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# **McKesson Bear Street**

# Data Usability Summary Report (DUSR)

SYRACUSE, NEW YORK

Volatile, Semivolatile and Methanol Analyses

SDG #s: 460-74504, 460-74604, and 460-74662

Analyses Performed By: TestAmerica Laboratories Edison, New Jersey

Report #: 21690R Review Level: Tier III Project: B0026003.2014.00010

#### SUMMARY

This data quality assessment summarizes the review of Sample Delivery Groups (SDGs) #s 460-74504, 460-74604, and 460-74662 for samples collected in association with the McKesson Bear Street site in Syracuse, New York. The review was conducted as a Tier III evaluation and included review of data package completeness. Only analytical data associated with constituents of concern were reviewed for this validation. Field documentation was not included in this review. Included with this assessment are the validation annotated sample result sheets, and chain of custody. Analyses were performed on the following samples:

		Lab ID		Sample	Parent	Analysis				
SDG	Sample ID		Matrix	Collection Date	Sample	voc	svoc	РСВ	METH	MISC
	TB041614	460-74504-1	Water	04/16/2014		Х				
	MW-4S	460-74504-2	Water	04/16/2014		Х	Х			
	PZ-4D	460-74504-3	Water	04/16/2014		Х	Х			
460-74504	MW-35	460-74504-4	Water	04/16/2014		Х			Х	
	MW-27	460-74504-5	Water	04/16/2014		Х	Х			
	MW-28	460-74504-6	Water	04/16/2014		Х	Х		Х	
	MW-8SR	460-74504-7	Water	04/16/2014		Х	Х			
	BD041614-01	460-74504-8	Water	04/16/2014	MW-8SR	Х	Х			
	TB041714	460-74604-1	Water	04/17/2014		Х				
	MW-31	460-74604-2	Water	04/17/2014		Х	Х		Х	
	MW-32	460-74604-3	Water	04/17/2014		Х	Х		Х	
	MW-33	460-74604-4	Water	04/17/2014		Х	Х			
460-74604	MW-34	460-74604-5	Water	04/17/2014		Х	Х		Х	
	MW-36R	460-74604-6	Water	04/17/2014		Х	Х			
	TW-02RRR	460-74604-7	Water	04/17/2014		Х	Х		Х	
	TW-01	460-74604-8	Water	04/17/2014		Х	Х		Х	
	BD041714-02	460-74604-9	Water	04/17/2014	TW-02RRR	Х	Х		Х	
460-74662	TB041814	460-74662-1	Water	04/18/2014		Х				
400-74002	MW-9S	460-74662-2	Water	04/18/2014		Х	Х		Х	

Notes:

1. METH - Methanol.

 The matrix spike /matrix spike duplicate (MS/MSD) analysis was performed on sample location MW-35.

3. MS/MSD analysis was performed for VOCs on sample location MW-31.

# ANALYTICAL DATA PACKAGE DOCUMENTATION

The table below is the evaluation of the data package completeness.

		Reported		Performance Acceptable		Not
	Items Reviewed	No	Yes	No	Yes	Required
1.	Sample receipt condition		Х		Х	
2.	Requested analyses and sample results		Х		Х	
3.	Master tracking list		Х		Х	
4.	Methods of analysis		Х		Х	
5.	Reporting limits		Х		Х	
6.	Sample collection date		Х		Х	
7.	Laboratory sample received date		Х		Х	
8.	Sample preservation verification (as applicable)		Х		Х	
9.	Sample preparation/extraction/analysis dates		Х		Х	
10.	Fully executed Chain-of-Custody (COC) form		Х		Х	
11.	Narrative summary of QA or sample problems provided		х		Х	
12.	Data Package Completeness and Compliance		Х		Х	

QA - Quality Assurance

# **ORGANIC ANALYSIS INTRODUCTION**

Analyses were performed according to United States Environmental Protection Agency (USEPA) SW-846 Methods 8260C, 8270D and 8015D as referenced in NYSDEC-ASP. Data were reviewed in accordance with USEPA National Functional Guidelines of October 1999 and USEPA Region II SOPs associated with USEPA SW-846 Validating Volatile Organic Compounds by GC/MS SW-846 Method 8260B (SOP HW-24 Revision 2, October 2006) and Validating Semivolatile Organic Compounds by GC/MS SW-846 Method 8270C (SOP HW-22 Revision 3, October 2006).

The data review process is an evaluation of data on a technical basis rather than a determination of contract compliance. As such, the standards against which the data are being weighed may differ from those specified in the analytical method. It is assumed that the data package represents the best efforts of the laboratory and had already been subjected to adequate and sufficient quality review prior to submission.

During the review process, laboratory qualified and unqualified data are verified against the supporting documentation. Based on this evaluation, qualifier codes may be added, deleted, or modified by the data reviewer. Results are qualified with the following codes in accordance with USEPA National Functional Guidelines:

- · Concentration (C) Qualifiers
  - U The compound was analyzed for but not detected. The associated value is the compound quantitation limit.
  - B The compound has been found in the sample as well as its associated blank, its presence in the sample may be suspect.
- · Quantitation (Q) Qualifiers
  - E The compound was quantitated above the calibration range.
  - D Concentration is based on a diluted sample analysis.
- · Validation Qualifiers
  - J The compound was positively identified; however, the associated numerical value is an estimated concentration only.
  - UJ The compound was not detected above the reported sample quantitation limit. However, the reported limit is approximate and may or may not represent the actual limit of quantitation.
  - JN The analysis indicates the presence of a compound for which there is presumptive evidence to make a tentative identification. The associated numerical value is an estimated concentration only.
  - UB Compound considered non-detect at the listed value due to associated blank contamination.
  - N The analysis indicates the presence of a compound for which there is presumptive evidence to make a tentative identification.
  - R The sample results are rejected as unusable. The compound may or may not be present in the sample.

Two facts should be noted by all data users. First, the "R" flag means that the associated value is unusable. In other words, due to significant quality control (QC) problems, the analysis is invalid and provides no information as to whether the compound is present or not. "R" values should not appear on data tables because they cannot be relied upon, even as a last resort. The second fact to keep in mind is that no compound concentration, even if it has passed all QC tests, is guaranteed to be accurate. Strict QC serves to increase confidence in data but any value potentially contains error.

# VOLATILE ORGANIC COMPOUND (VOC) ANALYSES

#### 1. Holding Times

The specified holding times for the following methods are presented in the following table.

Method	Matrix	Holding Time	Preservation
SW-846 8260C	Water	14 days from collection to analysis	Cool to <6 °C; preserved to a pH of less than 2 s.u.
300-040 02000	Soil	48 hours from collection to extraction and 14 days from collection to analysis	Cool to <6°C

s.u. Standard units

All samples were analyzed within the specified holding time criteria.

#### 2. Blank Contamination

Quality assurance (QA) blanks (i.e. laboratory method blanks, trip blanks, and equipment rinse blanks) are prepared to identify any contamination which may have been introduced into the samples during sample preparation or field activity. Method blanks measure laboratory contamination. Trip blanks measure sample storage contamination. Rinse blanks also measure contamination of samples during field operations.

A blank action level (BAL) of five times the concentration of a detected compound in an associated blank (common laboratory contaminant compounds are calculated at ten times) is calculated for QA blanks containing concentrations greater than the method detection limit (MDL). The BAL is compared to the associated sample results to determine the appropriate qualification of the sample results, if needed.

Compounds were not detected above the MDL in the associated blanks; therefore detected sample results were not associated with blank contamination.

#### 3. Mass Spectrometer Tuning

Mass spectrometer performance was acceptable and all analyses were performed within a 12-hour tune clock.

System performance and column resolution were acceptable.

#### 4. Calibration

Satisfactory instrument calibration is established to insure that the instrument is capable of producing acceptable quantitative data. An initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of an experimental sequence. The continuing calibration verifies that the instrument daily performance is satisfactory.

#### 4.1 Initial Calibration

The method specifies percent relative standard deviation (%RSD) and relative response factor (RRF) limits for select compounds only. A technical review of the data applies limits to all compounds with no exceptions.

All target compounds associated with the initial calibration standards must exhibit a %RSD less than the control limit (15%) or a correlation coefficient greater than 0.99, and a RRF value greater than control limit (0.05).

### 4.2 Continuing Calibration

All target compounds associated with the continuing calibration standard must exhibit a percent difference (%D) less than the control limit (20%) and RRF value greater than control limit (0.05).

All compounds associated with the calibrations were within the specified control limits.

#### 5. Surrogates/System Monitoring Compounds

All samples to be analyzed for organic compounds are spiked with surrogate compounds prior to sample preparation to evaluate overall laboratory performance and efficiency of the analytical technique. VOC analysis requires that all surrogates associated with the analysis exhibit recoveries within the laboratory-established acceptance limits.

All surrogate recoveries were within the control limits.

#### 6. Internal Standard Performance

Internal standard performance criteria insure that the GC/MS sensitivity and response are stable during every sample analysis. The criteria requires the internal standard compounds associated with the VOC analysis exhibit area counts that are not greater than two times (+100%) or less than one-half (-50%) of the area counts of the associated continuing calibration standard.

All internal standard responses were within control limits.

#### 7. Matrix Spike/Matrix Spike Duplicate (MS/MSD) Analysis

MS/MSD data are used to assess the precision and accuracy of the analytical method. The spiked compounds used in the MS/MSD analysis must exhibit recoveries within the laboratory-established acceptance limits. The relative percent difference (RPD) between the MS and MSD results must be within the laboratory-established acceptance limits.

Note: The MS/MSD recovery control limits do not apply for MS/MSDs performed on sample locations where the compound concentration detected in the parent sample exceeds the MS/MSD spiking concentration by a factor of four or greater. Sample results associated with MS/MSD exceedances where the parent samples are not site-specific are not qualified.

The MS/MSD exhibited acceptable recoveries. Sample locations associated with MS/MSD recoveries exhibiting an RPD greater than of the control limit presented in the following table.

Sample Locations	Compound
MW-35	Acetone

The criteria used to evaluate the RPD between the MS/MSD recoveries are presented in the following table. In the case of an RPD deviation, the sample results are qualified as documented in the table below.

Control Limit	Sample Result	Qualification
> UL	Non-detect	UJ
> OL	Detect	J

#### 8. Laboratory Control Sample (LCS) Analysis

The LCS analysis is used to assess the accuracy of the analytical method independent of matrix interferences. The spiked compounds used in the LCS analysis must exhibit recoveries within the laboratory-established acceptance limits.

All compounds associated with the LCS analyses exhibited recoveries within the control limits.

#### 9. Field Duplicate Analysis

The field duplicate analysis is used to assess the precision of the field sampling procedures and analytical method. A control limit of 50% for water matrices is applied to the RPD between the parent sample and the field duplicate. In the instance when the parent and/or duplicate sample concentrations are less than or equal to five times the reporting limit (RL), a control limit for the difference between the results of two times the RL is applied for water matrices.

Results for duplicate samples are summarized in the following table.

Sample ID/Duplicate ID	Compound	Sample Result	Duplicate Result	RPD
MW-8SR/ BD041614-01	Benzene	3.2	3.3	
	Ethylbenzene	0.25 J	0.27 J	AC
	Toluene	1.2	1.1	AC
	Xylenes, Total	13	13	
	Aceton	8.0 J	10	
	Benzene	1.2	1.2	
TW-02RRR/ BD041714-02	Ethylbenzene	0.51 J	0.44 J	AC
	Toluene	0.18 J	0.17 J	
	Xylenes, Total	1.0 J	0.96 J	

AC Acceptable

The calculated RPDs between the parent sample and field duplicate were acceptable.

#### 10. Compound Identification

Compounds are identified on the GC/MS by using the analytes relative retention time and ion spectra.

All identified compounds met the specified criteria.

#### 11. System Performance and Overall Assessment

Overall system performance was acceptable. Other than for those deviations specifically mentioned in this review, the overall data quality is within the guidelines specified in the method.

# DATA VALIDATION CHECKLIST FOR VOCs

VOCs: SW-846 8260C	Rep	orted		mance ptable	Not	
	No	Yes	No	Yes	Required	
GAS CHROMATOGRAPHY/MASS SPECTROMETR	Y (GC/MS	)				
Tier II Validation						
Holding times		Х		Х		
Reporting limits (units)		Х		Х		
Blanks					·	
A. Method blanks		Х		Х		
B. Equipment/Field blanks					Х	
C. Trip blanks		Х		Х		
Laboratory Control Sample (LCS) Accuracy (%R)		Х		Х		
Laboratory Control Sample Duplicate (LCSD) %R					Х	
LCS/LCSD Precision (RPD)					Х	
Matrix Spike (MS) %R		Х		Х		
Matrix Spike Duplicate (MSD) %R		Х		Х		
MS/MSD Precision RPD		Х	Х			
Field Duplicate RPD		Х		Х		
Surrogate Spike %R		Х		Х		
Dilution Factor		Х		Х		
Moisture Content					Х	
Tier III Validation						
System performance and column resolution		Х		Х		
Initial calibration %RSDs		Х		Х		
Continuing calibration RRFs		Х		Х		
Continuing calibration %Ds		Х		Х		
Instrument tune and performance check		Х		Х		
Ion abundance criteria for each instrument used		Х		Х		
Internal standard		Х		Х		
Compound identification and quantitation						
A. Reconstructed ion chromatograms	1	Х		Х		
B. Quantitation Reports		Х		Х		
C. RT of sample compounds within the established RT windows		х		х		
D. Transcription/calculations acceptable		Х		Х		
E. Reporting limits adjusted for sample dilutions		Х		Х		

%RPercent recoveryRPDRelative percent difference%RSDRelative standard deviation

%D Percent difference

# SEMIVOLATILE ORGANIC COMPOUND (SVOC) ANALYSES

#### 1. Holding Times

The specified holding times for the following methods are presented in the following table.

Method	Matrix	Holding Time	Preservation
SW-846 8270D	Water	7 days from collection to extraction and 40 days from extraction to analysis	Cool to <6°C
300-040 0270D	Soil	14 days from collection to extraction and 40 days from extraction to analysis	C00110 < 8 C

All samples were extracted and analyzed within the specified holding time criteria.

### 2. Blank Contamination

Quality assurance (QA) blanks (i.e. laboratory method blanks and equipment rinse blanks) are prepared to identify any contamination which may have been introduced into the samples during sample preparation or field activity. Method blanks measure laboratory contamination. Rinse blanks measure contamination of samples during field operations.

A blank action level (BAL) of five times the concentration of a detected compound in an associated blank (common laboratory contaminant compounds are calculated at ten times) is calculated for QA blanks containing concentrations greater than the method detection limit (MDL). The BAL is compared to the associated sample results to determine the appropriate qualification of the sample results, if needed.

Target compounds were not detected above the MDL in the associated blanks; therefore detected sample results were not associated with blank contamination.

## 3. Mass Spectrometer Tuning

Mass spectrometer performance was acceptable and all analyses were performed within a 12-hour tune clock.

System performance and column resolution are acceptable.

#### 4. Calibration

Satisfactory instrument calibration is established to insure that the instrument is capable of producing acceptable quantitative data. An initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of an experimental sequence. The continuing calibration verifies that the instrument daily performance is satisfactory.

#### 4.1 Initial Calibration

The method specifies percent relative standard deviation (%RSD) and relative response factor (RRF) limits for select compounds only. A technical review of the data applies limits to all compounds with no exceptions.

All target compounds associated with the initial calibration standards must exhibit a %RSD less than the control limit (15%) or a correlation coefficient greater than 0.99 and an RRF value greater than control limit (0.05).

#### 4.2 Continuing Calibration

All target compounds associated with the continuing calibration standard must exhibit a percent difference (%D) less than the control limit (20%) and RRF value greater than control limit (0.05).

All compounds associated with the calibrations were within the specified control limits.

#### 5. Surrogates/System Monitoring Compounds

All samples to be analyzed for organic compounds are spiked with surrogate compounds prior to sample preparation to evaluate overall laboratory performance and efficiency of the analytical technique. SVOC analysis requires that two of the three SVOC surrogate compounds within each fraction exhibit recoveries within the laboratory-established acceptance limits, and that all SVOC surrogate recoveries be greater than ten percent.

Sample locations associated with surrogates exhibiting recoveries outside of the control limits presented in the following table.

Sample Locations	Surrogate	Recovery
	Nitrobenzene-d5	
TW-02RRR BD041714-02	2-Fluorobiphenyl	D
	Terphenyl-d14	
	Nitrobenzene-d5	40
MW-34	2-Fluorobiphenyl	AC AC
	Terphenyl-d14	> UL
D Diluted		

AC Acceptable

The criteria used to evaluate the surrogate recoveries are presented in the following table. In the case of a surrogate deviation, the sample results associated with the deviant fraction are qualified as documented in the table below.

Control Limit	Sample Result	Qualification
s upper control limit (III)	Non-detect	No Action
> upper control limit (UL)	Detect	J
< lower control limit (LL) but > 10%	Non-detect	UJ
< lower control limit (LL) but > 10%	Detect	J
< 10%	Non-detect	R
< 10%	Detect	J
One surragete outside control limite but > 10%	Non-detect	No Action
One surrogate outside control limits but >10%	Detect	NO ACTION

Control Limit	Sample Result	Qualification
Surrogates diluted below the calibration curve due to the	Non-detect	$UJ^1$
high concentration of a target compounds	Detect	$J^1$

A more concentrated analysis was not performed with surrogate compounds within the calibration range; therefore, no determination of extraction efficiency could be made.

#### 6. Internal Standard Performance

Internal standard performance criteria insure that the GC/MS sensitivity and response are stable during every sample analysis. The criteria requires the internal standard compounds associated with the SVOC analysis exhibit area counts that are not greater than two times (+100%) or less than one-half (-50%) of the area counts of the associated continuing calibration standard.

All internal standard responses were within the control limits.

#### 7. Matrix Spike/Matrix Spike Duplicate (MS/MSD) Analysis

MS/MSD data are used to assess the precision and accuracy of the analytical method. The compounds used to perform the MS/MSD analysis must exhibit recoveries within the laboratory-established acceptance limits. The relative percent difference (RPD) between the MS and MSD results must be within the laboratory-established or analytical method-referenced acceptance limits.

Note: The MS/MSD recovery control limits do not apply for MS/MSD performed on sample locations where the compound concentration detected in the parent sample exceeds the MS/MSD concentration by a factor of four or greater. Sample results associated with MS/MSD exceedances where the parent samples are not site-specific are not qualified.

The MS/MSD exhibited acceptable recoveries and RPD between the MS/MSD recoveries.

#### 8. Laboratory Control Sample/Laboratory Control Sample Duplicate (LCS/LCSD) Analysis

The LCS/LCSD analysis is used to assess the precision and accuracy of the analytical method independent of matrix interferences. The compounds associated with the LCS/LCSD analysis must exhibit recoveries and relative percent differences (RPDs) between the LCS and LCSD results within the laboratory-established acceptance limits.

All compounds associated with the LCS/LCSD analyses exhibited recoveries and RPDs within the control limits.

#### 9. Field Duplicate Analysis

The field duplicate analysis is used to assess the precision of the field sampling procedures and analytical method. A control limit of 50% for water matrices is applied to the RPD between the parent sample and the field duplicate. In the instance when the parent and/or duplicate sample concentrations are less than or equal to five times the reporting limit (RL), a control limit for the difference between the results of two times the RL is applied for water matrices.

Results for duplicate samples are summarized in the following table.

Sample ID/Duplicate ID	Compound	Sample Result	Duplicate Result	RPD
MW-8SR/ BD041614-01	Aniline	3.9 J	5.9 J	AC
	n,n'-Dimethylaniline	1.4	1.9	AC
TW-02RRR/	Aniline	1300	1700	26.7%
BD041714-02	n,n'-Dimethylaniline	2.8 J	3.5 J	AC

AC Acceptable

The calculated RPDs between the parent sample and field duplicate were acceptable.

#### 10. Compound Identification

Compounds are identified on the GC/MS by using the analytes relative retention time and ion spectra.

All identified compounds met the specified criteria.

#### 11. System Performance and Overall Assessment

Overall system performance was acceptable. Other than for those deviations specifically mentioned in this review, the overall data quality is within the guidelines specified in the method.

DATA VALIDATION CHECKLIST FOR SVOCs
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SVOCs: SW-846 8270D	Rep	orted	Perfor Acce	Not	
	No	Yes	No	Yes	Required
GAS CHROMATOGRAPHY/MASS SPECTROMETRY (	GC/MS)				
Tier II Validation					
Holding Times		Х		Х	
Reporting Limits (units)		Х		Х	
Blanks					
A. Method Blanks		Х		Х	
B. Equipment/Field Blanks					Х
Laboratory Control Sample (LCS) Accuracy (%R)		Х		Х	
Laboratory Control Sample Duplicate (LCSD) %R		Х		Х	
LCS/LCSD Precision (RPD)		Х		Х	
Matrix Spike (MS) %R		Х		Х	
Matrix Spike Duplicate (MSD) %R		Х		Х	
MS/MSD RPD		Х		Х	
Field Duplicate RPD		Х		Х	
Surrogate Spike %R		Х	Х		
Dilution Factor		Х		Х	
Moisture Content					Х
Tier III Validation		•	•	•	•
System Performance and Column Resolution		Х		Х	
Initial Calibration %RSDs		Х		Х	
Continuing Calibration RRFs		Х		Х	
Continuing Calibration %Ds		Х		Х	
Instrument Tune and Performance Check		Х		Х	
Ion Abundance Criteria for Each Instrument Used		Х		Х	
Internal Standards		Х		Х	
Compound Identification and Quantitation					
A. Reconstructed Ion Chromatograms		Х		Х	
B. Quantitation Reports		Х		Х	
C. RT of Sample Compounds Within the Established RT Windows		х		х	
D. Transcription/calculations acceptable		Х		Х	
<ul><li>E. Reporting Limits Adjusted for Sample Dilutions</li><li>%R Percent Recovery</li></ul>		Х		Х	

%R

Percent Recovery Relative Percent Difference RPD

%RSDRelative Standard Deviation%DPercent Difference

# METHANOL ANALYSIS

#### 1. Holding Times

The specified holding times for the following methods are presented in the following table.

Method	Matrix	Holding Time	Preservation
Methanol	Soil	14 days from collection to analysis	Cool to <6°C
SW-846 8015D	Water	14 days from collection to analysis	C001 10 < 6 C

All samples were analyzed within the specified holding time criteria.

### 2. Blank Contamination

Quality assurance (QA) blanks (i.e. laboratory method blanks and equipment rinse blanks) are prepared to identify any contamination which may have been introduced into the samples during sample preparation or field activity. Method blanks measure laboratory contamination. Rinse blanks measure contamination of samples during field operations.

A blank action level (BAL) of five times the concentration of a detected analyte in an associated blank is calculated for QA blanks containing concentrations greater than the reporting limit (RL). The BAL is compared to the associated sample results to determine the appropriate qualification of the sample results, if needed.

Methanol was not detected above the MDL in the associated blanks; therefore detected sample results were not associated with blank contamination.

#### 3. System Performance

System performance and column resolution were acceptable.

#### 4. Calibration

Satisfactory instrument calibration is established to insure that the instrument is capable of producing acceptable quantitative data. An initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of an experimental sequence. The continuing calibration verifies that the instrument daily performance is satisfactory.

#### 4.1 Initial Calibration

A maximum RSD of 20% or a correlation coefficient of greater than 0.99 is allowed.

#### 4.2 Continuing Calibration

All target compounds associated with the continuing calibration standard must exhibit a percent difference (%D) less than the control limit (15%).

All calibration criteria were within the control limits.

#### 5. Surrogates/System Monitoring Compounds

All samples to be analyzed for organic compounds are spiked with surrogate compounds prior to sample preparation to evaluate overall laboratory performance and efficiency of the analytical technique. The analysis requires surrogate compounds exhibit recoveries within the laboratory-established acceptance limits.

All surrogate recoveries were within the control limits.

#### 6. Matrix Spike/Matrix Spike Duplicate (MS/MSD) Analysis

MS/MSD data are used to assess the precision and accuracy of the analytical method. The spiked analytes used in the MS/MSD analysis must exhibit recoveries within the laboratory-established acceptance limits. The relative percent difference (RPD) between the MS and MSD results must be within the laboratory-established acceptance limits.

Note: The MS/MSD recovery control limits do not apply for MS/MSDs performed on sample locations where the analyte concentration detected in the parent sample exceeds the MS/MSD concentration by a factor of four or greater. Sample results associated with MS/MSD exceedances where the parent samples are not site-specific are not qualified.

The MS/MSD exhibited acceptable recoveries and RPD between the MS/MSD recoveries.

#### 7. Laboratory Control Sample (LCS) Analysis

The LCS analysis is used to assess the accuracy of the analytical method independent of matrix interferences. The spiked compounds used in the LCS analysis must exhibit recoveries within the laboratory-established acceptance limits.

All compounds associated with the LCS analysis exhibited recoveries within the control limits.

#### 8. Field Duplicate Analysis

The field duplicate analysis is used to assess the precision of the field sampling procedures and analytical method. A control limit of 50% for water matrices is applied to the RPD between the parent sample and the field duplicate. In the instance when the parent and/or duplicate sample concentrations are less than or equal to five times the reporting limit (RL), a control limit for the difference between the results of two times the RL is applied for water matrices.

Results for duplicate samples are summarized in the following table.

Sample ID/Duplicate ID	Compound	Sample Result	Duplicate Result	RPD
TW-02RRR/ BD041714-02	Methanol	500 U	500 U	AC

AC Acceptable

The calculated RPDs between the parent sample and field duplicate were acceptable.

#### 9. Compound Identification

The retention times of all quantitated peaks must fall within the calculated retention time windows.

All identified compounds met the specified criteria.

#### 10. System Performance and Overall Assessment

Overall system performance was acceptable. Other than for those deviations specifically mentioned in this review, the overall data quality is within the guidelines specified in the method.

# DATA VALIDATION CHECKLIST FOR METHANOL

Methanol: SW-846 8015D	Rep	orted	Performance Acceptable		Not Required	
	No	Yes	No	Yes	Required	
GAS CHROMATOGRAPHY (GC/FID)						
Tier II Validation						
Holding Times		Х		Х		
Reporting Limits (Units)		Х		Х		
Blanks						
A. Method Blanks		Х		Х		
B. Equipment Blanks					Х	
C. Trip Blanks					Х	
Laboratory Control Sample (LCS) Accuracy (%R)		Х		Х		
Laboratory Control Sample Duplicate (LCSD) %R					Х	
LCS/LCSD Precision (RPD)					Х	
Matrix Spike (MS) %R		Х		Х		
Matrix Spike Duplicate (MSD) %R		Х		Х		
MS/MSD RPD		Х		Х		
Field Duplicate RPD		Х		Х		
Surrogate Spike %R		Х		Х		
Dilution Factor		Х		Х		
Moisture Content					Х	
Tier III Validation						
Initial Calibration %RSDs		Х		Х		
Continuing Calibration %Ds		Х		Х		
System Performance and Column Resolution		Х		Х		
Compound Identification and Quantitation						
A. Quantitation Reports		Х		Х		
B. RT of Sample Compounds Within Established RT Windows		х		х		
C. Pattern Identification					Х	
D. Transcription/calculations acceptable		Х		Х		
E. Reporting Limits adjusted for Sample Dilutions		Х		Х		

%RPercent RecoveryRPDRelative Percent Difference%RSDRelative Standard Deviation%DPercent Difference

# SAMPLE COMPLIANCE REPORT

Sample Delivery					Compliancy <sup>1</sup>			cy <sup>1</sup>		
Group (SDG)	Sampling Date	Protocol	Sample ID	Matrix	voc	svoc	РСВ	METH	MISC	Noncompliance
	04/16/2014		TB041614	Water	yes					
	04/16/2014		MW-4S	Water	yes	yes				
	04/16/2014		PZ-4D	Water	yes	yes				
460 74504	04/16/2014		MW-35	Water	No	yes		yes		MS/MSD RPD
460-74504	04/16/2014		MW-27	Water	yes	yes				
	04/16/2014 04/16/2014	MW-28	Water	yes	yes		yes			
			MW-8SR	Water	yes	yes				
0	04/16/2014		BD041614-01	Water	yes	yes				
	04/17/2014	TB041714	Water	yes						
	04/17/2014	SW846	MW-31	Water	yes	yes		yes		
	04/17/2014		MW-32	Water	yes	yes		yes		
	04/17/2014		MW-33	Water	yes	yes				
460-74604	04/17/2014		MW-34	Water	yes	no		yes		Surrogate %R
	04/17/2014		MW-36R	Water	yes	yes				-
	04/17/2014		TW-02RRR	Water	yes	no		yes		Surrogate %R
	04/17/2014		TW-01	Water	yes	yes		yes		-
	04/17/2014		BD041714-02	Water	yes	no		yes		Surrogate %R
400 74000	04/18/2014		TB041814	Water	yes					-
460-74662	04/18/2014		MW-9S	Water	yes	yes		yes		

1 Samples which are compliant with no added validation qualifiers are listed as "yes". Samples which are non-compliant or which have added qualifiers are listed as "no". A "no" designation does not necessarily indicate that the data have been rejected or are otherwise unusable.

Validation Performed By:	Lyndi Mott
Signature:	Sond us has
Date:	May 12, 2014
Peer Review:	Dennis Capria
Date:	May 16, 2014

# CHAIN OF CUSTODY/LABORAOTRY QUALIFIER DEFINITIONS/ CORRECTED SAMPLE ANALYSIS DATA SHEETS

	•	ab PM: Jhang, Grace		460-45501-28639.3
Phone: 382-4934		-Mail: )race.chang@testamericainc.com		Page: Page 3 of 5
				1034/1 #0.
Due Date Requested:				tion Cod
TAT Requested (days):	-			NaOH N - None Zn Acetate O - AsNaO2
Standar	¢.			Nitte Acid P - Naz045 NaHSO4 Q - Na2SO3 MeOH R - Na2S2SO3
Po #: Purchase Order Requeste	đ.	List		ic Acid
WO #		No) pound		
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36758		Cooler Temperature(s) <sup>o</sup> C and Other R	emarks: //{#5_0	.3/0.4°C
	Poison B X Unknown Duison B X Unknown Hill Le II4 Hill	$ \frac{24}{3} \frac{\text{foe}}{14} \frac{\text{Nicole (Nicole (Nontroe)}}{\text{Stau (dars)}} $	Image: Changer of the product of t	Image: Charge Grade     Career Trading Note:       Image: Charge Glessamericalinc.com     Image: Charge Glessamericalinc.com       Image: Charge Glessamericalinc.com     Image: Charg

**Chain of Custody Record** 

**TestAmerica** 

# DATA REPORTING QUALIFIERS

Lab Section	Qualifier	Description
GC/MS VOA		
	U	Indicates the analyte was analyzed for but not detected.
	J	Result is less than the RL but greater than or equal to the MDL and the concentration is an approximate value.
GC/MS Semi VOA		
	U	Indicates the analyte was analyzed for but not detected.
	J	Result is less than the RL but greater than or equal to the MDL and the concentration is an approximate value.
	Х	Surrogate is outside control limits
GC VOA		
	U	Indicates the analyte was analyzed for but not detected.

#### Client: ARCADIS U.S. Inc

# **Analytical Data**

Lab Sample ID: Client Matrix:	460-74504-1TB Water					Date Sampled: 04/16/2014 1600 Date Received: 04/17/2014 0920
		8260C Volatile Orgar	ic Compounds	s by GC/I	MS	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/18/2014 1725 04/18/2014 1725	Analysis Batch: Prep Batch:	460-219595 N/A		Instrument ID: Lab File ID: Initial Weight/Volur Final Weight/Volun	
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		10		U	2.7	10
Benzene		1.0		U	0.080	1.0
Ethylbenzene		1.0		U	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		1.0		U	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.0		U	0.13	3.0
Surrogate		%Rec		Qualifie	r Acc	eptance Limits
1,2-Dichloroethane-c	d4 (Surr)	103			70 -	- 130
Bromofluorobenzene		100			64 -	- 135
Dibromofluorometha	ne (Surr)	112			72 -	- 137
Toluene-d8 (Surr)		96			70 -	- 130

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-4S						
Lab Sample ID: Client Matrix:	460-74504-2 Water					Date Sampled: 04/16/ Date Received: 04/17/	
		8260C Volatile Orgar	ic Compounds	by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/18/2014 1813 04/18/2014 1813	Analysis Batch: Prep Batch:	460-219595 N/A		Instrument ID: Lab File ID: Initial Weight/Volu Final Weight/Volu		
Analyte		Result (u	g/L)	Qualifie	r MDL	RL	
Acetone		10	- /	U	2.7	10	
Benzene		1.0		U	0.080	1.0	
Ethylbenzene		1.0		U	0.10	1.0	
Methylene Chloride		1.0		U	0.18	1.0	
Toluene		1.0		U	0.15	1.0	
Trichloroethene		1.0		U	0.090	1.0	
Xylenes, Total		3.0		U	0.13	3.0	
Surrogate		%Rec		Qualifie	r Ac	ceptance Limits	
1,2-Dichloroethane-	d4 (Surr)	104			70	- 130	
Bromofluorobenzen	e	96			64	- 135	
Dibromofluorometha	ane (Surr)	111			72	- 137	
Toluene-d8 (Surr)		90			70	- 130	

Job Number: 460-74504-1

Client Sample ID:	PZ-4D					
Lab Sample ID:	460-74504-3				Dat	e Sampled: 04/16/2014 1140
Client Matrix:	Water				Dat	e Received: 04/17/2014 0920
		8260C Volatile Orgar	ic Compounds	by GC/I	MS	
Analysis Method:	8260C	Analysis Batch:	460-219595		Instrument ID:	CVOAMS9
Prep Method:	5030C	Prep Batch:	N/A		Lab File ID:	K26292.D
Dilution:	1.0				Initial Weight/Volume:	5 mL
Analysis Date:	04/18/2014 1837				Final Weight/Volume:	5 mL
Prep Date:	04/18/2014 1837					
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		10		U	2.7	10
Benzene		1.0		U	0.080	1.0
Ethylbenzene		1.0		U	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		1.0		U	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.0		U	0.13	3.0
Surrogate		%Rec		Qualifie	r Accept	ance Limits
1,2-Dichloroethane-	-d4 (Surr)	103			70 - 13	0
Bromofluorobenzen	e	96			64 - 13	5
Dibromofluorometha	ane (Surr)	110			72 - 13	7
Toluene-d8 (Surr)		87			70 - 13	0

Job Number: 460-74504-1

Client Sample ID:	MW-35					
Lab Sample ID: Client Matrix:	460-74504-4 Water					te Sampled: 04/16/2014 1330 te Received: 04/17/2014 0920
		8260C Volatile Orgar	nic Compounds	by GC/I	MS	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/18/2014 1902 04/18/2014 1902	Analysis Batch: Prep Batch:	460-219595 N/A		Instrument ID: Lab File ID: Initial Weight/Volume: Final Weight/Volume:	CVOAMS9 K26293.D 5 mL 5 mL
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		10		υJ	2.7	10
Benzene		1.0		U	0.080	1.0
Ethylbenzene		1.0		U	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		1.0		U	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.0		U	0.13	3.0
Surrogate		%Rec		Qualifie	r Accept	ance Limits
1,2-Dichloroethane-	-d4 (Surr)	105			70 - 13	0
Bromofluorobenzen	e	93			64 - 13	5
Dibromofluorometh	ane (Surr)	112			72 - 13	7
Toluene-d8 (Surr)		87			70 - 13	0

Job Number: 460-74504-1

Client Sample ID:	MW-27					
Lab Sample ID:	460-74504-5				D	ate Sampled: 04/16/2014 1440
Client Matrix:	Water				D	ate Received: 04/17/2014 0920
		8260C Volatile Orgar	nic Compounds	by GC/N	MS	
Analysis Method:	8260C	Analysis Batch:	460-219595		Instrument ID:	CVOAMS9
Prep Method:	5030C	Prep Batch:	N/A		Lab File ID:	K26294.D
Dilution:	1.0				Initial Weight/Volume	e: 5 mL
Analysis Date:	04/18/2014 1926				Final Weight/Volume	e: 5 mL
Prep Date:	04/18/2014 1926					
Analyte		Result (u	g/L)	Qualifier	r MDL	RL
Acetone		10		U	2.7	10
Benzene		1.0			0.080	1.0
Ethylbenzene		1.0		U	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		0.41		J	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		0.92		J	0.13	3.0
Surrogate		%Rec		Qualifier	r Acce	ptance Limits
1,2-Dichloroethane-	d4 (Surr)	104			70 - 1	30
Bromofluorobenzen	e	96			64 - 1	35
Dibromofluorometh	ane (Surr)	111			72 - 1	37
Toluene-d8 (Surr)		89			70 - 1	30

Job Number: 460-74504-1

Client Sample ID:	MW-28					
Lab Sample ID:	460-74504-6				Da	te Sampled: 04/16/2014 1545
Client Matrix:	Water				Da	te Received: 04/17/2014 0920
		8260C Volatile Orgar	nic Compounds	by GC/N	MS	
Analysis Method:	8260C	Analysis Batch:	460-219595		Instrument ID:	CVOAMS9
Prep Method:	5030C	Prep Batch:	N/A		Lab File ID:	K26295.D
Dilution:	1.0				Initial Weight/Volume	5 mL
Analysis Date:	04/18/2014 1950				Final Weight/Volume:	5 mL
Prep Date:	04/18/2014 1950					
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		13			2.7	10
Benzene		1.7			0.080	1.0
Ethylbenzene		1.0	I	U	0.10	1.0
Methylene Chloride	e	1.0	I	U	0.18	1.0
Toluene		0.29		J	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.0		U	0.13	3.0
Surrogate		%Rec	(	Qualifie	r Accep	ance Limits
1,2-Dichloroethane	e-d4 (Surr)	104			70 - 13	0
Bromofluorobenzei	ne	93			64 - 13	5
Dibromofluorometh	nane (Surr)	109			72 - 13	37
Toluene-d8 (Surr)		89			70 - 13	0

#### Client: ARCADIS U.S. Inc

# **Analytical Data**

Client Sample ID:	MW-8SR						
Lab Sample ID: Client Matrix:	460-74504-7 Water					•	04/16/2014 1600 04/17/2014 0920
		8260C Volatile Organ	ic Compounds	s by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/18/2014 2014 04/18/2014 2014	Analysis Batch: Prep Batch:	460-219595 N/A		Instrument ID: Lab File ID: Initial Weight/Volu Final Weight/Volu		
Analyte		Result (u	g/L)	Qualifie	r MDL	RL	
Acetone		10		U	2.7	10	
Benzene		3.2			0.080	1.0	
Ethylbenzene		0.25		J	0.10	1.0	
Methylene Chloride		1.0		U	0.18	1.0	
Toluene		1.2			0.15	1.0	
Trichloroethene		1.0		U	0.090	1.0	
Xylenes, Total		13			0.13	3.0	
Surrogate		%Rec		Qualifie	r Ac	ceptance Limits	
1,2-Dichloroethane-	d4 (Surr)	104			70	- 130	
Bromofluorobenzen	e	102			64	- 135	
Dibromofluorometha	ane (Surr)	113			72	- 137	
Toluene-d8 (Surr)		88			70	- 130	

#### Client: ARCADIS U.S. Inc

# **Analytical Data**

Client Sample ID:	BD041614-01						
Lab Sample ID: Client Matrix:	460-74504-8 Water						Campled: 04/16/2014 0930 Received: 04/17/2014 0920
		8260C Volatile Orgar	ic Compounds	by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/18/2014 2038 04/18/2014 2038	Analysis Batch: Prep Batch:	460-219595 N/A		Instrument ID: Lab File ID: Initial Weight/Volu Final Weight/Volu		CVOAMS9 K26297.D 5 mL 5 mL
Analyte		Result (u	g/L)	Qualifie	r MDL		RL
Acetone		10		U	2.7		10
Benzene		3.3			0.080		1.0
Ethylbenzene		0.27		J	0.10		1.0
Methylene Chloride		1.0		U	0.18		1.0
Toluene		1.1			0.15		1.0
Trichloroethene		1.0		U	0.090		1.0
Xylenes, Total		13			0.13		3.0
Surrogate		%Rec		Qualifie	r Ac	ceptanc	ce Limits
1,2-Dichloroethane-	d4 (Surr)	104			70	- 130	
Bromofluorobenzen	e	102			64	- 135	
Dibromofluorometha	ane (Surr)	114			72	- 137	
Toluene-d8 (Surr)		88			70	- 130	

#### Client: ARCADIS U.S. Inc

Client Sample ID:	MW-4S					
Lab Sample ID: Client Matrix:	460-74504-2 Water					e Sampled: 04/16/2014 1145 te Received: 04/17/2014 0920
		8270D Semivolatile Or	ganic Compou	nds (GC/M	IS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/03/2014 0351 04/17/2014 2035	Analysis Batch: Prep Batch:	460-222235 460-219569	L Ir F	nstrument ID: ab File ID: nitial Weight/Volume: inal Weight/Volume: njection Volume:	CBNAMS6 M79163.D 240 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		10		U	0.20	10
n,n'-Dimethylaniline		1.0		U	0.18	1.0
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Fluorobiphenyl		63			50 - 12	0
Nitrobenzene-d5 (S	urr)	66			60 - 11	4
Terphenyl-d14 (Sur	r)	77			72 - 13	0

#### Client: ARCADIS U.S. Inc

Client Sample ID:	PZ-4D					
Lab Sample ID: Client Matrix:	460-74504-3 Water					e Sampled: 04/16/2014 1140 e Received: 04/17/2014 0920
		8270D Semivolatile Or	ganic Compou	nds (GC/M	IS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/03/2014 0413 04/17/2014 2035	Analysis Batch: Prep Batch:	460-222235 460-219569	L: Ir F	nstrument ID: ab File ID: nitial Weight/Volume: inal Weight/Volume: njection Volume:	CBNAMS6 M79164.D 250 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		10		U	0.19	10
n,n'-Dimethylaniline		1.0		U	0.17	1.0
Surrogate		%Rec		Qualifier	Accepta	nce Limits
2-Fluorobiphenyl		61			50 - 120	
Nitrobenzene-d5 (S	urr)	62			60 - 114	ļ
Terphenyl-d14 (Sur	r)	74			72 - 130	

#### Client: ARCADIS U.S. Inc

Client Sample ID:	MW-35					
Lab Sample ID: Client Matrix:	460-74504-4 Water					e Sampled: 04/16/2014 1330 e Received: 04/17/2014 0920
		8270D Semivolatile Or	ganic Compou	nds (GC/N	NS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/02/2014 0347 04/17/2014 2035	Analysis Batch: Prep Batch:	460-222005 460-219569	L It F	nstrument ID: Lab File ID: nitial Weight/Volume: Final Weight/Volume: njection Volume:	CBNAMS6 M79106.D 240 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		10		U	0.20	10
n,n'-Dimethylaniline		1.6			0.18	1.0
Surrogate		%Rec		Qualifier	Accepta	ance Limits
2-Fluorobiphenyl		58			50 - 12	0
Nitrobenzene-d5 (S	urr)	68			60 - 11	4
Terphenyl-d14 (Sur	r)	89			72 - 130	0

#### Client: ARCADIS U.S. Inc

Client Sample ID:	MW-27					
Lab Sample ID: Client Matrix:	460-74504-5 Water					ate Sampled: 04/16/2014 1440 ate Received: 04/17/2014 0920
		8270D Semivolatile Or	ganic Compou	nds (GC/I	MS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/04/2014 1631 04/17/2014 2035	Analysis Batch: Prep Batch:	460-222416 460-219569		Instrument ID: Lab File ID: Initial Weight/Volume Final Weight/Volume Injection Volume:	
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		0.60		J	0.20	10
n,n'-Dimethylaniline		0.48		J	0.18	1.0
Surrogate		%Rec		Qualifier	Acce	ptance Limits
2-Fluorobiphenyl		57			50 - 1	120
Nitrobenzene-d5 (S	urr)	61			60 - 1	114
Terphenyl-d14 (Sur	r)	76			72 - 1	130

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-28					
Lab Sample ID: Client Matrix:	460-74504-6 Water					te Sampled: 04/16/2014 1545 te Received: 04/17/2014 0920
		8270D Semivolatile Or	ganic Compou	nds (GC/I	MS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/04/2014 1653 04/17/2014 2035	Analysis Batch: Prep Batch:	460-222416 460-219569		Instrument ID: Lab File ID: Initial Weight/Volume: Final Weight/Volume: Injection Volume:	CBNAMS6 M79223.D 250 mL 2 mL 5 uL
Analyte		Result (ug	g/L)	Qualifier	MDL	RL
Aniline		10		U	0.19	10
n,n'-Dimethylaniline		0.72		J	0.17	1.0
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Fluorobiphenyl		73			50 - 12	0
Nitrobenzene-d5 (S	urr)	75			60 - 11	4
Terphenyl-d14 (Sur	r)	103			72 - 13	0

# **Analytical Data**

Client Sample ID:	MW-8SR					
Lab Sample ID: Client Matrix:	460-74504-7 Water					e Sampled: 04/16/2014 1600 e Received: 04/17/2014 0920
		8270D Semivolatile Or	ganic Compou	nds (GC/M	S)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/04/2014 1715 04/17/2014 2035	Analysis Batch: Prep Batch:	460-222416 460-219569	La In Fi	strument ID: ab File ID: itial Weight/Volume: nal Weight/Volume: jection Volume:	CBNAMS6 M79224.D 240 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		3.9		J	0.20	10
n,n'-Dimethylaniline		1.4			0.18	1.0
Surrogate		%Rec		Qualifier	Accepta	nce Limits
2-Fluorobiphenyl		61			50 - 120	
Nitrobenzene-d5 (S	urr)	61			60 - 114	L.
Terphenyl-d14 (Sur	r)	75			72 - 130	)

# **Analytical Data**

Client Sample ID:	BD041614-01					
Lab Sample ID: Client Matrix:	460-74504-8 Water					te Sampled: 04/16/2014 0930 te Received: 04/17/2014 0920
		8270D Semivolatile Or	ganic Compou	nds (GC/M	IS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/04/2014 1737 04/17/2014 2035	Analysis Batch: Prep Batch:	460-222416 460-219569	L Ir F	nstrument ID: ab File ID: nitial Weight/Volume: inal Weight/Volume: njection Volume:	
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		5.6		J	0.20	10
n,n'-Dimethylaniline		1.9			0.18	1.0
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Fluorobiphenyl		78			50 - 12	20
Nitrobenzene-d5 (S	urr)	76			<b>60 - 1</b> 1	4
Terphenyl-d14 (Sur	r)	103			72 - 13	30

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-35					
Lab Sample ID:	460-74504-4				Dat	te Sampled: 04/16/2014 1330
Client Matrix:	Water				Da	te Received: 04/17/2014 0920
	8015D	Nonhalogenated Organic	c Compounds	- Direct lı	njection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volume:	1 mL
Dilution:	1.0				Final Weight/Volume:	
Analysis Date:	04/22/2014 1251				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifie	r Accept	ance Limits
2-Hexanone		111			62 - 12	9

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-28					
Lab Sample ID:	460-74504-6				Dat	e Sampled: 04/16/2014 1545
Client Matrix:	Water				Dat	te Received: 04/17/2014 0920
	8015D	Nonhalogenated Organi	c Compounds	- Direct Ir	ijection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volume:	1 mL
Dilution:	1.0				Final Weight/Volume:	
Analysis Date:	04/22/2014 1331				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Hexanone		113			62 - 12	9

Client Information         Binney Y, Rcg /N.         Maynets         Class State         Class State </th
Sample V. K.C. N. Monce         Change C
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tim     Sampler Y, Kec/N, Mon/Sc     Chang, Grace       Page:
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tion     Sampler V. K.e./N. Mon/Sc.     Chang, Grace       Phone     Phone     Phone     EMail:       Phone     Phone     Phone     EMail:       Due Date Requested:     Grace.chang@testamericain.co     Ana       Due Date Requested:     Pole:     Pole:       Down Pole:     Pole:     Pole:       Time     Sample     (wwww.etc.)       Sample     Cyce:     Pole:       Sample     Cyce:     Pole:       Sample     Cyce:     Pole:       Sample     Cyce:     Pole:       Sample     Cyce:     N       Sample     Cyce:
tion     Sampler X, Kee, N., Monvee     Chang, Grace       Phone:     Phone:     Phone:     E-Mail:       Phone:     Phone:     Phone:     E-Mail:       Juile 600     Purchase     Ouro Date Requested (days):     Int Requested (days):       Ininistration@arcadis-us.com     Porma     Porma     Porma       Porma     Sample     Porma     Porma       Sample Date     Time     Porma     Porma       Time     Sample Date     Preservation Code:     N       Value     N     N     N
tion     Sampler Y. R.C. N. Monice     Club PM: Chang. Grace       Physee     Physee     Physee     Physee       Ininistration@arcadis-us.com     Physee     Physee     Physee       Portinase     Order Requested     Physee     Physee       Portinase     Physee     Physee     Physee       Portinase     Physee     Physee     Physee       Portinase     Physee     Physee     Physee       Physee     Physee     Physee     Physee       Physee     Physee     Physee     Physee       Physee     Sample     (Warranting Strespecial Compound List     Physee       Physee     Physee     Physee     Physee     Physee       Physee     Physee     Physee <t< td=""></t<>
tion     Sampler Y. K.C. N. Mon/Sc.     Lus Pixer       Pione:     Pione:     Pione:     Pione:       Pione:     Sample:     Pione:     Pione:       Sample:     Carcomp:     Sample:     Matrix:       Sample:     Carcomp:     Sample:     Pione:       Sample:     Carcomp:     Sample:     Pione:       Sample:     Carcomp:     Sample:     Pione:       Sample:     Carcomp:     Sanal:     Pione:       Sample:
tion Sampler Y. Kc. / N. Monce Chang, Grace Physe: -382 - 473 4 E-Mai: Physe: -382 - 473 4 E-Mai: Physe: -382 - 473 4 E-Mai: International Control of Chang, Grace The Date Requested Clays): 
tion     Sampler Y. Kc/N. Monce     Lab PM. Chang. Grace       Phone:     Phone:     Phone:       Phone:     Phone:     Phone:       Phone:     Phone:     E-Mail:       Inte 600     Due Date Requested:     Image: Chang. Grace       Inte 600     TAT Requested (days):     Image: Chang. Grace       Inte 600     TAT Requested     Image: Chang. Grace       Inte 600     TAT Requested     Image: Chang. Grace       Inte 600     Pot::     Statundard       Inte 600     S
tion     Sampler Y. Kc/N. Monce     Lab PM. Chang, Grace       Phone:     Phone:     Phone:       Phone:     Phone:     F-383 - 473 4       Image:     Phone:     F-Mat:       Image:     Phone:     F-Mat:       Image:     Phone:     F-Mat:       Image:     Phone:     F-Mat:       Image:     Phone:     Phone:       Image:     Phone:     Phon
tion     Sampler Y. Kc/N.     Lab PM.       Phone:     Phone:     Chang, Grace       Phone:     Phone:     E-Mail:       Phone:     Phone:     E-Mail:       Starbulk     Image: Chang@testamericainc.co     Ana       Suite 600     TAT Requested:     Ana       Starbulk     Power     Ana       Starbulk     Image: Chang@testamericainc.co     Ana       WO #:     Image: Chang@testamericainc.co     Image: Chang@testamericainc.co       Image: Chang@testamericainc.co     Image: Chang@testamericaincainc.co     Image: Chang@testam
tion     Sampler Y. Kc/N.     Lab PM: Chang, Grace       Phone:     Phone:     Chang, Grace       Phone:     Phone:     E-Mail:       Inte 600     Due Date Requested:     Image: Chang@testamericainc.co       Inte 600     TAT Requested:     Image: Chang@testamericainc.co       Inte 600     Image: Chang@testamericainc.co     Image: Chang@testamericaincainc.co       Int
tion Sampler Y. Kce/N. Monree Lab PM: Phome: -382 - 473 4 Grace Chang, Grace Phome: -382 - 473 4 Grace.chang@testamericainc.co Inte 600 Due Date Requested (days): Due Date Requested (days):
tion Sampler X. Kce / N. Monrice Chang, Grace Chang, Grace Phone: - 383 - 4934 Grace.chang@testamericainc.com Analysis Due Date Requested: Analysis
Sampler Y. Kcc/N. Mon/cc     Lab PM: Chang, Grace       Phone:     Phone:       Phone:     - 383- 4934       grace.chang@testamericainc.com       Image:       Due Date Requested:
Sampler Y., Kcc / N., Mon/Sc. Chang, Grace Phone: Phone: - 323-4934 Grace.chang@testamericainc.com Analysis
Sampler Y. Kcc / N. Monroe Lab PM: Phone: Phone: - 383- 4934 E-Malt grace.chang@testameri
Sampler Y. Koc/N. Monroe Chang

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# **Analytical Data**

Client Sample ID:	TB041714						
Lab Sample ID: Client Matrix:	460-74604-1 Water						04/17/2014 1600 : 04/18/2014 0930
		8260C Volatile Orgar	ic Compounds	s by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/23/2014 0822 04/23/2014 0822	Analysis Batch: Prep Batch:	460-220323 N/A		Instrument ID: Lab File ID: Initial Weight/Volur Final Weight/Volur		
Analyte		Result (u	g/L)	Qualifie	r MDL	R	L
Acetone		10		U	2.7	1(	)
Benzene		1.0		U	0.080	1.	0
Ethylbenzene		1.0		U	0.10	1.	0
Methylene Chloride		1.0		U	0.18	1.	0
Toluene		1.0		U	0.15	1.	0
Trichloroethene		1.0		U	0.090	1.	0
Xylenes, Total		3.0		U	0.13	3.	0
Surrogate		%Rec		Qualifie	r Acc	eptance Limits	
1,2-Dichloroethane-	d4 (Surr)	97			70 -	- 130	
Bromofluorobenzen	e	96			64	- 135	
Dibromofluorometha	ane (Surr)	98			72 -	- 137	
Toluene-d8 (Surr)		96			70 -	- 130	

Job Number: 460-74604-1

Client Sample ID: Lab Sample ID: Client Matrix:	<b>MW-31</b> 460-74604-2 Water				[	Date Sampled: 04/17/2014 1600 Date Received: 04/18/2014 0930
		8260C Volatile Orgar	nic Compounds	s by GC/I	MS	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/23/2014 0910 04/23/2014 0910	Analysis Batch: Prep Batch:	460-220323 N/A		Instrument ID: Lab File ID: Initial Weight/Volum Final Weight/Volum	
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		10		U	2.7	10
Benzene		7.5			0.080	1.0
Ethylbenzene		0.22		J	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		0.93		J	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		4.6			0.13	3.0
Surrogate		%Rec		Qualifie	r Acce	eptance Limits
1,2-Dichloroethane	-d4 (Surr)	113			70 -	130
Bromofluorobenzen	e	107			64 -	135
Dibromofluorometh	ane (Surr)	112			72 -	137
Toluene-d8 (Surr)		108			70 -	130

Client: ARCADIS U.S. Inc

Job Number: 460-74604-1

Client Sample ID:	MW-32					
Lab Sample ID: Client Matrix:	460-74604-3 Water					Date Sampled: 04/17/2014 1305 Date Received: 04/18/2014 0930
		8260C Volatile Orgar	nic Compounds	by GC/I	MS	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/23/2014 0934 04/23/2014 0934	Analysis Batch: Prep Batch:	460-220323 N/A		Instrument ID: Lab File ID: Initial Weight/Volum Final Weight/Volum	
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		10		U	2.7	10
Benzene		1.0		U	0.080	1.0
Ethylbenzene		1.0		U	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		1.0		U	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.0		U	0.13	3.0
Surrogate		%Rec		Qualifie	r Acce	eptance Limits
1,2-Dichloroethane-	-d4 (Surr)	99			70 -	130
Bromofluorobenzen		96			64 -	135
Dibromofluorometha	ane (Surr)	101			72 -	137
Toluene-d8 (Surr)		97			70 -	130

Client: ARCADIS U.S. Inc

Client Sample ID:	MW-33					
Lab Sample ID:	460-74604-4					ate Sampled: 04/17/2014 1435
Client Matrix:	Water				D	ate Received: 04/18/2014 0930
		8260C Volatile Orgar	nic Compounds	s by GC/I	MS	
Analysis Method:	8260C	Analysis Batch:	460-220478		Instrument ID:	CVOAMS5
Prep Method:	5030C	Prep Batch:	N/A		Lab File ID:	E27817.D
Dilution:	1.0				Initial Weight/Volume	2: 5 mL
Analysis Date:	04/24/2014 0328				Final Weight/Volume	: 5 mL
Prep Date:	04/24/2014 0328					
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		10		U	2.7	10
Benzene		1.1			0.080	1.0
Ethylbenzene		1.0		U	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		1.0		U	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.0		U	0.13	3.0
Surrogate		%Rec		Qualifie	r Accep	otance Limits
1,2-Dichloroethane-	d4 (Surr)	123			70 - 1	30
Bromofluorobenzen	e	118			64 - 1	35
Dibromofluorometha	ane (Surr)	122			72 - 1	37
Toluene-d8 (Surr)		117			70 - 1	30

## Job Number: 460-74604-1

Client Sample ID: Lab Sample ID: Client Matrix:	<b>MW-34</b> 460-74604-5 Water					Date Sampled: 04/17/2014 1255 Date Received: 04/18/2014 0930
		8260C Volatile Orgar	nic Compounds	s by GC/I	MS	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/24/2014 0352 04/24/2014 0352	Analysis Batch: Prep Batch:	460-220478 N/A		Instrument ID: Lab File ID: Initial Weight/Volur Final Weight/Volun	
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		57			2.7	10
Benzene		1.4			0.080	1.0
Ethylbenzene		0.11		J	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		0.62		J	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.6			0.13	3.0
Surrogate		%Rec		Qualifie	r Acc	ceptance Limits
1,2-Dichloroethane-	-d4 (Surr)	97			70 -	- 130
Bromofluorobenzen	e	94			64 -	- 135
Dibromofluorometha	ane (Surr)	100			72 -	- 137
Toluene-d8 (Surr)		93			70 -	- 130

Client: ARCADIS U.S. Inc

# **Analytical Data**

Client Sample ID:	MW-36R						
Lab Sample ID: Client Matrix:	460-74604-6 Water					Date Sampled: 04/17 Date Received: 04/18	
		8260C Volatile Orgar	nic Compounds	s by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/24/2014 0416 04/24/2014 0416	Analysis Batch: Prep Batch:	460-220478 N/A		Instrument ID: Lab File ID: Initial Weight/Volur Final Weight/Volur		
Analyte		Result (u	g/L)	Qualifie	r MDL	RL	
Acetone		5.5		J	2.7	10	
Benzene		1.1			0.080	1.0	
Ethylbenzene		0.12		J	0.10	1.0	
Methylene Chloride		1.0		U	0.18	1.0	
Toluene		0.42		J	0.15	1.0	
Trichloroethene		1.0		U	0.090	1.0	
Xylenes, Total		1.6		J	0.13	3.0	
Surrogate		%Rec		Qualifie	r Acc	ceptance Limits	
1,2-Dichloroethane	-d4 (Surr)	103			70 -	- 130	
Bromofluorobenzer	e	98			64 -	- 135	
Dibromofluorometh	ane (Surr)	106			72 -	- 137	
Toluene-d8 (Surr)		97			70 -	- 130	

# **Analytical Data**

Client Sample ID:	TW-02RRR						
Lab Sample ID: Client Matrix:	460-74604-7 Water					Date Sampled: 04/1 Date Received: 04/1	
		8260C Volatile Orgar	nic Compounds	s by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/24/2014 0440 04/24/2014 0440	Analysis Batch: Prep Batch:	460-220478 N/A		Instrument ID: Lab File ID: Initial Weight/Volur Final Weight/Volur		
Analyte		Result (u	g/L)	Qualifie	r MDL	RL	
Acetone		8.0		J	2.7	10	
Benzene		1.2			0.080	1.0	
Ethylbenzene		0.51		J	0.10	1.0	
Methylene Chloride		1.0		U	0.18	1.0	
Toluene		0.18		J	0.15	1.0	
Trichloroethene		1.0		U	0.090	1.0	
Xylenes, Total		1.0		J	0.13	3.0	
Surrogate		%Rec		Qualifie	r Acc	ceptance Limits	
1,2-Dichloroethane-	d4 (Surr)	102			70	- 130	
Bromofluorobenzen	е	99			64	- 135	
Dibromofluorometha	ane (Surr)	103			72	- 137	
Toluene-d8 (Surr)		99			70	- 130	

Job Number: 460-74604-1

Client Sample ID:	TW-01					
Lab Sample ID:	460-74604-8				Date	e Sampled: 04/17/2014 1155
Client Matrix:	Water				Date	e Received: 04/18/2014 0930
		8260C Volatile Orgar	nic Compounds	by GC/N	IS	
Analysis Method:	8260C	Analysis Batch: 460-220478 Ir		Instrument ID:	CVOAMS5	
Prep Method:	5030C	Prep Batch:	N/A		Lab File ID:	E27821.D
Dilution:	1.0				Initial Weight/Volume:	5 mL
Analysis Date:	04/24/2014 0504				Final Weight/Volume:	5 mL
Prep Date:	04/24/2014 0504					
Analyte		Result (u	g/L) (	Qualifier	MDL	RL
Acetone		10		U	2.7	10
Benzene		1.0	I	U	0.080	1.0
Ethylbenzene		1.0	I	U	0.10	1.0
Methylene Chloride	•	1.0	I	U	0.18	1.0
Toluene		1.0	I	U	0.15	1.0
Trichloroethene		1.0	I	U	0.090	1.0
Xylenes, Total		3.0	I	U	0.13	3.0
Surrogate		%Rec	(	Qualifier	Accepta	ince Limits
1,2-Dichloroethane	-d4 (Surr)	107			70 - 130	)
Bromofluorobenzer	ne	104			64 - 135	5
Dibromofluorometh	ane (Surr)	110			72 - 137	7
Toluene-d8 (Surr)		103			70 - 130	)

Client: ARCADIS U.S. Inc

# **Analytical Data**

Client Sample ID:	BD041714-02						
Lab Sample ID: Client Matrix:	460-74604-9 Water						ampled: 04/17/2014 0900 eceived: 04/18/2014 0930
		8260C Volatile Orgar	ic Compounds	s by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/24/2014 0529 04/24/2014 0529	Analysis Batch: Prep Batch:	460-220478 N/A		Instrument ID: Lab File ID: Initial Weight/Volu Final Weight/Volu		CVOAMS5 E27822.D 5 mL 5 mL
Analyte		Result (u	g/L)	Qualifie	r MDL		RL
Acetone		10			2.7		10
Benzene		1.2			0.080		1.0
Ethylbenzene		0.44		J	0.10		1.0
Methylene Chloride		1.0		U	0.18		1.0
Toluene		0.17		J	0.15		1.0
Trichloroethene		1.0		U	0.090		1.0
Xylenes, Total		0.96		J	0.13		3.0
Surrogate		%Rec		Qualifie	r Ac	ceptance	e Limits
1,2-Dichloroethane	-d4 (Surr)	97			70	- 130	
Bromofluorobenzer	e	93			64	- 135	
Dibromofluorometh	ane (Surr)	99			72	- 137	
Toluene-d8 (Surr)		93			70	- 130	

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-31					
Lab Sample ID: Client Matrix:	460-74604-2 Water					e Sampled: 04/17/2014 1600 e Received: 04/18/2014 0930
		8270D Semivolatile Or	ganic Compou	nds (GC/I	MS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/03/2014 1743 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222297 460-219763		Instrument ID: Lab File ID: Initial Weight/Volume: Final Weight/Volume: Injection Volume:	CBNAMS6 M79198.D 250 mL 2 mL 5 uL
Analyte		Result (ug	g/L)	Qualifier	MDL	RL
Aniline		0.75		J	0.19	10
n,n'-Dimethylaniline		1.9			0.17	1.0
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Fluorobiphenyl		94			50 - 12	0
Nitrobenzene-d5 (S	urr)	93			60 - 11	4
Terphenyl-d14 (Sur	r)	122			72 - 13	0

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-32					
Lab Sample ID: Client Matrix:	460-74604-3 Water					Sampled: 04/17/2014 1305 Received: 04/18/2014 0930
		8270D Semivolatile Or	ganic Compou	nds (GC/M	S)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/03/2014 1805 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222297 460-219763	La In Fi	strument ID: ab File ID: itial Weight/Volume: nal Weight/Volume: jection Volume:	CBNAMS6 M79199.D 240 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		10		U	0.20	10
n,n'-Dimethylaniline		1.1			0.18	1.0
Surrogate		%Rec		Qualifier	Acceptar	nce Limits
2-Fluorobiphenyl		90			50 - 120	
Nitrobenzene-d5 (S	urr)	98			60 - 114	
Terphenyl-d14 (Sur	r)	127			72 - 130	

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-33					
Lab Sample ID: Client Matrix:	460-74604-4 Water					e Sampled: 04/17/2014 1435 e Received: 04/18/2014 0930
		8270D Semivolatile Or	ganic Compou	nds (GC/N	IS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/05/2014 0309 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222463 460-219763	L Ir F	nstrument ID: .ab File ID: nitial Weight/Volume: final Weight/Volume: njection Volume:	CBNAMS6 M79250.D 240 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		0.32		J	0.20	10
n,n'-Dimethylaniline		2.3			0.18	1.0
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Fluorobiphenyl		97			50 - 12	0
Nitrobenzene-d5 (S	urr)	98			60 - 11	4
Terphenyl-d14 (Sur	r)	124			72 - 13	D

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-34					
Lab Sample ID: Client Matrix:	460-74604-5 Water					Sampled: 04/17/2014 1255 Received: 04/18/2014 0930
		8270D Semivolatile Or	ganic Compou	nds (GC/M	S)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/05/2014 0330 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222463 460-219763	La In Fi	strument ID: ab File ID: itial Weight/Volume: inal Weight/Volume: jection Volume:	CBNAMS6 M79251.D 250 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		2.6		J	0.19	10
n,n'-Dimethylaniline		1.6			0.17	1.0
Surrogate		%Rec		Qualifier	Acceptar	nce Limits
2-Fluorobiphenyl		92			50 - 120	
Nitrobenzene-d5 (S	urr)	86			60 - 114	
Terphenyl-d14 (Sur	r)	137		Х	72 - 130	

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-36R					
Lab Sample ID: Client Matrix:	460-74604-6 Water					te Sampled: 04/17/2014 1410 te Received: 04/18/2014 0930
		8270D Semivolatile Or	ganic Compou	nds (GC/N	MS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/05/2014 0352 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222463 460-219763	L I F	Instrument ID: Lab File ID: Initial Weight/Volume: Final Weight/Volume: Injection Volume:	CBNAMS6 M79252.D 250 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		140			0.19	10
n,n'-Dimethylaniline		3.4			0.17	1.0
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Fluorobiphenyl		94			50 - 12	0
Nitrobenzene-d5 (S	urr)	90			60 - 11	4
Terphenyl-d14 (Sur	r)	128			72 - 13	0

# **Analytical Data**

Client Sample ID:	TW-02RRR					
Lab Sample ID: Client Matrix:	460-74604-7 Water					e Sampled: 04/17/2014 1120 e Received: 04/18/2014 0930
		8270D Semivolatile Or	ganic Compou	nds (GC/N	MS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 10 05/05/2014 0414 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222463 460-219763	L I F	Instrument ID: Lab File ID: Initial Weight/Volume: Final Weight/Volume: Injection Volume:	CBNAMS6 M79253.D 250 mL 2 mL 5 uL
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Aniline		1300		J	1.9	100
n,n'-Dimethylaniline		2.8		J	1.7	10
Surrogate		%Rec		Qualifier	Accepta	nce Limits
2-Fluorobiphenyl		0		Х	50 - 120	
Nitrobenzene-d5 (S	urr)	0		Х	60 - 114	
Terphenyl-d14 (Sur	r)	0		Х	72 - 130	)

## Client: ARCADIS U.S. Inc

Client Sample ID:	TW-01					
Lab Sample ID: Client Matrix:	460-74604-8 Water					ate Sampled: 04/17/2014 1155 ate Received: 04/18/2014 0930
		8270D Semivolatile Org	ganic Compou	nds (GC/I	MS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 05/05/2014 0436 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222463 460-219763	   	Instrument ID: Lab File ID: Initial Weight/Volume Final Weight/Volume Injection Volume:	
Analyte		Result (ug	g/L)	Qualifier	MDL	RL
Aniline		10		U	0.20	10
n,n'-Dimethylaniline		1.3			0.18	1.0
Surrogate		%Rec		Qualifier	Accep	tance Limits
2-Fluorobiphenyl		83			50 - 12	20
Nitrobenzene-d5 (S	urr)	83			60 - 1	14
Terphenyl-d14 (Sur	r)	120			72 - 1	30

# **Analytical Data**

Client Sample ID:	BD041714-02					
Lab Sample ID: Client Matrix:	460-74604-9 Water					te Sampled: 04/17/2014 0900 te Received: 04/18/2014 0930
		8270D Semivolatile Or	ganic Compou	nds (GC/	MS)	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 10 05/05/2014 0458 04/18/2014 2016	Analysis Batch: Prep Batch:	460-222463 460-219763		Instrument ID: Lab File ID: Initial Weight/Volume: Final Weight/Volume: Injection Volume:	
Analyte		Result (u	g/L)	Qualifie	MDL	RL
Aniline		1700		J	1.9	100
n,n'-Dimethylaniline		3.5		J	1.7	10
Surrogate		%Rec		Qualifie	Accept	ance Limits
2-Fluorobiphenyl		0		Х	50 - 12	20
Nitrobenzene-d5 (S	urr)	0		Х	60 - 11	4
Terphenyl-d14 (Sur	r)	0		Х	72 - 13	0

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-31					
Lab Sample ID:	460-74604-2					Date Sampled: 04/17/2014 1600
Client Matrix:	Water					Date Received: 04/18/2014 0930
	8015D	Nonhalogenated Organi	c Compounds	- Direct lı	njection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volur	ne: 1 mL
Dilution:	1.0				Final Weight/Volun	ne:
Analysis Date:	04/22/2014 1344				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifier	Acc	eptance Limits
2-Hexanone		111			62 -	129

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-32					
Lab Sample ID: Client Matrix:	460-74604-3 Water					Date Sampled: 04/17/2014 1305 Date Received: 04/18/2014 0930
	8015D	Nonhalogenated Organic	c Compounds	- Direct In	ijection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volur	me: 1 mL
Dilution:	1.0				Final Weight/Volun	me:
Analysis Date:	04/22/2014 1357				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifier	Acc	ceptance Limits
2-Hexanone		105			62 -	- 129

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-34					
Lab Sample ID:	460-74604-5				Da	ate Sampled: 04/17/2014 1255
Client Matrix:	Water				Da	ate Received: 04/18/2014 0930
	8015D	Nonhalogenated Organic	c Compounds	- Direct lı	njection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volume	e: 1 mL
Dilution:	1.0				Final Weight/Volume	:
Analysis Date:	04/22/2014 1410				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifie	- MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifie	- Accep	otance Limits
2-Hexanone		97			62 - 1	29

## Client: ARCADIS U.S. Inc

Job Number: 460-74604-1

62 - 129

Client Sample ID:	TW-02RRR					
Lab Sample ID:	460-74604-7				Date	Sampled: 04/17/2014 1120
Client Matrix:	Water				Date	Received: 04/18/2014 0930
	8015D	Nonhalogenated Organi	c Compounds	- Direct Inj	ection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397	Ir	nstrument ID:	HP5890-4
	N/A		N/A	Ir	nitial Weight/Volume:	1 mL
Dilution:	1.0			F	inal Weight/Volume:	
Analysis Date:	04/22/2014 1423			Ir	njection Volume:	1 uL
Prep Date:	N/A			F	esult Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifier	Acceptar	nce Limits

122

2-Hexanone

## Client: ARCADIS U.S. Inc

Client Sample ID:	TW-01					
Lab Sample ID:	460-74604-8				Date	e Sampled: 04/17/2014 1155
Client Matrix:	Water				Date	e Received: 04/18/2014 0930
	8015D	Nonhalogenated Organic	: Compounds	- Direct lı	njection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volume:	1 mL
Dilution:	1.0				Final Weight/Volume:	
Analysis Date:	04/22/2014 1436				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifier	Accepta	nce Limits
2-Hexanone		100			62 - 129	

## Client: ARCADIS U.S. Inc

Client Sample ID:	BD041714-02					
Lab Sample ID:	460-74604-9					Sampled: 04/17/2014 0900
Client Matrix:	Water				Date	Received: 04/18/2014 0930
	8015D	Nonhalogenated Organi	c Compounds	- Direct lı	njection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177397		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volume:	1 mL
Dilution:	1.0				Final Weight/Volume:	
Analysis Date:	04/22/2014 1450				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	ıg/L)	Qualifier	r MDL	RL

-				
Methanol	500	U	410	500
Surrogate	%Rec	Qualifier	Acceptance Li	mits
2-Hexanone	100		62 - 129	

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# **Analytical Data**

Client Sample ID:	TB041814					
Lab Sample ID: Client Matrix:	460-74662-1 Water					Date Sampled: 04/18/2014 1110 Date Received: 04/19/2014 1000
		8260C Volatile Orga	nic Compounds	s by GC/I	MS	
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/27/2014 1700 04/27/2014 1700	Analysis Batch: Prep Batch:	460-221116 N/A		Instrument ID: Lab File ID: Initial Weight/Volur Final Weight/Volun	
Analyte		Result (u	g/L)	Qualifie	r MDL	RL
Acetone		10		U	2.7	10
Benzene		1.0		U	0.080	1.0
Ethylbenzene		1.0		U	0.10	1.0
Methylene Chloride		1.0		U	0.18	1.0
Toluene		1.0		U	0.15	1.0
Trichloroethene		1.0		U	0.090	1.0
Xylenes, Total		3.0		U	0.13	3.0
Surrogate		%Rec		Qualifie	r Acc	eptance Limits
1,2-Dichloroethane	-d4 (Surr)	107			70 -	- 130
Bromofluorobenzer	e	101			64 -	- 135
Dibromofluorometh	ane (Surr)	109			72 -	- 137
Toluene-d8 (Surr)		98			70 -	- 130
. ,						

Job Number: 460-74662-1

Client Sample ID:	MW-9S						
Lab Sample ID: Client Matrix:	460-74662-2 Water						Sampled: 04/18/2014 1110 Received: 04/19/2014 1000
		8260C Volatile Organ	ic Compounds	s by GC/I	MS		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8260C 5030C 1.0 04/27/2014 1748 04/27/2014 1748	Analysis Batch: Prep Batch:	460-221116 N/A		Instrument ID: Lab File ID: Initial Weight/Volu Final Weight/Volu		CVOAMS5 E27992.D 5 mL 5 mL
Analyte		Result (u	g/L)	Qualifie	r MDL		RL
Acetone		10		U	2.7		10
Benzene		1.0			0.080		1.0
Ethylbenzene		19			0.10		1.0
Methylene Chloride		1.0		U	0.18		1.0
Toluene		2.2			0.15		1.0
Trichloroethene		1.0		U	0.090		1.0
Xylenes, Total		74			0.13		3.0
Surrogate		%Rec		Qualifie	r Ac	ceptan	ce Limits
1,2-Dichloroethane-	d4 (Surr)	104			70	- 130	
Bromofluorobenzen	e	100			64	- 135	
Dibromofluorometha	ane (Surr)	100			72	- 137	
Toluene-d8 (Surr)		96			70	- 130	

Client: ARCADIS U.S. Inc

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-9S						
Lab Sample ID: Client Matrix:	460-74662-2 Water						ed: 04/18/2014 1110 ed: 04/19/2014 1000
		8270D Semivolatile Or	ganic Compou	nds (GC/	MS)		
Analysis Method: Prep Method: Dilution: Analysis Date: Prep Date:	8270D 3510C 1.0 04/26/2014 1603 04/21/2014 2048	Analysis Batch: Prep Batch:	460-220995 460-220073		Instrument ID: Lab File ID: Initial Weight/Volu Final Weight/Volu Injection Volume:	M78 ime: 250	ıL
Analyte		Result (u	g/L)	Qualifier	r MDL		RL
Aniline		10		U	0.19		10
n,n'-Dimethylaniline		5.7			0.17		1.0
Surrogate		%Rec		Qualifier	Ac	ceptance Limi	its
2-Fluorobiphenyl		92			50	- 120	
Nitrobenzene-d5 (S	urr)	93			60	- 114	
Terphenyl-d14 (Sur	r)	115			72	- 130	

## Client: ARCADIS U.S. Inc

Client Sample ID:	MW-9S					
Lab Sample ID:	460-74662-2				Dat	e Sampled: 04/18/2014 1110
Client Matrix:	Water				Da	te Received: 04/19/2014 1000
	8015D	Nonhalogenated Organi	c Compounds	- Direct lı	njection (GC)	
Analysis Method:	8015D	Analysis Batch:	480-177672		Instrument ID:	HP5890-4
	N/A		N/A		Initial Weight/Volume:	1 mL
Dilution:	1.0				Final Weight/Volume:	
Analysis Date:	04/23/2014 1100				Injection Volume:	1 uL
Prep Date:	N/A				Result Type:	PRIMARY
Analyte		Result (u	g/L)	Qualifier	MDL	RL
Methanol		500		U	410	500
Surrogate		%Rec		Qualifier	Accept	ance Limits
2-Hexanone		106			62 - 12	9