.3	
Sample Wt/Vol:	30.06	Units:	g	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3541					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142511.D	1	05/20/25 09:45	05/22/25 15:36	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	210	U	27.9	210	ug/Kg
208-96-8	Acenaphthylene	210	U	35.5	210	ug/Kg
83-32-9	Acenaphthene	210	U	26.1	210	ug/Kg
86-73-7	Fluorene	210	U	31.1	210	ug/Kg
85-01-8	Phenanthrene	440		25.7	210	ug/Kg
120-12-7	Anthracene	210	U	40.9	210	ug/Kg
206-44-0	Fluoranthene	940		36.8	210	ug/Kg
129-00-0	Pyrene	490		44.2	210	ug/Kg
56-55-3	Benzo(a)anthracene	290		28.2	210	ug/Kg
218-01-9	Chrysene	320		24.4	210	ug/Kg
205-99-2	Benzo(b)fluoranthene	520		23.3	210	ug/Kg
207-08-9	Benzo(k)fluoranthene	180	J	27.5	210	ug/Kg
50-32-8	Benzo(a)pyrene	310		36.2	210	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	150	J	35.7	210	ug/Kg
53-70-3	Dibenz(a,h)anthracene	210	U	33.6	210	ug/Kg
191-24-2	Benzo(g,h,i)perylene	170	J	31.5	210	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	49.5		18 - 107	49%	SPK: 100
321-60-8	2-Fluorobiphenyl	43.1		20 - 109	43%	SPK: 100
1718-51-0	Terphenyl-d14	35.4		10 - 105	35%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	117000	6.898			
1146-65-2	Naphthalene-d8	418000	8.181			
15067-26-2	Acenaphthene-d10	191000	9.934			
1517-22-2	Phenanthrene-d10	271000	11.422			
1719-03-5	Chrysene-d12	225000	14.063			
1520-96-3	Perylene-d12	177000	15.557			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-910	SDG No.:	Q2075
Lab Sample ID:	Q2075-02	Matrix:	SOIL
Analytical Method:	8270E	% Solid:	81.3
Sample Wt/Vol:	30.06	Units:	g uL
Soil Aliquot Vol:		uL	Test: SVOCMS Group3
Extraction Type :		Decanted :	N Level : LOW
Injection Volume :		GPC Factor :	1.0 GPC Cleanup : N PH :
Prep Method :	SW3541		

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142511.D	1	05/20/25 09:45	05/22/25 15:36	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	05/15/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	SS-11			SDG No.:	Q2075	
Lab Sample ID:	Q2075-03			Matrix:	SOIL	
Analytical Method:	8270E			% Solid:	87.4	
Sample Wt/Vol:	30.04	Units:	g	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3541					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142486.D	1	05/20/25 09:45	05/21/25 14:37	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	190	U	25.9	190	ug/Kg
208-96-8	Acenaphthylene	190	U	33.0	190	ug/Kg
83-32-9	Acenaphthene	190	U	24.3	190	ug/Kg
86-73-7	Fluorene	190	U	28.9	190	ug/Kg
85-01-8	Phenanthrene	190	U	23.9	190	ug/Kg
120-12-7	Anthracene	190	U	38.1	190	ug/Kg
206-44-0	Fluoranthene	120	J	34.3	190	ug/Kg
129-00-0	Pyrene	190	U	41.1	190	ug/Kg
56-55-3	Benzo(a)anthracene	190	U	26.3	190	ug/Kg
218-01-9	Chrysene	190	U	22.7	190	ug/Kg
205-99-2	Benzo(b)fluoranthene	190	U	21.7	190	ug/Kg
207-08-9	Benzo(k)fluoranthene	190	U	25.6	190	ug/Kg
50-32-8	Benzo(a)pyrene	190	U	33.7	190	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	190	U	33.3	190	ug/Kg
53-70-3	Dibenz(a,h)anthracene	190	U	31.3	190	ug/Kg
191-24-2	Benzo(g,h,i)perylene	190	U	29.4	190	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	64.7		18 - 107	65%	SPK: 100
321-60-8	2-Fluorobiphenyl	56.2		20 - 109	56%	SPK: 100
1718-51-0	Terphenyl-d14	42.2		10 - 105	42%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	63600	6.898			
1146-65-2	Naphthalene-d8	239000	8.181			
15067-26-2	Acenaphthene-d10	132000	9.939			
1517-22-2	Phenanthrene-d10	251000	11.427			
1719-03-5	Chrysene-d12	246000	14.068			
1520-96-3	Perylene-d12	209000	15.557			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-11	SDG No.:	Q2075
Lab Sample ID:	Q2075-03	Matrix:	SOIL
Analytical Method:	8270E	% Solid:	87.4
Sample Wt/Vol:	30.04	Units:	g
Soil Aliquot Vol:		uL	
Extraction Type :		Decanted :	N
Injection Volume :		GPC Factor :	1.0
Prep Method :	SW3541	GPC Cleanup :	N
		Level :	LOW
		PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142486.D	1	05/20/25 09:45	05/21/25 14:37	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	05/16/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	SS-MW1-11.5			SDG No.:	Q2075	
Lab Sample ID:	Q2075-06			Matrix:	SOIL	
Analytical Method:	8270E			% Solid:	91	
Sample Wt/Vol:	30.06	Units:	g	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3541					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142489.D	1	05/20/25 09:45	05/21/25 16:03	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	190	U	24.9	190	ug/Kg
208-96-8	Acenaphthylene	190	U	31.7	190	ug/Kg
83-32-9	Acenaphthene	190	U	23.4	190	ug/Kg
86-73-7	Fluorene	190	U	27.7	190	ug/Kg
85-01-8	Phenanthrene	190	U	22.9	190	ug/Kg
120-12-7	Anthracene	190	U	36.5	190	ug/Kg
206-44-0	Fluoranthene	190	U	32.9	190	ug/Kg
129-00-0	Pyrene	190	U	39.5	190	ug/Kg
56-55-3	Benzo(a)anthracene	190	U	25.2	190	ug/Kg
218-01-9	Chrysene	190	U	21.8	190	ug/Kg
205-99-2	Benzo(b)fluoranthene	190	U	20.8	190	ug/Kg
207-08-9	Benzo(k)fluoranthene	190	U	24.6	190	ug/Kg
50-32-8	Benzo(a)pyrene	190	U	32.4	190	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	190	U	31.9	190	ug/Kg
53-70-3	Dibenz(a,h)anthracene	190	U	30.0	190	ug/Kg
191-24-2	Benzo(g,h,i)perylene	190	U	28.2	190	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	53.3		18 - 107	53%	SPK: 100
321-60-8	2-Fluorobiphenyl	48.6		20 - 109	49%	SPK: 100
1718-51-0	Terphenyl-d14	38.9		10 - 105	39%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	59300	6.898			
1146-65-2	Naphthalene-d8	207000	8.181			
15067-26-2	Acenaphthene-d10	104000	9.939			
1517-22-2	Phenanthrene-d10	179000	11.422			
1719-03-5	Chrysene-d12	153000	14.063			
1520-96-3	Perylene-d12	126000	15.557			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/16/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-MW1-11.5	SDG No.:	Q2075
Lab Sample ID:	Q2075-06	Matrix:	SOIL
Analytical Method:	8270E	% Solid:	91
Sample Wt/Vol:	30.06	Units:	g
Soil Aliquot Vol:		uL	
Extraction Type :		Decanted :	N
Injection Volume :		GPC Factor :	1.0
Prep Method :	SW3541	GPC Cleanup :	N
		Level :	LOW
		PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142489.D	1	05/20/25 09:45	05/21/25 16:03	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

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B = Analyte Found in Associated Method Blank

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() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Report of Analysis

Client:	CDM Smith			Date Collected:	05/15/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	FB-05152025			SDG No.:	Q2075	
Lab Sample ID:	Q2075-07			Matrix:	Water	
Analytical Method:	8270E			% Solid:	0	
Sample Wt/Vol:	810	Units:	mL	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3510C					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024758.D	1	05/21/25 08:40	05/22/25 14:19	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
91-20-3	Naphthalene	6.20	U	0.62	6.20	ug/L
208-96-8	Acenaphthylene	6.20	U	0.93	6.20	ug/L
83-32-9	Acenaphthene	6.20	U	0.68	6.20	ug/L
86-73-7	Fluorene	6.20	U	0.78	6.20	ug/L
85-01-8	Phenanthrene	6.20	U	0.62	6.20	ug/L
120-12-7	Anthracene	6.20	U	0.75	6.20	ug/L
206-44-0	Fluoranthene	6.20	U	1.00	6.20	ug/L
129-00-0	Pyrene	6.20	U	0.62	6.20	ug/L
56-55-3	Benzo(a)anthracene	6.20	U	0.56	6.20	ug/L
218-01-9	Chrysene	6.20	U	0.54	6.20	ug/L
205-99-2	Benzo(b)fluoranthene	6.20	U	0.60	6.20	ug/L
207-08-9	Benzo(k)fluoranthene	6.20	U	0.59	6.20	ug/L
50-32-8	Benzo(a)pyrene	6.20	U	0.68	6.20	ug/L
193-39-5	Indeno(1,2,3-cd)pyrene	6.20	U	0.73	6.20	ug/L
53-70-3	Dibenz(a,h)anthracene	6.20	U	0.83	6.20	ug/L
191-24-2	Benzo(g,h,i)perylene	6.20	U	0.85	6.20	ug/L
SURROGATES						
4165-60-0	Nitrobenzene-d5	66.0		49 - 133	66%	SPK: 100
321-60-8	2-Fluorobiphenyl	63.0		52 - 132	63%	SPK: 100
1718-51-0	Terphenyl-d14	83.4		48 - 125	83%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	104000	7.652			
1146-65-2	Naphthalene-d8	395000	10.422			
15067-26-2	Acenaphthene-d10	245000	14.287			
1517-22-2	Phenanthrene-d10	487000	17.086			
1719-03-5	Chrysene-d12	600000	21.522			
1520-96-3	Perylene-d12	736000	24.815			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	FB-05152025	SDG No.:	Q2075
Lab Sample ID:	Q2075-07	Matrix:	Water
Analytical Method:	8270E	% Solid:	0
Sample Wt/Vol:	810	Units:	mL
Soil Aliquot Vol:		uL	
Extraction Type :		Decanted :	N
Injection Volume :		GPC Factor :	1.0
Prep Method :	SW3510C	GPC Cleanup :	N
		Level :	LOW
		PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024758.D	1	05/21/25 08:40	05/22/25 14:19	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

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B = Analyte Found in Associated Method Blank

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D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Report of Analysis

Client:	CDM Smith			Date Collected:	05/16/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	FB-05162025			SDG No.:	Q2075	
Lab Sample ID:	Q2075-08			Matrix:	Water	
Analytical Method:	8270E			% Solid:	0	
Sample Wt/Vol:	940	Units:	mL	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3510C					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024748.D	1	05/21/25 08:40	05/21/25 22:18	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
91-20-3	Naphthalene	5.30	U	0.53	5.30	ug/L
208-96-8	Acenaphthylene	5.30	U	0.80	5.30	ug/L
83-32-9	Acenaphthene	5.30	U	0.59	5.30	ug/L
86-73-7	Fluorene	5.30	U	0.67	5.30	ug/L
85-01-8	Phenanthrene	5.30	U	0.53	5.30	ug/L
120-12-7	Anthracene	5.30	U	0.65	5.30	ug/L
206-44-0	Fluoranthene	5.30	U	0.87	5.30	ug/L
129-00-0	Pyrene	5.30	U	0.53	5.30	ug/L
56-55-3	Benzo(a)anthracene	5.30	U	0.48	5.30	ug/L
218-01-9	Chrysene	5.30	U	0.47	5.30	ug/L
205-99-2	Benzo(b)fluoranthene	5.30	U	0.52	5.30	ug/L
207-08-9	Benzo(k)fluoranthene	5.30	U	0.51	5.30	ug/L
50-32-8	Benzo(a)pyrene	5.30	U	0.59	5.30	ug/L
193-39-5	Indeno(1,2,3-cd)pyrene	5.30	U	0.63	5.30	ug/L
53-70-3	Dibenz(a,h)anthracene	5.30	U	0.71	5.30	ug/L
191-24-2	Benzo(g,h,i)perylene	5.30	U	0.73	5.30	ug/L
SURROGATES						
4165-60-0	Nitrobenzene-d5	73.7		49 - 133	74%	SPK: 100
321-60-8	2-Fluorobiphenyl	74.4		52 - 132	74%	SPK: 100
1718-51-0	Terphenyl-d14	89.5		48 - 125	90%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	87700	7.652			
1146-65-2	Naphthalene-d8	326000	10.422			
15067-26-2	Acenaphthene-d10	217000	14.286			
1517-22-2	Phenanthrene-d10	458000	17.092			
1719-03-5	Chrysene-d12	547000	21.51			
1520-96-3	Perylene-d12	672000	24.798			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/16/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	FB-05162025	SDG No.:	Q2075
Lab Sample ID:	Q2075-08	Matrix:	Water
Analytical Method:	8270E	% Solid:	0
Sample Wt/Vol:	940	Units:	mL
Soil Aliquot Vol:		uL	
Extraction Type :		Decanted :	N
Injection Volume :		GPC Factor :	1.0
Prep Method :	SW3510C	GPC Cleanup :	N
		Level :	LOW
		PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024748.D	1	05/21/25 08:40	05/21/25 22:18	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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B = Analyte Found in Associated Method Blank

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A = Aldol-Condensation Reaction Products



A
B
C
D
E
F
G

QC SUMMARY

Surrogate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: 8270E

Lab Sample ID	Client ID	Parameter	Spike (PPM)	Result (PPM)	Recovery (%)	Qual	Limits (%)	
							Low	High
PB168083BL	PB168083BL	Nitrobenzene-d5	100	67.4	67	67	18	107
		2-Fluorobiphenyl	100	69.4	69	69	20	109
		Terphenyl-d14	100	62.4	62	62	10	105
PB168083BS	PB168083BS	Nitrobenzene-d5	100	69.4	69	69	18	107
		2-Fluorobiphenyl	100	66.0	66	66	20	109
		Terphenyl-d14	100	81.6	82	82	10	105
Q2075-01	SS-10	Nitrobenzene-d5	100	43.6	44	44	18	107
		2-Fluorobiphenyl	100	37.5	38	38	20	109
		Terphenyl-d14	100	29.7	30	30	10	105
Q2075-02	SS-910	Nitrobenzene-d5	100	49.5	49	49	18	107
		2-Fluorobiphenyl	100	43.1	43	43	20	109
		Terphenyl-d14	100	35.4	35	35	10	105
Q2075-03	SS-11	Nitrobenzene-d5	100	64.7	65	65	18	107
		2-Fluorobiphenyl	100	56.2	56	56	20	109
		Terphenyl-d14	100	42.2	42	42	10	105
Q2075-04MS	SS-11MS	Nitrobenzene-d5	100	60.4	60	60	18	107
		2-Fluorobiphenyl	100	51.6	52	52	20	109
		Terphenyl-d14	100	41.5	41	41	10	105
Q2075-05MSD	SS-11MSD	Nitrobenzene-d5	100	59.6	60	60	18	107
		2-Fluorobiphenyl	100	54.4	54	54	20	109
		Terphenyl-d14	100	40.2	40	40	10	105
Q2075-06	SS-MW1-11.5	Nitrobenzene-d5	100	53.3	53	53	18	107
		2-Fluorobiphenyl	100	48.6	49	49	20	109
		Terphenyl-d14	100	38.9	39	39	10	105

Surrogate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: 8270E

Lab Sample ID	Client ID	Parameter	Spike (PPM)	Result (PPM)	Recovery (%)	Qual	Limits (%)	
							Low	High
PB168098BL	PB168098BL	Nitrobenzene-d5	100	75.0	75	75	49	133
		2-Fluorobiphenyl	100	77.3	77	77	52	132
		Terphenyl-d14	100	83.0	83	83	48	125
PB168098BS	PB168098BS	Nitrobenzene-d5	100	72.8	73	73	49	133
		2-Fluorobiphenyl	100	76.3	76	76	52	132
		Terphenyl-d14	100	77.9	78	78	48	125
PB168098BSD	PB168098BSD	Nitrobenzene-d5	100	73.9	74	74	49	133
		2-Fluorobiphenyl	100	76.8	77	77	52	132
		Terphenyl-d14	100	81.5	81	81	48	125
Q2075-07	FB-05152025	Nitrobenzene-d5	100	66.0	66	66	49	133
		2-Fluorobiphenyl	100	63.0	63	63	52	132
		Terphenyl-d14	100	83.4	83	83	48	125
Q2075-08	FB-05162025	Nitrobenzene-d5	100	73.7	74	74	49	133
		2-Fluorobiphenyl	100	74.4	74	74	52	132
		Terphenyl-d14	100	89.5	90	90	48	125

Matrix Spike/Matrix Spike Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8270E

Parameter	Spike	Sample Result	Result	Units	Rec	Rec Qual	RPD	RPD Qual	Limits Low	Limits High	RPD
Lab Sample ID:	Q2075-04MS	Client Sample ID:	SS-11MS				DataFile:	BF142487.D			
Naphthalene	1900	0	1600	ug/Kg	84				72	110	
Acenaphthylene	1900	0	1600	ug/Kg	84				79	118	
Acenaphthene	1900	0	1800	ug/Kg	95				70	121	
Fluorene	1900	0	1700	ug/Kg	89				68	116	
Phenanthrene	1900	0	1600	ug/Kg	84				52	128	
Anthracene	1900	0	1700	ug/Kg	89				62	124	
Fluoranthene	1900	120	2200	ug/Kg	109				44	125	
Pyrene	1900	0	1300	ug/Kg	68				26	142	
Benzo(a)anthracene	1900	0	1700	ug/Kg	89				71	114	
Chrysene	1900	0	1600	ug/Kg	84				57	121	
Benzo(b)fluoranthene	1900	0	1900	ug/Kg	100				67	121	
Benzo(k)fluoranthene	1900	0	1600	ug/Kg	84				57	134	
Benzo(a)pyrene	1900	0	1700	ug/Kg	89				70	142	
Indeno(1,2,3-cd)pyrene	1900	0	1400	ug/Kg	74				40	129	
Dibenz(a,h)anthracene	1900	0	1400	ug/Kg	74				43	123	
Benzo(g,h,i)perylene	1900	0	1300	ug/Kg	68				24	125	

Matrix Spike/Matrix Spike Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: SW8270E

Parameter	Spike	Sample Result	Result	Units	Rec	Rec Qual	RPD	RPD Qual	Limits Low	High	RPD
Lab Sample ID:	Q2075-05MSD	Client Sample ID:	SS-11MSD					DataFile:	BF142488.D		
Naphthalene	1900	0	1600	ug/Kg	84	0			72	110	20
Acenaphthylene	1900	0	1600	ug/Kg	84	0			79	118	20
Acenaphthene	1900	0	1800	ug/Kg	95	0			70	121	20
Fluorene	1900	0	1600	ug/Kg	84	6			68	116	20
Phenanthrene	1900	0	1700	ug/Kg	89	6			52	128	20
Anthracene	1900	0	1700	ug/Kg	89	0			62	124	20
Fluoranthene	1900	120	2300	ug/Kg	115	5			44	125	20
Pyrene	1900	0	1300	ug/Kg	68	0			26	142	20
Benzo(a)anthracene	1900	0	1600	ug/Kg	84	6			71	114	20
Chrysene	1900	0	1600	ug/Kg	84	0			57	121	20
Benzo(b)fluoranthene	1900	0	1700	ug/Kg	89	12			67	121	20
Benzo(k)fluoranthene	1900	0	1700	ug/Kg	89	6			57	134	20
Benzo(a)pyrene	1900	0	1700	ug/Kg	89	0			70	142	20
Indeno(1,2,3-cd)pyrene	1900	0	1400	ug/Kg	74	0			40	129	20
Dibenz(a,h)anthracene	1900	0	1300	ug/Kg	68	8			43	123	20
Benzo(g,h,i)perylene	1900	0	1300	ug/Kg	68	0			24	125	20

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: 8270E DataFile: BF142480.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Qual	Limits		RPD
									Low	High	
PB168083BS	Naphthalene	1700	1400	ug/Kg	82				62	100	
	Acenaphthylene	1700	1300	ug/Kg	76				63	101	
	Acenaphthene	1700	1500	ug/Kg	88				57	104	
	Fluorene	1700	1300	ug/Kg	76				61	101	
	Phenanthrene	1700	1400	ug/Kg	82				59	103	
	Anthracene	1700	1300	ug/Kg	76				61	105	
	Fluoranthene	1700	1500	ug/Kg	88				57	107	
	Pyrene	1700	1600	ug/Kg	94				59	103	
	Benzo(a)anthracene	1700	1400	ug/Kg	82				60	102	
	Chrysene	1700	1400	ug/Kg	82				59	101	
	Benzo(b)fluoranthene	1700	1500	ug/Kg	88				62	109	
	Benzo(k)fluoranthene	1700	1400	ug/Kg	82				62	109	
	Benzo(a)pyrene	1700	1500	ug/Kg	88				63	103	
	Indeno(1,2,3-cd)pyrene	1700	1600	ug/Kg	94				63	101	
	Dibenz(a,h)anthracene	1700	1600	ug/Kg	94				61	112	
	Benzo(g,h,i)perylene	1700	1600	ug/Kg	94				70	108	

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: 8270E DataFile: BP024755.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Qual	Limits		RPD
									Low	High	
PB168098BS	Naphthalene	50	43.0	ug/L	86				64	107	
	Acenaphthylene	50	42.4	ug/L	85				79	103	
	Acenaphthene	50	42.1	ug/L	84				59	113	
	Fluorene	50	44.1	ug/L	88				64	107	
	Phenanthrene	50	43.2	ug/L	86				62	109	
	Anthracene	50	44.5	ug/L	89				65	110	
	Fluoranthene	50	46.0	ug/L	92				64	110	
	Pyrene	50	44.5	ug/L	89				71	103	
	Benzo(a)anthracene	50	43.9	ug/L	88				62	107	
	Chrysene	50	44.1	ug/L	88				61	108	
	Benzo(b)fluoranthene	50	44.2	ug/L	88				77	113	
	Benzo(k)fluoranthene	50	44.0	ug/L	88				77	105	
	Benzo(a)pyrene	50	44.8	ug/L	90				72	131	
	Indeno(1,2,3-cd)pyrene	50	45.7	ug/L	91				72	105	
	Dibenz(a,h)anthracene	50	46.0	ug/L	92				78	115	
	Benzo(g,h,i)perylene	50	44.9	ug/L	90				75	118	

Laboratory Control Sample/Laboratory Control Sample Duplicate Summary

SW-846

SDG No.: Q2075

Client: CDM Smith

Analytical Method: 8270E DataFile: BP024756.D

Lab Sample ID	Parameter	Spike	Result	Unit	Rec	RPD	Qual	Qual	Limits			RPD
									Low	High	RPD	
PB168098BSD	Naphthalene	50	43.4	ug/L	87	1			64	107	20	
	Acenaphthylene	50	43.6	ug/L	87	3			79	103	20	
	Acenaphthene	50	43.5	ug/L	87	3			59	113	20	
	Fluorene	50	44.9	ug/L	90	2			64	107	20	
	Phenanthrene	50	44.0	ug/L	88	2			62	109	20	
	Anthracene	50	45.1	ug/L	90	1			65	110	20	
	Fluoranthene	50	45.8	ug/L	92	0			64	110	20	
	Pyrene	50	45.6	ug/L	91	2			71	103	20	
	Benzo(a)anthracene	50	45.5	ug/L	91	4			62	107	20	
	Chrysene	50	44.4	ug/L	89	1			61	108	20	
	Benzo(b)fluoranthene	50	46.7	ug/L	93	6			77	113	20	
	Benzo(k)fluoranthene	50	45.6	ug/L	91	4			77	105	20	
	Benzo(a)pyrene	50	46.1	ug/L	92	3			72	131	20	
	Indeno(1,2,3-cd)pyrene	50	44.8	ug/L	90	2			72	105	20	
	Dibenz(a,h)anthracene	50	45.2	ug/L	90	2			78	115	20	
	Benzo(g,h,i)perylene	50	43.9	ug/L	88	2			75	118	20	

4B

SEMIVOLATILE METHOD BLANK SUMMARY

EPA SAMPLE NO.

PB168083BL

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM Case No.: Q2075

SAS No.: Q2075 SDG No.: Q2075

Lab File ID: BF142479.D

Lab Sample ID: PB168083BL

Instrument ID: BNA_F

Date Extracted: 05/20/2025

Matrix: (soil/water) SOIL

Date Analyzed: 05/21/2025

Level: (low/med) LOW

Time Analyzed: 11:09

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
PB168083BS	PB168083BS	BF142480.D	05/21/2025
SS-11	Q2075-03	BF142486.D	05/21/2025
SS-11MS	Q2075-04MS	BF142487.D	05/21/2025
SS-11MSD	Q2075-05MSD	BF142488.D	05/21/2025
SS-MW1-11.5	Q2075-06	BF142489.D	05/21/2025
SS-10	Q2075-01	BF142510.D	05/22/2025
SS-910	Q2075-02	BF142511.D	05/22/2025

COMMENTS:

4B

SEMIVOLATILE METHOD BLANK SUMMARY

EPA SAMPLE NO.

PB168098BL

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM Case No.: Q2075

SAS No.: Q2075 SDG NO.: Q2075

Lab File ID: BP024754.D

Lab Sample ID: PB168098BL

Instrument ID: BNA_P

Date Extracted: 05/21/2025

Matrix: (soil/water) Water

Date Analyzed: 05/22/2025

Level: (low/med) LOW

Time Analyzed: 11:37

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED
PB168098BS	PB168098BS	BP024755.D	05/22/2025
PB168098BSD	PB168098BSD	BP024756.D	05/22/2025
FB-05152025	Q2075-07	BP024758.D	05/22/2025
FB-05162025	Q2075-08	BP024748.D	05/21/2025

COMMENTS:

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM

SAS No.: Q2075 SDG NO.: Q2075

Lab File ID: BF142465.D

DFTPP Injection Date: 05/20/2025

Instrument ID: BNA_F

DFTPP Injection Time: 11:13

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0% of mass 198	36.8
68	Less than 2.0% of mass 69	0.5 (1.9) 1
69	Mass 69 relative abundance	33.1
70	Less than 2.0% of mass 69	0.1 (0.5) 1
127	10.0 - 80.0% of mass 198	44.9
197	Less than 2.0% of mass 198	0.0
198	Base Peak, 100% relative abundance	100
199	5.0 to 9.0% of mass 198	6.6
275	10.0 - 60.0% of mass 198	29.9
365	Greater than 1% of mass 198	4.2
441	Present, but less than mass 443	19.9
442	Greater than 50% of mass 198	100
443	15.0 - 24.0% of mass 442	19 (19) 2

1-Value is % mass 69

2-Value is % mass 442

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
SSTDICC2.5	SSTDICC2.5	BF142467.D	05/20/2025	12:10
SSTDICC005	SSTDICC005	BF142468.D	05/20/2025	12:38
SSTDICC010	SSTDICC010	BF142469.D	05/20/2025	13:07
SSTDICC020	SSTDICC020	BF142470.D	05/20/2025	13:36
SSTDICCC040	SSTDICCC040	BF142471.D	05/20/2025	14:05
SSTDICC050	SSTDICC050	BF142472.D	05/20/2025	14:34
SSTDICC060	SSTDICC060	BF142473.D	05/20/2025	15:03
SSTDICC080	SSTDICC080	BF142474.D	05/20/2025	15:31

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM

SAS No.: Q2075 SDG NO.: Q2075

Lab File ID: BF142477.D

DFTPP Injection Date: 05/21/2025

Instrument ID: BNA_F

DFTPP Injection Time: 09:41

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0% of mass 198	40.0
68	Less than 2.0% of mass 69	0.6 (2) 1
69	Mass 69 relative abundance	34.9
70	Less than 2.0% of mass 69	0.2 (0.7) 1
127	10.0 - 80.0% of mass 198	46.9
197	Less than 2.0% of mass 198	0.0
198	Base Peak, 100% relative abundance	100
199	5.0 to 9.0% of mass 198	6.8
275	10.0 - 60.0% of mass 198	29.1
365	Greater than 1% of mass 198	3.8
441	Present, but less than mass 443	18.3
442	Greater than 50% of mass 198	100
443	15.0 - 24.0% of mass 442	19.4 (19.4) 2

1-Value is % mass 69

2-Value is % mass 442

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
SSTDCCC040	SSTDCCC040	BF142478.D	05/21/2025	10:40
PB168083BL	PB168083BL	BF142479.D	05/21/2025	11:09
PB168083BS	PB168083BS	BF142480.D	05/21/2025	11:37
SS-11	Q2075-03	BF142486.D	05/21/2025	14:37
SS-11MS	Q2075-04MS	BF142487.D	05/21/2025	15:06
SS-11MSD	Q2075-05MSD	BF142488.D	05/21/2025	15:34
SS-MW1-11.5	Q2075-06	BF142489.D	05/21/2025	16:03

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM

SAS No.: Q2075 SDG NO.: Q2075

Lab File ID: BF142501.D

DFTPP Injection Date: 05/22/2025

Instrument ID: BNA_F

DFTPP Injection Time: 10:43

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0% of mass 198	37.5
68	Less than 2.0% of mass 69	0.5 (1.8) 1
69	Mass 69 relative abundance	33.6
70	Less than 2.0% of mass 69	0.2 (0.6) 1
127	10.0 - 80.0% of mass 198	46.8
197	Less than 2.0% of mass 198	0.0
198	Base Peak, 100% relative abundance	100
199	5.0 to 9.0% of mass 198	6.8
275	10.0 - 60.0% of mass 198	30.3
365	Greater than 1% of mass 198	4.1
441	Present, but less than mass 443	20.4
442	Greater than 50% of mass 198	100
443	15.0 - 24.0% of mass 442	18.7 (18.7) 2

1-Value is % mass 69

2-Value is % mass 442

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
SSTDCCC040	SSTDCCC040	BF142502.D	05/22/2025	11:11
SS-10	Q2075-01	BF142510.D	05/22/2025	15:07
SS-910	Q2075-02	BF142511.D	05/22/2025	15:36

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM

SAS No.: Q2075 SDG NO.: Q2075

Lab File ID: BP024613.D

DFTPP Injection Date: 05/13/2025

Instrument ID: BNA_P

DFTPP Injection Time: 10:00

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0% of mass 198	33.5
68	Less than 2.0% of mass 69	0.2 (0.6) 1
69	Mass 69 relative abundance	36.2
70	Less than 2.0% of mass 69	0.2 (0.6) 1
127	10.0 - 80.0% of mass 198	49.3
197	Less than 2.0% of mass 198	0.0
198	Base Peak, 100% relative abundance	100
199	5.0 to 9.0% of mass 198	6.6
275	10.0 - 60.0% of mass 198	33.1
365	Greater than 1% of mass 198	5.4
441	Present, but less than mass 443	19.4
442	Greater than 50% of mass 198	100
443	15.0 - 24.0% of mass 442	19.4 (19.4) 2

1-Value is % mass 69

2-Value is % mass 442

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
SSTDICC2.5	SSTDICC2.5	BP024614.D	05/13/2025	10:41
SSTDICC005	SSTDICC005	BP024615.D	05/13/2025	11:22
SSTDICC010	SSTDICC010	BP024616.D	05/13/2025	12:03
SSTDICC020	SSTDICC020	BP024617.D	05/13/2025	12:43
SSTDICCC040	SSTDICCC040	BP024618.D	05/13/2025	13:24
SSTDICC050	SSTDICC050	BP024619.D	05/13/2025	14:05
SSTDICC060	SSTDICC060	BP024620.D	05/13/2025	14:45
SSTDICC080	SSTDICC080	BP024621.D	05/13/2025	15:26

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM

SAS No.: Q2075 SDG NO.: Q2075

Lab File ID: BP024736.D

DFTPP Injection Date: 05/21/2025

Instrument ID: BNA_P

DFTPP Injection Time: 12:54

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0% of mass 198	31.7
68	Less than 2.0% of mass 69	0.0 (0.0) 1
69	Mass 69 relative abundance	33.9
70	Less than 2.0% of mass 69	0.1 (0.4) 1
127	10.0 - 80.0% of mass 198	47.3
197	Less than 2.0% of mass 198	0.0
198	Base Peak, 100% relative abundance	100
199	5.0 to 9.0% of mass 198	6.9
275	10.0 - 60.0% of mass 198	33.3
365	Greater than 1% of mass 198	5.6
441	Present, but less than mass 443	22.0
442	Greater than 50% of mass 198	100
443	15.0 - 24.0% of mass 442	19.9 (19.9) 2

1-Value is % mass 69

2-Value is % mass 442

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
SSTDCCC040	SSTDCCC040	BP024737.D	05/21/2025	14:48
FB-05162025	Q2075-08	BP024748.D	05/21/2025	22:18

5B

SEMIVOLATILE ORGANIC INSTRUMENT PERFORMANCE CHECK
DECAFLUOROTRIPHENYLPHOSPHINE (DFTPP)

Lab Name: CHEMTECH

Contract: CAMP02

Lab Code: CHEM

SAS No.: Q2075 SDG NO.: Q2075

Lab File ID: BP024752.D

DFTPP Injection Date: 05/22/2025

Instrument ID: BNA_P

DFTPP Injection Time: 09:35

m/e	ION ABUNDANCE CRITERIA	% RELATIVE ABUNDANCE
51	10.0 - 80.0% of mass 198	30.1
68	Less than 2.0% of mass 69	0.3 (1.3) 1
69	Mass 69 relative abundance	32.0
70	Less than 2.0% of mass 69	0.1 (0.5) 1
127	10.0 - 80.0% of mass 198	46.0
197	Less than 2.0% of mass 198	0.0
198	Base Peak, 100% relative abundance	100
199	5.0 to 9.0% of mass 198	7.0
275	10.0 - 60.0% of mass 198	33.7
365	Greater than 1% of mass 198	5.4
441	Present, but less than mass 443	22.4
442	Greater than 50% of mass 198	100
443	15.0 - 24.0% of mass 442	19.6 (19.6) 2

1-Value is % mass 69

2-Value is % mass 442

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

EPA SAMPLE NO.	LAB SAMPLE ID	LAB FILE ID	DATE ANALYZED	TIME ANALYZED
SSTDCCC040	SSTDCCC040	BP024753.D	05/22/2025	10:56
PB168098BL	PB168098BL	BP024754.D	05/22/2025	11:37
PB168098BS	PB168098BS	BP024755.D	05/22/2025	12:18
PB168098BSD	PB168098BSD	BP024756.D	05/22/2025	12:58
FB-05152025	Q2075-07	BP024758.D	05/22/2025	14:19



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8B

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: CHEMTECH
Lab Code: CHEM Case No.: Q2075 SAS No.: Q2075 SDG No.: Q2075
EPA Sample No.: SSTDCCC040 Date Analyzed: 05/21/2025
Lab File ID: BF142478.D Time Analyzed: 10:40
Instrument ID: BNA_F GC Column: DB-UI ID: 0.18 (mm)

	IS1 (DCB) AREA #	RT #	IS2 (NPT) AREA #	RT #	IS3 (ANT) AREA #	RT #
12 HOUR STD	124102	6.898	475212	8.19	258765	9.94
UPPER LIMIT	248204	7.398	950424	8.686	517530	10.439
LOWER LIMIT	62051	6.398	237606	7.686	129383	9.439
EPA SAMPLE NO.						
01 PB168083BL	127003	6.90	492428	8.18	264579	9.94
02 PB168083BS	124824	6.90	495438	8.19	277044	9.94
03 SS-11	63588	6.90	238505	8.18	131818	9.94
04 SS-11MS	54835 *	6.90	200654 *	8.18	113830 *	9.94
05 SS-11MSD	53473 *	6.90	192497 *	8.18	95696 *	9.94
06 SS-MW1-11.5	59263 *	6.90	206974 *	8.18	103588 *	9.94

IS1 (DCB) = 1,4-Dichlorobenzene-d4

IS2 (NPT) = Naphthalene-d8

IS3 (ANT) = Acenaphthene-d10

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = -50% of internal standard area

RT UPPER LIMIT = +0.50 minutes of internal standard RT

RT LOWER LIMIT = -0.50 minutes of internal standard RT

Column used to flag values outside QC limits with an asterisk.

* Values outside of QC limits.

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SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name:	CHEMTECH						
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075	SDG NO.:	Q2075
EPA Sample No.:	SSTDCCC040		Date Analyzed:	05/21/2025			
Lab File ID:	BF142478.D		Time Analyzed:	10:40			
Instrument ID:	BNA_F		GC Column:	DB-UI	ID:	0.18	(mm)

	IS4 (PHN) AREA #	RT #	IS5 (CRY) AREA #	RT #	IS6 (PRY) AREA #	RT #
12 HOUR STD	454448	11.427	231233	14.068	233183	15.562
	908896	11.927	462466	14.568	466366	16.062
	227224	10.927	115617	13.568	116592	15.062
EPA SAMPLE NO.						
01 PB168083BL	511963	11.43	324877	14.07	234653	15.56
02 PB168083BS	468851	11.43	214548	14.07	229705	15.56
03 SS-11	250730	11.43	246437	14.07	209013	15.56
04 SS-11MS	229012	11.43	210926	14.07	176162	15.56
05 SS-11MSD	179294 *	11.43	174160	14.07	147670	15.56
06 SS-MW1-11.5	179364 *	11.42	153166	14.06	125915	15.56

IS4 (PHN) = Phenanthrene-d10

IS5 (CRY) = Chrysene-d12

IS6 (PRY) = Perylene-d12

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = -50% of internal standard area

RT UPPER LIMIT = +0.50 minutes of internal standard RT

RT LOWER LIMIT = -0.50 minutes of internal standard RT

Column used to flag values outside QC limits with an asterisk.

* Values outside of QC limits.



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8B

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: CHEMTECH
Lab Code: CHEM Case No.: Q2075 SAS No.: Q2075 SDG NO.: Q2075
EPA Sample No.: SSTDCCC040 Date Analyzed: 05/22/2025
Lab File ID: BF142502.D Time Analyzed: 11:11
Instrument ID: BNA_F GC Column: DB-UI ID: 0.18 (mm)

	IS1 (DCB) AREA #	RT #	IS2 (NPT) AREA #	RT #	IS3 (ANT) AREA #	RT #
12 HOUR STD	96659	6.898	372701	8.19	197309	9.94
UPPER LIMIT	193318	7.398	745402	8.687	394618	10.439
LOWER LIMIT	48329.5	6.398	186351	7.687	98654.5	9.439
EPA SAMPLE NO.						
01 SS-10	120361	6.90	452635	8.18	223139	9.93
02 SS-910	116551	6.90	418491	8.18	190753	9.93

IS1 (DCB) = 1,4-Dichlorobenzene-d4

IS2 (NPT) = Naphthalene-d8

IS3 (ANT) = Acenaphthene-d10

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.

8C

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name:	CHEMTECH			
Lab Code:	CHEM	Case No.:	Q2075	
SAS No.:	Q2075		SDG NO.:	Q2075
EPA Sample No.:	SSTDCCC040		Date Analyzed:	05/22/2025
Lab File ID:	BF142502.D		Time Analyzed:	11:11
Instrument ID:	BNA_F		GC Column:	DB-U1
			ID:	0.18 (mm)

	IS4 (PHN) AREA #	RT #	IS5 (CRY) AREA #	RT #	IS6 (PRY) AREA #	RT #
12 HOUR STD	315055	11.428	157895	14.069	195527	15.557
	630110	11.928	315790	14.569	391054	16.057
	157528	10.928	78947.5	13.569	97763.5	15.057
EPA SAMPLE NO.						
01 SS-10	308296	11.42	228905	14.06	198076	15.56
02 SS-910	271357	11.42	225312	14.06	176642	15.56

IS4 (PHN) = Phenanthrene-d10

IS5 (CRY) = Chrysene-d12

IS6 (PRY) = Perylene-d12

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = -50% of internal standard area

RT UPPER LIMIT = +0.50 minutes of internal standard RT

RT LOWER LIMIT = -0.50 minutes of internal standard RT

Column used to flag values outside QC limits with an asterisk.

* Values outside of QC limits.



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Fax : 908 789 8922

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8B

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: CHEMTECH
Lab Code: CHEM Case No.: Q2075 SAS No.: Q2075 SDG NO.: Q2075
EPA Sample No.: SSTDCCC040 Date Analyzed: 05/21/2025
Lab File ID: BP024737.D Time Analyzed: 14:48
Instrument ID: BNA_P GC Column: ZB-GR ID: 0.25 (mm)

	IS1 (DCB) AREA #	RT #	IS2 (NPT) AREA #	RT #	IS3 (ANT) AREA #	RT #
12 HOUR STD	128557	7.646	504282	10.42	338214	14.29
UPPER LIMIT	257114	8.146	1008560	10.916	676428	14.787
LOWER LIMIT	64278.5	7.146	252141	9.916	169107	13.787
EPA SAMPLE NO.						
01 FB-05162025	87679	7.65	325514	10.42	217493	14.29

IS1 (DCB) = 1,4-Dichlorobenzene-d4

IS2 (NPT) = Naphthalene-d8

IS3 (ANT) = Acenaphthene-d10

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.

8C

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name:	CHEMTECH			
Lab Code:	CHEM	Case No.:	Q2075	
SAS No.:	Q2075		SDG NO.:	Q2075
EPA Sample No.:	SSTDCCC040		Date Analyzed:	05/21/2025
Lab File ID:	BP024737.D		Time Analyzed:	14:48
Instrument ID:	BNA_P		GC Column:	ZB-GR
			ID:	0.25 (mm)

	IS4 (PHN) AREA #	RT #	IS5 (CRY) AREA #	RT #	IS6 (PRY) AREA #	RT #
12 HOUR STD	643332	17.081	728456	21.51	831576	24.786
	1286660	17.581	1456910	22.01	1663150	25.286
	321666	16.581	364228	21.01	415788	24.286
EPA SAMPLE NO.						
01 FB-05162025	457611	17.09	547413	21.51	671507	24.80

IS4 (PHN) = Phenanthrene-d10

IS5 (CRY) = Chrysene-d12

IS6 (PRY) = Perylene-d12

AREA UPPER LIMIT = +100% of internal standard area

AREA LOWER LIMIT = -50% of internal standard area

RT UPPER LIMIT = +0.50 minutes of internal standard RT

RT LOWER LIMIT = -0.50 minutes of internal standard RT

Column used to flag values outside QC limits with an asterisk.

* Values outside of QC limits.



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6

8B

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name: CHEMTECH
Lab Code: CHEM Case No.: Q2075 SAS No.: Q2075 SDG No.: Q2075
EPA Sample No.: SSTDCCC040 Date Analyzed: 05/22/2025
Lab File ID: BP024753.D Time Analyzed: 10:56
Instrument ID: BNA_P GC Column: ZB-GR ID: 0.25 (mm)

	IS1 (DCB) AREA #	RT #	IS2 (NPT) AREA #	RT #	IS3 (ANT) AREA #	RT #
12 HOUR STD	103714	7.652	424752	10.42	258861	14.28
UPPER LIMIT	207428	8.152	849504	10.922	517722	14.781
LOWER LIMIT	51857	7.152	212376	9.922	129431	13.781
EPA SAMPLE NO.						
01 PB168098BL	115351	7.65	453620	10.42	285428	14.29
02 PB168098BS	114889	7.65	451979	10.42	283120	14.29
03 PB168098BSD	126327	7.65	523024	10.42	338733	14.29
04 FB-05152025	103638	7.65	395386	10.42	244728	14.29

IS1 (DCB) = 1,4-Dichlorobenzene-d4

IS2 (NPT) = Naphthalene-d8

IS3 (ANT) = Acenaphthene-d10

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.

8C

SEMIVOLATILE INTERNAL STANDARD AREA AND RT SUMMARY

Lab Name:	CHEMTECH						
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075	SDG NO.:	Q2075
EPA Sample No.:	SSTDCCC040		Date Analyzed:	05/22/2025			
Lab File ID:	BP024753.D		Time Analyzed:	10:56			
Instrument ID:	BNA_P		GC Column:	ZB-GR	ID:	0.25	(mm)

	IS4 (PHN) AREA #	RT #	IS5 (CRY) AREA #	RT #	IS6 (PRY) AREA #	RT #
12 HOUR STD	515151	17.075	613668	21.516	759397	24.792
	1030300	17.575	1227340	22.016	1518790	25.292
	257576	16.575	306834	21.016	379699	24.292
EPA SAMPLE NO.						
01 PB168098BL	613349	17.09	715536	21.52	855120	24.82
02 PB168098BS	568580	17.09	655197	21.52	754804	24.82
03 PB168098BSD	686645	17.09	768545	21.51	835982	24.79
04 FB-05152025	486633	17.09	599957	21.52	736321	24.82

IS4 (PHN) = Phenanthrene-d10

IS5 (CRY) = Chrysene-d12

IS6 (PRY) = Perylene-d12

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.



QC SAMPLE

DATA

A
B
C
D
E
F
G

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168083BL			SDG No.:	Q2075
Lab Sample ID:	PB168083BL			Matrix:	SOIL
Analytical Method:	8270E			% Solid:	100
Sample Wt/Vol:	30.01	Units:	g	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142479.D	1	05/20/25 09:45	05/21/25 11:09	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	170	U	22.7	170	ug/Kg
208-96-8	Acenaphthylene	170	U	28.9	170	ug/Kg
83-32-9	Acenaphthene	170	U	21.3	170	ug/Kg
86-73-7	Fluorene	170	U	25.3	170	ug/Kg
85-01-8	Phenanthrene	170	U	20.9	170	ug/Kg
120-12-7	Anthracene	170	U	33.3	170	ug/Kg
206-44-0	Fluoranthene	170	U	30.0	170	ug/Kg
129-00-0	Pyrene	170	U	36.0	170	ug/Kg
56-55-3	Benzo(a)anthracene	170	U	23.0	170	ug/Kg
218-01-9	Chrysene	170	U	19.9	170	ug/Kg
205-99-2	Benzo(b)fluoranthene	170	U	19.0	170	ug/Kg
207-08-9	Benzo(k)fluoranthene	170	U	22.4	170	ug/Kg
50-32-8	Benzo(a)pyrene	170	U	29.5	170	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	170	U	29.1	170	ug/Kg
53-70-3	Dibenz(a,h)anthracene	170	U	27.4	170	ug/Kg
191-24-2	Benzo(g,h,i)perylene	170	U	25.7	170	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	67.4		18 - 107	67%	SPK: 100
321-60-8	2-Fluorobiphenyl	69.4		20 - 109	69%	SPK: 100
1718-51-0	Terphenyl-d14	62.4		10 - 105	62%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	127000	6.898			
1146-65-2	Naphthalene-d8	492000	8.181			
15067-26-2	Acenaphthene-d10	265000	9.939			
1517-22-2	Phenanthrene-d10	512000	11.428			
1719-03-5	Chrysene-d12	325000	14.069			
1520-96-3	Perylene-d12	235000	15.563			

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168083BL			SDG No.:	Q2075
Lab Sample ID:	PB168083BL			Matrix:	SOIL
Analytical Method:	8270E			% Solid:	100
Sample Wt/Vol:	30.01	Units:	g	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142479.D	1	05/20/25 09:45	05/21/25 11:09	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168098BL			SDG No.:	Q2075
Lab Sample ID:	PB168098BL			Matrix:	Water
Analytical Method:	8270E			% Solid:	0
Sample Wt/Vol:	1000	Units:	mL	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :				GPC Factor :	1.0
				GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024754.D	1	05/21/25 08:40	05/22/25 11:37	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
91-20-3	Naphthalene	5.00	U	0.50	5.00	ug/L
208-96-8	Acenaphthylene	5.00	U	0.75	5.00	ug/L
83-32-9	Acenaphthene	5.00	U	0.55	5.00	ug/L
86-73-7	Fluorene	5.00	U	0.63	5.00	ug/L
85-01-8	Phenanthrene	5.00	U	0.50	5.00	ug/L
120-12-7	Anthracene	5.00	U	0.61	5.00	ug/L
206-44-0	Fluoranthene	5.00	U	0.82	5.00	ug/L
129-00-0	Pyrene	5.00	U	0.50	5.00	ug/L
56-55-3	Benzo(a)anthracene	5.00	U	0.45	5.00	ug/L
218-01-9	Chrysene	5.00	U	0.44	5.00	ug/L
205-99-2	Benzo(b)fluoranthene	5.00	U	0.49	5.00	ug/L
207-08-9	Benzo(k)fluoranthene	5.00	U	0.48	5.00	ug/L
50-32-8	Benzo(a)pyrene	5.00	U	0.55	5.00	ug/L
193-39-5	Indeno(1,2,3-cd)pyrene	5.00	U	0.59	5.00	ug/L
53-70-3	Dibenz(a,h)anthracene	5.00	U	0.67	5.00	ug/L
191-24-2	Benzo(g,h,i)perylene	5.00	U	0.69	5.00	ug/L
SURROGATES						
4165-60-0	Nitrobenzene-d5	75.0		49 - 133	75%	SPK: 100
321-60-8	2-Fluorobiphenyl	77.3		52 - 132	77%	SPK: 100
1718-51-0	Terphenyl-d14	83.0		48 - 125	83%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	115000	7.652			
1146-65-2	Naphthalene-d8	454000	10.422			
15067-26-2	Acenaphthene-d10	285000	14.287			
1517-22-2	Phenanthrene-d10	613000	17.092			
1719-03-5	Chrysene-d12	716000	21.522			
1520-96-3	Perylene-d12	855000	24.821			

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168098BL			SDG No.:	Q2075
Lab Sample ID:	PB168098BL			Matrix:	Water
Analytical Method:	8270E			% Solid:	0
Sample Wt/Vol:	1000	Units:	mL	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024754.D	1	05/21/25 08:40	05/22/25 11:37	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168083BS			SDG No.:	Q2075
Lab Sample ID:	PB168083BS			Matrix:	SOIL
Analytical Method:	8270E			% Solid:	100
Sample Wt/Vol:	30.03	Units:	g	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142480.D	1	05/20/25 09:45	05/21/25 11:37	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	1400	22.7		170	ug/Kg
208-96-8	Acenaphthylene	1300	28.9		170	ug/Kg
83-32-9	Acenaphthene	1500	21.3		170	ug/Kg
86-73-7	Fluorene	1300	25.3		170	ug/Kg
85-01-8	Phenanthrene	1400	20.9		170	ug/Kg
120-12-7	Anthracene	1300	33.3		170	ug/Kg
206-44-0	Fluoranthene	1500	30.0		170	ug/Kg
129-00-0	Pyrene	1600	36.0		170	ug/Kg
56-55-3	Benzo(a)anthracene	1400	23.0		170	ug/Kg
218-01-9	Chrysene	1400	19.9		170	ug/Kg
205-99-2	Benzo(b)fluoranthene	1500	19.0		170	ug/Kg
207-08-9	Benzo(k)fluoranthene	1400	22.4		170	ug/Kg
50-32-8	Benzo(a)pyrene	1500	29.5		170	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	1600	29.1		170	ug/Kg
53-70-3	Dibenz(a,h)anthracene	1600	27.4		170	ug/Kg
191-24-2	Benzo(g,h,i)perylene	1600	25.7		170	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	69.4	18 - 107		69%	SPK: 100
321-60-8	2-Fluorobiphenyl	66.0	20 - 109		66%	SPK: 100
1718-51-0	Terphenyl-d14	81.6	10 - 105		82%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	125000	6.898			
1146-65-2	Naphthalene-d8	495000	8.186			
15067-26-2	Acenaphthene-d10	277000	9.939			
1517-22-2	Phenanthrene-d10	469000	11.427			
1719-03-5	Chrysene-d12	215000	14.068			
1520-96-3	Perylene-d12	230000	15.562			

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168083BS			SDG No.:	Q2075
Lab Sample ID:	PB168083BS			Matrix:	SOIL
Analytical Method:	8270E			% Solid:	100
Sample Wt/Vol:	30.03	Units:	g	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142480.D	1	05/20/25 09:45	05/21/25 11:37	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168098BS			SDG No.:	Q2075
Lab Sample ID:	PB168098BS			Matrix:	Water
Analytical Method:	8270E			% Solid:	0
Sample Wt/Vol:	1000	Units:	mL	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :				GPC Factor :	1.0
				GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024755.D	1	05/21/25 08:40	05/22/25 12:18	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
91-20-3	Naphthalene	43.0	0.50		5.00	ug/L
208-96-8	Acenaphthylene	42.4	0.75		5.00	ug/L
83-32-9	Acenaphthene	42.1	0.55		5.00	ug/L
86-73-7	Fluorene	44.1	0.63		5.00	ug/L
85-01-8	Phenanthrene	43.2	0.50		5.00	ug/L
120-12-7	Anthracene	44.5	0.61		5.00	ug/L
206-44-0	Fluoranthene	46.0	0.82		5.00	ug/L
129-00-0	Pyrene	44.5	0.50		5.00	ug/L
56-55-3	Benzo(a)anthracene	43.9	0.45		5.00	ug/L
218-01-9	Chrysene	44.1	0.44		5.00	ug/L
205-99-2	Benzo(b)fluoranthene	44.2	0.49		5.00	ug/L
207-08-9	Benzo(k)fluoranthene	44.0	0.48		5.00	ug/L
50-32-8	Benzo(a)pyrene	44.8	0.55		5.00	ug/L
193-39-5	Indeno(1,2,3-cd)pyrene	45.7	0.59		5.00	ug/L
53-70-3	Dibenz(a,h)anthracene	46.0	0.67		5.00	ug/L
191-24-2	Benzo(g,h,i)perylene	44.9	0.69		5.00	ug/L
SURROGATES						
4165-60-0	Nitrobenzene-d5	72.8	49 - 133		73%	SPK: 100
321-60-8	2-Fluorobiphenyl	76.3	52 - 132		76%	SPK: 100
1718-51-0	Terphenyl-d14	77.9	48 - 125		78%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	115000	7.652			
1146-65-2	Naphthalene-d8	452000	10.422			
15067-26-2	Acenaphthene-d10	283000	14.287			
1517-22-2	Phenanthrene-d10	569000	17.087			
1719-03-5	Chrysene-d12	655000	21.522			
1520-96-3	Perylene-d12	755000	24.821			

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168098BS			SDG No.:	Q2075
Lab Sample ID:	PB168098BS			Matrix:	Water
Analytical Method:	8270E			% Solid:	0
Sample Wt/Vol:	1000	Units:	mL	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024755.D	1	05/21/25 08:40	05/22/25 12:18	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168098BSD			SDG No.:	Q2075
Lab Sample ID:	PB168098BSD			Matrix:	Water
Analytical Method:	8270E			% Solid:	0
Sample Wt/Vol:	1000	Units:	mL	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024756.D	1	05/21/25 08:40	05/22/25 12:58	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
91-20-3	Naphthalene	43.4	0.50		5.00	ug/L
208-96-8	Acenaphthylene	43.6	0.75		5.00	ug/L
83-32-9	Acenaphthene	43.5	0.55		5.00	ug/L
86-73-7	Fluorene	44.9	0.63		5.00	ug/L
85-01-8	Phenanthrene	44.0	0.50		5.00	ug/L
120-12-7	Anthracene	45.1	0.61		5.00	ug/L
206-44-0	Fluoranthene	45.8	0.82		5.00	ug/L
129-00-0	Pyrene	45.6	0.50		5.00	ug/L
56-55-3	Benzo(a)anthracene	45.5	0.45		5.00	ug/L
218-01-9	Chrysene	44.4	0.44		5.00	ug/L
205-99-2	Benzo(b)fluoranthene	46.7	0.49		5.00	ug/L
207-08-9	Benzo(k)fluoranthene	45.6	0.48		5.00	ug/L
50-32-8	Benzo(a)pyrene	46.1	0.55		5.00	ug/L
193-39-5	Indeno(1,2,3-cd)pyrene	44.8	0.59		5.00	ug/L
53-70-3	Dibenz(a,h)anthracene	45.2	0.67		5.00	ug/L
191-24-2	Benzo(g,h,i)perylene	43.9	0.69		5.00	ug/L
SURROGATES						
4165-60-0	Nitrobenzene-d5	73.9	49 - 133		74%	SPK: 100
321-60-8	2-Fluorobiphenyl	76.8	52 - 132		77%	SPK: 100
1718-51-0	Terphenyl-d14	81.5	48 - 125		81%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	126000	7.652			
1146-65-2	Naphthalene-d8	523000	10.416			
15067-26-2	Acenaphthene-d10	339000	14.287			
1517-22-2	Phenanthrene-d10	687000	17.092			
1719-03-5	Chrysene-d12	769000	21.51			
1520-96-3	Perylene-d12	836000	24.786			

Report of Analysis

Client:	CDM Smith			Date Collected:	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	
Client Sample ID:	PB168098BSD			SDG No.:	Q2075
Lab Sample ID:	PB168098BSD			Matrix:	Water
Analytical Method:	8270E			% Solid:	0
Sample Wt/Vol:	1000	Units:	mL	Final Vol:	1000 uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3
Extraction Type :				Decanted :	N
Injection Volume :				Level :	LOW
Prep Method :	GPC Factor : 1.0			GPC Cleanup :	N
				PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BP024756.D	1	05/21/25 08:40	05/22/25 12:58	PB168098

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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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

Report of Analysis

Client:	CDM Smith			Date Collected:	05/15/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	SS-11MS			SDG No.:	Q2075	
Lab Sample ID:	Q2075-04MS			Matrix:	SOIL	
Analytical Method:	8270E			% Solid:	87.4	
Sample Wt/Vol:	30.07	Units:	g	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3541					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142487.D	1	05/20/25 09:45	05/21/25 15:06	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	1600	25.9		190	ug/Kg
208-96-8	Acenaphthylene	1600	33.0		190	ug/Kg
83-32-9	Acenaphthene	1800	24.3		190	ug/Kg
86-73-7	Fluorene	1700	28.9		190	ug/Kg
85-01-8	Phenanthrene	1600	23.9		190	ug/Kg
120-12-7	Anthracene	1700	38.0		190	ug/Kg
206-44-0	Fluoranthene	2200	34.2		190	ug/Kg
129-00-0	Pyrene	1300	41.1		190	ug/Kg
56-55-3	Benzo(a)anthracene	1700	26.3		190	ug/Kg
218-01-9	Chrysene	1600	22.7		190	ug/Kg
205-99-2	Benzo(b)fluoranthene	1900	21.7		190	ug/Kg
207-08-9	Benzo(k)fluoranthene	1600	25.6		190	ug/Kg
50-32-8	Benzo(a)pyrene	1700	33.7		190	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	1400	33.2		190	ug/Kg
53-70-3	Dibenz(a,h)anthracene	1400	31.3		190	ug/Kg
191-24-2	Benzo(g,h,i)perylene	1300	29.3		190	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	60.4	18 - 107		60%	SPK: 100
321-60-8	2-Fluorobiphenyl	51.6	20 - 109		52%	SPK: 100
1718-51-0	Terphenyl-d14	41.5	10 - 105		41%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	54800	6.898			
1146-65-2	Naphthalene-d8	201000	8.181			
15067-26-2	Acenaphthene-d10	114000	9.939			
1517-22-2	Phenanthrene-d10	229000	11.427			
1719-03-5	Chrysene-d12	211000	14.068			
1520-96-3	Perylene-d12	176000	15.563			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-11MS	SDG No.:	Q2075
Lab Sample ID:	Q2075-04MS	Matrix:	SOIL
Analytical Method:	8270E	% Solid:	87.4
Sample Wt/Vol:	30.07 Units: g	Final Vol:	1000 uL
Soil Aliquot Vol:	uL	Test:	SVOCMS Group3
Extraction Type :	Decanted : N	Level :	LOW
Injection Volume :	GPC Factor : 1.0	GPC Cleanup :	N PH :
Prep Method :	SW3541		

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142487.D	1	05/20/25 09:45	05/21/25 15:06	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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U = Not Detected

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LOD = Limit of Detection

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Q = indicates LCS control criteria did not meet requirements

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* = Values outside of QC limits

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A = Aldol-Condensation Reaction Products

Report of Analysis

Client:	CDM Smith			Date Collected:	05/15/25	
Project:	Con Ed UTEN Mount Vernon, NY			Date Received:	05/16/25	
Client Sample ID:	SS-11MSD			SDG No.:	Q2075	
Lab Sample ID:	Q2075-05MSD			Matrix:	SOIL	
Analytical Method:	8270E			% Solid:	87.4	
Sample Wt/Vol:	30.03	Units:	g	Final Vol:	1000	uL
Soil Aliquot Vol:	uL			Test:	SVOCMS Group3	
Extraction Type :				Decanted :	N	Level :
Injection Volume :				GPC Factor :	1.0	GPC Cleanup : N PH :
Prep Method :	SW3541					

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142488.D	1	05/20/25 09:45	05/21/25 15:34	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units(Dry Weight)
TARGETS						
91-20-3	Naphthalene	1600		25.9	190	ug/Kg
208-96-8	Acenaphthylene	1600		33.0	190	ug/Kg
83-32-9	Acenaphthene	1800		24.3	190	ug/Kg
86-73-7	Fluorene	1600		28.9	190	ug/Kg
85-01-8	Phenanthrene	1700		23.9	190	ug/Kg
120-12-7	Anthracene	1700		38.1	190	ug/Kg
206-44-0	Fluoranthene	2300		34.3	190	ug/Kg
129-00-0	Pyrene	1300		41.1	190	ug/Kg
56-55-3	Benzo(a)anthracene	1600		26.3	190	ug/Kg
218-01-9	Chrysene	1600		22.7	190	ug/Kg
205-99-2	Benzo(b)fluoranthene	1700		21.7	190	ug/Kg
207-08-9	Benzo(k)fluoranthene	1700		25.6	190	ug/Kg
50-32-8	Benzo(a)pyrene	1700		33.7	190	ug/Kg
193-39-5	Indeno(1,2,3-cd)pyrene	1400		33.3	190	ug/Kg
53-70-3	Dibenz(a,h)anthracene	1300		31.3	190	ug/Kg
191-24-2	Benzo(g,h,i)perylene	1300		29.4	190	ug/Kg
SURROGATES						
4165-60-0	Nitrobenzene-d5	59.6		18 - 107	60%	SPK: 100
321-60-8	2-Fluorobiphenyl	54.4		20 - 109	54%	SPK: 100
1718-51-0	Terphenyl-d14	40.2		10 - 105	40%	SPK: 100
INTERNAL STANDARDS						
3855-82-1	1,4-Dichlorobenzene-d4	53500	6.898			
1146-65-2	Naphthalene-d8	192000	8.181			
15067-26-2	Acenaphthene-d10	95700	9.939			
1517-22-2	Phenanthrene-d10	179000	11.427			
1719-03-5	Chrysene-d12	174000	14.068			
1520-96-3	Perylene-d12	148000	15.557			

Report of Analysis

Client:	CDM Smith	Date Collected:	05/15/25
Project:	Con Ed UTEN Mount Vernon, NY	Date Received:	05/16/25
Client Sample ID:	SS-11MSD	SDG No.:	Q2075
Lab Sample ID:	Q2075-05MSD	Matrix:	SOIL
Analytical Method:	8270E	% Solid:	87.4
Sample Wt/Vol:	30.03	Units:	g
Soil Aliquot Vol:		uL	
Extraction Type :		Decanted :	N
Injection Volume :		GPC Factor :	1.0
Prep Method :	SW3541	GPC Cleanup :	N
		Level :	LOW
		PH :	

File ID/Qc Batch:	Dilution:	Prep Date	Date Analyzed	Prep Batch ID
BF142488.D	1	05/20/25 09:45	05/21/25 15:34	PB168083

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
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U = Not Detected

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LOD = Limit of Detection

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J = Estimated Value

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A
B
C
D
E
F
G

CALIBRATION

SUMMARY

Method Path : Z:\svoasrv\HPCHEM1\BNA_F\Methods\
 Method File : 8270-BF052025.M
 Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION
 Last Update : Tue May 20 16:26:47 2025
 Response Via : Initial Calibration

Calibration Files

2.5 =BF142467.D 5 =BF142468.D 10 =BF142469.D 20 =BF142470.D 40 =BF142471.D 50 =BF142472.D 60 =BF142473.D 80 =BF142474.D

	Compound	2.5	5	10	20	40	50	60	80	Avg	%RSD
<hr/>											
1) I	1,4-Dichlorobenzene					-----ISTD-----					
2)	1,4-Dioxane	0.493	0.460	0.474	0.458	0.500	0.486	0.456	0.475	3.79	
3)	Pyridine	1.237	1.150	1.212	1.170	1.273	1.234	1.190	1.210	3.52	
4)	n-Nitrosodimethylamine	0.612	0.593	0.627	0.619	0.673	0.652	0.621	0.628	4.21	
5) S	2-Fluorophenol	1.264	1.214	1.225	1.125	1.220	1.164	1.098	1.187	5.03	
6)	Aniline	2.044	1.889	1.963	1.844	1.993	1.910	1.808	1.921	4.35	
7) S	Phenol-d6	1.530	1.449	1.459	1.367	1.467	1.403	1.328	1.429	4.77	
8)	2-Chlorophenol	1.345	1.293	1.315	1.252	1.338	1.285	1.223	1.293	3.44	
9)	Benzaldehyde	1.035	0.975	0.969	0.817	0.872	0.758	0.591	0.859	17.81	
10) C	Phenol	1.716	1.621	1.646	1.530	1.657	1.597	1.487	1.608	4.85	
11)	bis(2-Chloroethyl)ether	1.202	1.155	1.168	1.108	1.209	1.156	1.105	1.157	3.52	
12)	1,3-Dichlorobenzene	1.562	1.470	1.473	1.389	1.482	1.407	1.317	1.443	5.48	
13) C	1,4-Dichlorobenzene	1.540	1.476	1.491	1.407	1.495	1.430	1.335	1.453	4.70	
14)	1,2-Dichlorobenzene	1.495	1.405	1.436	1.327	1.432	1.357	1.284	1.391	5.20	
15)	Benzyl Alcohol	1.059	1.024	1.058	1.021	1.131	1.073	1.026	1.056	3.69	
16)	2,2'-oxybis(1,4-phenylene)	2.082	1.978	1.983	1.868	2.011	1.898	1.786	1.944	5.11	
17)	2-Methylphenol	1.040	0.992	1.026	0.976	1.053	1.015	0.965	1.010	3.27	
18)	Hexachloroethane	0.533	0.503	0.523	0.489	0.529	0.497	0.477	0.507	4.23	
19) P	n-Nitroso-di-n-butylamine	0.923	0.941	0.880	0.900	0.843	0.912	0.866	0.819	0.886	4.69
20)	3+4-Methylphenols	1.412	1.319	1.337	1.246	1.337	1.250	1.149	1.293	6.59	
21) I	Naphthalene-d8				-----ISTD-----						
22)	Acetophenone	0.480	0.453	0.459	0.429	0.452	0.428	0.399	0.443	5.98	
23) S	Nitrobenzene-d5	0.376	0.365	0.379	0.356	0.382	0.363	0.347	0.367	3.51	
24)	Nitrobenzene	0.338	0.328	0.338	0.323	0.343	0.331	0.316	0.331	2.94	
25)	Isophorone	0.636	0.615	0.620	0.593	0.638	0.607	0.585	0.613	3.26	
26) C	2-Nitrophenol	0.167	0.170	0.180	0.175	0.190	0.182	0.173	0.177	4.44	
27)	2,4-Dimethylphenol	0.315	0.315	0.318	0.303	0.325	0.312	0.295	0.312	3.22	
28)	bis(2-Chloroethyl)ether	0.406	0.394	0.394	0.364	0.391	0.375	0.358	0.383	4.63	
29) C	2,4-Dichlorophenol	0.288	0.282	0.290	0.276	0.300	0.283	0.266	0.283	3.74	
30)	1,2,4-Trichlorobenzene	0.325	0.313	0.317	0.295	0.320	0.300	0.284	0.308	4.89	
31)	Naphthalene	1.061	1.021	1.020	0.941	1.007	0.953	0.891	0.985	5.94	
32)	Benzoic acid		0.153	0.176	0.188	0.211	0.209	0.201	0.190	11.73	
33)	4-Chloroaniline	0.424	0.409	0.416	0.389	0.415	0.397	0.343	0.399	6.87	
34) C	Hexachlorobutane	0.203	0.192	0.197	0.187	0.198	0.192	0.176	0.192	4.52	
35)	Caprolactam	0.081	0.076	0.083	0.079	0.085	0.078	0.076	0.080	4.29	
36) C	4-Chloro-3-methylphenol	0.304	0.290	0.299	0.282	0.301	0.287	0.271	0.291	4.02	
37)	2-Methylnaphthalene	0.679	0.638	0.646	0.590	0.631	0.598	0.556	0.620	6.62	
38)	1-Methylnaphthalene	0.703	0.668	0.672	0.615	0.650	0.611	0.566	0.641	7.19	

Method Path : Z:\svoasrv\HPCHEM1\BNA_F\Methods\
 Method File : 8270-BF052025.M

		ISTD-----										
39)	I	Acenaphthene-d10	0.592	0.580	0.588	0.552	0.592	0.573	0.544	0.574	3.39	
40)		1,2,4,5-Tetrac...	0.318	0.350	0.383	0.393	0.430	0.425	0.411	0.387	10.57	A
41)	P	Hexachlorocycl...	0.230	0.225	0.234	0.211	0.233	0.218	0.202	0.222	5.39	B
42)	S	2,4,6-Tribromo...	0.387	0.373	0.406	0.371	0.406	0.388	0.379	0.387	3.69	C
43)	C	2,4,6-Trichlor...	0.410	0.411	0.416	0.395	0.437	0.408	0.381	0.408	4.27	D
44)		2,4,5-Trichlor...	1.726	1.619	1.558	1.387	1.480	1.396	1.268	1.490	10.46	E
45)	S	2-Fluorobiphenyl	1.672	1.605	1.595	1.480	1.595	1.507	1.408	1.552	5.82	F
46)		1,1'-Biphenyl	1.218	1.170	1.177	1.094	1.183	1.122	1.061	1.146	4.84	G
47)		2-Chloronaphth...	0.333	0.318	0.338	0.324	0.354	0.332	0.320	0.331	3.72	
48)		2-Nitroaniline	2.064	1.982	2.027	1.851	1.998	1.885	1.745	1.936	5.85	
49)		Acenaphthylene	1.441	1.340	1.367	1.255	1.366	1.256	1.211	1.320	6.16	
50)		Dimethylphthalate	0.290	0.283	0.289	0.280	0.302	0.281	0.268	0.285	3.74	
51)		2,6-Dinitrotol...	1.259	1.222	1.227	1.120	1.223	1.146	1.077	1.182	5.72	
52)	C	Acenaphthene	0.320	0.309	0.327	0.299	0.330	0.305	0.287	0.311	5.02	
53)		3-Nitroaniline	0.110	0.137	0.149	0.169	0.160	0.158	0.147	14.36		
54)	P	2,4-Dinitrophenol	1.886	1.766	1.768	1.606	1.736	1.635	1.509	1.701	7.38	
55)		Dibenzofuran	0.221	0.217	0.242	0.227	0.252	0.229	0.220	0.230	5.57	
56)	P	4-Nitrophenol	0.372	0.367	0.387	0.364	0.398	0.372	0.344	0.372	4.62	
57)		2,4-Dinitrotol...	1.477	1.399	1.372	1.231	1.343	1.238	1.140	1.314	8.85	
58)		Fluorene	0.347	0.336	0.360	0.333	0.361	0.333	0.317	0.341	4.71	
59)		2,3,4,6-Tetrac...	1.422	1.310	1.359	1.231	1.343	1.218	1.140	1.289	7.51	
60)		Diethylphthalate	0.724	0.678	0.676	0.609	0.653	0.612	0.566	0.646	8.25	
61)		4-Chlorophenyl...	0.303	0.283	0.303	0.277	0.294	0.265	0.251	0.282	6.95	
62)		4-Nitroaniline	1.239	1.194	1.188	1.107	1.197	1.123	1.055	1.158	5.55	
63)		Azobenzene										
64)	I	Phenanthrene-d10	ISTD-----									
65)		4,6-Dinitro-2....	0.086	0.094	0.110	0.115	0.129	0.125	0.123	0.112	14.72	
66)	c	n-Nitrosodiphe...	0.712	0.682	0.696	0.649	0.697	0.694	0.660	0.684	3.30	
67)		4-Bromophenyl....	0.240	0.235	0.239	0.222	0.246	0.243	0.230	0.236	3.45	
68)		Hexachlorobenzene	0.265	0.267	0.263	0.251	0.273	0.262	0.250	0.262	3.19	
69)		Atrazine	0.184	0.174	0.186	0.178	0.196	0.181	0.174	0.182	4.29	
70)	C	Pentachlorophenol	0.130	0.135	0.149	0.147	0.162	0.157	0.154	0.148	7.88	
71)		Phenanthrene	1.196	1.094	1.100	1.012	1.085	1.033	0.964	1.069	7.00	
72)		Anthracene	1.191	1.132	1.124	1.036	1.110	1.048	0.996	1.091	6.15	
73)		Carbazole	1.030	0.968	0.982	0.889	0.950	0.877	0.832	0.933	7.39	
74)		Di-n-butylphth...	1.108	1.039	1.083	0.976	1.059	0.974	0.908	1.021	6.95	
75)	C	Fluoranthene	1.177	1.081	1.052	0.938	0.997	0.901	0.846	0.999	11.41	
76)	I	Chrysene-d12	ISTD-----									
77)		Benzidine	0.670	0.782	0.903	0.797	0.805	0.698	0.556	0.744	15.12	
78)		Pyrene	1.929	1.967	2.066	1.859	1.959	1.773	1.507	1.866	9.79	
79)	S	Terphenyl-d14	1.624	1.582	1.660	1.423	1.492	1.337	1.126	1.464	12.80	
80)		Butylbenzylpht...	0.462	0.453	0.525	0.526	0.575	0.550	0.517	0.516	8.54	
81)		Benzo(a)anthra...	1.424	1.287	1.403	1.278	1.391	1.327	1.238	1.336	5.36	
82)		3,3'-Dichlorob...	0.360	0.364	0.402	0.407	0.444	0.442	0.417	0.405	8.30	
83)		Chrysene	1.222	1.204	1.193	1.145	1.255	1.223	1.158	1.200	3.20	
84)		Bis(2-ethylhex...	0.510	0.548	0.618	0.673	0.765	0.778	0.727	0.660	15.91	
85)	c	Di-n-octyl pht...	0.958	1.108	1.266	1.432	1.542	1.437	1.290	17.30		

Method Path : Z:\svoasrv\HPCHEM1\BNA_F\Methods\

Method File : 8270-BF052025.M

86)	I	Perylene-d12	-----ISTD-----								
87)		Indeno(1,2,3-c...)	1.421	1.495	1.583	1.448	1.583	1.546	1.434	1.501	4.64
88)		Benzo(b)fluora...	1.317	1.105	1.293	1.126	1.209	1.122	1.114	1.184	7.62
89)		Benzo(k)fluora...	1.191	1.166	1.032	1.042	1.178	1.128	1.023	1.109	6.68
90)	C	Benzo(a)pyrene	1.154	1.081	1.114	1.081	1.185	1.133	1.062	1.116	4.00
91)		Dibenzo(a,h)an...	1.152	1.228	1.275	1.184	1.290	1.245	1.149	1.218	4.67
92)		Benzo(g,h,i)pe...	1.154	1.220	1.270	1.180	1.299	1.245	1.158	1.218	4.64

(#) = Out of Range

A
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Method Path : Z:\svoasrv\HPCHEM1\BNA_P\Methods\
 Method File : 8270E-BP051325.M
 Title : ASP BNA STANDARDS FOR 5 POINT CALIBRATION
 Last Update : Tue May 13 16:21:39 2025
 Response Via : Initial Calibration

Calibration Files

2.5 =BP024614.D 5 =BP024615.D 10 =BP024616.D 20 =BP024617.D 40 =BP024618.D 50 =BP024619.D 60 =BP024620.D 80 =BP0246
21.D

	Compound	2.5	5	10	20	40	50	60	80	Avg	%RSD	
<hr/>												
1) I	1,4-Dichlorobenzene					ISTD						
2)	1,4-Dioxane	0.641	0.557	0.535	0.505	0.562	0.530	0.513	0.549	8.31		
3)	Pyridine	1.054	1.135	1.202	1.198	1.363	1.307	1.263	1.218	8.57		
4)	n-Nitrosodimethylamine	0.536	0.507	0.527	0.517	0.580	0.557	0.537	0.538	4.61		
5) S	2-Fluorophenol	1.134	1.096	1.162	1.105	1.272	1.223	1.193	1.169	5.50		
6)	Aniline	1.717	1.703	1.932	1.917	2.130	2.091	1.999	1.927	8.67		
7) S	Phenol-d6	1.301	1.344	1.486	1.472	1.671	1.620	1.552	1.492	9.09		
8)	2-Chlorophenol	1.200	1.204	1.311	1.303	1.465	1.433	1.376	1.327	7.83		
9)	Benzaldehyde	0.991	1.008	1.046	0.955	1.043	0.954	0.766	0.966	9.91		
10) C	Phenol	1.373	1.401	1.526	1.509	1.707	1.670	1.597	1.540	8.24		
11)	bis(2-Chloroethyl)ether	1.317	1.170	1.257	1.213	1.357	1.299	1.247	1.266	5.04		
12)	1,3-Dichlorobenzene	1.596	1.487	1.520	1.440	1.622	1.546	1.480	1.527	4.29		
13) C	1,4-Dichlorobenzene	1.558	1.505	1.525	1.470	1.637	1.567	1.491	1.536	3.67		
14)	1,2-Dichlorobenzene	1.530	1.467	1.498	1.426	1.579	1.529	1.449	1.497	3.56		
15)	Benzyl Alcohol	0.883	0.950	1.128	1.149	1.306	1.288	1.236	1.134	14.43		
16)	2,2'-oxybis(1-phenylpropane)	1.543	1.445	1.520	1.467	1.617	1.564	1.454	1.516	4.22		
17)	2-Methylphenol	0.966	0.949	1.105	1.099	1.246	1.213	1.154	1.105	10.30		
18)	Hexachloroethane	0.606	0.591	0.582	0.551	0.623	0.597	0.570	0.589	4.04		
19) P	n-Nitroso-di-n-butylamine	0.880	0.977	0.967	1.086	1.048	1.142	1.132	1.026	1.032	8.63	
20)	3+4-Methylphenols	1.221	1.302	1.503	1.515	1.702	1.692	1.584	1.503	12.21		
<hr/>												
21) I	Naphthalene-d8				ISTD							
22)	Acetophenone	0.486	0.481	0.511	0.483	0.534	0.512	0.491	0.500	3.97		
23) S	Nitrobenzene-d5	0.381	0.376	0.401	0.382	0.429	0.408	0.394	0.396	4.67		
24)	Nitrobenzene	0.333	0.340	0.355	0.337	0.373	0.356	0.344	0.348	4.03		
25)	Isophorone	0.644	0.640	0.695	0.673	0.753	0.721	0.682	0.687	5.91		
26) C	2-Nitrophenol	0.135	0.143	0.168	0.172	0.196	0.192	0.189	0.171	14.05		
27)	2,4-Dimethylphenol	0.273	0.279	0.300	0.298	0.329	0.318	0.309	0.301	6.60		
28)	bis(2-Chloroethyl)ether	0.403	0.391	0.411	0.399	0.443	0.419	0.405	0.410	4.15		
29) C	2,4-Dichlorophenol	0.250	0.258	0.290	0.291	0.327	0.316	0.309	0.292	9.96		
30)	1,2,4-Trichlorobenzene	0.346	0.322	0.327	0.306	0.347	0.329	0.323	0.329	4.33		
31)	Naphthalene	1.064	1.026	1.049	0.980	1.091	1.037	1.007	1.036	3.54		
32)	Benzoic acid		0.089	0.146	0.191	0.215	0.220	0.225	0.181	29.55		
33)	4-Chloroaniline	0.348	0.371	0.420	0.418	0.465	0.462	0.442	0.418	10.62		
34) C	Hexachlorobutane	0.231	0.214	0.211	0.198	0.222	0.209	0.206	0.213	5.08		
35)	Caprolactam	0.082	0.092	0.104	0.109	0.121	0.121	0.110	0.106	13.80		
36) C	4-Chloro-3-methylphenol	0.316	0.325	0.358	0.356	0.391	0.384	0.359	0.356	7.78		
37)	2-Methylnaphthalene	0.673	0.654	0.670	0.640	0.713	0.693	0.654	0.671	3.73		
38)	1-Methylnaphthalene	0.741	0.700	0.720	0.690	0.751	0.733	0.690	0.718	3.47		

Method Path : Z:\svoasrv\HPCHEM1\BNA_P\Methods\
 Method File : 8270E-BP051325.M

39) I	Acenaphthene-d10	-----ISTD-----	
40)	1,2,4,5-Tetrac...	0.593 0.558 0.574 0.549 0.625 0.580 0.586 0.581	4.26
41) P	Hexachlorocycl...	0.250 0.276 0.350 0.356 0.420 0.391 0.421 0.352	19.02
42) S	2,4,6-Tribromo...	0.324 0.321 0.335 0.323 0.369 0.357 0.346 0.339	5.49
43) C	2,4,6-Trichlor...	0.305 0.341 0.375 0.371 0.428 0.410 0.402 0.376	11.29
44)	2,4,5-Trichlor...	0.376 0.375 0.418 0.404 0.471 0.447 0.440 0.419	8.68
45) S	2-Fluorobiphenyl	1.601 1.498 1.553 1.432 1.602 1.515 1.485 1.527	4.10
46)	1,1'-Biphenyl	1.524 1.434 1.492 1.383 1.586 1.473 1.456 1.478	4.40
47)	2-Chloronaphth...	1.147 1.106 1.131 1.065 1.196 1.132 1.113 1.127	3.56
48)	2-Nitroaniline	0.279 0.304 0.331 0.335 0.381 0.364 0.343 0.334	10.28
49)	Acenaphthylene	1.845 1.818 1.920 1.804 2.029 1.930 1.856 1.886	4.19
50)	Dimethylphthalate	1.570 1.527 1.518 1.450 1.611 1.535 1.460 1.524	3.73
51)	2,6-Dinitrotol...	0.299 0.308 0.326 0.312 0.354 0.337 0.319 0.322	5.75
52) C	Acenaphthene	1.105 1.054 1.094 1.028 1.159 1.099 1.052 1.084	4.02
53)	3-Nitroaniline	0.258 0.275 0.329 0.340 0.379 0.370 0.350 0.329	14.04
54) P	2,4-Dinitrophenol	0.098 0.143 0.172 0.201 0.201 0.201 0.169	24.79
55)	Dibenzofuran	1.810 1.716 1.740 1.633 1.816 1.733 1.658 1.730	4.00
56) P	4-Nitrophenol	0.159 0.222 0.257 0.299 0.288 0.274 0.250	20.82
57)	2,4-Dinitrotol...	0.400 0.422 0.456 0.454 0.508 0.483 0.454 0.454	7.87
58)	Fluorene	1.466 1.415 1.410 1.336 1.494 1.410 1.341 1.410	4.14
59)	2,3,4,6-Tetrac...	0.367 0.364 0.381 0.379 0.428 0.406 0.394 0.388	5.89
60)	Diethylphthalate	1.593 1.542 1.537 1.444 1.639 1.563 1.468 1.541	4.40
61)	4-Chlorophenyl...	0.752 0.697 0.700 0.656 0.741 0.709 0.668 0.703	5.01
62)	4-Nitroaniline	0.228 0.259 0.326 0.330 0.375 0.363 0.341 0.317	17.06
63)	Azobenzene	1.327 1.299 1.356 1.250 1.407 1.343 1.243 1.318	4.45
64) I	Phenanthrene-d10	-----ISTD-----	
65)	4,6-Dinitro-2....	0.094 0.120 0.129 0.149 0.143 0.141 0.129	15.64
66) c	n-Nitrosodiphe...	0.603 0.589 0.627 0.593 0.667 0.621 0.600 0.614	4.41
67)	4-Bromophenyl....	0.231 0.223 0.237 0.226 0.252 0.242 0.240 0.236	4.31
68)	Hexachlorobenzene	0.303 0.288 0.298 0.282 0.321 0.304 0.300 0.299	4.25
69)	Atrazine	0.210 0.209 0.227 0.218 0.248 0.237 0.226 0.225	6.32
70) C	Pentachlorophenol	0.124 0.137 0.163 0.172 0.198 0.191 0.192 0.168	17.12
71)	Phenanthrene	1.144 1.085 1.125 1.051 1.170 1.125 1.083 1.112	3.68
72)	Anthracene	1.097 1.064 1.142 1.075 1.205 1.139 1.108 1.119	4.31
73)	Carbazole	0.923 0.944 1.039 1.001 1.114 1.053 1.013 1.012	6.43
74)	Di-n-butylphth...	1.182 1.218 1.332 1.260 1.447 1.363 1.313 1.302	6.97
75) C	Fluoranthene	1.271 1.254 1.314 1.250 1.399 1.337 1.272 1.300	4.18
76) I	Chrysene-d12	-----ISTD-----	
77)	Benzidine	0.362 0.543 0.560 0.657 0.599 0.548 0.545	18.22
78)	Pyrene	1.184 1.137 1.197 1.153 1.273 1.222 1.223 1.198	3.85
79) S	Terphenyl-d14	1.192 1.113 1.165 1.125 1.226 1.175 1.167 1.166	3.30
80)	Butylbenzylpht...	0.458 0.465 0.537 0.527 0.604 0.571 0.569 0.533	10.28
81)	Benzo(a)anthra...	1.251 1.218 1.253 1.208 1.336 1.275 1.250 1.256	3.36
82)	3,3'-Dichlorob...	0.357 0.392 0.474 0.477 0.539 0.511 0.514 0.466	14.45
83)	Chrysene	1.214 1.180 1.185 1.131 1.255 1.188 1.180 1.190	3.16
84)	Bis(2-ethylhex...	0.673 0.683 0.801 0.766 0.884 0.836 0.841 0.783	10.32
85) c	Di-n-octyl pht...	1.048 1.116 1.272 1.280 1.458 1.376 1.417 1.281	11.95

Method Path : Z:\svoasrv\HPCHEM1\BNA_P\Methods\

Method File : 8270E-BP051325.M

86)	I	Perylene-d12	- - - - - ISTD - - - - -											
87)		Indeno(1,2,3-c...)	1.341	1.357	1.447	1.424	1.608	1.549	1.494	1.460				6.70
88)		Benzo(b)fluora...	1.028	1.070	1.152	1.103	1.288	1.184	1.189	1.145				7.58
89)		Benzo(k)fluora...	1.147	1.106	1.182	1.140	1.258	1.238	1.185	1.180				4.59
90)	C	Benzo(a)pyrene	0.995	1.001	1.112	1.064	1.221	1.167	1.136	1.099				7.68
91)		Dibenzo(a,h)an...	1.068	1.066	1.186	1.167	1.321	1.271	1.236	1.188				8.18
92)		Benzo(g,h,i)pe...	1.101	1.106	1.167	1.147	1.294	1.242	1.212	1.181				6.07

(#) = Out of Range

A B C D E F G

7C

SEMOVOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH		Contract:	CAMP02	
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075
Instrument ID:	BNA_F		Calibration Date/Time:	05/21/2025	10:40
Lab File ID:	BF142478.D		Init. Calib. Date(s):	05/20/2025	05/20/2025
EPA Sample No.:	SSTDCCCC040		Init. Calib. Time(s):	12:10	15:31
GC Column:	DB-UI	ID: 0.18	(mm)		

COMPOUND	RRF	RRF040	MIN RRF	%D	MAX%D
2-Fluorophenol	1.187	1.087		-8.4	
Phenol-d6	1.429	1.358		-5.0	
Nitrobenzene-d5	0.367	0.343		-6.5	
Naphthalene	0.985	0.928		-5.8	
2-Fluorobiphenyl	1.490	1.373		-7.9	
Acenaphthylene	1.936	1.795		-7.3	
Acenaphthene	1.182	1.125		-4.8	20.0
Fluorene	1.314	1.229		-6.5	
2,4,6-Tribromophenol	0.222	0.225		1.4	
Phenanthrene	1.069	0.989		-7.5	
Anthracene	1.091	1.022		-6.3	
Fluoranthene	0.999	0.984		-1.5	20.0
Pyrene	1.866	1.932		3.5	
Terphenyl-d14	1.464	1.466		0.1	
Benzo(a)anthracene	1.336	1.289		-3.5	
Chrysene	1.200	1.116		-7.0	
Benzo(b)fluoranthene	1.184	1.216		2.7	
Benzo(k)fluoranthene	1.109	0.996		-10.2	
Benzo(a)pyrene	1.116	1.061		-4.9	20.0
Indeno(1,2,3-cd)pyrene	1.501	1.409		-6.1	
Dibenzo(a,h)anthracene	1.218	1.156		-5.1	
Benzo(g,h,i)perylene	1.218	1.202		-1.3	

All other compounds must meet a minimum RRF of 0.010.

7C

SEMOVOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH		Contract:	CAMP02	
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075
Instrument ID:	BNA_F		Calibration Date/Time:	05/22/2025	11:11
Lab File ID:	BF142502.D		Init. Calib. Date(s):	05/20/2025	05/20/2025
EPA Sample No.:	SSTDCCC040		Init. Calib. Time(s):	12:10	15:31
GC Column:	DB-UI	ID: 0.18	(mm)		

COMPOUND	RRF	RRF040	MIN RRF	%D	MAX%D
2-Fluorophenol	1.187	1.158		-2.4	
Phenol-d6	1.429	1.379		-3.5	
Nitrobenzene-d5	0.367	0.356		-3.0	
Naphthalene	0.985	0.939		-4.7	
2-Fluorobiphenyl	1.490	1.389		-6.8	
Acenaphthylene	1.936	1.864		-3.7	
Acenaphthene	1.182	1.118		-5.4	20.0
Fluorene	1.314	1.233		-6.2	
2,4,6-Tribromophenol	0.222	0.208		-6.3	
Phenanthrene	1.069	1.009		-5.6	
Anthracene	1.091	1.031		-5.5	
Fluoranthene	0.999	0.878		-12.1	20.0
Pyrene	1.866	1.705		-8.6	
Terphenyl-d14	1.464	1.259		-14.0	
Benzo(a)anthracene	1.336	1.258		-5.8	
Chrysene	1.200	1.136		-5.3	
Benzo(b)fluoranthene	1.184	1.087		-8.2	
Benzo(k)fluoranthene	1.109	1.044		-5.9	
Benzo(a)pyrene	1.116	1.078		-3.4	20.0
Indeno(1,2,3-cd)pyrene	1.501	1.416		-5.7	
Dibenzo(a,h)anthracene	1.218	1.143		-6.2	
Benzo(g,h,i)perylene	1.218	1.144		-6.1	

All other compounds must meet a minimum RRF of 0.010.

7C

SEMOVOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH	Contract:	CAMP02				
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075	SDG No.:	Q2075
Instrument ID:	BNA_P	Calibration Date/Time:			05/21/2025	14:48	
Lab File ID:	BP024737.D	Init. Calib. Date(s):			05/13/2025	05/13/2025	
EPA Sample No.:	SSTDCCC040	Init. Calib. Time(s):			10:41	15:26	
GC Column:	ZB-GR	ID:	0.25	(mm)			

COMPOUND	RRF	RRF040	MIN RRF	%D	MAX%D
2-Fluorophenol	1.169	1.095		-6.3	
Phenol-d6	1.492	1.334		-10.6	
Nitrobenzene-d5	0.396	0.360		-9.1	
Naphthalene	1.036	1.000		-3.5	
2-Fluorobiphenyl	1.527	1.480		-3.1	
Acenaphthylene	1.886	1.807		-4.2	
Acenaphthene	1.084	1.033		-4.7	20.0
Fluorene	1.410	1.271		-9.9	
2,4,6-Tribromophenol	0.339	0.356		5.0	
Phenanthrene	1.112	1.063		-4.4	
Anthracene	1.119	1.090		-2.6	
Fluoranthene	1.300	1.252		-3.7	20.0
Pyrene	1.198	1.164		-2.8	
Terphenyl-d14	1.166	1.170		0.3	
Benzo(a)anthracene	1.256	1.204		-4.1	
Chrysene	1.190	1.132		-4.9	
Benzo(b)fluoranthene	1.145	1.108		-3.2	
Benzo(k)fluoranthene	1.180	1.122		-4.9	
Benzo(a)pyrene	1.099	1.059		-3.6	20.0
Indeno(1,2,3-cd)pyrene	1.460	1.437		-1.6	
Dibenzo(a,h)anthracene	1.188	1.179		-0.8	
Benzo(g,h,i)perylene	1.181	1.143		-3.2	

All other compounds must meet a minimum RRF of 0.010.

7C

SEMOVOLATILE CONTINUING CALIBRATION CHECK

Lab Name:	CHEMTECH		Contract:	CAMP02	
Lab Code:	CHEM	Case No.:	Q2075	SAS No.:	Q2075
Instrument ID:	BNA_P		Calibration Date/Time:	05/22/2025	10:56
Lab File ID:	BP024753.D		Init. Calib. Date(s):	05/13/2025	05/13/2025
EPA Sample No.:	SSTDCCCC040		Init. Calib. Time(s):	10:41	15:26
GC Column:	ZB-GR	ID: 0.25	(mm)		

COMPOUND	RRF	RRF040	MIN RRF	%D	MAX%D
2-Fluorophenol	1.169	1.135		-2.9	
Phenol-d6	1.492	1.407		-5.7	
Nitrobenzene-d5	0.396	0.368		-7.1	
Naphthalene	1.036	0.988		-4.6	
2-Fluorobiphenyl	1.527	1.502		-1.6	
Acenaphthylene	1.886	1.818		-3.6	
Acenaphthene	1.084	1.055		-2.7	20.0
Fluorene	1.410	1.365		-3.2	
2,4,6-Tribromophenol	0.339	0.364		7.4	
Phenanthrene	1.112	1.056		-5.0	
Anthracene	1.119	1.093		-2.3	
Fluoranthene	1.300	1.270		-2.3	20.0
Pyrene	1.198	1.114		-7.0	
Terphenyl-d14	1.166	1.110		-4.8	
Benzo(a)anthracene	1.256	1.196		-4.8	
Chrysene	1.190	1.132		-4.9	
Benzo(b)fluoranthene	1.145	1.090		-4.8	
Benzo(k)fluoranthene	1.180	1.078		-8.6	
Benzo(a)pyrene	1.099	1.059		-3.6	20.0
Indeno(1,2,3-cd)pyrene	1.460	1.467		0.5	
Dibenzo(a,h)anthracene	1.188	1.197		0.8	
Benzo(g,h,i)perylene	1.181	1.175		-0.5	

All other compounds must meet a minimum RRF of 0.010.



SHIPPING DOCUMENTS



284 Sheffield Street, Mountainside, NJ 07092
 (908) 789-8900 • Fax (908) 789-8922
www.chemtech.net

ALLIANCE PROJECT NO.

QUOTE NO.

COC Number

Q2075

7

2045940

CLIENT INFORMATION

CLIENT PROJECT INFORMATION

CLIENT BILLING INFORMATION

REPORT TO BE SENT TO:

COMPANY: CDM Smith

ADDRESS: 110 Fieldcrest Ave. #8 6th Flr

CITY Edison STATE: NJ ZIP: 08837

ATTENTION: M. Encinas

PHONE: 732 590 4679 FAX:

PROJECT NAME: UTEN UST

PROJECT NO.: LOCATION: Mt. Vernon NY

PROJECT MANAGER: M. Encinas

e-mail: Encinas.Ma@cdmsmith.com

PHONE: 732 590 4679 FAX:

BILL TO: CDM Smith

PO#:

ADDRESS: 110 Fieldcrest Ave #8 6th Flr

CITY Edison STATE: NJ ZIP: 08837

ATTENTION: M. Encinas PHONE: 732 590 4679

ANALYSIS

DATA TURNAROUND INFORMATION

FAX (RUSH) 5 business days*

HARDCOPY (DATA PACKAGE): 5 business days*

EDD: 5 business days*

*TO BE APPROVED BY CHEMTECH

STANDARD HARDCOPY TURNAROUND TIME IS 10 BUSINESS

DATA DELIVERABLE INFORMATION

- Level 1 (Results Only) Level 4 (QC + Full Raw Data)
 Level 2 (Results + QC) NJ Reduced US EPA CLP
 Level 3 (Results + QC) NYS ASP A NYS ASP B
 + Raw Data Other
 EDD FORMAT

1. *Method 51 Task 3*
 2. *Method 51 Task 3*
 3. *Method 51 Task 3*
 4. *Method 51 Task 3*
 5. *Method 51 Task 3*
 6. *Method 51 Task 3*
 7. *Method 51 Task 3*
 8. *Method 51 Task 3*
 9. *Method 51 Task 3*

PRESERVATIVES

COMMENTS

← Specify Preservatives
 A-HCl D-NaOH
 B-HNO3 E-ICE
 C-H₂SO₄ F-OTHER

CP-51 Task 3
 per email and
 see attached

ALLIANCE SAMPLE ID	PROJECT SAMPLE IDENTIFICATION	SAMPLE MATRIX	SAMPLE TYPE		SAMPLE COLLECTION DATE	TIME	# OF BOTTLES	PRESERVATIVES									COMMENTS			
			COMP	GRAB				A-A	E	F	1	2	3	4	5	6	7	8	9	
1.	FB-05152025	Ag			✓ 5/15/25	1230	3		V	V										
2.	SS-10	Soil			✓ 5/15/25	1310	5		J	J										
3.	SS-910	Soil			✓ 5/15/25	1315	5		J	J										
4.	SS-1L (ME/MED)	Soil			✓ 5/15/25	1400	15		V	V										
5.	SS-Merit-11.5	Soil			✓ 5/16/25	0835	5		V	V										
6.	FB-05162025	Ag			✓ 5/16/25	1535	3		V	V										
7.																				
8.																				
9.																				
10.																				

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

RELINQUISHED BY SAMPLER:

DATE/TIME: 5/16/25 1555

RECEIVED BY: 1. *CR*Conditions of bottles or coolers at receipt: COMPLIANT NON COMPLIANT COOLER TEMP

2.4 °C

Comments:

RELINQUISHED BY SAMPLER:

DATE/TIME:

RECEIVED BY: 2.

2.

DATE/TIME:

RECEIVED BY: 3.

DATE/TIME:

RECEIVED BY: 3.

Page ____ of _____

CLIENT: Hand Delivered Other

Shipment Complete

 YES NO

Laboratory Certification

Certified By	License No.
CAS EPA CLP Contract	68HERH20D0011
Connecticut	PH-0830
DOD ELAP (ANAB)	L2219
Maine	2024021
Maryland	296
New Hampshire	255424 Rev 1
New Jersey	20012
New York	11376
Pennsylvania	68-00548
Soil Permit	525-24-234-08441
Texas	T104704488

LOGIN REPORT/SAMPLE TRANSFER

Order ID : Q2075	CAMP02	Order Date : 5/16/2025 4:10:00 PM	Project Mgr :
Client Name : CDM Smith		Project Name : Con Ed UTEN Mount Verm	Report Type : NYS ASP B
Client Contact : Marcie Ann Encinas		Receive DateTime : 5/16/2025 3:55:00 PM	EDD Type : EQUIS
Invoice Name : CDM Smith		Purchase Order :	Hard Copy Date :
Invoice Contact : Marcie Ann Encinas			Date Signoff :

LAB ID	CLIENT ID	MATRIX	SAMPLE DATE	SAMPLE TIME	TEST	TEST GROUP	METHOD	FAX DATE	DUE DATES
Q2075-01	SS-10	Solid	05/15/2025	12:30 13:10	VOCMS Group3		8260D	10-Bus. Days	5 days
Q2075-02	SS-910	Solid	05/15/2025	13:10 13:15	VOCMS Group3		8260D	10-Bus. Days	
Q2075-03	SS-11	Solid	05/15/2025	13:15 14:00	VOCMS Group3		8260D	10-Bus. Days	
Q2075-04	Q2075-03MS	Solid	05/15/2025	13:15 14:00	VOCMS Group3		8260D	10-Bus. Days	
Q2075-05	Q2075-03MSD	Solid	05/15/2025	13:15 14:00	VOCMS Group3		8260D	10-Bus. Days	
Q2075-06	SS-MW1-11.5	Solid	05/16/2025	08:35	VOCMS Group3		8260D	10-Bus. Days	
Q2075-07	FB-05152025	Water	05/15/2025	12:30	VOCMS Group3		8260-Low	10 Bus. Days	
Q2075-08	FB-05162025	Water	05/16/2025	15:35					

LOGIN REPORT/SAMPLE TRANSFER

Order ID :	Q2075	CAMP02	Order Date :	5/16/2025 4:10:00 PM	Project Mgr :
Client Name :	CDM Smith		Project Name :	Con Ed UTEN Mount Vern	
Client Contact :	Marcie Ann Encinas		Receive Date/Time :	5/16/2025 3:55:00 PM	Report Type : NYS ASP B
Invoice Name :	CDM Smith		Purchase Order :		EDD Type : EQUIS
Invoice Contact :	Marcie Ann Encinas				Hard Copy Date :
					Date Signoff :

LAB ID	CLIENT ID	MATRIX	SAMPLE DATE	SAMPLE TIME	TEST	TEST GROUP	METHOD	FAX DATE	DUE DATES
					VOCMS Group3		8260-Low	40 Bus. Days	5 Days

Relinquished By : CL
 Date / Time : 5/19/25 11:30

Received By : Jam
 Date / Time : 05/19/25 11:30 DS 6 522
 Storage Area : VOA Refrigerator Room Bay # 4