

DATA PACKAGE

GENERAL CHEMISTRY
METALS
SEMI-VOLATILE ORGANICS
VOLATILE ORGANICS

PROJECT NAME: PVSC MONTHLY 2025

ARDMORE CHEMICAL

29 Riverside Avenue

Newark, NJ - 07104-

Phone No: 973-481-2406

ORDER ID: Q2006

ATTENTION: Michael Sharphouse







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Cover Page

Order ID:	Q2006
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Project ID: PVSC Monthly 2025

Client: Ardmore Chemical

Lab Sample Number Client Sample Number

Q2006-01 EFF-WW Q2006-02 EFF-WW

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

Signature :		
Signature .	———— Date:	5/23/2025

NYDOH CERTIFICATION NO - 11376 NJDEP CERTIFICATION NO - 20012

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CASE NARRATIVE

Ardmore Chemical

Project Name: PVSC Monthly 2025

Project # N/A Order ID # Q2006 Test Name: VOC-PP

A. Number of Samples and Date of Receipt:

2 Water samples were received on 05/09/2025.

B. Parameters

According to the Chain of Custody document, the following analyses were requested: BOD5, Cyanide, Mercury, Metals Group3, SVOCMS Group1, TSS and VOC-PP. This data package contains results for VOC-PP.

C. Analytical Techniques:

The analysis performed on instrument MSVOA_N were done using GC column Rxi-624SIL MS 30m, 0.25mm, 1.4 um, Cat. #13868.The analysis of VOC-PP was based on method 624.1.

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 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 %RSD is greater than 35% in the Initial Calibration method (624N042325W.M)

for Acrolein, this compound is passing on Linear Regression.

The Continuous Calibration met the requirements.

The Tuning criteria met requirements.

Sample EFF-WW was directly analyzed with 5X due to foamy Sample.

E. Additional Comments:

As per method, MS/MSD is required to be performed with the sample analysis. However, Lab did not receive sufficient volume to perform the MS/MSD therefore MS/MSD were not performed for this project. However, Lab has performed LCS/LCSD instead.

Trip Blank was not provided with this set of samples.

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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 <35% 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 > 35% 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.

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Signature		

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CASE NARRATIVE

Ardmore Chemical

Project Name: PVSC Monthly 2025

Project # N/A Order ID # Q2006

Test Name: SVOCMS Group1

A. Number of Samples and Date of Receipt:

2 Water samples were received on 05/09/2025.

B. Parameters

According to the Chain of Custody document, the following analyses were requested: BOD5, Cyanide, Mercury, Metals Group3, SVOCMS Group1, TSS and VOC-PP. This data package contains results for SVOCMS Group1.

C. Analytical Techniques:

The 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 Group1 was based on method 625.1 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 except for EFF-WW [2-

Fluorophenol - 45%, Phenol-d6 - 28%]. Due to matrix interference, which can be observed from the abnormal chromatogram. Hence this analysis will be final.

The Internal Standards Areas met the acceptable requirements.

The Retention Times were acceptable for all samples.

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 % RSD is greater than 20% in the Initial Calibration (8270-BP051325.M) for 2,4-

Dinitrophenol, 4-Nitrophenol, these compounds are passing on Linear Regression.

The Continuous Calibration met the requirements.

The Tuning criteria met requirements.

E. Additional Comments:

As per method, MS/MSD is required to be performed with the sample analysis. However, Lab did not receive sufficient volume to perform the MS/MSD therefore MS/MSD were not performed for this project. However, Lab has performed LCS/LCSD instead. 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

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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.

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Signature		

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CASE NARRATIVE

Ardmore Chemical

Project Name: PVSC Monthly 2025

Project # N/A Order ID # Q2006

Test Name: Mercury, Metals Group3

A. Number of Samples and Date of Receipt:

2 Water samples were received on 05/09/2025.

B. Parameters:

According to the Chain of Custody document, the following analyses were requested: BOD5, Cyanide, Mercury, Metals Group3, SVOCMS Group1, TSS and VOC-PP. This data package contains results for Mercury, Metals Group3.

C. Analytical Techniques:

The analysis and digestion of Metals Group3 was based on 200.7 and The analysis and digestion of Mercury was based on 245.1.

D. QA/ QC Samples:

The Holding Times were met for all analysis.

The Blank Spike met requirements for all samples.

The Duplicate (EFF-WWDUP) analysis met criteria for all samples except for Zinc due to sample matrix interference. The Duplicate (EFF-WWMSD) analysis met criteria for all samples except for Zinc due to Chemical Interference during Digestion Process.

The Matrix Spike analysis met criteria for all samples.

The Matrix Spike Duplicate (EFF-WWMSD) analysis met criteria for all samples except for Zinc due to Chemical Interference during Digestion Process.

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

The Calibration met the requirements.

The Serial Dilution (EFF-WWL) met criteria for all samples except for Zinc due to sample matrix interference.

E. Additional Comments:

The Post Digest Spike (EFF-WWA) analysis met criteria for all samples except for Zinc due to unknown chemical interferences of matrix with the addition of spike amount after digestion and before analysis, matrix has suppression effect during addition of spike.

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CASE NARRATIVE

Ardmore Chemical

Project Name: PVSC Monthly 2025

Project # N/A Order ID # O2006

Test Name: BOD5, Cyanide, TSS

A. Number of Samples and Date of Receipt:

2 Water samples were received on 05/09/2025.

B. Parameters:

According to the Chain of Custody document, the following analyses were requested: BOD5, Cyanide, Mercury, Metals Group3, SVOCMS Group1, TSS and VOC-PP. This data package contains results for BOD5, Cyanide, TSS.

C. Analytical Techniques:

The analysis of TSS was based on method SM2540 D, The analysis of Cyanide was based on method SM4500-CN C,E and The analysis of BOD5 was based on method SM5210 B.

D. QA/ QC Samples:

The Holding Times were met for all analysis.

The Blank Spike met requirements for all samples.

The Duplicate analysis met criteria for all samples.

The Matrix Spike analysis met criteria for all samples.

The Matrix Spike Duplicate analysis met criteria for all samples.

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

The Calibration met the requirements.

E. Additional Comments:

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.

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DATA REPORTING QUALIFIERS- INORGANIC

For reporting results, the following "Results Qualifiers" are used:

- J Indicates the reported value was obtained from a reading that was less than the Contract Required Detection Limit (CRDL), but greater than or equal to the Instrument Detection Limit (IDL).
- U Indicates the analyte was analyzed for, but not detected.
- ND Indicates the analyte was analyzed for, but not detected
- E Indicates the reported value is estimated because of the presence of interference
- M Indicates Duplicate injection precision not met.
- N Indicates the spiked sample recovery is not within control limits.
- S Indicates the reported value was determined by the Method of Standard Addition (MSA).
- * Indicates that the duplicate analysis is not within control limits.
- + Indicates the correlation coefficient for the MSA is less than 0.995.
- D Indicates the reported value is from a secondary analysis with a dilution factor. The original analysis exceeded the calibration range.
- M Method qualifiers
 - **"P"** for ICP instrument
 - "PM" for ICP when Microwave Digestion is used
 - "CV" for Manual Cold Vapor AA
 "AV" for automated Cold Vapor AA
 - "CA" for MIDI-Distillation Spectrophotometric
 - "AS" for Semi –Automated Spectrophotometric
 - "C" for Manual Spectrophotometric
 - **"T"** for Titrimetric
 - "NR" for analyte not required to be analyzed
- OR Indicates the analyte's concentration exceeds the calibrated range of the instrument for that specific analysis.
 - instrument for that speeme analysis.
- Q Indicates the LCS did not meet the control limits requirements
- H Sample Analysis Out Of Hold Time



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 is flag is used when similar situation arise on any organic parameter i.e. Pest, PCB and others. 						
В	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						

Aliance
TECHNICAL GROUP

APPENDIX A

QA REVIEW GENERAL DOCUMENTATION

Project #: Q2006

	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)	<u> </u>
Check chain-of-custody for proper relinquish/return of samples	<u> </u>
Is the chain of custody signed and complete	<u> </u>
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	<u> </u>
COVER PAGE:	
Do numbers of samples correspond to the number of samples in the Chain of Custody on login page	<u> </u>
Do lab numbers and client Ids on cover page agree with the Chain of Custody	<u> </u>
CHAIN OF CUSTODY:	
Do requested analyses on Chain of Custody agree with form I results	<u> </u>
Do requested analyses on Chain of Custody agree with the log-in page	<u>*</u> <u>*</u> <u>*</u>
Were the correct method log-in for analysis according to the Analytical Request and Chain of Castody	<u> </u>
Were the samples received within hold time	<u> </u>
Were any problems found with the samples at arrival recorded in the Sample Management Laboratory Chronicle	<u> </u>
ANALYTICAL:	
Was method requirement followed?	<u> </u>
Was client requirement followed?	<u> </u>
Does the case narrative summarize all QC failure?	√ √ √
All runlogs and manual integration are reviewed for requirements	<u> </u>
All manual calculations and /or hand notations verified	

QA Review Signature: SOHIL JODHANI Date: 05/23/2025

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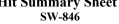


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Hit Summary Sheet

SDG No.: Q2006

Client: Ardmore Chemical





Sample ID	Client ID	Matrix	Parameter	Concentration	C MDL	RDL	Units
Client ID: Q2006-01	EFF-WW EFF-WW	Water	Chloroform	31.9	2.80	25.0	ug/L
			Total Voc:	31.9			
			Total Concentration:	31.9			

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SAMPLE DATA



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Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 Project: Date Received: PVSC Monthly 2025 05/09/25 Client Sample ID: SDG No.: Q2006 EFF-WW Q2006-01 Matrix: Lab Sample ID: Water Analytical Method: E624.1 % Solid: Sample Wt/Vol: 5 Final Vol: uL Units: mL5000 Soil Aliquot Vol: uL Test: VOC-PP GC Column: RXI-624 ID: 0.25 Level: LOW

Prep Method:

File ID/Qc Batch: Dilution: Prep Date Date Analyzed Prep Batch ID VN086665.D 5 05/16/25 13:54 VN051625

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
74-87-3	Chloromethane	3.20	U	3.20	25.0	ug/L
75-01-4	Vinyl Chloride	4.20	U	4.20	25.0	ug/L
74-83-9	Bromomethane	4.00	U	4.00	25.0	ug/L
75-00-3	Chloroethane	11.6	U	11.6	25.0	ug/L
75-69-4	Trichlorofluoromethane	4.00	U	4.00	25.0	ug/L
75-35-4	1,1-Dichloroethene	3.80	U	3.80	25.0	ug/L
107-02-8	Acrolein	33.1	U	33.1	130	ug/L
107-13-1	Acrylonitrile	14.0	U	14.0	130	ug/L
75-09-2	Methylene Chloride	4.30	U	4.30	25.0	ug/L
156-60-5	trans-1,2-Dichloroethene	4.10	U	4.10	25.0	ug/L
75-34-3	1,1-Dichloroethane	3.40	U	3.40	25.0	ug/L
56-23-5	Carbon Tetrachloride	3.70	U	3.70	25.0	ug/L
67-66-3	Chloroform	31.9		2.80	25.0	ug/L
71-55-6	1,1,1-Trichloroethane	3.20	U	3.20	25.0	ug/L
71-43-2	Benzene	2.30	U	2.30	25.0	ug/L
107-06-2	1,2-Dichloroethane	2.50	U	2.50	25.0	ug/L
79-01-6	Trichloroethene	2.50	U	2.50	25.0	ug/L
78-87-5	1,2-Dichloropropane	2.30	U	2.30	25.0	ug/L
75-27-4	Bromodichloromethane	3.20	U	3.20	25.0	ug/L
108-88-3	Toluene	2.30	U	2.30	25.0	ug/L
10061-02-6	t-1,3-Dichloropropene	3.60	U	3.60	25.0	ug/L
10061-01-5	cis-1,3-Dichloropropene	3.40	U	3.40	25.0	ug/L
79-00-5	1,1,2-Trichloroethane	2.30	U	2.30	25.0	ug/L
110-75-8	2-Chloroethyl vinyl ether	23.2	U	23.2	130	ug/L
124-48-1	Dibromochloromethane	3.30	U	3.30	25.0	ug/L
127-18-4	Tetrachloroethene	4.20	U	4.20	25.0	ug/L
108-90-7	Chlorobenzene	2.40	U	2.40	25.0	ug/L
100-41-4	Ethyl Benzene	2.80	U	2.80	25.0	ug/L
179601-23-1	m/p-Xylenes	6.50	U	6.50	50.0	ug/L
95-47-6	o-Xylene	3.40	U	3.40	25.0	ug/L

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Test:

VOC-PP

Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 Date Received: Project: PVSC Monthly 2025 05/09/25 Client Sample ID: SDG No.: Q2006 EFF-WW Lab Sample ID: Q2006-01 Matrix: Water Analytical Method: E624.1 % Solid: 5 Final Vol: Sample Wt/Vol: Units: mL5000 uL

GC Column: RXI-624 ID: 0.25 Level: LOW

uL

Prep Method:

Soil Aliquot Vol:

File ID/Qc Batch: Dilution: Prep Date Date Analyzed Prep Batch ID VN086665.D 5 05/16/25 13:54 VN051625

CAS Number	Parameter	Conc.	Conc. Qualifier MDL		LOQ / CRQL	Units
75-25-2	Bromoform	4.70	U	4.70	25.0	ug/L
79-34-5	1,1,2,2-Tetrachloroethane	2.20	U	2.20	25.0	ug/L
541-73-1	1,3-Dichlorobenzene	3.40	U	3.40	25.0	ug/L
106-46-7	1,4-Dichlorobenzene	4.10	U	4.10	25.0	ug/L
95-50-1	1,2-Dichlorobenzene	3.40	U	3.40	25.0	ug/L
SURROGATES						
17060-07-0	1,2-Dichloroethane-d4	29.1		91 - 110	97%	SPK: 30
2037-26-5	Toluene-d8	29.5		91 - 112	98%	SPK: 30
460-00-4	4-Bromofluorobenzene	27.2		63 - 112	91%	SPK: 30
INTERNAL STA	ANDARDS					
74-97-5	Bromochloromethane	29000	7.812			
540-36-3	1,4-Difluorobenzene	167000	9.1			
3114-55-4	Chlorobenzene-d5	152000	11.865			

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

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LAB CHRONICLE

OrderID: Q2006

Client: Ardmore Chemical
Contact: Michael Sharphouse

OrderDate: 5/9/2025 2:30:00 PM

Project: PVSC Monthly 2025 Location: L41,VOA Ref. #3 Water

ClientID Sample Date **Prep Date** Received LabID Matrix Test Method **Anal Date** Q2006-01 05/09/25 05/09/25 **EFF-WW** Water 05/16/25 VOC-PP 624.1

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Hit Summary Sheet SW-846

SDG No.: Q2006

Client: Ardmore Chemical

Sample ID Client ID Matrix Parameter Concentration C MDL RDL Units

Client ID:

0.000

Total Svoc: 0.00
Total Concentration: 0.00

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SAMPLE DATA

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Test:

Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 Project: PVSC Monthly 2025 Date Received: 05/09/25

Client Sample ID: **EFF-WW** SDG No.: Q2006 Lab Sample ID: Q2006-02 Matrix: Water % Solid: 0 Analytical Method: 625.1

Sample Wt/Vol: 960 Units: mL Final Vol: 1000 uL SVOCMS Group1

Extraction Type: Decanted: N Level: LOW

uL

GPC Cleanup: PH: Injection Volume: GPC Factor: Ν

3510C Prep Method:

Soil Aliquot Vol:

File ID/Qc Batch: Dilution: Prep Date Date Analyzed Prep Batch ID BP024656.D 1 05/15/25 08:55 PB168019 05/16/25 14:14

B1 02 100 0.B	•	337.237.23		00/10/20 11.11	12100017	
CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
TARGETS						
62-75-9	n-Nitrosodimethylamine	0.90	U	0.90	10.4	ug/L
108-95-2	Phenol	0.95	U	0.95	5.20	ug/L
111-44-4	bis(2-Chloroethyl)ether	0.84	U	0.84	5.20	ug/L
95-57-8	2-Chlorophenol	0.60	U	0.60	5.20	ug/L
108-60-1	2,2-oxybis(1-Chloropropane)	1.30	U	1.30	5.20	ug/L
621-64-7	n-Nitroso-di-n-propylamine	1.50	U	1.50	5.20	ug/L
67-72-1	Hexachloroethane	0.68	U	0.68	5.20	ug/L
98-95-3	Nitrobenzene	0.79	U	0.79	5.20	ug/L
78-59-1	Isophorone	0.78	U	0.78	5.20	ug/L
88-75-5	2-Nitrophenol	1.80	U	1.80	5.20	ug/L
105-67-9	2,4-Dimethylphenol	1.90	U	1.90	5.20	ug/L
111-91-1	bis(2-Chloroethoxy)methane	0.71	U	0.71	5.20	ug/L
120-83-2	2,4-Dichlorophenol	0.54	U	0.54	5.20	ug/L
120-82-1	1,2,4-Trichlorobenzene	0.56	U	0.56	5.20	ug/L
91-20-3	Naphthalene	0.52	U	0.52	5.20	ug/L
87-68-3	Hexachlorobutadiene	0.56	U	0.56	5.20	ug/L
59-50-7	4-Chloro-3-methylphenol	0.61	U	0.61	5.20	ug/L
77-47-4	Hexachlorocyclopentadiene	3.80	U	3.80	10.4	ug/L
88-06-2	2,4,6-Trichlorophenol	0.53	U	0.53	5.20	ug/L
91-58-7	2-Chloronaphthalene	0.64	U	0.64	5.20	ug/L
131-11-3	Dimethylphthalate	0.64	U	0.64	5.20	ug/L
208-96-8	Acenaphthylene	0.78	U	0.78	5.20	ug/L
606-20-2	2,6-Dinitrotoluene	0.96	U	0.96	5.20	ug/L
83-32-9	Acenaphthene	0.57	U	0.57	5.20	ug/L
51-28-5	2,4-Dinitrophenol	6.20	U	6.20	10.4	ug/L
100-02-7	4-Nitrophenol	2.50	U	2.50	10.4	ug/L
121-14-2	2,4-Dinitrotoluene	1.30	U	1.30	5.20	ug/L
84-66-2	Diethylphthalate	0.72	U	0.72	5.20	ug/L
7005-72-3	4-Chlorophenyl-phenylether	0.71	U	0.71	5.20	ug/L

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Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 Project: PVSC Monthly 2025 Date Received: 05/09/25

Client Sample ID: **EFF-WW** SDG No.: Q2006 Lab Sample ID: Q2006-02 Matrix: Water % Solid: 0 Analytical Method: 625.1

Sample Wt/Vol: 960 Units: mL Final Vol: 1000 uL

SVOCMS Group1 Soil Aliquot Vol: uL Test:

Extraction Type: Decanted: N Level: LOW

GPC Cleanup: PH: Injection Volume: GPC Factor: Ν

3510C Prep Method:

File ID/Qc Batch: Dilution: Prep Date Date Analyzed Prep Batch ID BP024656.D 1 05/15/25 08:55 PB168019 05/16/25 14:14

CAS Number	Parameter	Conc.	Qualifier	MDL	LOQ / CRQL	Units
86-73-7	Fluorene	0.66	U	0.66	5.20	ug/L
534-52-1	4,6-Dinitro-2-methylphenol	3.00	U	3.00	10.4	ug/L
86-30-6	n-Nitrosodiphenylamine	0.60	U	0.60	5.20	ug/L
103-33-3	Azobenzene	0.84	U	0.84	5.20	ug/L
101-55-3	4-Bromophenyl-phenylether	0.42	U	0.42	5.20	ug/L
118-74-1	Hexachlorobenzene	0.54	U	0.54	5.20	ug/L
87-86-5	Pentachlorophenol	1.60	U	1.60	10.4	ug/L
85-01-8	Phenanthrene	0.52	U	0.52	5.20	ug/L
120-12-7	Anthracene	0.64	U	0.64	5.20	ug/L
84-74-2	Di-n-butylphthalate	1.30	U	1.30	5.20	ug/L
206-44-0	Fluoranthene	0.85	U	0.85	5.20	ug/L
92-87-5	Benzidine	4.50	U	4.50	10.4	ug/L
129-00-0	Pyrene	0.52	U	0.52	5.20	ug/L
85-68-7	Butylbenzylphthalate	2.00	U	2.00	5.20	ug/L
91-94-1	3,3-Dichlorobenzidine	0.97	U	0.97	10.4	ug/L
56-55-3	Benzo(a)anthracene	0.47	U	0.47	5.20	ug/L
218-01-9	Chrysene	0.46	U	0.46	5.20	ug/L
117-81-7	Bis(2-ethylhexyl)phthalate	1.70	U	1.70	5.20	ug/L
117-84-0	Di-n-octyl phthalate	2.40	U	2.40	10.4	ug/L
205-99-2	Benzo(b)fluoranthene	0.51	U	0.51	5.20	ug/L
207-08-9	Benzo(k)fluoranthene	0.50	U	0.50	5.20	ug/L
50-32-8	Benzo(a)pyrene	0.57	U	0.57	5.20	ug/L
193-39-5	Indeno(1,2,3-cd)pyrene	0.61	U	0.61	5.20	ug/L
53-70-3	Dibenzo(a,h)anthracene	0.70	U	0.70	5.20	ug/L
191-24-2	Benzo(g,h,i)perylene	0.72	U	0.72	5.20	ug/L
SURROGATES						
367-12-4	2-Fluorophenol	45.3	*	60 - 140	45%	SPK: 100
13127-88-3	Phenol-d6	28.1	*	60 - 140	28%	SPK: 100
4165-60-0	Nitrobenzene-d5	75.1		60 - 140	75%	SPK: 100
321-60-8	2-Fluorobiphenyl	73.0		60 - 140	73%	SPK: 100

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Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 Project: PVSC Monthly 2025 Date Received: 05/09/25 Client Sample ID: **EFF-WW** SDG No.: Q2006 Lab Sample ID: Q2006-02 Matrix: Water Analytical Method: 625.1 % Solid: 0 Sample Wt/Vol: 960 Units: Final Vol: 1000 uL mLSVOCMS Group1 Soil Aliquot Vol: иL Test: Level: Extraction Type: Decanted: Ν LOW

Prep Method: 3510C

Injection Volume:

 File ID/Qc Batch:
 Dilution:
 Prep Date
 Date Analyzed
 Prep Batch ID

 BP024656.D
 1
 05/15/25 08:55
 05/16/25 14:14
 PB168019

GPC Factor: 1.0

CAS Number	Parameter	Conc.	Qualifier MDL	LOQ / CRQL	Units
118-79-6	2,4,6-Tribromophenol	70.5	60 - 140	71%	SPK: 100
1718-51-0	Terphenyl-d14	75.2	60 - 140	75%	SPK: 100
INTERNAL STA	ANDARDS				
3855-82-1	1,4-Dichlorobenzene-d4	157000	7.658		
1146-65-2	Naphthalene-d8	630000	10.434		
15067-26-2	Acenaphthene-d10	402000	14.293		
1517-22-2	Phenanthrene-d10	750000	17.11		
1719-03-5	Chrysene-d12	879000	21.533		
1520-96-3	Pervlene-d12	1060000	24.851		

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

GPC Cleanup:

Ν

PH:

N = Presumptive Evidence of a Compound

* = Values outside of QC limits

D = Dilution

() = Laboratory InHouse Limit

A = Aldol-Condensation Reaction Products

Q2006



LAB CHRONICLE

OrderID: Q2006 OrderDate: 5/9/2025 2:30:00 PM

Client:Ardmore ChemicalProject:PVSC Monthly 2025Contact:Michael SharphouseLocation:L41,VOA Ref. #3 Water

Sample Date **Prep Date** Received LabID ClientID Matrix Test Method **Anal Date** Q2006-02 05/09/25 05/09/25 **EFF-WW** Water SVOCMS Group1 625.1 05/15/25 05/16/25

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Q2006

SDG No.:

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

Hit Summary Sheet SW-846

Order ID: Q2006

Client: Ardmore Chemical Project ID: PVSC Monthly 2025

Chent.	Ardinore Chemical		1 TOJECT ID	1 v Se Wollding 2023				
Sample ID	Client ID	Matrix	Parameter	Concentration	C	MDL	RDL	Units
Client ID:	EFF-WW							
Q2006-02	EFF-WW	Water	Lead	1.77	J	1.21	6.00	ug/L
Q2006-02	EFF-WW	Water	Zinc	231		2.00	20.0	ug/L

Q2006 **25 of 36**











SAMPLE DATA

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

Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 Project: PVSC Monthly 2025 Date Received: 05/09/25 Client Sample ID: EFF-WW SDG No.: Q2006 Lab Sample ID: Q2006-02 Matrix: Water % Solid: Level (low/med): 0 low

Cas	Parameter	Conc.	Qua.	DF	MDL	LOQ / CRQL	Units	Prep Date	Date Ana.	Ana Met.	Prep Met.
7439-92-1	Lead	1.77	J	1	1.21	6.00	ug/L	05/13/25 15:05	05/15/25 18:05	EPA 200.7	_
7439-97-6	Mercury	0.027	U	1	0.027	0.20	ug/L	05/22/25 09:00	05/22/25 14:49	E245.1	
7440-66-6	Zinc	231	N*	1	2.00	20.0	ug/L	05/13/25 15:05	05/15/25 18:05	EPA 200.7	

Color Before:

Colorless

Mercury

Clarity Before: Clarity After: Cloudy

Clear

Texture:
Artifacts:

Color After: Colorless

Comments:

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

D = Dilution

Q = indicates LCS control criteria did not meet requirements

J = Estimated Value

B = Analyte Found in Associated Method Blank

* = indicates the duplicate analysis is not within control limits.

E = Indicates the reported value is estimated because of the presence

of interference.

OR = Over Range

N =Spiked sample recovery not within control limits

Q2006



LAB CHRONICLE

OrderID: Q2006 **OrderDate:** 5/9/2025 2:30:00 PM

Client:Ardmore ChemicalProject:PVSC Monthly 2025Contact:Michael SharphouseLocation:L41,VOA Ref. #3 Water

Sample Date **Prep Date** Received LabID ClientID Matrix Test Method **Anal Date** Q2006-02 05/09/25 05/09/25 **EFF-WW** Water Mercury 245.1 05/22/25 05/22/25 Metals Group3 200.7 05/13/25 05/15/25

Q2006 **28 of 36**



8



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SAMPLE DATA

Q2006 **29 of 36**



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

Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 09:00

Project: PVSC Monthly 2025 Date Received: 05/09/25

Client Sample ID: EFF-WW SDG No.: Q2006

Lab Sample ID: Q2006-01 Matrix: WATER

% Solid: 0

Parameter	Conc. Qua.	DF MDL	LOQ / CRQL	Units	Prep Date	Date Ana.	Ana Met.
Cyanide	0.0012 U	1 0.0012	0.0050	mg/L	05/14/25 08:15	05/14/25 12:31	SM 4500-CN
							C-16 plus E-16

Comments:

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

D = Dilution

Q = indicates LCS control criteria did not meet requirements

H = Sample Analysis Out Of Hold Time

J = Estimated Value

B = Analyte Found in Associated Method Blank

* = indicates the duplicate analysis is not within control limits.

E = Indicates the reported value is estimated because of the presence of interference.

OR = Over Range

N =Spiked sample recovery not within control limits

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

Report of Analysis

Client: Ardmore Chemical Date Collected: 05/09/25 09:00

Project: PVSC Monthly 2025 Date Received: 05/09/25

Client Sample ID: EFF-WW SDG No.: Q2006

Lab Sample ID: Q2006-02 Matrix: WATER

% Solid: 0

Parameter	Conc. Qua.	DF	MDL	LOQ / CRQL	Units	Prep Date	Date Ana.	Ana Met.
BOD5	1250	1	0.20	2.00	mg/L		05/09/25 16:40	SM 5210 B-16
TSS	98.6	1	1.00	4.00	mg/L		05/13/25 10:00	SM 2540 D-15

Comments:

U = Not Detected

LOQ = Limit of Quantitation

MDL = Method Detection Limit

LOD = Limit of Detection

D = Dilution

Q = indicates LCS control criteria did not meet requirements

H = Sample Analysis Out Of Hold Time

J = Estimated Value

B = Analyte Found in Associated Method Blank

* = indicates the duplicate analysis is not within control limits.

E = Indicates the reported value is estimated because of the presence of interference.

OR = Over Range

N =Spiked sample recovery not within control limits

Q2006 **31**



LAB CHRONICLE

OrderID: Q2006

Client: Ardmore Chemical
Contact: Michael Sharphouse

OrderDate: 5/9/2025 2:30:00 PM

Project: PVSC Monthly 2025 Location: L41,VOA Ref. #3 Water

LabID	ClientID	Matrix	Test	Method S	ample Date	Prep Date	Anal Date	Received
Q2006-01	EFF-WW	WATER			05/09/25			05/09/25
•					09:00			
			Cyanide	SM4500-CN		05/14/25	05/14/25	
				C,E			12:31	
Q2006-02	EFF-WW	WATER			05/09/25			05/09/25
					09:00			
			BOD5	SM5210 B			05/09/25	
							16:40	
			TSS	SM2540 D			05/13/25	
							10:00	

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SHIPPING DOCUMENTS

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

CHEMT	ECH	PROJ	ECT	NO

QUOTE NO.

COC Number 2041889

CLIENT INFORMATION	ALC: NO PERSONS ASSESSMENT	CLIENT PROJECT INFORMATION		CLIENT BILLING INFORMATION		
REPORT TO BE SENT TO: COMPANY: FIRD MORE INC	PROJECT NAM	1E:	BILL TO:	PO#:		
ADDRESS: 29 RIVERSIDE AVE BIg#14	PROJECT NO.:	LOCATION:	ADDRESS:			
CITY Newark STATE: NJ ZIP: 07405	PROJECT MANA	GER:	CITY	STATE: ZIP:		
ATTENTION: MICHAEL SHARPHOOSE	e-mail:		ATTENTION:	PHONE:		
PHONE: 973 481 2406 FAX:	PHONE:	FAX:	1000	ANALYSIS		
DATA TURNAROUND INFORMATION FAX (RUSH)DAYS*		A DELIVERABLE INFORMATION S Only) Level 4 (QC + Full Raw Data)	///	10/0//////		
HARDCOPY (DATA PACKAGE): STANDARD DAYS* EDD: DAYS* *TO BE APPROVED BY CHEMTECH	☐ Level 2 (Results☐ Level 3 (Results		100 3 5 VOR 3 V	CTP 8 9		
STANDARD HARDCOPY TURNAROUND TIME IS 10 BUSINESS DAYS	□ EDD FORMAT_	4	PRESERVATIVE			
CHEMTECH SAMPLE SAMPLE IDENTIFICATION	SAMPLE TYPE MATRIX W W W W W W W W W W W W W W W W W W W	E SAMPLE COLLECTION LD DATE TIME DATE TIME DATE TIME DATE TIME DATE DATE DATE DATE DATE DATE DATE DAT		← Specify Preservatives A-HCI D-NaOH B-HN03 E-ICE		
1. EFF WW	ww X	5/9/2 9:00 X X		PH 10		
2. EFF WW	wwx	5/5/25 9:00	XXX	рн 1.3		
3.				,		
4.						
6.						
7.						
8.						
9.						
10.	I I I I I I I I I I I I I I I I I I I			^		
RELINQUISHED BY SAMPLER: DATE/TIME: 1416 RECEIVED BY: 1. Bleet Shaphoat 5/9/35 1. RELINQUISHED BY SAMPLER: DATE/TIME: RECEIVED BY: 2. 2.)	Conditions of bottles or coolers at receipt: Comments: METALS	□ COMPLIANT □ NON COMPLIANT	□ COOLER TEMP 4.3 °C		
RELINQUISHED BY SAMPLER: DATE/TIME: RECEIVED BY: 3. 3. WHITE - CHEMIT	CH COPY FOR RETURN 1	Page of CLIENT: CHEMTECH TO CLIENT 34, Q. [3,6] CHEMTECH COPY	☐ Hand Delivered ☐ Other ☐ Picked Up ☐ Field Sa	Shipment Complete ampling		

PINK - SAMPLER COPY



Laboratory Certification

Certified By	License No.
CAS EPA CLP Contract	68HERH20D0011
O constituent	DIL 2022
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

QA Control Code: A2070148



LOGIN REPORT/SAMPLE TRANSFER

Order ID: Q2006

ARDM01

Order Date: 5/9/2025 2:30:00 PM

Project Mgr:

Client Name: Ardmore Chemical

Project Name: PVSC Monthly 2025

Report Type: Level 1

Client Contact: Michael Sharphouse

Receive DateTime: 5/9/2025 2:16:00 PM

EDD Type: NONE

14:40 Re\$5

Invoice Name: Ardmore Chemical

Purchase Order:

Hard Copy Date:

Invoice Contact: Michael Sharphouse

Date Signoff:

LAB ID	CLIENT ID	MATRIX SAMPLE SAMPLE DATE TIME	TEST	TEST GROUP	METHOD	FAX DATE	DUE DATES
Q2006-01	EFF-WW	Water 05/09/2025 09:00					
			VOC-PP		624.1	10 Bus. Days	

Relinguished By:

Date / Time:

Storage Area: VOA Refridgerator Room

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