

Hard Copy Required

Page 2 of 2

Thursday, February 02, 2012

REQUEST NUMBER: 12-700

PRIORITY	METHOD CODE	CNTNR	SAMPLE ID	SAMPLE MATRIX	DATE SAMPLED	SPECIAL INSTRUCTIONS
	SW-846:8260B	1	CAAN-12-2032	WG	2/1/2012	
		2	CAAN-12-2032	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		1	CAAN-12-2201	WG	2/1/2012	
		2	CAAN-12-2201	WG	2/1/2012	
	SW-846:8270C	1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		3	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		3	CAAN-12-2199	WG	2/1/2012	
		1	CAAN-12-2201	WG	2/1/2012	
		2	CAAN-12-2201	WG	2/1/2012	
		3	CAAN-12-2201	WG	2/1/2012	
	SW-846:8321A_MOD	1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		3	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		3	CAAN-12-2199	WG	2/1/2012	
	SW-846:9060	1	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	

Final Page of REQUEST NUMBER 12-700

Thursday, February 02, 2012

REQUEST NUMBER: 12-700

LOS ALAMOS
NATIONAL LABORATORY

ATTN: Valerie Davis

General Engineering Laboratories, Inc., Charleston, SC.
2040 Savage Rd
Charleston, SC 29407

These Samples are on:

LANL Request Number: 12-700


Per Agreement Number: 126310011

Project Cost Code: MR1A015AGWJ0

Please analyse the enclosed samples
according to the schedule indicated:

SHIP DATE: 2/2/2012**TURNAROUND/REPORT DUE: 3/3/2012****TURNAROUND REQ'D: 30 Days****RAD SCREENING: Yes, Below Background****LAB REQUEST COMMENTS:**

LANL ER SMO CONTACT:

Signature: 

PRIORITY	METHOD CODE	CNTNR	SAMPLE ID	SAMPLE MATRIX	DATE SAMPLED	SPECIAL INSTRUCTIONS
	EPA:350.1	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	EPA:351.2	1	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
	EPA:353.2	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	EPA:365.4	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	SW-846:8260B	1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	

Thursday, February 02, 2012

LAB CHAIN OF CUSTODY DOCUMENT NUMBER: 12-700C

LOS ALAMOS

REQUEST NUMBER: 12-700

NATIONAL LABORATORY

ATTN: Valerie Davis

TURNAROUND/REPORT DUE: 3/3/2012

General Engineering Laboratories, Inc., Charleston, SC.

TURNAROUND REQ'D: 30

2040 Savage Rd

Charleston, SC 29407

LAB REQUEST COMMENTS:

SAMPLE ID	CTNR	CTNR DESC	ORDER	PRESERV	MATRIX
CAAN-12-2199	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2199	1	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	2	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	3	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	1	AMBER GLASS	WSP-TKN+TOC	Sulfuric Acid (H2SO4)	WG
CAAN-12-2200	1	AMBER GLASS	WSP-NH3+NO3/NO2+ PO4	Sulfuric Acid (H2SO4)	WG
CAAN-12-2201	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2201	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2201	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2201	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2201	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG

Relinquished By:

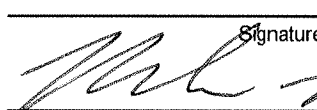
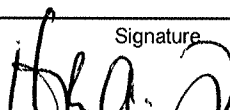

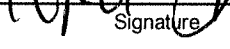
Date

Time

Received By:

Date

Time

	
Signature	Signature
	
Signature	Signature

Received for DISPOSAL By: Date

Time

Remarks:

Signature

Thursday, February 02, 2012

LAB CHAIN OF CUSTODY DOCUMENT NUMBER: 12-700C

LOS ALAMOS

REQUEST NUMBER: 12-700

NATIONAL LABORATORY

ATTN: Valerie Davis

TURNAROUND/REPORT DUE: 3/3/2012

General Engineering Laboratories, Inc., Charleston, SC.

TURNAROUND REQ'D: 30

2040 Savage Rd

Charleston, SC 29407

LAB REQUEST COMMENTS:

295270

SAMPLE ID	CTNR	CTNR DESC	ORDER	PRESERV	MATRIX
CAAN-12-2030	1	AMBER GLASS	WSP-NH3+NO3/NO2+PO4	Sulfuric Acid (H2SO4)	WG
CAAN-12-2031	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2031	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2031	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	1	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	2	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	3	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	1	AMBER GLASS	WSP-TKN+TOC	Sulfuric Acid (H2SO4)	WG
CAAN-12-2032	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2032	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2199	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY

EVENT ID: 3734

EVENT NAME: Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011
Interim Plan rev. 1

SAMPLE ID: CAAN-12-2030

WORK ORDER:

AS PLANNED		AS COLLECTED	AS PLANNED		AS COLLECTED
DATE COLLECTED(MM/DD/YYYY):		2/1/2012	MEDIA:	WGR	OK
TIME COLLECTED (HH:MM)		1245	SUB-MEDIA:	UA	
PRS ID:	Ancho	OK	SAMPLE TECH CODE:	GSP	
LOCATION ID:	R-30		FIELD QC TYPE:	NA	
LOCATION TYPE:	MON		FIELD PREP:	F	
PORT:	SINGLE COMPLETION		SAMPLE USAGE:	INV	
			SCREEN/PORT DESC:		
FIELD MATRIX:	WG		EXCAVATED: YES/NO/NA		
COMPOSITE TYPE:	NA		COMPOSITE TIME INTERVAL:	NA	
			WATER FLOWING: YES/NO/NA		
BOREHOLE: YES/NO/NA	NA		BOREHOLE DECLINATION:	NA	
			BOREHOLE DIRECTION:	NA	

#	PRIORITY	ORDER	CNTNR	PRESERVATIVE	COLLECTED Y/N	SPECIAL INSTRUCTIONS
1	NA	WSP-GENINORG	1 LITER POLY	Ice	Y	NA
1		WSP-Met+B+SN +SR+U	1 LITER POLY	Nitric Acid (HNO3)		
1		WSP-NH3+NO3/ NO2+PO4	500 ML AMBER GLASS	Sulfuric Acid (H2SO4)		

SAMPLE DESC:

SAMPLE COMMENTS:

LOCATION DESC:

FIELD SCREENING/MEASUREMENT RESULTS:

COLLECTED BY (PRINT)

D. Woody

REVIEWED BY (PRINT)

W. Shan

RELINQUISHED BY (Printed Name) D Woody (Signature) D Woody	Date/Time 02/01/12 1415	RECEIVED BY (Printed Name) Dean Sheppard (Signature) Dean Sheppard	Date/Time 02/01/12 1415
RELINQUISHED BY (Printed Name) (Signature)	Date/Time	RECEIVED BY (Printed Name) (Signature)	Date/Time

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY

EVENT ID: 3734

EVENT NAME: Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011
Interim Plan rev. 1

SAMPLE ID: CAAN-12-2031

WORK ORDER:

AS PLANNED		AS COLLECTED		AS PLANNED		AS COLLECTED	
DATE COLLECTED(MM/DD/YYYY):		2 / 1 / 2012		MEDIA:		WGR	
TIME COLLECTED (HH:MM)		1245		SUB-MEDIA:		UA	
PRS ID: Ancho		OK		SAMPLE TECH CODE:		GSP	
LOCATION ID: R-30				FIELD QC TYPE:		NA	
LOCATION TYPE: MON				FIELD PREP:		UF	
PORT: SINGLE COMPLETION				SAMPLE USAGE:		INV	
				SCREEN/PORT DESC:			
FIELD MATRIX: WG				EXCAVATED: YES/NO/NA		NA	
COMPOSITE TYPE: NA				COMPOSITE TIME INTERVAL: NA		WATER FLOWING: YES/NO/NA	
BOREHOLE: YES/NO/NA		BOREHOLE DECLINATION: NA		BOREHOLE DIRECTION: NA			

#	PRIORITY	ORDER	CNTNR	PRESERVATIVE	COLLECTED Y/N	SPECIAL INSTRUCTIONS
2	NA	WSP-8260B-VOA	40 ML SEPTUM AMBER GLASS	Hydrochloric Acid (HCL)	Y	NA
3		WSP-8270C-SVOA	1 LITER AMBER GLASS	Ice		
3		WSP-8321A-NMED HEXP	1 LITER AMBER GLASS	Ice		
1		WSP-GrossA/B	1 LITER POLY	None		
2		WSP-HEXMOD	1 LITER AMBER GLASS	Ice		
1		WSP-LL-H-3	1 LITER POLY	None		
1		WSP-RAD	1 GAL POLY	Nitric Acid (HNO3)		
1		WSP-TKN+TOC	500 ML AMBER GLASS	Sulfuric Acid (H2SO4)		
1		Re226+228	1 GAL POLY	Nitric Acid (HNO3)	NA 1/30/12	

SAMPLE DESC: NA

SAMPLE COMMENTS:

Deisel generator running ~50' away during sampling

LOCATION DESC:

NA

FIELD SCREENING/MEASUREMENT RESULTS:

pH	TC	SC (mg/L)	DO (mg/L)	ORP (mV)	Q (gpm)	Turb (NTU)
7.97	20.25	117	7.98	176.6	1.25 ^{1.25} ₅ ^{1/2}	1.25

COLLECTED BY (PRINT)

D Woody

REVIEWED BY (PRINT)

W. Shaw

RELINQUISHED BY

Date/Time

RECEIVED BY

Date/Time

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY**EVENT ID:** 3734**EVENT NAME:** Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011

Interim Plan rev 1

(Printed Name) <i>D Woody</i>	<i>02/01/12</i>	(Printed Name) <i>Sherrin Greenwood</i>	<i>02/01/12</i>
(Signature) <i>D Woody</i>	<i>1415</i>	(Signature) <i>Sherrin Greenwood</i>	<i>1415</i>
RELINQUISHED BY	Date/Time	RECEIVED BY	Date/Time
(Printed Name)		(Printed Name)	
(Signature)		(Signature)	

3734

CAAN-12-2031

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY

EVENT ID: 3734

EVENT NAME: Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011
Interim Plan rev. 1

SAMPLE ID: CAAN-12-2032

WORK ORDER:

AS PLANNED		AS COLLECTED		AS PLANNED		AS COLLECTED	
DATE COLLECTED(MM/DD/YYYY):		2/1/2012		MEDIA:		WGR	
TIME COLLECTED (HH:MM)		1245		SUB-MEDIA:		UA	
PRS ID: Ancho		OK		SAMPLE TECH CODE:		NC	
LOCATION ID: R-30				FIELD QC TYPE:		ETB	
LOCATION TYPE: MON				FIELD PREP:		UF	
PORT: SINGLE COMPLETION				SAMPLE USAGE:		QC	
				SCREEN/PORT DESC:			
FIELD MATRIX: WG				EXCAVATED: YES/NO/NA		NA	
COMPOSITE TYPE: NA				COMPOSITE TIME INTERVAL: NA		WATER FLOWING: YES/NO (NA)	
BOREHOLE: YES/NO/NA		BOREHOLE DECLINATION: NA		BOREHOLE DIRECTION: NA			

#	PRIORITY	ORDER	CNTNR	PRESERVATIVE	COLLECTED Y/N	SPECIAL INSTRUCTIONS
1	1/31/12	WSP-8260B-VOA	40 ML SEPTUM AMBER GLASS	Hydrochloric Acid (HCL)	Y	NA

SAMPLE DESC: QC Sample of CAAN-12-2031

SAMPLE COMMENTS:

LOCATION DESC:

FIELD SCREENING/MEASUREMENT RESULTS:

COLLECTED BY (PRINT)

W. Shaw

REVIEWED BY (PRINT)

D. Woody

RELINQUISHED BY (Printed Name) D Woody (Signature) D Woody	Date/Time 02/01/12 1415	RECEIVED BY (Printed Name) Sheri Sherwood (Signature) Sheri Sherwood	Date/Time 02/01/12 1415
RELINQUISHED BY (Printed Name) (Signature)	Date/Time	RECEIVED BY (Printed Name) (Signature)	Date/Time

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY

EVENT ID: 3734

EVENT NAME: Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011
Interim Plan rev. 1

SAMPLE ID: CAAN-12-2199

WORK ORDER:

AS PLANNED		AS COLLECTED		AS PLANNED		AS COLLECTED	
DATE COLLECTED(MM/DD/YYYY):		2/1/2012		MEDIA:	WGR		OK
TIME COLLECTED (HH:MM)		1245		SUB-MEDIA:	UA		
PRS ID:	Ancho	OK		SAMPLE TECH CODE:	GSP		
LOCATION ID:	R-30			FIELD QC TYPE:	ED		
LOCATION TYPE:	MON			FIELD PREP:	UE		
PORT:	SINGLE COMPLETION			SAMPLE USAGE:	QC		
				SCREEN/PORT DESC:			
FIELD MATRIX:	WG			EXCAVATED: YES/NO/NA	NA		
COMPOSITE TYPE:	NA			COMPOSITE TIME INTERVAL:	NA		
				WATER FLOWING: YES/NO/NA	NA		
BOREHOLE: YES/NO/NA	NA			BOREHOLE DECLINATION:	NA		
				BOREHOLE DIRECTION:	NA		

#	PRIORITY	ORDER	CNTNR	PRESERVATIVE	COLLECTED Y/N	SPECIAL INSTRUCTIONS
2	NA	WSP-8260B-VOA	40 ML SEPTUM AMBER GLASS	Hydrochloric Acid (HCL)	Y	NA
1	NA	WSP-8270C-SVOA	1 LITER AMBER GLASS	Ice	Y	NA
1	NA	WSP-8321A-NMED HEXP	1 LITER AMBER GLASS	Ice	Y	NA
1	NA	WSP-GrossA/B	1 LITER POLY	None	Y	NA
2	NA	WSP-HEXMOD	1 LITER AMBER GLASS	Ice	Y	NA
1	NA	WSP-LL-H-3	1 LITER POLY	None	Y	NA
1	NA	WSP-RAD	1 GAL POLY	Nitric Acid (HNO3)	Y	NA
1	NA	WSP-TKN+TOC	500 ML AMBER GLASS	Sulfuric Acid (H2SO4)	Y	NA

SAMPLE DESC: QC Sample of CAAN-12-2031

SAMPLE COMMENTS:

LOCATION DESC:

FIELD SCREENING/MEASUREMENT RESULTS:

COLLECTED BY (PRINT)

J. Romero

REVIEWED BY (PRINT)

W. Shaw

RELINQUISHED BY

(Printed Name)

Date/Time

RECEIVED BY

(Printed Name)

Date/Time

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY

EVENT ID: 3734

EVENT NAME: Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011
Interim Plan rev. 1

(Signature)		(Signature)	ONS
RELINQUISHED BY	Date/Time	RECEIVED BY	Date/Time
(Printed Name) D. Woody	02/01/12	(Printed Name) JERRI SHERWOOD	12/01/12
(Signature) D. Woody	1415	(Signature) JERRI SHERWOOD	1415

3734

CAAN-12-2199

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY

EVENT ID: 3734

EVENT NAME: Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011
Interim Plan rev. 1

SAMPLE ID: CAAN-12-2200

WORK ORDER:

AS PLANNED		AS COLLECTED		AS PLANNED		AS COLLECTED	
DATE COLLECTED(MM/DD/YYYY):		2/1/2012		MEDIA:	WGR		OK
TIME COLLECTED (HH:MM)		1245		SUB-MEDIA:	UA		
PRS ID:	Ancho			SAMPLE TECH CODE:	GSP		
LOCATION ID:	R-30			FIELD QC TYPE:	FD		
LOCATION TYPE:	MON			FIELD PREP:	E		
PORT:	SINGLE COMPLETION			SAMPLE USAGE:	QC		
				SCREEN/PORT DESC:			
FIELD MATRIX:	WG			EXCAVATED: YES/NO/NA	NA		
COMPOSITE TYPE:	NA			COMPOSITE TIME INTERVAL:	NA		
				WATER FLOWING: YES/NO/NA	NA		
BOREHOLE: YES/NO/NA	NA			BOREHOLE DECLINATION:	NA		
				BOREHOLE DIRECTION:	NA		

#	PRIORITY	ORDER	CNTNR	PRESERVATIVE	COLLECTED Y/N	SPECIAL INSTRUCTIONS
1	NA	WSP-GENINORG	1 LITER POLY	Ice	Y	NA
1		WSP-Met+B+SN +SR+U	1 LITER POLY	Nitric Acid (HNO3)	I	
1		WSP-NH3+NO3/ NO2+PO4	500 ML AMBER GLASS	Sulfuric Acid (H2SO4)	I	

SAMPLE DESC: QC Sample of CAAN-12-2030

SAMPLE COMMENTS:

LOCATION DESC:

FIELD SCREENING/MEASUREMENT RESULTS:

COLLECTED BY (PRINT)

D. Woody

REVIEWED BY (PRINT)

W. Shaw

RELINQUISHED BY	Date/Time	RECEIVED BY	Date/Time
(Printed Name) D Woody	01/02/12	(Printed Name) D. Woody	01/02/12
(Signature) D Woody	1415	(Signature) D. Woody	1415
RELINQUISHED BY	Date/Time	RECEIVED BY	Date/Time
(Printed Name)		(Printed Name)	
(Signature)		(Signature)	

SAMPLE COLLECTION LOG/FIELD CHAIN OF CUSTODY

EVENT ID: 3734

EVENT NAME: Ancho, MDA AB Mon. Group Sampling Q2, January 2012, 2011
Interim Plan rev. 1

SAMPLE ID: CAAN-12-2201

WORK ORDER:

AS PLANNED		AS COLLECTED		AS PLANNED		AS COLLECTED	
DATE COLLECTED(MM/DD/YYYY):		2/1/2012		MEDIA:	WGR		OK
TIME COLLECTED (HH:MM)		1245		SUB-MEDIA:	UA		
PRS ID:	Ancho	OK		SAMPLE TECH CODE:	02		
LOCATION ID:	R-30			FIELD QC TYPE:	FB		
LOCATION TYPE:	MON			FIELD PREP:	UF		
PORT:	SINGLE COMPLETION			SAMPLE USAGE:	QC		
				SCREEN/PORT DESC:			
FIELD MATRIX:	WG			EXCAVATED: YES/NO/NA			
COMPOSITE TYPE:	N/A			COMPOSITE TIME INTERVAL:	N/A		
BOREHOLE: YES/NO/NA				WATER FLOWING: YES/NO/NA			
BOREHOLE DECLINATION:	N/A			BOREHOLE DIRECTION:	N/A		

#	PRIORITY	ORDER	CNTNR	PRESERVATIVE	COLLECTED Y/N	SPECIAL INSTRUCTIONS
2	N/A	WSP-8260B-VOA	40 ML SEPTUM AMBER GLASS	Hydrochloric Acid (HCL)	Y	N/A
1	1/3/12 vs	WSP-8270C-SVOA	1 LITER AMBER GLASS	Ice	J	J

SAMPLE DESC: QC Sample of CAAN-12-2031

SAMPLE COMMENTS:

OT from TA-54 Building 54 basement lab 18.2 MD/an

LOCATION DESC:

FIELD SCREENING/MEASUREMENT RESULTS:

see CAAN-12-2031

COLLECTED BY (PRINT)

W. J. Shaw

REVIEWED BY (PRINT)

D. Woody

RELINQUISHED BY (Printed Name) D Woody (Signature) D Woody	Date/Time 02/01/12 1415	RECEIVED BY (Printed Name) Sheri Sherwood (Signature) Sheri Sherwood	Date/Time 02/01/12 1415
RELINQUISHED BY (Printed Name) (Signature)	Date/Time	RECEIVED BY (Printed Name) (Signature)	Date/Time

DATA VALIDATION COVER SHEET**5114-1****Data Validation Cover Sheet**

Records Use only

**Section I.**REQUEST NUMBER: 12-700 VALIDATION DATE: 3/6/12 LAB CODE: GELCONTRACT LABORATORY NAME: GEL Laboratories LLCVALIDATOR: Larry Fukui ORGANIZATION: Analytical Quality Associates, Inc.

ANALYTICAL SUITE (CHECK ALL THAT APPLY):

- | | | | |
|--|--|---|---|
| <input type="checkbox"/> TPH-GRO | <input type="checkbox"/> HIGH EXPLOSIVES | <input type="checkbox"/> DIOXIN FURANS | <input type="checkbox"/> LCMSMS PERCHLORATES |
| <input type="checkbox"/> TPH-DRO | <input type="checkbox"/> METALS | <input type="checkbox"/> PCB CONGENERS | <input type="checkbox"/> ORGANOCHLORINE |
| <input type="checkbox"/> GENERAL CHEMISTRY | <input type="checkbox"/> RADIOCHEMISTRY | <input type="checkbox"/> LCMSMS HIGH EXPLOSIVES | <input type="checkbox"/> PESTICIDES/POLYCHLORINATED BIPHENYLS |
- ☒ OTHER (DESCRIBE): GC/MS VOC

Section II. Completeness Check

- | YES | NO | N/A | (CHECK ONE) | YES | NO | N/A | (CHECK ONE) |
|-------------------------------------|--------------------------|-------------------------------------|-----------------------------|-------------------------------------|--------------------------|--------------------------|--------------------------|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 1. CHAIN-OF-CUSTODY FORM(S) | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 6. RAW/BSS DATA |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 2. CASE NARRATIVE | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 7. QUALITY CONTROL FORMS |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 3. SAMPLE RESULT FORMS | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 8. QUANTITATION REPORTS |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 4. SAMPLE CHROMATOGRAMS | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 9. TICS FORMS |
| <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 5. STANDARD CHROMATOGRAMS | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 10. TICS MASS SPECTRA |

Comments/problems noted (include information about requests for further information submitted to the contract laboratory and agreed-upon date of resolution and contract laboratory point of contact):

- The ICAL %RSDs were >15% for bromoform and 1,2-dibromo-3-chloropropane. The associated sample results were NDs and, thus, were qualified UJ,V7a.
- The ICAL and/or ICV/CCV RRFs were <0.05 but ≥0.01 for acetonitrile; n-butyl alcohol; propionitrile; and isobutyl alcohol. The associated sample results were NDs and, thus, were qualified UJ,V7b.
- The ICV and/or CCV %Ds were >20% for dichlorodifluoromethane; carbon disulfide; acrolein; trichlorotrifluoroethane; and 2-chloro-1,3-butadiene. The associated sample results were NDs and, thus, were qualified UJ,V7c.
- It should be noted that ten target analytes were not present in the MS/MSD spiking solution. Since MS/MSD analyses were not required, no sample results were qualified.

Reviewed by: Eric T. MinkLevel: 1Date: 3/7/12

VALIDATOR'S SIGNATURE: _____

DATE: 3/6/12

VOLATILE ORGANIC COMPOUND (VOC) ANALYTICAL DATA VALIDATION CHECKLIST

5114-2

Volatile Organic Compound (VOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	1. The holding time was >1 and ≤2 times the applicable holding time requirement.	UJ, V9	J-, V9
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	2. The holding time was >2 times the applicable holding time requirement.	R, V9a	J-, V9a
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	3. The instrument performance sample did not pass method acceptance criteria.	R, V16	R, V16
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	4. Samples were analyzed outside specific method tune time criteria.	N/A	J, V16b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	5. The required instrument performance sample information is missing. Contact the SMO or external laboratory for information.	R, V16c	R, V16c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	6. The affected results were not analyzed with a valid 5-point calibration curve and/or a standard at the reporting limit.	UJ or R, V7	J, V7
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	7. The affected analytes were analyzed with an initial calibration curve that exceeded the %RSD criteria and/or the associated multipoint calibration correlation coefficient is <0.995.	UJ, V7a	J, V7a
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	8. The affected analytes were analyzed with an RRF of <0.05 in the initial calibration and/or CCV.	R, V7b	J, V7b
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	9. The Initial Calibration Verification (ICV) and/or Continuing Calibration Verification (CCV) were recovered outside the method-specific limits.	UJ, V7c	J, V7c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	10. The ICV and/or CCV were not analyzed at the appropriate method frequency.	UJ, V7d	J, V7d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	11. Required calibration information is missing or samples were analyzed on an expired calibration. Contact the SMO or external laboratory for information.	R, V7f	R, V7f

VOLATILE ORGANIC COMPOUND (VOC) ANALYTICAL DATA VALIDATION CHECKLIST

5114-2

Volatile Organic Compound (VOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A (Check One)				Assign Qualifier Listed Below If Criterion = Yes	
				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	12. The sample result is $\leq 5X$ (10X for common organic laboratory contaminants) the concentration of the related analyte in the method blank.	U, V4	N/A
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	13. The affected analytes are considered estimated and biased high because this analyte was identified in the method blank but was $> 5X$ (10X for common laboratory contaminants).	N/A	J, V4a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	14. The sample result is $\leq 5X$ the concentration of the related analyte in the trip blank, rinsate blank, and/or equipment blank.	U, V4d	N/A
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	15. Required method blank information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, V4e	R, V4e
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	16. The IS retention time has shifted by more than 30 seconds.	UJ, V0	J, V0
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	17. Analyte is positively confirmed but outside the IS retention time window; however, spectral matches must be provided.	N/A	J, V0a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	18. Required IS retention time documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, V0b	R, V0b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	19. The quantitating IS are count is $< 10\%$ of the expected value, which indicates increased potential for false negative results and other possible problems with sample quantitation. Follow method-specific windows.	R, V1a	J, V1a
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	20. The IS area count for the quantitating IS is $< 50\%$ but $> 10\%$ for organics window relation to the previous continuing calibration. Follow the method-specific windows.	UJ, V1b	J, V1b

VOLATILE ORGANIC COMPOUND (VOC) ANALYTICAL DATA VALIDATION CHECKLIST

5114-2

Volatile Organic Compound (VOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	21. The IS area count for the quantitating IS is >200% of the area count for the previous organic continuing calibration. Follow the method-specific windows.	UJ, V1c	J, V1c
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	22. Required IS information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, V1d	R, V1d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	23. The surrogate is <10%R. Follow the external laboratory limits located within the associated data package.	R, V3	J-, V3
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	24. The surrogate is < the Lower Acceptance Limit (LAL) but ≥10%R. Follow the external laboratory limits located within the associated data package.	UJ, V3a	J-, V3a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	25. The surrogate %R is > the Upper Acceptance Limit (UAL) Follow the external laboratory limits located within the associated data package.	N/A	J+, V3b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	26. At least one surrogate is > the UAL and one surrogate is < the LAL. Follow the external laboratory limits located within the associated data package.	UJ, V3c	J, V3c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	27. Required surrogate information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, V3d	R, V3d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	28. The LCS percent recovery was <10%. Follow the external laboratory limits located within the associated data package.	R, V12	J-, V12
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	29. The LCS percent recovery was < the LAL but > 10%. Follow the external laboratory limits located within the associated data package.	UV, V12a	J-, V12a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	30. The LCS percent recover was > the UAL. Follow the external laboratory limits located within the associated data package.	N/A	J+, V12b

VOLATILE ORGANIC COMPOUND (VOC) ANALYTICAL DATA VALIDATION CHECKLIST

5114-2

Volatile Organic Compound (VOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A (Check One)				Assign Qualifier Listed Below If Criterion = Yes	
				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	31. The LCS documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, V12c	R, V12c
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	32. The affected analyte is considered not detected because mass spectrum did not meet specifications.	N/A	U, V8
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	33. The mass spectrum column documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, V8a	R, V8a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	34. Duplicate, dilution, or reanalysis.	UJ, V88	J, V88
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	35. The affected analytes have elevated detection limits and may not meet project DQOs because the sample was diluted without any target analytes identified due to matrix interference. Qualify as Reject if the analytical laboratory cannot provide proof for matrix interference.	UJ, R, V15	R, V15
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	36. Qualification of data via data validation did not occur based on Quality Control requirements in this procedure. Adhere to the external laboratory qualifiers found within the Form I analytical data summary sheets generated by the external laboratory.	U, U_LAB	J, J_LAB, NQ, NQ
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	37. The LANL project chemist identified quality deficiencies in the reported data that requires further qualification. This code can only be used and/or under advisement by the LANL project chemist.	UJ, R, V19	J, R, V19

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2031

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 15:30

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 15:30

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y711.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00 UJ,V7c
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0 UJ,V7b
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00 UJ,V7c
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0 UJ,V7b
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

LMF
3/6/12

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2031

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 15:30

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 15:30

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y711.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ	
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00	
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00	
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00	
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00	
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00	
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00	
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00	
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00	
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00	
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00	UJ,V7a
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00	
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00	
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00	
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00	
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00	
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00	
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00	
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00	
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00	
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00	UJ,V7a
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00	
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00	
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00	
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00	UJ,V7c
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00	UJ,V7c
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00	
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00	
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00	UJ,V7c
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00	UJ,V7b
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00	
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0	UJ,V7b
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00	

LMF
3/6/12

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	295270002	Date Received:	02/03/2012 08:45		
		Client:	ARSL001	Project:	ESHL00210
Client ID:	CAAN-12-2031	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 15:30	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 15:30				
Data File:	021212V9\9Y711.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	44.6	50.0	ug/L 89.2	(76%-127%)
Bromofluorobenzene	50.8	50.0	ug/L 102	(80%-120%)
Toluene-d8	49.6	50.0	ug/L 99.1	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.33	75	ug/L	0	J
	unknown siloxane	12.41	5.54	ug/L	0	J
	unknown siloxane	14.77	6.68	ug/L	0	J

LMF
3/6/12

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2199

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 15:56

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 15:56

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y712.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00 UJ,V7c
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0 UJ,V7b
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00 UJ,V7c
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0 UJ,V7b
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

LMF
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Volatile
Certificate of Analysis
Sample Summary

Page 2 of 3

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2199

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 15:56

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 15:56

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y712.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ	
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00	
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00	
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00	
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00	
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00	
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00	
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00	
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00	
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00	
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00	UJ,V7a
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00	
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00	
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00	
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00	
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00	
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00	
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00	
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00	
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00	
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00	UJ,V7a
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00	
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00	
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00	
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00	UJ,V7c
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00	UJ,V7c
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00	
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00	
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00	UJ,V7c
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00	UJ,V7b
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00	
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0	UJ,V7b
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00	

LMF
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**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 295270004	Date Received: 02/03/2012 08:45	
	Client: ARSL001	Project: ESHL00210
Client ID: CAAN-12-2199	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 15:56	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 15:56		
Data File: 021212V9\9Y712.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	46.4	50.0	ug/L 92.8	(76%-127%)
Bromofluorobenzene	49.8	50.0	ug/L 99.5	(80%-120%)
Toluene-d8	47.4	50.0	ug/L 94.8	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.33	78.6	ug/L	0	J
	unknown siloxane	14.77	5.03	ug/L	0	J

LMF
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**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270007

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2201

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 16:23

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 16:23

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y713.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00 UJ,V7c
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0 UJ,V7b
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00 UJ,V7c
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0 UJ,V7b
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

**Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700
Lab Sample ID: 295270007

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2201

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 16:23

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 16:23

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y713.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ	
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00	
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00	
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00	
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00	
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00	
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00	
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00	
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00	
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00	
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00	UJ,V7a
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00	
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00	
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00	
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00	
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00	
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00	
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00	
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00	
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00	
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00	UJ,V7a
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00	
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00	
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00	
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00	UJ,V7c
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00	UJ,V7c
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00	
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00	
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00	UJ,V7c
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00	UJ,V7b
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00	
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0	UJ,V7b
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00	

LMF
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**Volatile
Certificate of Analysis
Sample Summary**

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	295270007	Date Received:	02/03/2012 08:45		
		Client:	ARSL001	Project:	ESHL00210
Client ID:	CAAN-12-2201	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 16:23	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 16:23				
Data File:	021212V9\9Y713.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	44.2	50.0	ug/L 88.5	(76%-127%)
Bromofluorobenzene	53.3	50.0	ug/L 107	(80%-120%)
Toluene-d8	45.9	50.0	ug/L 91.7	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.29	37.3	ug/L	0	J
	unknown siloxane	14.77	7.93	ug/L	0	J

LMF
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**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270008

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2032

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 16:49

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 16:49

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y714.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00 UJ,V7c
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0 UJ,V7b
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00 UJ,V7c
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0 UJ,V7b
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

LMF
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Volatile
Certificate of Analysis
Sample Summary

Page 2 of 3

SDG Number: 12-700
Lab Sample ID: 295270008

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2032

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 16:49

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 16:49

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y714.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ	
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00	
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00	
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00	
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00	
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00	
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00	
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00	
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00	
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00	
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00	UJ,V7a
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00	
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00	
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00	
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00	
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00	
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00	
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00	
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00	
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00	
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00	
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00	
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00	
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00	UJ,V7a
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00	
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00	
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00	
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00	UJ,V7c
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00	UJ,V7c
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00	
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00	
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00	UJ,V7c
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00	UJ,V7b
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00	
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0	UJ,V7b
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00	

LMF
3/6/12

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 295270008	Date Received: 02/03/2012 08:45	
Client ID: CAAN-12-2032	Client: ARSL001	Project: ESHL00210
Batch ID: 1187820	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Run Date: 02/12/2012 16:49	Inst: VOA9.I	Dilution: 1
Prep Date: 02/12/2012 16:49	Analyst: GRB2	Purge Vol: 5 mL
Data File: 021212V9\9Y714.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	45.7	50.0	ug/L 91.4	(76%-127%)
Bromofluorobenzene	53.0	50.0	ug/L 106	(80%-120%)
Toluene-d8	49.1	50.0	ug/L 98.2	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.33	60.8	ug/L	0	J
	unknown siloxane	16.74	7.92	ug/L	0	J

LMF
3/6/12

DATA VALIDATION COVER SHEET**5115-1****Data Validation Cover Sheet**

Records Use only

**Section I.**REQUEST NUMBER: 12-700 VALIDATION DATE: 3/6/12 LAB CODE: GELCONTRACT LABORATORY NAME: GEL Laboratories LLCVALIDATOR: Larry Fukui ORGANIZATION: Analytical Quality Associates, Inc.

ANALYTICAL SUITE (CHECK ALL THAT APPLY):

- | | | | |
|--|--|---|---|
| <input type="checkbox"/> TPH-GRO | <input type="checkbox"/> HIGH EXPLOSIVES | <input type="checkbox"/> DIOXIN FURANS | <input type="checkbox"/> LCMSMS PERCHLORATES |
| <input type="checkbox"/> TPH-DRO | <input type="checkbox"/> METALS | <input type="checkbox"/> PCB CONGENERS | <input type="checkbox"/> ORGANOCHLORINE |
| <input type="checkbox"/> GENERAL CHEMISTRY | <input type="checkbox"/> RADIOCHEMISTRY | <input type="checkbox"/> LCMSMS HIGH EXPLOSIVES | <input type="checkbox"/> PESTICIDES/POLYCHLORINATED BIPHENYLS |
- ☒ OTHER (DESCRIBE): GC/MS SVOC

Section II. Completeness Check

- | YES | NO | N/A | (CHECK ONE) | YES | NO | N/A | (CHECK ONE) |
|-------------------------------------|--------------------------|-------------------------------------|-----------------------------|-------------------------------------|--------------------------|-------------------------------------|--------------------------|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 1. CHAIN-OF-CUSTODY FORM(S) | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 6. RAW/BSS DATA |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 2. CASE NARRATIVE | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 7. QUALITY CONTROL FORMS |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 3. SAMPLE RESULT FORMS | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 8. QUANTITATION REPORTS |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 4. SAMPLE CHROMATOGRAMS | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 9. TICS FORMS |
| <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 5. STANDARD CHROMATOGRAMS | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 10. TICS MASS SPECTRA |

Comments/problems noted (include information about requests for further information submitted to the contract laboratory and agreed-upon date of resolution and contract laboratory point of contact):

- The ICAL RRF was <0.05 but ≥ 0.01 for atrazine. The associated sample results were NDs and, thus, were qualified UJ,SV7b.
- The ICV and/or CCV %Ds were $>20\%$ for hexachlorocyclopentadiene; 4-nitrophenol; and 4-nitroaniline. The associated sample results were NDs and, thus, were qualified UJ,SV7c.
- The surrogate %R for 2-fluorophenol was $>$ the laboratory UAL for the MSD. This was a QC sample and, thus, no sample data were qualified as a result.
- The LCS %Rs were $<$ the laboratory LAL but $\geq 10\%$ for benzoic acid and benzidine. The associated sample results were NDs and, thus, were qualified UJ,SV12a.
- It should be noted that dinoseb; n-nitrosodiethylamine; n-nitrosodi-n-butylamine; and pentachlorobenzene were not present in the LCS, the MS, or the MSD. No sample results were qualified as a result.
- The MSD %Rs and MS/MSD RPDs did not meet laboratory acceptance criteria for several compounds. Since MS/MSD analyses were not required for SVOC, no sample data were qualified.

Reviewed by: Eric T. MinkLevel: 1Date: 3/7/12

VALIDATOR'S SIGNATURE: _____

DATE: 3/6/12

DATA VALIDATION COVER SHEET**5115-1****Data Validation Cover Sheet**

Records Use only



Form 5115-1, Revision 0.0

LOS ALAMOS

Environmental Restoration Project

SEMIVOLATILE ORGANIC COMPOUND (SVOC) ANALYTICAL DATA VALIDATION CHECKLIST

5115-2

Semivolatile Organic Compound (SVOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A (Check One)				Assign Qualifier Listed Below If Criterion = Yes	
				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	1. The holding time was >1 and ≤2 times the applicable holding time requirement.	UJ, SV9	J-, SV9
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	2. The holding time was >2 times the applicable holding time requirement.	R, SV9a	J-, SV9a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	3. The affected analytes are regarded as rejected because the analytical holding time was exceeded.	R, SV9b	R, SV9b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	4. The instrument performance sample did not pass method acceptance criteria.	R, SV16	R, SV16
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	5. Samples were analyzed outside specific method tune time criteria.	N/A	J, SV16b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	6. The required instrument performance sample information is missing. Contact the SMO or external laboratory for information.	R, SV16c	R, SV16c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	7. The affected results were not analyzed with a valid 5-point calibration curve and/or a standard at the reporting limit.	UJ, R, SV7	J, SV7
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	8. The affected analytes were analyzed with an initial calibration curve that exceeded the %RSD criteria and/or the associated multipoint calibration correlation coefficient is <0.995.	UJ, SV7a	J, SV7a
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	9. The affected analytes were analyzed with an RRF of <0.05 in the initial calibration and/or Continuing Calibration Verification (CCV).	R, SV7b	J, SV7b
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	10. The Initial Calibration Verification (ICV) and/or CCV were recovered outside the method-specific limits.	UJ, SV7c	J, SV7c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	11. The ICV and/or CCV were not analyzed at the appropriate method frequency.	UJ, SV7d	J, SV7d

SEMIVOLATILE ORGANIC COMPOUND (SVOC) ANALYTICAL DATA VALIDATION CHECKLIST

5115-2

Semivolatile Organic Compound (SVOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	12. Required calibration information is missing or samples were analyzed on an expired calibration. Contact the SMO or external laboratory for information.	R, SV7f	R, SV7f
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	13. The sample result is $\leq 5X$ (10X for common organic laboratory contaminants) the concentration of the related analyte in the method blank.	U, SV4	J, V4a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	14. The affected analytes are considered estimated and biased high because this analyte was identified in the method blank but was greater than 5X (10X for common laboratory contaminants).	N/A	J, SV4a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	15. The sample result is $\leq 5X$ the concentration of the related analyte in the trip blank, rinsate blank, or equipment blank.	UJ, SV4d	N/A
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	16. Required method blank information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, SV4e	R, SV4e
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	17. The IS retention time has shifted by more than 30 seconds.	UJ, SV0	J, SV0
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	18. Analyte is positively confirmed but outside the IS retention time window; however, spectral matches must be provided.	N/A	J, SV0a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	19. Required IS retention time documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, SV0b	R, SV0b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	20. The quantitating IS area count is $<10\%$ of the expected value, which indicates increased potential for false negative results and other possible problems with sample quantitation. Follow method-specific windows.	R, SV1a	J, SV1a

SEMIVOLATILE ORGANIC COMPOUND (SVOC) ANALYTICAL DATA VALIDATION CHECKLIST

5115-2

Semivolatile Organic Compound (SVOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A (Check One)				Assign Qualifier Listed Below If Criterion = Yes	
				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	21. The IS area count for the quantitating IS is <50% but >10% for organics window relation to the previous continuing calibration. Follow method-specific windows.	UJ, SV1b	J, SV1b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	22. The IS area count for the quantitating IS is >200% of the area count for the previous continuing calibration. Follow method-specific windows.	UJ, SV1c	J, SV1c
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	23. Required IS information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, SV1d	R, SV1d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	24. The surrogate is <10%R. Follow the external laboratory limits located within the associated data package.	R, SV3	J-, SV3
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	25. The surrogate is < the Lower Acceptance Level (LAL) but ≥10%R. Follow the external laboratory limits located within the associated data package.	UJ, SV3a	J-, SV3a
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	26. The surrogate %R value is > the UAL. Follow the external laboratory limits located within the associated data package.	N/A	J+, SV3b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	27. At least one surrogate is > the Upper Acceptance Limit (UAL) and one surrogate is < the LAL. Follow the external laboratory limits located within the associated data package.	UJ, SV3c	J, SV3c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	28. Required surrogate information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, SV3d	R, SV3d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	29. The LCS percent recovery was <10%. Follow the external laboratory limits located within the associated data package.	R, SV12	J-, SV12

SEMIVOLATILE ORGANIC COMPOUND (SVOC) ANALYTICAL DATA VALIDATION CHECKLIST

5115-2

Semivolatile Organic Compound (SVOC) Analytical Data Validation Checklist

Records Use only



Yes No N/A (Check One)				Assign Qualifier Listed Below If Criterion = Yes	
				Non-detected Analyte	Detected Analyte
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	30. The LCS percent recovery was < the LAL but >10%. Follow the external laboratory limits located within the associated data package.	UJ, SV12a	J-, SV12a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	31. The LCS percent recovery was > the UAL. Follow the external laboratory limits located within the associated data package.	N/A	J+, SV12b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	32. The LCS documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, SV12c	R, SV12c
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	33. The affected analyte is considered not detected because mass spectrum did not meet specifications.	N/A	U, SV8
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	34. The mass spectrum column documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, SV8a	R, SV8a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	35. Duplicate, dilution, or reanalysis.	UJ, SV88	J, SV88
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	36. The affected analytes have elevated detection limits and may not meet project DQOs because the sample was diluted without any target analytes identified due to matrix interference. Qualify as Reject if the analytical laboratory cannot provide proof for matrix interference.	UJ, R, SV15	R, SV15
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	37. Qualification of data via data validation did not occur based on Quality Control requirements in this procedure. Adhere to the external laboratory qualifiers found within the Form I analytical data summary sheets generated by the external laboratory.	U, U_LAB	J, J_LAB, NQ, NQ
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	38. The LANL project chemist identified quality deficiencies in the reported data that requires further qualification. This code can only be used and/or under advisement by the LANL project chemist.	UJ, R, SV19	J, R, SV19

**Semi-Volatile
Certificate of Analysis
Sample Summary**

Page 1 of 3

SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 1000 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2031
Batch ID: 1186837
Run Date: 02/09/2012 10:28
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0907.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether	U	10.0	ug/L	3.00	10.0
62-75-9	N-Methyl-N-nitrosomethylamine	U	10.0	ug/L	3.00	10.0
110-86-1	Pyridine	U	10.0	ug/L	3.00	10.0
62-53-3	Aniline	U	10.0	ug/L	3.00	10.0
108-95-2	Phenol	U	10.0	ug/L	3.00	10.0
111-44-4	bis(2-Chloroethyl) ether	U	10.0	ug/L	3.00	10.0
95-57-8	2-Chlorophenol	U	10.0	ug/L	3.00	10.0
541-73-1	1,3-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
106-46-7	1,4-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
95-50-1	1,2-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
100-51-6	Benzyl alcohol	U	10.0	ug/L	3.00	10.0
95-48-7	o-Cresol	U	10.0	ug/L	3.00	10.0
65794-96-9	m,p-Cresols	U	10.0	ug/L	3.00	10.0
621-64-7	N-Nitrosodi--n-propylamine	U	10.0	ug/L	3.00	10.0
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane	U	10.0	ug/L	3.00	10.0
98-95-3	Nitrobenzene	U	10.0	ug/L	3.00	10.0
78-59-1	Isophorone	U	10.0	ug/L	3.00	10.0
88-75-5	2-Nitrophenol	U	10.0	ug/L	3.00	10.0
105-67-9	2,4-Dimethylphenol	U	10.0	ug/L	3.00	10.0
111-91-1	bis(2-Chloroethoxy)methane	U	10.0	ug/L	3.00	10.0
120-83-2	2,4-Dichlorophenol	U	10.0	ug/L	3.00	10.0
65-85-0	Benzoic acid	U	20.0	ug/L	6.00	20.0 UJ,SV12a
106-47-8	4-Chloroaniline	U	10.0	ug/L	3.00	10.0
87-68-3	Hexachlorobutadiene	U	10.0	ug/L	3.00	10.0
59-50-7	Parachlorometa cresol	U	10.0	ug/L	3.00	10.0
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.300	1.00
90-12-0	1-Methylnaphthalene	U	1.00	ug/L	0.300	1.00
77-47-4	Hexachlorocyclopentadiene	U	10.0	ug/L	3.00	10.0 UJ,SV7c
88-06-2	2,4,6-Trichlorophenol	U	10.0	ug/L	3.00	10.0
95-95-4	2,4,5-Trichlorophenol	U	10.0	ug/L	3.00	10.0
91-58-7	2-Chloronaphthalene	U	1.00	ug/L	0.300	1.00
88-74-4	2-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate	U	10.0	ug/L	3.00	10.0
606-20-2	2,6-Dinitrotoluene	U	10.0	ug/L	3.00	10.0

LMF
3/6/12

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 1000 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2031
Batch ID: 1186837
Run Date: 02/09/2012 10:28
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0907.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene	U	10.0	ug/L	3.00	10.0
208-96-8	Acenaphthylene	U	1.00	ug/L	0.300	1.00
83-32-9	Acenaphthene	U	1.00	ug/L	0.300	1.00
51-28-5	2,4-Dinitrophenol	U	20.0	ug/L	5.00	20.0
132-64-9	Dibenzofuran	U	10.0	ug/L	3.00	10.0
58-90-2	2,3,4,6-Tetrachlorophenol	U	10.0	ug/L	3.00	10.0
84-66-2	Diethylphthalate	U	10.0	ug/L	3.00	10.0
100-02-7	4-Nitrophenol	U	10.0	ug/L	3.00	10.0 UJ,SV7c
86-73-7	Fluorene	U	1.00	ug/L	0.300	1.00
7005-72-3	4-Chlorophenylphenylether	U	10.0	ug/L	3.00	10.0
100-01-6	4-Nitroaniline	U	10.0	ug/L	3.00	10.0 UJ,SV7c
	<i>p-Nitroaniline</i>					
534-52-1	2-Methyl-4,6-dinitrophenol	U	10.0	ug/L	3.00	10.0
122-39-4	Diphenylamine	U	10.0	ug/L	3.00	10.0
122-66-7	Azobenzene	U	10.0	ug/L	3.00	10.0
	<i>1,2-Diphenylhydrazine</i>					
101-55-3	4-Bromophenylphenylether	U	10.0	ug/L	3.00	10.0
118-74-1	Hexachlorobenzene	U	10.0	ug/L	3.00	10.0
87-86-5	Pentachlorophenol	U	10.0	ug/L	3.00	10.0
88-85-7	Dinoseb	U	10.0	ug/L	3.00	10.0
85-01-8	Phenanthrene	U	1.00	ug/L	0.300	1.00
120-12-7	Anthracene	U	1.00	ug/L	0.300	1.00
84-74-2	Di-n-butylphthalate	U	10.0	ug/L	3.00	10.0
206-44-0	Fluoranthene	U	1.00	ug/L	0.300	1.00
129-00-0	Pyrene	U	1.00	ug/L	0.300	1.00
85-68-7	Butylbenzylphthalate	U	10.0	ug/L	3.00	10.0
117-81-7	bis(2-Ethylhexyl)phthalate	U	10.0	ug/L	3.00	10.0
56-55-3	Benzo(a)anthracene	U	1.00	ug/L	0.300	1.00
218-01-9	Chrysene	U	1.00	ug/L	0.300	1.00
117-84-0	Di-n-octylphthalate	U	10.0	ug/L	3.00	10.0
205-99-2	Benzo(b)fluoranthene	U	1.00	ug/L	0.300	1.00
207-08-9	Benzo(k)fluoranthene	U	1.00	ug/L	0.300	1.00
50-32-8	Benzo(a)pyrene	U	1.00	ug/L	0.300	1.00
193-39-5	Indeno(1,2,3-cd)pyrene	U	1.00	ug/L	0.300	1.00
53-70-3	Dibenzo(a,h)anthracene	U	1.00	ug/L	0.300	1.00
191-24-2	Benzo(ghi)perylene	U	1.00	ug/L	0.300	1.00
123-91-1	1,4-Dioxane	U	10.0	ug/L	3.00	10.0
55-18-5	N-Nitrosodiethylamine	U	10.0	ug/L	3.00	10.0
930-55-2	N-Nitrosopyrrolidine	U	10.0	ug/L	3.00	10.0

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**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270002

Client ID: CAAN-12-2031
Batch ID: 1186837
Run Date: 02/09/2012 10:28
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0907.D

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 1000 mL
Column: DB-5ms

Matrix: WG

Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.0	ug/L	3.00	10.0
95-94-3	1,2,4,5-Tetrachlorobenzene	U	10.0	ug/L	3.00	10.0
608-93-5	Pentachlorobenzene	U	10.0	ug/L	3.00	10.0
1912-24-9	Atrazine	U	10.0	ug/L	3.00	10.0 UJ,SV7b
92-87-5	Benzidine	U	10.0	ug/L	3.00	10.0 UJ,SV12a
91-94-1	3,3'-Dichlorobenzidine	U	10.0	ug/L	3.00	10.0
120-82-1	1,2,4-Trichlorobenzene	U	10.0	ug/L	3.00	10.0

Surrogate/Tracer recovery	Result	Nominal		Recovery%	Acceptable Limits
2,4,6-Tribromophenol	75.0	100	ug/L	75.0	(30%-128%)
2-Fluorobiphenyl	34.0	50.0	ug/L	67.9	(34%-98%)
2-Fluorophenol	41.1	100	ug/L	41.1	(21%-78%)
Nitrobenzene-d5	37.3	50.0	ug/L	74.6	(39%-117%)
Phenol-d5	26.3	100	ug/L	26.3	(14%-80%)
p-Terphenyl-d14	37.3	50.0	ug/L	74.6	(39%-129%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	No Tentatively Identified Compounds Found			ug/L		

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2199
Batch ID: 1186837
Run Date: 02/09/2012 11:43
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0910.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether	U	10.5	ug/L	3.16	10.5
62-75-9	N-Methyl-N-nitrosomethylamine	U	10.5	ug/L	3.16	10.5
110-86-1	Pyridine	U	10.5	ug/L	3.16	10.5
62-53-3	Aniline	U	10.5	ug/L	3.16	10.5
108-95-2	Phenol	U	10.5	ug/L	3.16	10.5
111-44-4	bis(2-Chloroethyl) ether	U	10.5	ug/L	3.16	10.5
95-57-8	2-Chlorophenol	U	10.5	ug/L	3.16	10.5
541-73-1	1,3-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
106-46-7	1,4-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
95-50-1	1,2-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
100-51-6	Benzyl alcohol	U	10.5	ug/L	3.16	10.5
95-48-7	o-Cresol	U	10.5	ug/L	3.16	10.5
65794-96-9	m,p-Cresols	U	10.5	ug/L	3.16	10.5
621-64-7	N-Nitrosodi--n-propylamine	U	10.5	ug/L	3.16	10.5
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane	U	10.5	ug/L	3.16	10.5
98-95-3	Nitrobenzene	U	10.5	ug/L	3.16	10.5
78-59-1	Isophorone	U	10.5	ug/L	3.16	10.5
88-75-5	2-Nitrophenol	U	10.5	ug/L	3.16	10.5
105-67-9	2,4-Dimethylphenol	U	10.5	ug/L	3.16	10.5
111-91-1	bis(2-Chloroethoxy)methane	U	10.5	ug/L	3.16	10.5
120-83-2	2,4-Dichlorophenol	U	10.5	ug/L	3.16	10.5
65-85-0	Benzoic acid	U	21.1	ug/L	6.32	21.1 UJ,SV12a
106-47-8	4-Chloroaniline	U	10.5	ug/L	3.16	10.5
87-68-3	Hexachlorobutadiene	U	10.5	ug/L	3.16	10.5
59-50-7	Parachlorometa cresol	U	10.5	ug/L	3.16	10.5
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
91-20-3	Naphthalene	U	1.05	ug/L	0.316	1.05
90-12-0	1-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
77-47-4	Hexachlorocyclopentadiene	U	10.5	ug/L	3.16	10.5 UJ,SV7c
88-06-2	2,4,6-Trichlorophenol	U	10.5	ug/L	3.16	10.5
95-95-4	2,4,5-Trichlorophenol	U	10.5	ug/L	3.16	10.5
91-58-7	2-Chloronaphthalene	U	1.05	ug/L	0.316	1.05
88-74-4	2-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate	U	10.5	ug/L	3.16	10.5
606-20-2	2,6-Dinitrotoluene	U	10.5	ug/L	3.16	10.5

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**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2199
Batch ID: 1186837
Run Date: 02/09/2012 11:43
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0910.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene	U	10.5	ug/L	3.16	10.5
208-96-8	Acenaphthylene	U	1.05	ug/L	0.316	1.05
83-32-9	Acenaphthene	U	1.05	ug/L	0.316	1.05
51-28-5	2,4-Dinitrophenol	U	21.1	ug/L	5.26	21.1
132-64-9	Dibenzofuran	U	10.5	ug/L	3.16	10.5
58-90-2	2,3,4,6-Tetrachlorophenol	U	10.5	ug/L	3.16	10.5
84-66-2	Diethylphthalate	U	10.5	ug/L	3.16	10.5
100-02-7	4-Nitrophenol	U	10.5	ug/L	3.16	10.5 UJ,SV7c
86-73-7	Fluorene	U	1.05	ug/L	0.316	1.05
7005-72-3	4-Chlorophenylphenylether	U	10.5	ug/L	3.16	10.5
100-01-6	4-Nitroaniline	U	10.5	ug/L	3.16	10.5 UJ,SV7c
	<i>p-Nitroaniline</i>					
534-52-1	2-Methyl-4,6-dinitrophenol	U	10.5	ug/L	3.16	10.5
122-39-4	Diphenylamine	U	10.5	ug/L	3.16	10.5
122-66-7	Azobenzene	U	10.5	ug/L	3.16	10.5
	<i>1,2-Diphenylhydrazine</i>					
101-55-3	4-Bromophenylphenylether	U	10.5	ug/L	3.16	10.5
118-74-1	Hexachlorobenzene	U	10.5	ug/L	3.16	10.5
87-86-5	Pentachlorophenol	U	10.5	ug/L	3.16	10.5
88-85-7	Dinoseb	U	10.5	ug/L	3.16	10.5
85-01-8	Phenanthrene	U	1.05	ug/L	0.316	1.05
120-12-7	Anthracene	U	1.05	ug/L	0.316	1.05
84-74-2	Di-n-butylphthalate	U	10.5	ug/L	3.16	10.5
206-44-0	Fluoranthene	U	1.05	ug/L	0.316	1.05
129-00-0	Pyrene	U	1.05	ug/L	0.316	1.05
85-68-7	Butylbenzylphthalate	U	10.5	ug/L	3.16	10.5
117-81-7	bis(2-Ethylhexyl)phthalate	U	10.5	ug/L	3.16	10.5
56-55-3	Benzo(a)anthracene	U	1.05	ug/L	0.316	1.05
218-01-9	Chrysene	U	1.05	ug/L	0.316	1.05
117-84-0	Di-n-octylphthalate	U	10.5	ug/L	3.16	10.5
205-99-2	Benzo(b)fluoranthene	U	1.05	ug/L	0.316	1.05
207-08-9	Benzo(k)fluoranthene	U	1.05	ug/L	0.316	1.05
50-32-8	Benzo(a)pyrene	U	1.05	ug/L	0.316	1.05
193-39-5	Indeno(1,2,3-cd)pyrene	U	1.05	ug/L	0.316	1.05
53-70-3	Dibenzo(a,h)anthracene	U	1.05	ug/L	0.316	1.05
191-24-2	Benzo(ghi)perylene	U	1.05	ug/L	0.316	1.05
123-91-1	1,4-Dioxane	U	10.5	ug/L	3.16	10.5
55-18-5	N-Nitrosodiethylamine	U	10.5	ug/L	3.16	10.5
930-55-2	N-Nitrosopyrrolidine	U	10.5	ug/L	3.16	10.5

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**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2199
Batch ID: 1186837
Run Date: 02/09/2012 11:43
Prep Date: 02/08/2012 17:46
Data File: S020912.B\3b0910.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.5	ug/L	3.16	10.5
95-94-3	1,2,4,5-Tetrachlorobenzene	U	10.5	ug/L	3.16	10.5
608-93-5	Pentachlorobenzene	U	10.5	ug/L	3.16	10.5
1912-24-9	Atrazine	U	10.5	ug/L	3.16	10.5 UJ,SV7b
92-87-5	Benzidine	U	10.5	ug/L	3.16	10.5 UJ,SV12a
91-94-1	3,3'-Dichlorobenzidine	U	10.5	ug/L	3.16	10.5
120-82-1	1,2,4-Trichlorobenzene	U	10.5	ug/L	3.16	10.5

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
2,4,6-Tribromophenol	87.6	105	ug/L 83.3	(30%-128%)
2-Fluorobiphenyl	40.4	52.6	ug/L 76.8	(34%-98%)
2-Fluorophenol	58.3	105	ug/L 55.4	(21%-78%)
Nitrobenzene-d5	48.1	52.6	ug/L 91.5	(39%-117%)
Phenol-d5	37.2	105	ug/L 35.4	(14%-80%)
p-Terphenyl-d14	41.7	52.6	ug/L 79.2	(39%-129%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
No Tentatively Identified Compounds Found				ug/L		

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**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700
Lab Sample ID: 295270007

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2201
Batch ID: 1186837
Run Date: 02/09/2012 12:08
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0911.D

Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether	U	10.5	ug/L	3.16	10.5
62-75-9	N-Methyl-N-nitrosomethylamine	U	10.5	ug/L	3.16	10.5
110-86-1	Pyridine	U	10.5	ug/L	3.16	10.5
62-53-3	Aniline	U	10.5	ug/L	3.16	10.5
108-95-2	Phenol	U	10.5	ug/L	3.16	10.5
111-44-4	bis(2-Chloroethyl) ether	U	10.5	ug/L	3.16	10.5
95-57-8	2-Chlorophenol	U	10.5	ug/L	3.16	10.5
541-73-1	1,3-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
106-46-7	1,4-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
95-50-1	1,2-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
100-51-6	Benzyl alcohol	U	10.5	ug/L	3.16	10.5
95-48-7	o-Cresol	U	10.5	ug/L	3.16	10.5
65794-96-9	m,p-Cresols	U	10.5	ug/L	3.16	10.5
621-64-7	N-Nitrosodi--n-propylamine	U	10.5	ug/L	3.16	10.5
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane	U	10.5	ug/L	3.16	10.5
98-95-3	Nitrobenzene	U	10.5	ug/L	3.16	10.5
78-59-1	Isophorone	U	10.5	ug/L	3.16	10.5
88-75-5	2-Nitrophenol	U	10.5	ug/L	3.16	10.5
105-67-9	2,4-Dimethylphenol	U	10.5	ug/L	3.16	10.5
111-91-1	bis(2-Chloroethoxy)methane	U	10.5	ug/L	3.16	10.5
120-83-2	2,4-Dichlorophenol	U	10.5	ug/L	3.16	10.5
65-85-0	Benzoic acid	U	21.1	ug/L	6.32	21.1 UJ,SV12a
106-47-8	4-Chloroaniline	U	10.5	ug/L	3.16	10.5
87-68-3	Hexachlorobutadiene	U	10.5	ug/L	3.16	10.5
59-50-7	Parachlorometa cresol	U	10.5	ug/L	3.16	10.5
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
91-20-3	Naphthalene	U	1.05	ug/L	0.316	1.05
90-12-0	1-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
77-47-4	Hexachlorocyclopentadiene	U	10.5	ug/L	3.16	10.5 UJ,SV7c
88-06-2	2,4,6-Trichlorophenol	U	10.5	ug/L	3.16	10.5
95-95-4	2,4,5-Trichlorophenol	U	10.5	ug/L	3.16	10.5
91-58-7	2-Chloronaphthalene	U	1.05	ug/L	0.316	1.05
88-74-4	2-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate	U	10.5	ug/L	3.16	10.5
606-20-2	2,6-Dinitrotoluene	U	10.5	ug/L	3.16	10.5

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**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270007

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2201
Batch ID: 1186837
Run Date: 02/09/2012 12:08
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0911.D

Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ	
121-14-2	2,4-Dinitrotoluene	U	10.5	ug/L	3.16	10.5	
208-96-8	Acenaphthylene	U	1.05	ug/L	0.316	1.05	
83-32-9	Acenaphthene	U	1.05	ug/L	0.316	1.05	
51-28-5	2,4-Dinitrophenol	U	21.1	ug/L	5.26	21.1	
132-64-9	Dibenzofuran	U	10.5	ug/L	3.16	10.5	
58-90-2	2,3,4,6-Tetrachlorophenol	U	10.5	ug/L	3.16	10.5	
84-66-2	Diethylphthalate	U	10.5	ug/L	3.16	10.5	
100-02-7	4-Nitrophenol	U	10.5	ug/L	3.16	10.5	UJ,SV7c
86-73-7	Fluorene	U	1.05	ug/L	0.316	1.05	
7005-72-3	4-Chlorophenylphenylether	U	10.5	ug/L	3.16	10.5	
100-01-6	4-Nitroaniline	U	10.5	ug/L	3.16	10.5	UJ,SV7c
	<i>p</i> -Nitroaniline						
534-52-1	2-Methyl-4,6-dinitrophenol	U	10.5	ug/L	3.16	10.5	
122-39-4	Diphenylamine	U	10.5	ug/L	3.16	10.5	
122-66-7	Azobenzene	U	10.5	ug/L	3.16	10.5	
	<i>1,2</i> -Diphenylhydrazine						
101-55-3	4-Bromophenylphenylether	U	10.5	ug/L	3.16	10.5	
118-74-1	Hexachlorobenzene	U	10.5	ug/L	3.16	10.5	
87-86-5	Pentachlorophenol	U	10.5	ug/L	3.16	10.5	
88-85-7	Dinoseb	U	10.5	ug/L	3.16	10.5	
85-01-8	Phenanthrene	U	1.05	ug/L	0.316	1.05	
120-12-7	Anthracene	U	1.05	ug/L	0.316	1.05	
84-74-2	Di-n-butylphthalate	U	10.5	ug/L	3.16	10.5	
206-44-0	Fluoranthene	U	1.05	ug/L	0.316	1.05	
129-00-0	Pyrene	U	1.05	ug/L	0.316	1.05	
85-68-7	Butylbenzylphthalate	U	10.5	ug/L	3.16	10.5	
117-81-7	bis(2-Ethylhexyl)phthalate	U	10.5	ug/L	3.16	10.5	
56-55-3	Benzo(a)anthracene	U	1.05	ug/L	0.316	1.05	
218-01-9	Chrysene	U	1.05	ug/L	0.316	1.05	
117-84-0	Di-n-octylphthalate	U	10.5	ug/L	3.16	10.5	
205-99-2	Benzo(b)fluoranthene	U	1.05	ug/L	0.316	1.05	
207-08-9	Benzo(k)fluoranthene	U	1.05	ug/L	0.316	1.05	
50-32-8	Benzo(a)pyrene	U	1.05	ug/L	0.316	1.05	
193-39-5	Indeno(1,2,3-cd)pyrene	U	1.05	ug/L	0.316	1.05	
53-70-3	Dibenzo(a,h)anthracene	U	1.05	ug/L	0.316	1.05	
191-24-2	Benzo(ghi)perylene	U	1.05	ug/L	0.316	1.05	
123-91-1	1,4-Dioxane	U	10.5	ug/L	3.16	10.5	
55-18-5	N-Nitrosodiethylamine	U	10.5	ug/L	3.16	10.5	
930-55-2	N-Nitrosopyrrolidine	U	10.5	ug/L	3.16	10.5	

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**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270007

Client ID: CAAN-12-2201
Batch ID: 1186837
Run Date: 02/09/2012 12:08
Prep Date: 02/08/2012 17:46
Data File: S020912.B\3b0911.D

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Matrix: WG

Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.5	ug/L	3.16	10.5
95-94-3	1,2,4,5-Tetrachlorobenzene	U	10.5	ug/L	3.16	10.5
608-93-5	Pentachlorobenzene	U	10.5	ug/L	3.16	10.5
1912-24-9	Atrazine	U	10.5	ug/L	3.16	10.5 UJ,SV7b
92-87-5	Benzidine	U	10.5	ug/L	3.16	10.5 UJ,SV12a
91-94-1	3,3'-Dichlorobenzidine	U	10.5	ug/L	3.16	10.5
120-82-1	1,2,4-Trichlorobenzene	U	10.5	ug/L	3.16	10.5

Surrogate/Tracer recovery	Result	Nominal		Recovery%	Acceptable Limits
2,4,6-Tribromophenol	74.3	105	ug/L	70.6	(30%-128%)
2-Fluorobiphenyl	38.9	52.6	ug/L	73.9	(34%-98%)
2-Fluorophenol	50.4	105	ug/L	47.9	(21%-78%)
Nitrobenzene-d5	44.8	52.6	ug/L	85.2	(39%-117%)
Phenol-d5	31.2	105	ug/L	29.7	(14%-80%)
p-Terphenyl-d14	42.5	52.6	ug/L	80.8	(39%-129%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	No Tentatively Identified Compounds Found			ug/L		

LMF
3/6/12

DATA VALIDATION COVER SHEET**5122-1****Data Validation Cover Sheet**

Records Use only

**Section I.**REQUEST NUMBER: 12-700 VALIDATION DATE: 3/6/12 LAB CODE: GELCONTRACT LABORATORY NAME: GEL Laboratories LLCVALIDATOR: Larry Fukui ORGANIZATION: Analytical Quality Associates, Inc.

ANALYTICAL SUITE (CHECK ALL THAT APPLY):

- | | | | |
|--|--|--|--|
| <input type="checkbox"/> TPH-GRO | <input type="checkbox"/> HIGH EXPLOSIVES | <input type="checkbox"/> DIOXIN FURANS | <input type="checkbox"/> LCMSMS PERCHLORATES |
| <input type="checkbox"/> TPH-DRO | <input type="checkbox"/> METALS | <input type="checkbox"/> PCB CONGENERS | <input type="checkbox"/> ORGANOCHLORINE |
| <input type="checkbox"/> GENERAL CHEMISTRY | <input type="checkbox"/> RADIOCHEMISTRY | <input checked="" type="checkbox"/> LCMSMS HIGH EXPLOSIVES | PESTICIDES/POLYCHLORINATED BIPHENYLS |
| <input type="checkbox"/> OTHER (DESCRIBE): _____ | | | |

Section II. Completeness Check

- | YES | NO | N/A | (CHECK ONE) | YES | NO | N/A | (CHECK ONE) |
|-------------------------------------|--------------------------|-------------------------------------|-----------------------------|-------------------------------------|--------------------------|-------------------------------------|--------------------------|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 1. CHAIN-OF-CUSTODY FORM(S) | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 6. RAW/BSS DATA |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 2. CASE NARRATIVE | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 7. QUALITY CONTROL FORMS |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 3. SAMPLE RESULT FORMS | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 8. QUANTITATION REPORTS |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 4. SAMPLE CHROMATOGRAMS | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 9. TICS FORMS |
| <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 5. STANDARD CHROMATOGRAMS | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 10. TICS MASS SPECTRA |

Comments/problems noted (include information about requests for further information submitted to the contract laboratory and agreed-upon date of resolution and contract laboratory point of contact):

1. It should be noted that the raw ICAL data from the instrument used for the secondary HE analysis were not reported in the data package. Thus, the surrogate retention time criteria could not be evaluated. No sample data were qualified as a result.

Reviewed by: Eric T. MinkLevel: 1Date: 3/7/12

VALIDATOR'S SIGNATURE: _____

DATE: 3/6/12

LC/MS/MS HIGH EXPLOSIVE ANALYTICAL DATA VALIDATION CHECKLIST**5122-2****LC/MS/MS High Explosive Analytical Data Validation Checklist**

Records Use only



Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	1. The IS retention time has shifted by more than 30 seconds.	R, UJ, HE0	J, HE0
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	2. Required IS retention time documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, HE0b	R, HE0b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	3. The quantitating IS area count is <25% of the expected value, which indicates increased potential for false negative results and other possible problems with sample quantitation. Follow the method-specific windows.	R, HE1a	J, HE1a
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	4. The IS area count for the quantitating IS is <70% but >25% of the average of that obtained from the calibration standards.	UJ, HE1b	J+, HE1b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	5. The IS area count for the quantitating IS is >130% of the average of that obtained from the calibration standards.	UJ, HE1c	J-, HE1c
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	6. Required IS information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, HE1d	R, HE1d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	7. The surrogate is <10%R. Follow the external laboratory limits.	R, HE3	J-, HE3
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	8. The surrogate is < the Lower Acceptance Limit but ≥10% recovery. Follow the external laboratory limits.	UJ, HE3a	J-, HE3a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	9. The surrogate %R value is > the Upper Acceptance Limit. Follow the external laboratory limits.	N/A	J+, HE3b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	10. At least one surrogate is > the Upper Acceptance Limit and one surrogate is < the Lower Acceptance Limit. Follow the external laboratory limits.	UJ, HE3c	J, HE3c

LC/MS/MS HIGH EXPLOSIVE ANALYTICAL DATA VALIDATION CHECKLIST**5122-2****LC/MS/MS High Explosive Analytical Data Validation Checklist**

Records Use only



Yes No N/A (Check One)				Assign Qualifier Listed Below If Criterion = Yes	
				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	11. Required surrogate information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, HE3d	R, HE3d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	12. The sample result is ≤ 5 times the concentration of the related analyte in the method blank.	U, HE4	N/A
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	13. The affected analytes are considered estimated and biased high because this analyte was identified in the method blank but was $>5x$.	N/A	J, HE4a
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	14. The sample result is ≤ 5 times the concentration of the related analyte in the trip blank, rinsate blank, and/or equipment blank.	U, HE4d	N/A
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	15. Required method blank information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, HE4e	R, HE4e
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	16. The absence of sample carry-over must be determined and verified.	N/A	R, N, HE4f
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	17. The affected results were not analyzed with a valid 5-point calibration curve and/or a standard at the reporting limit.	UJ, HE7	J, HE7
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	18. The affected analytes were analyzed with an initial calibration curve that exceeded the %RSD criteria and/or the associated multipoint calibration correlation coefficient is less < 0.99 .	UJ, R, HE7a	J, HE7a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	19. The affected analytes were analyzed with a RRF of < 0.05 in the initial calibration and/or CCV.	UJ, R, HE7b	J, HE7b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	20. The ICV and/or CCV were recovered outside the method limits.	UJ, R, HE7c	J, HE7c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	21. The ICV and/or CCV were not analyzed at the appropriate method frequency.	UJ, R, HE7d	J, HE7d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	22. Required calibration information is missing or samples were analyzed on an expired calibration. Contact the SMO or external laboratory for information.	R, HE7f	R, HE7f

LC/MS/MS HIGH EXPLOSIVE ANALYTICAL DATA VALIDATION CHECKLIST**5122-2****LC/MS/MS High Explosive Analytical Data Validation Checklist**

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Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	23. The mass spectral documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, HE8a	R, HE8a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	24. The holding time was >1 and ≤2 times the applicable holding time requirement.	UJ, HE9	J-, HE9
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	25. The holding time was >2 times the applicable holding time requirement.	R, HE9a	J-, HE9a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	26. The LCS percent recovery was <10%. Follow the external laboratory limits.	R, HE12	J-, HE12
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	27. The LCS percent recovery was < the Lower Acceptance Limit but >10%. Follow the external laboratory limits.	UJ, HE12a	J-, HE12a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	28. The LCS percent recovery was > the Upper Acceptance Limit. Follow the external laboratory limits.	N/A	J+, HE12b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	29. The LCS documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, HE12c	R, HE12c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	30. The MS/MSD percent recovery was <10%.	R, HE12d	R, HE12d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	31. The MS/MSD percent recovery was >10% but <70%.	UJ, HE12e	J, HE12e
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	32. The MS/MSD percent recover was >70%.	N/A	J+, HE12f
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	33. The MS/MSD relative percent difference was >30%.	UJ, HE12g	J, HE12g
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	34. The affected analytes are considered suspect because the sample was diluted without any target analytes identified due to matrix interference. (Qualify as Reject if the analytical laboratory cannot provide proof for matrix interference.)	UJ, R, HE15	R, HE15
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	35. The sample was diluted because target analytes were > the initial verification calibration.	UJ, HE15a	J, HE15a

LC/MS/MS HIGH EXPLOSIVE ANALYTICAL DATA VALIDATION CHECKLIST**5122-2****LC/MS/MS High Explosive Analytical Data Validation Checklist**

Records Use only



Yes No N/A (Check One)				Assign Qualifier Listed Below If Criterion = Yes	
				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	36. The Contract Required Detection Limit Check Standard (CRI) sample did not pass method acceptance criteria.	UJ, R, HE16	J, HE16
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	37. The required CRI sample information is missing. Contact the SMO or external laboratory for information.	R, HE16c	R, HE16c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	38. The LANL project chemist identified quality deficiencies in the reported data that requires further qualification. This code can only be used and/or under advisement by the LANL project chemist.	UJ, R, HE19	J, R, HE19
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	39. Duplicate, dilution, or reanalysis.	UJ, HE88	J, HE88

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270003

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture:

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227015a

Date Analyzed: 28-FEB-12 01:51

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
118-96-7	2,4,6-Trinitrotoluene	0.263	U	0.0842	0.263
<i>118-96-7</i>	<i>2,4,6-Trinitrotoluene</i>				
121-14-2	2,4-Dinitrotoluene	0.263	U	0.0842	0.263
<i>121-14-2</i>	<i>2,4-Dinitrotoluene</i>				
121-82-4	RDX	0.263	U	0.0842	0.263
<i>121-82-4</i>	<i>RDX</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				
2691-41-0	HMX	0.263	U	0.0842	0.263
<i>2691-41-0</i>	<i>HMX</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
606-20-2	2,6-Dinitrotoluene	0.263	U	0.0842	0.263
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
88-72-2	o-Nitrotoluene	0.263	U	0.0863	0.263
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
98-95-3	Nitrobenzene	0.263	U	0.0842	0.263
<i>98-95-3</i>	<i>Nitrobenzene</i>				
99-08-1	m-Nitrotoluene	0.263	U	0.0842	0.263
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	0.263	U	0.0842	0.263
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
99-65-0	m-Dinitrobenzene	0.263	U	0.0842	0.263
<i>99-65-0</i>	<i>m-Dinitrobenzene</i>				
479-45-8	Tetryl	0.526	U	0.0842	0.526
<i>479-45-8</i>	<i>Tetryl</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270003

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
78-11-5	PETN	0.526	U	0.105	0.526
<i>78-11-5</i>	<i>PETN</i>				
99-99-0	p-Nitrotoluene	0.526	U	0.158	0.526
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270003

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160016.wiff

Date Analyzed: 16-FEB-12 18:19

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
3058-38-6	TATB	1.05	U	0.316	1.05
<i>3058-38-6</i>	<i>TATB</i>				
618-87-1	3,5-Dinitroaniline	1.05	U	0.316	1.05
<i>618-87-1</i>	<i>3,5-Dinitroaniline</i>				
78-30-8	tris(o-cresyl) phosphate	1.05	U	0.316	1.05
<i>78-30-8</i>	<i>tris(o-cresyl) phosphate</i>				
59229-75-3	2,6-Diamino-4-nitrotoluene	2.63	U	0.526	2.63
<i>59229-75-3</i>	<i>2,6-Diamino-4-nitrotoluene</i>				
6629-29-4	2,4-Diamino-6-nitrotoluene	2.63	U	0.526	2.63
<i>6629-29-4</i>	<i>2,4-Diamino-6-nitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2199

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270005

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227018a

Date Analyzed: 28-FEB-12 03:20

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
118-96-7	2,4,6-Trinitrotoluene	0.263	U	0.0842	0.263
<i>118-96-7</i>	<i>2,4,6-Trinitrotoluene</i>				
121-14-2	2,4-Dinitrotoluene	0.263	U	0.0842	0.263
<i>121-14-2</i>	<i>2,4-Dinitrotoluene</i>				
121-82-4	RDX	0.263	U	0.0842	0.263
<i>121-82-4</i>	<i>RDX</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				
2691-41-0	HMX	0.263	U	0.0842	0.263
<i>2691-41-0</i>	<i>HMX</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
606-20-2	2,6-Dinitrotoluene	0.263	U	0.0842	0.263
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
88-72-2	o-Nitrotoluene	0.263	U	0.0863	0.263
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
98-95-3	Nitrobenzene	0.263	U	0.0842	0.263
<i>98-95-3</i>	<i>Nitrobenzene</i>				
99-08-1	m-Nitrotoluene	0.263	U	0.0842	0.263
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	0.263	U	0.0842	0.263
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
99-65-0	m-Dinitrobenzene	0.263	U	0.0842	0.263
<i>99-65-0</i>	<i>m-Dinitrobenzene</i>				
479-45-8	Tetryl	0.526	U	0.0842	0.526
<i>479-45-8</i>	<i>Tetryl</i>				

LMF
3/6/12

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2199

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270005

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
78-11-5	PETN	0.526	U	0.105	0.526
<i>78-11-5</i>	<i>PETN</i>				
99-99-0	p-Nitrotoluene	0.526	U	0.158	0.526
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2199

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270005

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160019.wiff

Date Analyzed: 16-FEB-12 19:09

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
3058-38-6	TATB	1.05	U	0.316	1.05
<i>3058-38-6</i>	<i>TATB</i>				
618-87-1	3,5-Dinitroaniline	1.05	U	0.316	1.05
<i>618-87-1</i>	<i>3,5-Dinitroaniline</i>				
78-30-8	tris(o-cresyl) phosphate	1.05	U	0.316	1.05
<i>78-30-8</i>	<i>tris(o-cresyl) phosphate</i>				
59229-75-3	2,6-Diamino-4-nitrotoluene	2.63	U	0.526	2.63
<i>59229-75-3</i>	<i>2,6-Diamino-4-nitrotoluene</i>				
6629-29-4	2,4-Diamino-6-nitrotoluene	2.63	U	0.526	2.63
<i>6629-29-4</i>	<i>2,4-Diamino-6-nitrotoluene</i>				

DATA VALIDATION COVER SHEET**5120-1****Data Validation Cover Sheet**

Records Use only

**Section I.**REQUEST NUMBER: 12-700 VALIDATION DATE: 3/6/12 LAB CODE: GELCONTRACT LABORATORY NAME: GEL Laboratories LLCVALIDATOR: Larry Fukui ORGANIZATION: Analytical Quality Associates, Inc.

ANALYTICAL SUITE (CHECK ALL THAT APPLY):

- | | | | |
|---|--|---|--|
| <input type="checkbox"/> TPH-GRO | <input type="checkbox"/> HIGH EXPLOSIVES | <input type="checkbox"/> DIOXIN FURANS | <input type="checkbox"/> LCMSMS PERCHLORATES |
| <input type="checkbox"/> TPH-DRO | <input type="checkbox"/> METALS | <input type="checkbox"/> PCB CONGENERS | <input type="checkbox"/> ORGANOCHLORINE |
| <input checked="" type="checkbox"/> GENERAL CHEMISTRY | <input type="checkbox"/> RADIOCHEMISTRY | <input type="checkbox"/> LCMSMS HIGH EXPLOSIVES | PESTICIDES/POLYCHLORINATED BIPHENYLS |

☐ OTHER (DESCRIBE):
_____**Section II. Completeness Check**

- | YES | NO | N/A | (CHECK ONE) | YES | NO | N/A | (CHECK ONE) |
|-------------------------------------|--------------------------|-------------------------------------|-----------------------------|-------------------------------------|--------------------------|-------------------------------------|--------------------------|
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 1. CHAIN-OF-CUSTODY FORM(S) | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 6. RAW/BSS DATA |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 2. CASE NARRATIVE | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 7. QUALITY CONTROL FORMS |
| <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 3. SAMPLE RESULT FORMS | <input checked="" type="checkbox"/> | <input type="checkbox"/> | <input type="checkbox"/> | 8. QUANTITATION REPORTS |
| <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 4. SAMPLE CHROMATOGRAMS | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 9. TICS FORMS |
| <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 5. STANDARD CHROMATOGRAMS | <input type="checkbox"/> | <input type="checkbox"/> | <input checked="" type="checkbox"/> | 10. TICS MASS SPECTRA |

Comments/problems noted (include information about requests for further information submitted to the contract laboratory and agreed-upon date of resolution and contract laboratory point of contact):

- The MS %R was < the laboratory LAL but $\geq 10\%$ for TKN. The associated sample results were NDs and, thus, were qualified UJ,I6a.
- It should be noted that the matrix QC for all analyses except ammonia as nitrogen and TKN were performed on parent samples from other LANL RNs. No sample data were qualified as a result.

Reviewed by: Eric T. MinkLevel: 1Date: 3/7/12

VALIDATOR'S SIGNATURE: _____

DATE: 3/6/12

GENERAL CHEMISTRY ANALYTICAL DATA VALIDATION CHECKLIST

5120-2

General Chemistry Analytical Data Validation Checklist

Records Use only



Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	1. The holding time was >1 and ≤2 times the applicable holding time requirement.	UJ, I9	J-, I9
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	2. The holding time was >2 times the applicable holding time requirement.	R, I9a	J-, I9a
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	3. The affected analytes are regarded as rejected because the analytical holding time was exceeded.	R, I9b	R, I9b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	4. The affected results were not analyzed with a valid 5-point calibration curve and/or a standard at the reporting limit.	UJ, R, I7	J, I7
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	5. The affected analytes were analyzed with an initial calibration curve that exceeded the %RSD criteria and/or the associated multipoint calibration correlation coefficient is <0.995.	UJ, I7a	J, I7a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	6. The ICV and/or CCV were recovered outside the method specific limits.	UJ, I7c	J, I7c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	7. The ICV and/or CCV were not analyzed at the appropriate method frequency.	UJ, I7d	J, I7d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	8. Required calibration information is missing or samples were analyzed on an expired calibration. Contact the SMO or external laboratory for information.	R, I7f	R, I7f
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	9. The interference check sample percent recovery value is <50%.	R, I2	J-, I2
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	10. The interference check sample percent recovery value is ≥50% and <80%.	UJ, I2a	J-, I2a
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	11. The interference check sample percent recovery value is >120%.	N/A	J+, I2b
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	12. The interference check sample was not analyzed with the samples.	R, I2c	R, I2c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	13. The sample result is ≤5X the concentration of the related analyte in the method blank.	U, I4	N/A

GENERAL CHEMISTRY ANALYTICAL DATA VALIDATION CHECKLIST

5120-2

General Chemistry Analytical Data Validation Checklist

Records Use only



Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	14. The affected analytes are considered estimated and biased high because this analyte was identified in the method blank but was >5X.	N/A	J, I4a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	15. The sample result is ≤5X the concentration of the related analyte in the instrument blank and continuing calibration blank.	U, I4b	N/A
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	16. Continuing calibration blanks were not analyzed at the appropriate method frequency.	UJ, I4c	J, I4c
<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	17. The sample result is ≤5X the concentration of the related analyte in the trip blank, rinsate blank, or equipment blank.	U, I4d	N/A
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	18. Required method blank information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	R, I4e	R, I4e
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	19. The associate matrix spike recovery was <10%. Follow the external laboratory limits located within the associated data package.	R, I6	R, I6
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	20. The associated matrix spike recovery was below the Lower Acceptance Limit (LAL) but >10%. Follow the external laboratory limits located within the associated data package.	UJ, I6a	J-, I6a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	21. The associated matrix spike recovery was above the Upper Acceptance Limit (UAL). Follow the external laboratory limits located within the associated data package.	UJ, I6b	J+, I6b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	22. Required matrix spike information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information. If LCS information is present, do not reject. Qualify data based on LCS information.	R, I6c	R, I6c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	23. The sample and/or the duplicate sample results RPD is not within the acceptance limits. Follow the external laboratory limits located within the associated data package.	UJ, I10b	J, I10b

GENERAL CHEMISTRY ANALYTICAL DATA VALIDATION CHECKLIST

5120-2

General Chemistry Analytical Data Validation Checklist

Records Use only



Yes No N/A				Assign Qualifier Listed Below If Criterion = Yes	
(Check One)				Non-detected Analyte	Detected Analyte
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	24. The duplicate sample was not prepared and/or analyzed with the samples for unspecified reasons. The duplicate information is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information.	UJ, I10d	J, I10d
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	25. The LCS percent recovery was <10%. Follow the external laboratory limits located within the associated data package.	R, I12	R, I12
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	26. The LCS percent recover was < the LAL but >10%. Follow the external laboratory limits located within the associated data package.	UJ, I12a	J-, I12a
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	27. The LCS percent recovery was > the UAL. Follow the external laboratory limits located within the associated data package.	N/A	J+, I12b
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	28. The LCS documentation is missing. Data may not be acceptable for use. Contact the SMO or external laboratory for information. Do not Reject if MS/MSD information is present. Qualify according to MS/MSD criteria.	R, I12c	R, I12c
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	29. Duplicate, dilution, or reanalysis	UJ, I88	J, I88
<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	30. The LANL project chemist identified quality deficiencies in the reported data that require further qualification. This code can ONLY be used and/or under advisement by the LANL project chemist.	UJ, R, I19	J, R, I19
<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	31. Qualification of data via data validation does not occur based on Quality Control requirements in this procedure. Adhere to the external laboratory qualifiers found within the Form I analytical data summary sheets generated by the external laboratory.	U, U_LAB	J, J_LAB NQ, NQ (no qualification)

GEL LABORATORIES LLC

2040 Savage Road Charleston SC 29407 - (843) 556-8171 - www.gel.com

Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2030
Sample ID: 295270001
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method
Nutrient Analysis											
EPA 350.1 Nitrogen, Ammonia L "As Received"											
Nitrogen, Ammonia	U	ND	0.016	0.050	mg/L	1	KLP1	02/08/12	1542	1186023	1
EPA 353.2 Nitrogen, Nitrate/Nitrite "As Received"											
Nitrogen, Nitrate/Nitrite		0.304	0.050	0.250	mg/L	5	KLP1	02/14/12	1237	1186031	2
EPA 365.4 Phosphorus, Total in "As Received"											
Phosphorus, Total as P		0.0562	0.015	0.050	mg/L	1	KLP1	02/08/12	1417	1186020	3

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 350.2 Prep	EPA 350.1 Ammonia Nitrogen Prep	AXS5	02/07/12	1658	1186022
EPA 365.4 Prep	EPA 365.4 Phosphorus, Total in liquid PR	AXS5	02/08/12	1159	1186019

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	EPA 350.1	
2	EPA 353.2	
3	EPA 365.4	

LMF
3/6/12

GEL LABORATORIES LLC

2040 Savage Road Charleston SC 29407 - (843) 556-8171 - www.gel.com

Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2031
Sample ID: 295270003
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method	
Carbon Analysis												
SW 9060 Total Organic Carbon "As Received"												
Total Organic Carbon Average	J	0.349	0.330	1.00	mg/L	1	TSM	02/07/12	1841	1186182	1	
Nutrient Analysis												
Nitrogen, Total Kjeldahl (TKN) "As Received"												
Nitrogen, Total Kjeldahl	U	ND	UJ.l6a	0.035	0.100	mg/L	1	KLP1	02/17/12	1038	1186025	2

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 351.2 Prep	EPA 351.2 Total Kjeldahl Nitrogen Prep	AXS5	02/14/12	1649	1186024

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	SW846 9060	
2	EPA 351.2	

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Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2199
Sample ID: 295270005
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method	
Carbon Analysis												
SW 9060 Total Organic Carbon "As Received"												
Total Organic Carbon Average	J	0.359	0.330	1.00	mg/L	1	TSM	02/07/12	1915	1186182	1	
Nutrient Analysis												
Nitrogen, Total Kjeldahl (TKN) "As Received"												
Nitrogen, Total Kjeldahl	U	ND	UJ,l6a	0.175	0.500	mg/L	5	KLP1	02/17/12	1049	1186025	2

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 351.2 Prep	EPA 351.2 Total Kjeldahl Nitrogen Prep	AXS5	02/14/12	1649	1186024

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	SW846 9060	
2	EPA 351.2	

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3/6/12

GEL LABORATORIES LLC

2040 Savage Road Charleston SC 29407 - (843) 556-8171 - www.gel.com

Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2200
Sample ID: 295270006
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method
Nutrient Analysis											
EPA 350.1 Nitrogen, Ammonia L "As Received"											
Nitrogen, Ammonia	U	ND	0.016	0.050	mg/L	1	KLP1	02/08/12	1554	1186023	1
EPA 353.2 Nitrogen, Nitrate/Nitrite "As Received"											
Nitrogen, Nitrate/Nitrite		0.304	0.050	0.250	mg/L	5	KLP1	02/14/12	1238	1186031	2
EPA 365.4 Phosphorus, Total in "As Received"											
Phosphorus, Total as P	J	0.0356	0.015	0.050	mg/L	1	KLP1	02/08/12	1418	1186020	3

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 350.2 Prep	EPA 350.1 Ammonia Nitrogen Prep	AXS5	02/07/12	1658	1186022
EPA 365.4 Prep	EPA 365.4 Phosphorus, Total in liquid PR	AXS5	02/08/12	1159	1186019

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	EPA 350.1	
2	EPA 353.2	
3	EPA 365.4	

LMF
3/6/12

Thursday, February 02, 2012

LAB CHAIN OF CUSTODY DOCUMENT NUMBER: 12-700C

LOS ALAMOS

REQUEST NUMBER: 12-700

NATIONAL LABORATORY

ATTN: Valerie Davis

TURNAROUND/REPORT DUE: 3/3/2012

General Engineering Laboratories, Inc., Charleston, SC.

TURNAROUND REQ'D: 30

2040 Savage Rd

Charleston, SC 29407

LAB REQUEST COMMENTS:

SAMPLE ID	CTNR	CTNR DESC	ORDER	PRESERV	MATRIX
CAAN-12-2030	1	AMBER GLASS	WSP-NH3+NO3/NO2+PO4	Sulfuric Acid (H2SO4)	WG
CAAN-12-2031	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2031	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2031	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	1	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	2	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	3	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	1	AMBER GLASS	WSP-TKN+TOC	Sulfuric Acid (H2SO4)	WG
CAAN-12-2032	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2032	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2199	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG

Thursday, February 02, 2012

LAB CHAIN OF CUSTODY DOCUMENT NUMBER: 12-700C

LOS ALAMOS

REQUEST NUMBER: 12-700

NATIONAL LABORATORY

ATTN: Valerie Davis

TURNAROUND/REPORT DUE: 3/3/2012

General Engineering Laboratories, Inc., Charleston, SC.

TURNAROUND REQ'D: 30

2040 Savage Rd

Charleston, SC 29407

LAB REQUEST COMMENTS:

SAMPLE ID	CTNR	CTNR DESC	ORDER	PRESERV	MATRIX
CAAN-12-2199	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2199	1	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	2	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	3	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	1	AMBER GLASS	WSP-TKN+TOC	Sulfuric Acid (H2SO4)	WG
CAAN-12-2200	1	AMBER GLASS	WSP-NH3+NO3/NO2+ PO4	Sulfuric Acid (H2SO4)	WG
CAAN-12-2201	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2201	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2201	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2201	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2201	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG

Relinquished By:

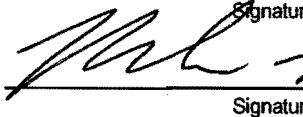
Date

Time

Received By:

Date

Time

Signature	Signature
	
Signature	Signature
Signature	Signature

Received for DISPOSAL By: Date

Time

Remarks:

Signature

Thursday, February 02, 2012

REQUEST NUMBER: 12-700

PRIORITY	METHOD CODE	CNTNR	SAMPLE ID	SAMPLE MATRIX	DATE SAMPLED	SPECIAL INSTRUCTIONS
	SW-846:8260B	1	CAAN-12-2032	WG	2/1/2012	
		2	CAAN-12-2032	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		1	CAAN-12-2201	WG	2/1/2012	
		2	CAAN-12-2201	WG	2/1/2012	
	SW-846:8270C	1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		3	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		3	CAAN-12-2199	WG	2/1/2012	
		1	CAAN-12-2201	WG	2/1/2012	
		2	CAAN-12-2201	WG	2/1/2012	
		3	CAAN-12-2201	WG	2/1/2012	
	SW-846:8321A_MOD	1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		3	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		3	CAAN-12-2199	WG	2/1/2012	
	SW-846:9060	1	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	

Thursday, February 02, 2012

REQUEST NUMBER: 12-700

LOS ALAMOS
NATIONAL LABORATORY

ATTN: Valerie Davis

General Engineering Laboratories, Inc., Charleston, SC.
2040 Savage Rd
Charleston, SC 29407

These Samples are on:

LANL Request Number: 12-700

Per Agreement Number: 126310011

Project Cost Code: MR1A015AGWJ0

Please analyse the enclosed samples
according to the schedule indicated:

SHIP DATE: 2/2/2012**TURNAROUND/REPORT DUE: 3/3/2012****TURNAROUND REQ'D: 30 Days****RAD SCREENING: Yes, Below Background****LAB REQUEST COMMENTS:**

LANL ER SMO CONTACT:

Signature: 

PRIORITY	METHOD CODE	CNTNR	SAMPLE ID	SAMPLE MATRIX	DATE SAMPLED	SPECIAL INSTRUCTIONS
	EPA:350.1	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	EPA:351.2	1	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
	EPA:353.2	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	EPA:365.4	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	SW-846:8260B	1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG		



February 07, 2012

www.gel.com

Ms. Joylene Valdez
Los Alamos National Laboratory
TA-03, SM271, Drop Pt. 02U, Rm111
Los Alamos, New Mexico 87545

Re: LANL-WQH Water Samples
Work Order: 295270
SDG: 12-700

Dear Ms. Valdez:

GEL Laboratories, LLC (GEL) appreciates the opportunity to provide the following analytical results for the sample(s) we received on February 03, 2012, and analyzed for Explosives by LCMSMS, GC/MS Semivolatile, GC/MS Volatile and General Chemistry. This original data report has been prepared and reviewed in accordance with GEL's standard operating procedures.

Our policy is to provide high quality, personalized analytical services to enable you to meet your analytical needs on time every time. We trust that you will find everything in order and to your satisfaction. If you have any questions, please do not hesitate to call me at (843) 556-8171, ext. 4485.

Sincerely,

Hope Taylor for
Valerie Davis
Project Manager

Purchase Order: 63641-10
Chain of Custody: 12-700 and
Enclosures



ARS International (63641-10)
LANL-WQH Water Samples
Work Order #: 295270
SDG: 12-700

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Case Narrative

**Case Narrative for
ARS International (63641-10)
LANL-WQH Water Samples
Workorder #: 295270
SDG # : 12-700**

February 07, 2012

Laboratory Identification:

GEL Laboratories LLC
2040 Savage Road
Charleston, South Carolina 29407
(843) 556-8171

Summary

Sample receipt The samples arrived at GEL Laboratories LLC, Charleston, South Carolina on February 03, 2012 for analysis. The samples were prepared/analyzed within the required holding time. Shipping container temperatures were checked, documented, and within specifications. The samples were screened according to GEL Standard Operating Procedure. Please see attached email for discrepancies. All sample containers arrived without any visible signs of tampering or breakage. Containers were checked for pH, where appropriate, and matched the preservative as documented on the accompanying chain of custody. Shipping container temperature was within specification (0 - 6C).

Sample Identification The laboratory received the following samples:

<u>Laboratory ID</u>	<u>Client ID</u>
295270001	CAAN-12-2030
295270002	CAAN-12-2031
295270003	CAAN-12-2031
295270004	CAAN-12-2199
295270005	CAAN-12-2199
295270006	CAAN-12-2200
295270007	CAAN-12-2201
295270008	CAAN-12-2032

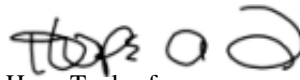
Case Narrative

Sample analyses were conducted using methodology as outlined in GEL Laboratories, LLC (GEL) Standard Operating Procedures. Any technical or administrative problems during analysis, data review, and reduction are contained in the analytical case narratives in the enclosed data package.

Data Package

The enclosed data package contains the following sections: Case Narrative, Chain of Custody, Cooler Receipt Checklist, Data Package Qualifier Definitions and data from the following fractions: Explosives by LCMSMS, GC/MS Semivolatile, GC/MS Volatile and General Chemistry.

I certify that this data report is in compliance with the terms and conditions of the subcontract and task order, both technically and for completeness, for other than the conditions detailed in the attached case narrative.

A handwritten signature in black ink, appearing to read 'Hope Taylor'.

Hope Taylor for
Valerie Davis
Project Manager

List of current GEL Certifications as of 07 February 2012

State	Certification
Arizona	AZ0766
Arkansas	88-0651
CLIA	42D0904046
California NELAP	01151CA
Colorado	SC00012
Connecticut	PH-0169
Delaware	SC00012
DoD ELAP A2LA ISO 17025	2567.01
Florida NELAP	E87156
Foreign Soils Permit	P330-09-00191
Georgia	SC00012
Georgia SDWA	967
Hawaii	SC00012
Idaho	SC00012
Illinois NELAP	200029
Indiana	C-SC-01
Kansas NELAP	E-10332
Kentucky	90129
Louisiana NELAP	03046 (AI33904)
Louisiana SDWA	LA110008
Maryland	270
Massachusetts	M-SC012
Mississippi	SC00012
Nevada	SC000122011-1
New Hampshire NELAP	2054
New Jersey NELAP	SC002
New Mexico	SC00012
New York NELAP	11501
North Carolina	233
North Carolina SDWA	45709
Oklahoma	9904
Pennsylvania NELAP	68-00485
South Carolina Chemistry	10120001
South Carolina Radiochemi	10120002
Tennessee	TN 02934
Texas NELAP	T104704235-12-7
Utah NELAP	SC00012
Vermont	VT87156
Virginia NELAP	460202
Washington	C780
Wisconsin	999887790

Chain of Custody and Supporting Documentation

Thursday, February 02, 2012

LAB CHAIN OF CUSTODY DOCUMENT NUMBER: 12-700C

LOS ALAMOS

REQUEST NUMBER: 12-700

NATIONAL LABORATORY

ATTN: Valerie Davis

TURNAROUND/REPORT DUE: 3/3/2012

General Engineering Laboratories, Inc., Charleston, SC.

TURNAROUND REQ'D: 30

2040 Savage Rd

Charleston, SC 29407

LAB REQUEST COMMENTS:

295270

SAMPLE ID	CTNR	CTNR DESC	ORDER	PRESERV	MATRIX
CAAN-12-2030	1	AMBER GLASS	WSP-NH3+NO3/NO2+PO4	Sulfuric Acid (H2SO4)	WG
CAAN-12-2031	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2031	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2031	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2031	1	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	2	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	3	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2031	1	AMBER GLASS	WSP-TKN+TOC	Sulfuric Acid (H2SO4)	WG
CAAN-12-2032	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2032	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2199	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2199	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG

Thursday, February 02, 2012

LAB CHAIN OF CUSTODY DOCUMENT NUMBER: 12-700C

LOS ALAMOS

REQUEST NUMBER: 12-700

NATIONAL LABORATORY

ATTN: Valerie Davis

TURNAROUND/REPORT DUE: 3/3/2012

General Engineering Laboratories, Inc., Charleston, SC.

TURNAROUND REQ'D: 30

2040 Savage Rd

Charleston, SC 29407

LAB REQUEST COMMENTS:

SAMPLE ID	CTNR	CTNR DESC	ORDER	PRESERV	MATRIX
CAAN-12-2199	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2199	1	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	2	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	3	AMBER GLASS	WSP-8321A-NMED HEXP	Ice	WG
CAAN-12-2199	1	AMBER GLASS	WSP-TKN+TOC	Sulfuric Acid (H2SO4)	WG
CAAN-12-2200	1	AMBER GLASS	WSP-NH3+NO3/NO2+ PO4	Sulfuric Acid (H2SO4)	WG
CAAN-12-2201	1	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2201	2	SEPTUM AMBER GLASS	WSP-8260B-VOA	Hydrochloric Acid (HCL)	WG
CAAN-12-2201	1	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2201	2	AMBER GLASS	WSP-8270C-SVOA	Ice	WG
CAAN-12-2201	3	AMBER GLASS	WSP-8270C-SVOA	Ice	WG

Relinquished By:

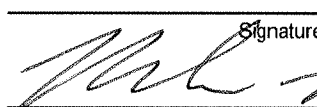
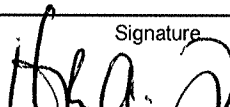

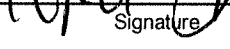
Date

Time

Received By:

Date

Time

	
Signature	Signature
	
Signature	Signature


2/2/12 3:00 02/01/12 0845

	
Signature	Signature

Received for DISPOSAL By: Date

Time

Remarks:


Signature

Thursday, February 02, 2012

**LOS ALAMOS
NATIONAL LABORATORY**

ATTN: Valerie Davis

General Engineering Laboratories, Inc., Charleston, SC.
2040 Savage Rd
Charleston, SC 29407

Please analyse the enclosed samples
according to the schedule indicated:

SHIP DATE: 2/2/2012

TURNAROUND/REPORT DUE: 3/3/2012

TURNAROUND REQ'D: 30 Days

RAD SCREENING: Yes, Below Background

LAB REQUEST COMMENTS:

LANL ER SMO CONTACT:

Signature:



These Samples are on:

LANL Request Number: 12-700
Per Agreement Number: 126310011
Project Cost Code: MR1A015AGWJ0

REQUEST NUMBER: 12-700

PRIORITY	METHOD CODE	CNTNR	SAMPLE ID	SAMPLE MATRIX	DATE SAMPLED	SPECIAL INSTRUCTIONS
	EPA:350.1	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	EPA:351.2	1	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
	EPA:353.2	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	EPA:365.4	1	CAAN-12-2030	WG	2/1/2012	
		1	CAAN-12-2200	WG	2/1/2012	
	SW-846:8260B	1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	

Hard Copy Required

Page 2 of 2

Thursday, February 02, 2012

REQUEST NUMBER: 12-700

PRIORITY	METHOD CODE	CNTNR	SAMPLE ID	SAMPLE MATRIX	DATE SAMPLED	SPECIAL INSTRUCTIONS
	SW-846:8260B	1	CAAN-12-2032	WG	2/1/2012	
		2	CAAN-12-2032	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		1	CAAN-12-2201	WG	2/1/2012	
		2	CAAN-12-2201	WG	2/1/2012	
		1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		3	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
	SW-846:8270C	2	CAAN-12-2199	WG	2/1/2012	
		3	CAAN-12-2199	WG	2/1/2012	
		1	CAAN-12-2201	WG	2/1/2012	
		2	CAAN-12-2201	WG	2/1/2012	
		3	CAAN-12-2201	WG	2/1/2012	
		1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		3	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
	SW-846:8321A_MOD	3	CAAN-12-2201	WG	2/1/2012	
		1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		3	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
		2	CAAN-12-2199	WG	2/1/2012	
		3	CAAN-12-2199	WG	2/1/2012	
		1	CAAN-12-2031	WG	2/1/2012	
		2	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	
	SW-846:9060	1	CAAN-12-2031	WG	2/1/2012	
		1	CAAN-12-2199	WG	2/1/2012	

Final Page of REQUEST NUMBER 12-700



SAMPLE RECEIPT & REVIEW FORM

Client: LANL		SDG/AR/COC/Work Order: 12-700	
Received By: HOPE TAYLOR		Date Received: 03 February 2012	
Suspected Hazard Information	Yes	No	*If Net Counts > 100cpm on samples not marked "radioactive", contact the Radiation Safety Group for further investigation.
COC/Samples marked as radioactive?		X	Maximum Net Counts Observed* (Observed Counts - Area Background Counts): 20cpm
Classified Radioactive II or III by RSO?		X	If yes, Were swipes taken of sample containers < action levels?
COC/Samples marked containing PCBs?		X	
Shipped as a DOT Hazardous?		X	Hazard Class Shipped: UN#:
Samples identified as Foreign Soil?		X	

Sample Receipt Criteria	Yes	NA	No	Comments/Qualifiers (Required for Non-Conforming Items)
1 Shipping containers received intact and sealed?	X			Circle Applicable: Seals broken Damaged container Leaking container Other (describe)
2 Samples requiring cold preservation within (0 ≤ 6 deg. C)?*	X			Preservation Method: Ice bags Blue ice Dry ice None Other (describe) *all temperatures are recorded in Celsius 4-5, 16
2a Daily check performed and passed on IR temperature gun?	X			Temperature Device Serial #: 61524646 Secondary Temperature Device Serial # (If Applicable):
3 Chain of custody documents included with shipment?	X			
4 Sample containers intact and sealed?	X			Circle Applicable: Seals broken Damaged container Leaking container Other (describe)
5 Samples requiring chemical preservation at proper pH?	X			Sample ID's, containers affected and observed pH: If Preservation added, Lot#:
6 VOA vials free of headspace (defined as < 6mm bubble)?		X		Sample ID's and containers affected:
7 Are Encore containers present?			X	(If yes, immediately deliver to Volatiles laboratory)
8 Samples received within holding time?	X			ID's and tests affected:
9 Sample ID's on COC match ID's on bottles?	X			Sample ID's and containers affected:
10 Date & time on COC match date & time on bottles?			X	Sample ID's affected: TIME NOT NOTATED ON CHAIN OF CUSTODY
11 Number of containers received match number indicated on COC?			X	Sample ID's affected: Lab received one container each for IDs CAAN-12-2199 and 2201 for HEXP and SVOA chain indicates three each. Also, for ID CAAN12-2032 received one container for 8260 chain indicates two.
12 Are sample containers identifiable as GEL provided?			X	
13 COC form is properly signed in relinquished/received sections?	X			
14 Carrier and tracking number.	X			Circle Applicable: FedEx Air FedEx Ground UPS Field Services Courier Other 7209 7856 3090 4C 7209 7856 3104 4C 7209 7856 3115 5C 7209 7856 3089 16C

Comments (Use Continuation Form if needed):

Subject: Sample Receipt for 020312

From: Hope Taylor <Hope.Taylor@gel.com>

Date: 2/3/2012 3:27 PM

To: "Keith R. Greene" <kgreene@lanl.gov>, Joylene Valdez <joylenev@lanl.gov>, LANL@amrad.com, "team.davis" <team.davis@gel.com>

Good afternoon all,

The containers for Gross A/B were preserved prior to analysis.

RN 12-699 Lab did not receive a container for Ra226+Ra228 for ID CAAN-12-2031. Please advise

RN 12-705 Lab received one vial for WSTLA-12-1558 chain indicates two.

RN 12-704 Lab received two containers for CAAN-12-2027 and 2035 for HEXP chain indicates three.

RN 12-700 Lab received one container each for IDs CAAN-12-2199 and 2201 for HEXP and SVOA chain indicates three each. Also, for ID CAAN-12-2032 received one container for 8260 chain indicates two.

Thanks

--

Hope Taylor
Project Manager Assistant
GEL Laboratories, LLC
2040 Savage Road
Charleston, SC 29407
Direct: 843.769.7376 ext. 4778
Main: 843.556.8171
Fax: 843.766.1178
E-mail: hop01200@gel.com
Web: www.gel.com

Data Review Qualifier Flag Definition Sheet

Data Review Qualifier Definitions

Qualifier Explanation

*	A quality control analyte recovery is outside of specified acceptance criteria
**	Analyte is a surrogate compound
<	Result is less than value reported
>	Result is greater than value reported
^	RPD of sample and duplicate evaluated using +/-RL. Concentrations are <5X the RL
A	The TIC is a suspected aldol-condensation product
B	Target analyte was detected in the associated blank
B	Metals-Either presence of analyte detected in the associated blank, or MDL/IDL < sample value < PQL
BD	Results are either below the MDC or tracer recovery is low
C	Analyte has been confirmed by GC/MS analysis
D	Results are reported from a diluted aliquot of the sample
d	5-day BOD-The 2:1 depletion requirement was not met for this sample
E	Organics-Concentration of the target analyte exceeds the instrument calibration range
E	Metals-%difference of sample and SD is >10%. Sample concentration must meet flagging criteria
H	Analytical holding time was exceeded
h	Preparation or preservation holding time was exceeded
J	Value is estimated
N	Metals-The Matrix spike sample recovery is not within specified control limits
N	Organics-Presumptive evidence based on mass spectral library search to make a tentative identification of the analyte (TIC). Quantitation is based on nearest internal standard response factor
N/A	Spike recovery limits do not apply. Sample concentration exceeds spike concentration by 4X or more
ND	Analyte concentration is not detected above the reporting limit
UI	Gamma Spectroscopy-Uncertain identification
X	Consult Case Narrative, Data Summary package, or Project Manager concerning this qualifier
Y	QC Samples were not spiked with this compound
Z	Paint Filter Test-Particulates passed through the filter, however no free liquids were observed.

Volatile Analysis

Case Narrative

**ChemStation Case Narrative
ARS International (ARSL)
SDG 12-700**

Method/Analysis Information

Procedure: Volatile Organic Compounds (VOC) by Gas Chromatograph/Mass Spectrometer

Analytical Method: SW846 8260B DOE-AL

Analytical Batch Number: 1187820

Sample Analysis

The following client and quality control samples were analyzed to complete this SDG using the methods referenced in the Analysis Information section:

Sample ID	Client ID
295270002	CAAN-12-2031
295270004	CAAN-12-2199
295270007	CAAN-12-2201
295270008	CAAN-12-2032
1202597760	Method Blank (MB)
1202597761	295270002(CAAN-12-2031) Post Spike (PS)
1202597762	295270002(CAAN-12-2031) Post Spike Duplicate (PSD)
1202597763	Laboratory Control Sample (LCS)
1202597764	Laboratory Control Sample (LCS)

NOTE: For volatile organic analyses the matrix spike designations may be indicated as "PS" or "PSD". The "PS" designation (post spike) indicates that the matrix was fortified prior to analysis but after applying any prep factors, such as a dilution. The laboratory considers the MS/MSD and PS/PSD designations interchangeable.

The samples in this SDG were analyzed on an "as received" basis.

Preparation/Analytical Method Verification

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-OA-E-038 REV# 16.

Raw data reports are processed and reviewed by the analyst using the Chemstation software package. False positives have been removed from the quantitation reports per standard operating procedures (SOP) section 19.1.2. False positive analytes are designated on the quantitation report with a 'd' qualifier.

Calibration Information

Please note that the 'Cal Date' indicated on each quantitation report reflects the date and time of the most recent calibrated analyte(s) in the processing method. Since the laboratory may calibrate with multiple solutions on

different days using the same processing method, the software will update the 'Cal Date' to the last calibration file, date and time. The correct dates and times for all calibration files are located on the Calibration History report in the Standard Data section in the data package.

The surrogate compounds were calibrated using a minimum five-point calibration curve. The surrogates were added by the auto sampler at a concentration of 50 ug/L. GEL Laboratories LLC will not have surrogate recoveries reported for Dibromofluoromethane. This is due to increased regulations for this analyte and an industry shortage.

A complete list of the initial calibration data files are shown in the Calibration History report located in the Standard Data section of the data package.

Initial Calibration

All initial calibration requirements have been met for this sample delivery groups (SDG). A second source initial calibration verification (ICV) was included in the standard section directly behind the initial calibration.

Continuing Calibration Verification Requirements

All associated calibration verification standard(s) (ICV or CCV) met the acceptance criteria.

Quality Control (QC) Information

Method Blank (MB) Statement

The MB analyzed with this SDG met the acceptance criteria.

Surrogate Recoveries

Surrogate recoveries in all client and quality control samples were within the acceptance limits.

Laboratory Control Sample (LCS) Recovery

The LCS spike recoveries met the acceptance limits.

QC Sample Designation

Sample 295270002 (CAAN-12-2031) was designated for spike analysis.

Matrix Spike (PS) Recovery Statement

The spike recoveries were within the required acceptance limits.

Matrix Spike Duplicate (PSD) Recovery Statement

The spike duplicate recoveries were within the required acceptance limits.

Relative Percent Difference (RPD) Statement

The RPDs between the matrix spike pair met the acceptance limits.

Internal Standard (ISTD) Acceptance

The internal standard responses in all client and quality control samples met the required acceptance criteria.

Technical Information

Holding Time Specifications

GEL assigns holding times based on the associated methodology, which assigns the date and time from sample collection or sample receipt. Those holding times expressed in hours are calculated in the ALPHALIMS system. Those holding times expressed as days expire at midnight on the day of expiration. All samples in this SDG met the specified holding time.

Sample Preservation and Integrity

All samples met the sample preservation and integrity requirements.

Sample Dilutions/Methanol Dilutions

The samples in this SDG did not require dilutions.

Sample Re-extraction/Re-analysis

Samples 1202597761 (CAAN-12-2031) and 1202597762 (CAAN-12-2031) were re-analyzed due to unacceptable recoveries in the initial analysis.

Miscellaneous Information**Electronic Packaging Comment**

This data package was generated using an electronic data processing program referred to as virtual packaging. In an effort to increase quality and efficiency, the laboratory has developed systems to generate all data packages electronically. The following change from traditional packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. An electronic signature page inserted after the case narrative will include the data validator's signature and title. The signature page also includes the data qualifiers used in the fractional package. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

Data Exception (DER) Documentation

A Data Exception Document was not required for this SDG.

Manual Integrations

Data files associated with the initial calibration, continuing calibration check, and samples did not require manual integrations.

TIC Comment

Tentatively identified compounds (TIC) were required for this sample delivery group/work order. The tentatively identified compounds may include some silanols. These compounds were due to column or septum bleed and were not native to the affected samples. Please note that non-requested target analytes that are reported on the quantitation reports will not be present on the Form I. These detected analytes are included in the calibrated method and as a result cannot be reported on the TIC quantitation report.

Additional Comments

Additional comments were not required for this SDG.

Residual Chlorine

Residual Chlorine was not detected in any of the samples in this SDG.

System Configuration

The Volatile-GC/MS analysis was performed on the following instrument configuration:

Instrument ID	Instrument	System Configuration	Column ID	Column Description	P & T Trap
VOA9.I	Agilent 6890/5973 GC/MS w/ OI Eclipse/Archon Autosampler	HP6890/HP5973	DB-624	J&W, 60m x 0.25mm x 1.4um	Trap 10

Certification Statement

Where the analytical method has been performed under NELAP certification, the analysis has met all the requirements of the NELAC standard unless otherwise noted in the analytical case narrative.

GEL LABORATORIES LLC

2040 Savage Road Charleston SC 29407 - (843) 556-8171 - www.gel.com

Qualifier Definition Report for

ARSL001 ARS International (63641-10)

Client SDG: 12-700 GEL Work Order: 295270

The Qualifiers in this report are defined as follows:

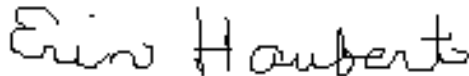
- * A quality control analyte recovery is outside of specified acceptance criteria
- ** Analyte is a surrogate compound
- U Analyte was analyzed for, but not detected above the MDL, MDA, or LOD.
- DL Indicates that sample is diluted.
- RA Indicates that sample is re-analyzed without re-extraction.
- RE Indicates that sample is re-extracted.

Review/Validation

GEL requires all analytical data to be verified by a qualified data reviewer. In addition, all CLP-like deliverables receive a third level review of the fractional data package.

The following data validator verified the information presented in this data report:

Signature:



Name: Erin Haubert

Date: 27 FEB 2012

Title: Data Validator

Sample Data Summary

Volatile
Certificate of Analysis
Sample Summary

SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2031

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 15:30

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 15:30

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y711.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

Volatile
Certificate of Analysis
Sample Summary

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	295270002	Date Received:	02/03/2012 08:45		
		Client:	ARSL001	Project:	ESHL00210
Client ID:	CAAN-12-2031	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 15:30	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 15:30				
Data File:	021212V9\9Y711.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	295270002	Date Received:	02/03/2012 08:45		
		Client:	ARSL001	Project:	ESHL00210
Client ID:	CAAN-12-2031	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 15:30	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 15:30				
Data File:	021212V9\9Y711.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	44.6	50.0	ug/L 89.2	(76%-127%)
Bromofluorobenzene	50.8	50.0	ug/L 102	(80%-120%)
Toluene-d8	49.6	50.0	ug/L 99.1	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.33	75	ug/L	0	J
	unknown siloxane	12.41	5.54	ug/L	0	J
	unknown siloxane	14.77	6.68	ug/L	0	J

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2199

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 15:56

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 15:56

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y712.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	295270004	Date Received:	02/03/2012 08:45		
		Client:	ARSL001	Project:	ESHL00210
Client ID:	CAAN-12-2199	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 15:56	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 15:56				
Data File:	021212V9\9Y712.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 295270004	Date Received: 02/03/2012 08:45	
	Client: ARSL001	Project: ESHL00210
Client ID: CAAN-12-2199	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 15:56	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 15:56		
Data File: 021212V9\9Y712.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	46.4	50.0	ug/L 92.8	(76%-127%)
Bromofluorobenzene	49.8	50.0	ug/L 99.5	(80%-120%)
Toluene-d8	47.4	50.0	ug/L 94.8	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.33	78.6	ug/L	0	J
	unknown siloxane	14.77	5.03	ug/L	0	J

Volatile
Certificate of Analysis
Sample Summary

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 295270007	Date Received: 02/03/2012 08:45	
	Client: ARSL001	Project: ESHL00210
Client ID: CAAN-12-2201	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 16:23	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 16:23		
Data File: 021212V9\9Y713.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

Volatile
Certificate of Analysis
Sample Summary

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 295270007	Date Received: 02/03/2012 08:45	
	Client: ARSL001	Project: ESHL00210
Client ID: CAAN-12-2201	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 16:23	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 16:23		
Data File: 021212V9\9Y713.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 295270007	Date Received: 02/03/2012 08:45	
	Client: ARSL001	Project: ESHL00210
Client ID: CAAN-12-2201	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 16:23	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 16:23		
Data File: 021212V9\9Y713.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	44.2	50.0	ug/L 88.5	(76%-127%)
Bromofluorobenzene	53.3	50.0	ug/L 107	(80%-120%)
Toluene-d8	45.9	50.0	ug/L 91.7	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.29	37.3	ug/L	0	J
	unknown siloxane	14.77	7.93	ug/L	0	J

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270008

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2032

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 16:49

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 16:49

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y714.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

Volatile
Certificate of Analysis
Sample Summary

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SDG Number: 12-700
Lab Sample ID: 295270008

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2032

Client: ARSL001

Project: ESHL00210

Batch ID: 1187820

Method: SW846 8260B DOE-AL

SOP Ref: GL-OA-E-038

Run Date: 02/12/2012 16:49

Inst: VOA9.I

Dilution: 1

Prep Date: 02/12/2012 16:49

Analyst: GRB2

Purge Vol: 5 mL

Data File: 021212V9\9Y714.D

Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 295270008	Date Received: 02/03/2012 08:45	
	Client: ARSL001	Project: ESHL00210
Client ID: CAAN-12-2032	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 16:49	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 16:49		
Data File: 021212V9\9Y714.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	45.7	50.0	ug/L 91.4	(76%-127%)
Bromofluorobenzene	53.0	50.0	ug/L 106	(80%-120%)
Toluene-d8	49.1	50.0	ug/L 98.2	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.33	60.8	ug/L	0	J
	unknown siloxane	16.74	7.92	ug/L	0	J

Quality Control Summary

Volatile
Surrogate Recovery Report

Page 1 of 1

SDG Number: 12-700**Matrix Type: LIQUID**

Sample ID	Client ID	DCED4 %REC	TOL %REC	BFB %REC
1202597763	LCS for batch 1187820	88	91	101
1202597764	LCS for batch 1187820	86	92	107
1202597760	MB for batch 1187820	88	96	106
295270002	CAAN-12-2031	89	99	102
295270004	CAAN-12-2199	93	95	100
295270007	CAAN-12-2201	88	92	107
295270008	CAAN-12-2032	91	98	106
1202597761	CAAN-12-2031PS	85	93	102
1202597762	CAAN-12-2031PSD	84	94	97

Surrogate**Acceptance Limits**

DCED4 = 1,2-Dichloroethane-d4

(76%-127%)

TOL = Toluene-d8

(80%-120%)

BFB = Bromofluorobenzene

(80%-120%)

* Recovery outside Acceptance Limits

Column to be used to flag recovery values

D Sample Diluted

Volatile
Quality Control Summary
Spike Recovery Report

Page 1 of 8

SDG Number: 12-700

Sample Type: Post Spike

Client ID: CAAN-12-2031PS

Matrix: WG

Lab Sample ID: 1202597761

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:20

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
75-71-8	PS Dichlorodifluoromethane	50.0	0.00 U	37.3	75	33-133
74-87-3	PS Chloromethane	50.0	0.00 U	51.2	102	52-142
75-01-4	PS Vinyl chloride	50.0	0.00 U	48.2	96	54-132
74-83-9	PS Bromomethane	50.0	0.00 U	50.2	100	63-125
75-00-3	PS Chloroethane	50.0	0.00 U	49.6	99	71-133
75-69-4	PS Trichlorofluoromethane	50.0	0.00 U	44.9	90	65-139
60-29-7	PS Ethyl ether	50.0	0.00 U	44.8	90	70-114
67-64-1	PS Acetone	250	0.00 U	162	65	30-162
75-05-8	PS Acetonitrile	1250	0.00 U	1220	98	57-126
75-35-4	PS 1,1-Dichloroethylene	50.0	0.00 U	54.3	109	68-123
74-88-4	PS Iodomethane	250	0.00 U	298	119	71-122
75-09-2	PS Methylene chloride	50.0	0.00 U	47.2	94	74-128
75-15-0	PS Carbon disulfide	250	0.00 U	314	126	69-130
1634-04-4	PS tert-Butyl methyl ether	50.0	0.00 U	48.3	97	69-126
156-60-5	PS trans-1,2-Dichloroethylene	50.0	0.00 U	54.2	108	64-122
108-05-4	PS Vinyl acetate	250	0.00 U	261	104	49-155
75-34-3	PS 1,1-Dichloroethane	50.0	0.00 U	53.3	107	74-122
78-93-3	PS 2-Butanone	250	0.00 U	197	79	16-146
156-59-2	PS cis-1,2-Dichloroethylene	50.0	0.00 U	53.7	107	66-137
594-20-7	PS 2,2-Dichloropropane	50.0	0.00 U	58.0	116	68-134
67-66-3	PS Chloroform	50.0	0.00 U	51.7	103	75-126
74-97-5	PS Bromochloromethane	50.0	0.00 U	49.4	99	79-126

Volatile
Quality Control Summary
Spike Recovery Report

Page 2 of 8

SDG Number: 12-700

Sample Type: Post Spike

Client ID: CAAN-12-2031PS

Matrix: WG

Lab Sample ID: 1202597761

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:20

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
71-55-6	PS 1,1,1-Trichloroethane	50.0	0.00 U	55.3	111	70-137
563-58-6	PS 1,1-Dichloropropene	50.0	0.00 U	56.9	114	74-125
71-36-3	PS n-Butyl alcohol	5000	0.00 U	4940	99	59-136
56-23-5	PS Carbon tetrachloride	50.0	0.00 U	57.9	116	70-143
107-06-2	PS 1,2-Dichloroethane	50.0	0.00 U	48.9	98	70-126
71-43-2	PS Benzene	50.0	0.00 U	54.6	109	74-119
79-01-6	PS Trichloroethylene	50.0	0.00 U	56.5	113	69-126
78-87-5	PS 1,2-Dichloropropane	50.0	0.00 U	55.7	111	76-121
75-27-4	PS Bromodichloromethane	50.0	0.00 U	50.8	102	77-131
74-95-3	PS Dibromomethane	50.0	0.00 U	48.4	97	79-123
108-10-1	PS 4-Methyl-2-pentanone	250	0.00 U	247	99	65-128
10061-01-5	PS cis-1,3-Dichloropropylene	50.0	0.00 U	52.7	105	77-125
108-88-3	PS Toluene	50.0	0.00 U	51.9	104	69-119
10061-02-6	PS trans-1,3-Dichloropropylene	50.0	0.00 U	51.8	104	75-125
79-00-5	PS 1,1,2-Trichloroethane	50.0	0.00 U	49.0	98	74-119
591-78-6	PS 2-Hexanone	250	0.00 U	191	76	31-144
142-28-9	PS 1,3-Dichloropropane	50.0	0.00 U	49.7	99	76-117
127-18-4	PS Tetrachloroethylene	50.0	0.00 U	56.1	112	65-126
124-48-1	PS Dibromochloromethane	50.0	0.00 U	52.5	105	74-126
106-93-4	PS 1,2-Dibromoethane	50.0	0.00 U	49.0	98	78-121
108-90-7	PS Chlorobenzene	50.0	0.00 U	53.1	106	74-119
100-41-4	PS Ethylbenzene	50.0	0.00 U	54.0	108	71-121

Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Post Spike

Client ID: CAAN-12-2031PS

Matrix: WG

Lab Sample ID: 1202597761

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:20

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
179601-23-1	PS m,p-Xylenes	100	0.00 U	109	109	70-123
95-47-6	PS o-Xylene	50.0	0.00 U	53.9	108	71-125
100-42-5	PS Styrene	50.0	0.00 U	52.9	106	71-128
75-25-2	PS Bromoform	50.0	0.00 U	51.4	103	71-126
79-34-5	PS 1,1,2,2-Tetrachloroethane	50.0	0.00 U	48.0	96	67-126
96-18-4	PS 1,2,3-Trichloropropane	50.0	0.00 U	48.0	96	76-124
108-86-1	PS Bromobenzene	50.0	0.00 U	52.1	104	72-120
103-65-1	PS n-Propylbenzene	50.0	0.00 U	55.3	111	64-125
95-49-8	PS 2-Chlorotoluene	50.0	0.00 U	54.8	110	68-126
98-82-8	PS Isopropylbenzene	50.0	0.00 U	54.5	109	68-127
108-67-8	PS 1,3,5-Trimethylbenzene	50.0	0.00 U	54.2	108	67-128
106-43-4	PS 4-Chlorotoluene	50.0	0.00 U	52.8	106	67-123
98-06-6	PS tert-Butylbenzene	50.0	0.00 U	54.3	109	67-130
95-63-6	PS 1,2,4-Trimethylbenzene	50.0	0.00 U	52.7	105	67-127
135-98-8	PS sec-Butylbenzene	50.0	0.00 U	54.4	109	66-129
99-87-6	PS 4-Isopropyltoluene	50.0	0.00 U	54.7	109	63-134
541-73-1	PS 1,3-Dichlorobenzene	50.0	0.00 U	51.7	103	68-122
106-46-7	PS 1,4-Dichlorobenzene	50.0	0.00 U	50.8	102	67-121
104-51-8	PS n-Butylbenzene	50.0	0.00 U	53.6	107	59-133
96-12-8	PS 1,2-Dibromo-3-chloropropane	50.0	0.00 U	48.7	97	62-129
87-68-3	PS Hexachlorobutadiene	50.0	0.00 U	54.1	108	52-137
91-20-3	PS Naphthalene	50.0	0.00 U	43.0	86	63-129

Volatile

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Quality Control Summary
Spike Recovery Report

SDG Number: 12-700

Sample Type: Post Spike

Client ID: CAAN-12-2031PS

Matrix: WG

Lab Sample ID: 1202597761

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:20

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
87-61-6	PS 1,2,3-Trichlorobenzene	50.0	0.00 U	44.7	89	59-131
630-20-6	PS 1,1,1,2-Tetrachloroethane	50.0	0.00 U	54.4	109	76-128
120-82-1	PS 1,2,4-Trichlorobenzene	50.0	0.00 U	47.9	96	59-127
95-50-1	PS 1,2-Dichlorobenzene	50.0	0.00 U	49.7	99	72-119

Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Post Spike Duplicate

Client ID: CAAN-12-2031PSD

Matrix: WG

Lab Sample ID: 1202597762

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:46

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
75-71-8	PSD Dichlorodifluoromethane	50.0	0.00	U 36.1	72	33-133	3	0-20
74-87-3	PSD Chloromethane	50.0	0.00	U 50.0	100	52-142	2	0-20
75-01-4	PSD Vinyl chloride	50.0	0.00	U 47.6	95	54-132	1	0-20
74-83-9	PSD Bromomethane	50.0	0.00	U 48.6	97	63-125	3	0-20
75-00-3	PSD Chloroethane	50.0	0.00	U 50.5	101	71-133	2	0-20
75-69-4	PSD Trichlorofluoromethane	50.0	0.00	U 44.0	88	65-139	2	0-20
60-29-7	PSD Ethyl ether	50.0	0.00	U 45.6	91	70-114	2	0-20
67-64-1	PSD Acetone	250	0.00	U 153	61	30-162	5	0-21
75-05-8	PSD Acetonitrile	1250	0.00	U 1160	93	57-126	5	0-20
75-35-4	PSD 1,1-Dichloroethylene	50.0	0.00	U 51.7	103	68-123	5	0-20
74-88-4	PSD Iodomethane	250	0.00	U 289	116	71-122	3	0-20
75-09-2	PSD Methylene chloride	50.0	0.00	U 47.9	96	74-128	2	0-20
75-15-0	PSD Carbon disulfide	250	0.00	U 302	121	69-130	4	0-20
1634-04-4	PSD tert-Butyl methyl ether	50.0	0.00	U 48.2	96	69-126	0	0-20
156-60-5	PSD trans-1,2-Dichloroethylene	50.0	0.00	U 53.2	106	64-122	2	0-20
108-05-4	PSD Vinyl acetate	250	0.00	U 250	100	49-155	4	0-20
75-34-3	PSD 1,1-Dichloroethane	50.0	0.00	U 52.5	105	74-122	2	0-20
78-93-3	PSD 2-Butanone	250	0.00	U 190	76	16-146	3	0-20
156-59-2	PSD cis-1,2-Dichloroethylene	50.0	0.00	U 52.4	105	66-137	3	0-20
594-20-7	PSD 2,2-Dichloropropane	50.0	0.00	U 56.5	113	68-134	3	0-20
67-66-3	PSD Chloroform	50.0	0.00	U 50.5	101	75-126	2	0-20
74-97-5	PSD Bromochloromethane	50.0	0.00	U 49.6	99	79-126	0	0-20

Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Post Spike Duplicate

Client ID: CAAN-12-2031PSD

Matrix: WG

Lab Sample ID: 1202597762

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:46

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
71-55-6	PSD 1,1,1-Trichloroethane	50.0	0.00	U 54.7	109	70-137	1	0-20
563-58-6	PSD 1,1-Dichloropropene	50.0	0.00	U 55.1	110	74-125	3	0-20
71-36-3	PSD n-Butyl alcohol	5000	0.00	U 4790	96	59-136	3	0-22
56-23-5	PSD Carbon tetrachloride	50.0	0.00	U 56.1	112	70-143	3	0-20
107-06-2	PSD 1,2-Dichloroethane	50.0	0.00	U 48.0	96	70-126	2	0-20
71-43-2	PSD Benzene	50.0	0.00	U 53.1	106	74-119	3	0-20
79-01-6	PSD Trichloroethylene	50.0	0.00	U 54.6	109	69-126	3	0-20
78-87-5	PSD 1,2-Dichloropropane	50.0	0.00	U 53.8	108	76-121	3	0-20
75-27-4	PSD Bromodichloromethane	50.0	0.00	U 51.3	103	77-131	1	0-20
74-95-3	PSD Dibromomethane	50.0	0.00	U 48.2	96	79-123	0	0-20
108-10-1	PSD 4-Methyl-2-pentanone	250	0.00	U 239	96	65-128	3	0-20
10061-01-5	PSD cis-1,3-Dichloropropylene	50.0	0.00	U 53.1	106	77-125	1	0-20
108-88-3	PSD Toluene	50.0	0.00	U 51.7	103	69-119	0	0-20
10061-02-6	PSD trans-1,3-Dichloropropylene	50.0	0.00	U 51.2	102	75-125	1	0-20
79-00-5	PSD 1,1,2-Trichloroethane	50.0	0.00	U 47.6	95	74-119	3	0-20
591-78-6	PSD 2-Hexanone	250	0.00	U 182	73	31-144	5	0-20
142-28-9	PSD 1,3-Dichloropropane	50.0	0.00	U 49.1	98	76-117	1	0-20
127-18-4	PSD Tetrachloroethylene	50.0	0.00	U 55.5	111	65-126	1	0-20
124-48-1	PSD Dibromochloromethane	50.0	0.00	U 50.9	102	74-126	3	0-20
106-93-4	PSD 1,2-Dibromoethane	50.0	0.00	U 47.9	96	78-121	2	0-20
108-90-7	PSD Chlorobenzene	50.0	0.00	U 51.9	104	74-119	2	0-20
100-41-4	PSD Ethylbenzene	50.0	0.00	U 52.5	105	71-121	3	0-20

Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Post Spike Duplicate

Client ID: CAAN-12-2031PSD

Matrix: WG

Lab Sample ID: 1202597762

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:46

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
179601-23-1	PSD m,p-Xylenes	100	0.00	U 107	107	70-123	2	0-23
95-47-6	PSD o-Xylene	50.0	0.00	U 51.5	103	71-125	5	0-20
100-42-5	PSD Styrene	50.0	0.00	U 50.9	102	71-128	4	0-20
75-25-2	PSD Bromoform	50.0	0.00	U 49.2	98	71-126	4	0-20
79-34-5	PSD 1,1,2,2-Tetrachloroethane	50.0	0.00	U 46.5	93	67-126	3	0-20
96-18-4	PSD 1,2,3-Trichloropropane	50.0	0.00	U 46.7	93	76-124	3	0-20
108-86-1	PSD Bromobenzene	50.0	0.00	U 50.8	102	72-120	3	0-20
103-65-1	PSD n-Propylbenzene	50.0	0.00	U 52.8	106	64-125	5	0-23
95-49-8	PSD 2-Chlorotoluene	50.0	0.00	U 53.6	107	68-126	2	0-20
98-82-8	PSD Isopropylbenzene	50.0	0.00	U 51.2	102	68-127	6	0-20
108-67-8	PSD 1,3,5-Trimethylbenzene	50.0	0.00	U 52.6	105	67-128	3	0-21
106-43-4	PSD 4-Chlorotoluene	50.0	0.00	U 49.4	99	67-123	7	0-20
98-06-6	PSD tert-Butylbenzene	50.0	0.00	U 54.3	109	67-130	0	0-20
95-63-6	PSD 1,2,4-Trimethylbenzene	50.0	0.00	U 51.5	103	67-127	2	0-22
135-98-8	PSD sec-Butylbenzene	50.0	0.00	U 52.8	106	66-129	3	0-20
99-87-6	PSD 4-Isopropyltoluene	50.0	0.00	U 53.6	107	63-134	2	0-22
541-73-1	PSD 1,3-Dichlorobenzene	50.0	0.00	U 49.8	100	68-122	4	0-20
106-46-7	PSD 1,4-Dichlorobenzene	50.0	0.00	U 50.3	101	67-121	1	0-20
104-51-8	PSD n-Butylbenzene	50.0	0.00	U 50.8	102	59-133	5	0-20
96-12-8	PSD 1,2-Dibromo-3-chloropropane	50.0	0.00	U 46.9	94	62-129	4	0-20
87-68-3	PSD Hexachlorobutadiene	50.0	0.00	U 52.8	106	52-137	2	0-20
91-20-3	PSD Naphthalene	50.0	0.00	U 43.5	87	63-129	1	0-20

Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Post Spike Duplicate

Client ID: CAAN-12-2031PSD

Matrix: WG

Lab Sample ID:1202597762

Instrument: VOA9.I

Analysis Date: 02/12/2012 20:46

Dilution: 1

Analyst: GRB2

Prep Batch ID 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
87-61-6	PSD 1,2,3-Trichlorobenzene	50.0	0.00 U	47.3	95	59-131	6	0-22
630-20-6	PSD 1,1,1,2-Tetrachloroethane	50.0	0.00 U	54.3	109	76-128	0	0-20
120-82-1	PSD 1,2,4-Trichlorobenzene	50.0	0.00 U	46.2	92	59-127	4	0-22
95-50-1	PSD 1,2-Dichlorobenzene	50.0	0.00 U	49.6	99	72-119	0	0-20

Volatile
Quality Control Summary
Spike Recovery Report

Page 1 of 4

SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1187820

Matrix: WATER

Lab Sample ID: 1202597763

Instrument: VOA9.I

Analysis Date: 02/12/2012 12:51

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
75-71-8	LCS Dichlorodifluoromethane	50.0	0.0	34.0	68	40-136
74-87-3	LCS Chloromethane	50.0	0.0	48.5	97	62-136
75-01-4	LCS Vinyl chloride	50.0	0.0	46.8	94	64-128
74-83-9	LCS Bromomethane	50.0	0.0	47.1	94	69-122
75-00-3	LCS Chloroethane	50.0	0.0	48.1	96	79-131
75-69-4	LCS Trichlorofluoromethane	50.0	0.0	42.3	85	74-138
60-29-7	LCS Ethyl ether	50.0	0.0	43.2	86	73-120
67-64-1	LCS Acetone	250	0.0	324	130	42-164
75-05-8	LCS Acetonitrile	1250	0.0	1320	106	62-120
75-35-4	LCS 1,1-Dichloroethylene	50.0	0.0	49.7	99	74-122
74-88-4	LCS Iodomethane	250	0.0	277	111	76-120
75-09-2	LCS Methylene chloride	50.0	0.0	45.1	90	80-124
75-15-0	LCS Carbon disulfide	250	0.0	300	120	74-130
1634-04-4	LCS tert-Butyl methyl ether	50.0	0.0	47.6	95	73-122
156-60-5	LCS trans-1,2-Dichloroethylene	50.0	0.0	50.9	102	67-123
108-05-4	LCS Vinyl acetate	250	0.0	279	111	70-150
75-34-3	LCS 1,1-Dichloroethane	50.0	0.0	50.8	102	80-120
78-93-3	LCS 2-Butanone	250	0.0	296	118	45-153
156-59-2	LCS cis-1,2-Dichloroethylene	50.0	0.0	51.0	102	71-134
594-20-7	LCS 2,2-Dichloropropane	50.0	0.0	55.1	110	75-137
67-66-3	LCS Chloroform	50.0	0.0	49.4	99	80-122
74-97-5	LCS Bromochloromethane	50.0	0.0	47.9	96	80-123

Volatile
Quality Control Summary
Spike Recovery Report

Page 2 of 4

SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1187820

Matrix: WATER

Lab Sample ID: 1202597763

Instrument: VOA9.I

Analysis Date: 02/12/2012 12:51

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
71-55-6	LCS 1,1,1-Trichloroethane	50.0	0.0	52.7	105	75-136
563-58-6	LCS 1,1-Dichloropropene	50.0	0.0	53.7	107	80-124
71-36-3	LCS n-Butyl alcohol	5000	0.0	5540	111	64-133
56-23-5	LCS Carbon tetrachloride	50.0	0.0	53.0	106	75-144
107-06-2	LCS 1,2-Dichloroethane	50.0	0.0	47.7	95	74-121
71-43-2	LCS Benzene	50.0	0.0	50.5	101	80-120
79-01-6	LCS Trichloroethylene	50.0	0.0	51.4	103	80-120
78-87-5	LCS 1,2-Dichloropropane	50.0	0.0	51.6	103	80-120
75-27-4	LCS Bromodichloromethane	50.0	0.0	48.8	98	80-128
74-95-3	LCS Dibromomethane	50.0	0.0	48.0	96	80-120
108-10-1	LCS 4-Methyl-2-pentanone	250	0.0	267	107	70-125
10061-01-5	LCS cis-1,3-Dichloropropylene	50.0	0.0	51.1	102	80-124
108-88-3	LCS Toluene	50.0	0.0	47.2	94	77-120
10061-02-6	LCS trans-1,3-Dichloropropylene	50.0	0.0	49.1	98	80-124
79-00-5	LCS 1,1,2-Trichloroethane	50.0	0.0	46.6	93	77-120
591-78-6	LCS 2-Hexanone	250	0.0	296	118	53-149
142-28-9	LCS 1,3-Dichloropropane	50.0	0.0	47.9	96	78-120
127-18-4	LCS Tetrachloroethylene	50.0	0.0	50.3	101	75-124
124-48-1	LCS Dibromochloromethane	50.0	0.0	48.3	97	77-125
106-93-4	LCS 1,2-Dibromoethane	50.0	0.0	46.6	93	80-120
108-90-7	LCS Chlorobenzene	50.0	0.0	48.8	98	80-120
100-41-4	LCS Ethylbenzene	50.0	0.0	49.6	99	80-120

Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1187820

Matrix: WATER

Lab Sample ID: 1202597763

Instrument: VOA9.I

Analysis Date: 02/12/2012 12:51

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
179601-23-1	LCS m,p-Xylenes	100	0.0	101	101	80-120
95-47-6	LCS o-Xylene	50.0	0.0	49.4	99	80-121
100-42-5	LCS Styrene	50.0	0.0	49.6	99	79-125
75-25-2	LCS Bromoform	50.0	0.0	49.2	98	73-126
79-34-5	LCS 1,1,2,2-Tetrachloroethane	50.0	0.0	47.1	94	73-122
96-18-4	LCS 1,2,3-Trichloropropane	50.0	0.0	48.5	97	78-121
108-86-1	LCS Bromobenzene	50.0	0.0	48.9	98	79-120
103-65-1	LCS n-Propylbenzene	50.0	0.0	51.2	102	76-121
95-49-8	LCS 2-Chlorotoluene	50.0	0.0	50.3	101	80-120
98-82-8	LCS Isopropylbenzene	50.0	0.0	49.2	98	78-124
108-67-8	LCS 1,3,5-Trimethylbenzene	50.0	0.0	49.7	99	79-123
106-43-4	LCS 4-Chlorotoluene	50.0	0.0	49.7	99	78-120
98-06-6	LCS tert-Butylbenzene	50.0	0.0	49.4	99	79-125
95-63-6	LCS 1,2,4-Trimethylbenzene	50.0	0.0	49.7	99	79-121
135-98-8	LCS sec-Butylbenzene	50.0	0.0	50.0	100	79-124
99-87-6	LCS 4-Isopropyltoluene	50.0	0.0	50.0	100	80-128
541-73-1	LCS 1,3-Dichlorobenzene	50.0	0.0	48.6	97	80-120
106-46-7	LCS 1,4-Dichlorobenzene	50.0	0.0	48.7	97	79-120
104-51-8	LCS n-Butylbenzene	50.0	0.0	49.8	100	78-127
96-12-8	LCS 1,2-Dibromo-3-chloropropane	50.0	0.0	47.6	95	66-126
87-68-3	LCS Hexachlorobutadiene	50.0	0.0	48.6	97	71-132
91-20-3	LCS Naphthalene	50.0	0.0	43.8	88	68-127

Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1187820

Matrix: WATER

Lab Sample ID: 1202597763

Instrument: VOA9.I

Analysis Date: 02/12/2012 12:51

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
87-61-6	LCS 1,2,3-Trichlorobenzene	50.0	0.0	45.7	91	70-129
630-20-6	LCS 1,1,1,2-Tetrachloroethane	50.0	0.0	49.8	100	80-125
120-82-1	LCS 1,2,4-Trichlorobenzene	50.0	0.0	47.3	95	74-126
95-50-1	LCS 1,2-Dichlorobenzene	50.0	0.0	46.6	93	80-120

Volatile
Quality Control Summary
Spike Recovery Report

Page 1 of 1

SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1187820

Matrix: WATER

Lab Sample ID: 1202597764

Instrument: VOA9.I

Analysis Date: 02/12/2012 13:17

Dilution: 1

Analyst: GRB2

Prep Batch ID: 1187820

Purge Vol: 5 mL

Batch ID: 1187820

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
107-02-8	LCS Acrolein	250	0.0	248	99	60-137
76-13-1	LCS Trichlorotrifluoroethane	250	0.0	293	117	66-140
107-05-1	LCS Allyl chloride	250	0.0	241	96	59-137
107-13-1	LCS Acrylonitrile	250	0.0	254	102	69-120
126-99-8	LCS 2-Chloro-1,3-butadiene	50.0	0.0	41.9	84	61-138
107-12-0	LCS Propionitrile	250	0.0	274	110	69-120
126-98-7	LCS Methacrylonitrile	250	0.0	259	104	67-120
78-83-1	LCS Isobutyl alcohol	2500	0.0	2580	103	60-126
80-62-6	LCS Methyl methacrylate	250	0.0	256	102	73-120
97-63-2	LCS Ethyl methacrylate	250	0.0	247	99	71-120

Method Blank Summary

Page 1 of 1

SDG Number:	12-700	Client:	ARSL001	Matrix:	WATER
Client ID:	MB for batch 1187820	Instrument ID:	VOA9.I	Data File:	021212V9\9Y710BAR.D
Lab Sample ID:	1202597760	Prep Date:	02/12/2012 14:50	Analyzed:	02/12/12 14:50
Column:	DB-624				

This method blank applies to the following samples and quality control samples:

Client Sample ID	Lab Sample ID	File ID	Date Analyzed	Time Analyzed
01 LCS for batch 1187820	1202597763	021212V9\9Y706LAR.D	02/12/12	1251
02 LCS for batch 1187820	1202597764	021212V9\9Y707SHAR.D	02/12/12	1317
03 CAAN-12-2031	295270002	021212V9\9Y711.D	02/12/12	1530
04 CAAN-12-2199	295270004	021212V9\9Y712.D	02/12/12	1556
05 CAAN-12-2201	295270007	021212V9\9Y713.D	02/12/12	1623
06 CAAN-12-2032	295270008	021212V9\9Y714.D	02/12/12	1649
07 CAAN-12-2031PS	1202597761	021212V9\9Y722.D	02/12/12	2020
08 CAAN-12-2031PSD	1202597762	021212V9\9Y723.D	02/12/12	2046

Quality Control Data

Volatile
Certificate of Analysis
Sample Summary

SDG Number: 12-700		Matrix:	WATER
Lab Sample ID: 1202597760			
Client Sample: QC for batch 1187820	Client: ARSL001	Project:	QC
Client ID: MB for batch 1187820	Method: SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution:	1
Run Date: 02/12/2012 14:50	Analyst: GRB2	Purge Vol:	5 mL
Prep Date: 02/12/2012 14:50			
Data File: 021212V9\9Y710BAR.D	Column: DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

Volatile
Certificate of Analysis
Sample Summary

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SDG Number: 12-700	Matrix: WATER
Lab Sample ID: 1202597760	
Client Sample: QC for batch 1187820	Client: ARSL001
Client ID: MB for batch 1187820	Method: SW846 8260B DOE-AL
Batch ID: 1187820	Project: QC
Run Date: 02/12/2012 14:50	SOP Ref: GL-OA-E-038
Prep Date: 02/12/2012 14:50	Dilution: 1
Data File: 021212V9\9Y710BAR.D	Purge Vol: 5 mL
	Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

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Certificate of Analysis
Sample Summary

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SDG Number:	12-700	Matrix:	WATER
Lab Sample ID:	1202597760		
Client Sample:	QC for batch 1187820	Client:	ARSL001
Client ID:	MB for batch 1187820	Method:	SW846 8260B DOE-AL
Batch ID:	1187820	Inst:	VOA9.I
Run Date:	02/12/2012 14:50	Analyst:	GRB2
Prep Date:	02/12/2012 14:50	Purge Vol:	5 mL
Data File:	021212V9\9Y710BAR.D	Column:	DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	44.1	50.0	ug/L 88.1	(76%-127%)
Bromofluorobenzene	53.2	50.0	ug/L 106	(80%-120%)
Toluene-d8	48.1	50.0	ug/L 96.3	(80%-120%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	unknown	4.29	25.4	ug/L	0	J

Volatile
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Sample Summary

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 1202597761	Date Received: 02/03/2012 08:45	
Client Sample: QC for batch 1187820	Client: ARSL001	Project: QC
Client ID: CAAN-12-2031PS	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 20:20	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 20:20		
Data File: 021212V9\9Y722.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane		37.3	ug/L	0.300	1.00
74-87-3	Chloromethane		51.2	ug/L	0.300	1.00
75-01-4	Vinyl chloride		48.2	ug/L	0.500	1.00
74-83-9	Bromomethane		50.2	ug/L	0.300	1.00
75-00-3	Chloroethane		49.6	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane		44.9	ug/L	0.300	1.00
60-29-7	Ethyl ether		44.8	ug/L	0.300	1.00
67-64-1	Acetone		162	ug/L	3.50	10.0
75-05-8	Acetonitrile		1220	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene		54.3	ug/L	0.300	1.00
74-88-4	Iodomethane		298	ug/L	1.25	5.00
75-09-2	Methylene chloride		47.2	ug/L	3.00	10.0
75-15-0	Carbon disulfide		314	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether		48.3	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene		54.2	ug/L	0.300	1.00
108-05-4	Vinyl acetate		261	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane		53.3	ug/L	0.300	1.00
78-93-3	2-Butanone		197	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene		53.7	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane		58.0	ug/L	0.300	1.00
67-66-3	Chloroform		51.7	ug/L	0.250	1.00
74-97-5	Bromochloromethane		49.4	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane		55.3	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene		56.9	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol		4940	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride		57.9	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane		48.9	ug/L	0.250	1.00
71-43-2	Benzene		54.6	ug/L	0.300	1.00
79-01-6	Trichloroethylene		56.5	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane		55.7	ug/L	0.250	1.00
75-27-4	Bromodichloromethane		50.8	ug/L	0.250	1.00
74-95-3	Dibromomethane		48.4	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone		247	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene		52.7	ug/L	0.250	1.00
108-88-3	Toluene		51.9	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene		51.8	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane		49.0	ug/L	0.250	1.00
591-78-6	2-Hexanone		191	ug/L	1.25	5.00

Volatile
Certificate of Analysis
Sample Summary

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SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 1202597761	Date Received: 02/03/2012 08:45	
Client Sample: QC for batch 1187820	Client: ARSL001	Project: QC
Client ID: CAAN-12-2031PS	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 20:20	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 20:20		
Data File: 021212V9\9Y722.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane		49.7	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene		56.1	ug/L	0.300	1.00
124-48-1	Dibromochloromethane		52.5	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane		49.0	ug/L	0.250	1.00
108-90-7	Chlorobenzene		53.1	ug/L	0.250	1.00
100-41-4	Ethylbenzene		54.0	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes		109	ug/L	0.500	2.00
95-47-6	o-Xylene		53.9	ug/L	0.300	1.00
100-42-5	Styrene		52.9	ug/L	0.250	1.00
75-25-2	Bromoform		51.4	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane		48.0	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane		48.0	ug/L	0.300	1.00
108-86-1	Bromobenzene		52.1	ug/L	0.250	1.00
103-65-1	n-Propylbenzene		55.3	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene		54.8	ug/L	0.250	1.00
98-82-8	Isopropylbenzene		54.5	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene		54.2	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene		52.8	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene		54.3	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene		52.7	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene		54.4	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene		54.7	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene		51.7	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene		50.8	ug/L	0.250	1.00
104-51-8	n-Butylbenzene		53.6	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane		48.7	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene		54.1	ug/L	0.300	1.00
91-20-3	Naphthalene		43.0	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene		44.7	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	1202597761	Date Received:	02/03/2012 08:45		
Client Sample:	QC for batch 1187820	Client:	ARSL001	Project:	QC
Client ID:	CAAN-12-2031PS	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 20:20	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 20:20				
Data File:	021212V9\9Y722.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane		54.4	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene		47.9	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene		49.7	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	42.6	50.0	85.2	(76%-127%)
Bromofluorobenzene	50.9	50.0	102	(80%-120%)
Toluene-d8	46.5	50.0	93.1	(80%-120%)

**Volatile
Certificate of Analysis
Sample Summary**

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	1202597762	Date Received:	02/03/2012 08:45		
Client Sample:	QC for batch 1187820	Client:	ARSL001	Project:	QC
Client ID:	CAAN-12-2031PSD	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 20:46	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 20:46				
Data File:	021212V9\9Y723.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane		36.1	ug/L	0.300	1.00
74-87-3	Chloromethane		50.0	ug/L	0.300	1.00
75-01-4	Vinyl chloride		47.6	ug/L	0.500	1.00
74-83-9	Bromomethane		48.6	ug/L	0.300	1.00
75-00-3	Chloroethane		50.5	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane		44.0	ug/L	0.300	1.00
60-29-7	Ethyl ether		45.6	ug/L	0.300	1.00
67-64-1	Acetone		153	ug/L	3.50	10.0
75-05-8	Acetonitrile		1160	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene		51.7	ug/L	0.300	1.00
74-88-4	Iodomethane		289	ug/L	1.25	5.00
75-09-2	Methylene chloride		47.9	ug/L	3.00	10.0
75-15-0	Carbon disulfide		302	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether		48.2	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene		53.2	ug/L	0.300	1.00
108-05-4	Vinyl acetate		250	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane		52.5	ug/L	0.300	1.00
78-93-3	2-Butanone		190	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene		52.4	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane		56.5	ug/L	0.300	1.00
67-66-3	Chloroform		50.5	ug/L	0.250	1.00
74-97-5	Bromochloromethane		49.6	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane		54.7	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene		55.1	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol		4790	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride		56.1	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane		48.0	ug/L	0.250	1.00
71-43-2	Benzene		53.1	ug/L	0.300	1.00
79-01-6	Trichloroethylene		54.6	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane		53.8	ug/L	0.250	1.00
75-27-4	Bromodichloromethane		51.3	ug/L	0.250	1.00
74-95-3	Dibromomethane		48.2	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone		239	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene		53.1	ug/L	0.250	1.00
108-88-3	Toluene		51.7	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene		51.2	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane		47.6	ug/L	0.250	1.00
591-78-6	2-Hexanone		182	ug/L	1.25	5.00

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SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 1202597762	Date Received: 02/03/2012 08:45	
Client Sample: QC for batch 1187820	Client: ARSL001	Project: QC
Client ID: CAAN-12-2031PSD	Method: SW846 8260B DOE-AL	SOP Ref: GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution: 1
Run Date: 02/12/2012 20:46	Analyst: GRB2	Purge Vol: 5 mL
Prep Date: 02/12/2012 20:46		
Data File: 021212V9\9Y723.D	Column: DB-624	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane		49.1	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene		55.5	ug/L	0.300	1.00
124-48-1	Dibromochloromethane		50.9	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane		47.9	ug/L	0.250	1.00
108-90-7	Chlorobenzene		51.9	ug/L	0.250	1.00
100-41-4	Ethylbenzene		52.5	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes		107	ug/L	0.500	2.00
95-47-6	o-Xylene		51.5	ug/L	0.300	1.00
100-42-5	Styrene		50.9	ug/L	0.250	1.00
75-25-2	Bromoform		49.2	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane		46.5	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane		46.7	ug/L	0.300	1.00
108-86-1	Bromobenzene		50.8	ug/L	0.250	1.00
103-65-1	n-Propylbenzene		52.8	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene		53.6	ug/L	0.250	1.00
98-82-8	Isopropylbenzene		51.2	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene		52.6	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene		49.4	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene		54.3	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene		51.5	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene		52.8	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene		53.6	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene		49.8	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene		50.3	ug/L	0.250	1.00
104-51-8	n-Butylbenzene		50.8	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane		46.9	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene		52.8	ug/L	0.300	1.00
91-20-3	Naphthalene		43.5	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene		47.3	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

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SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	1202597762	Date Received:	02/03/2012 08:45		
Client Sample:	QC for batch 1187820	Client:	ARSL001	Project:	QC
Client ID:	CAAN-12-2031PSD	Method:	SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID:	1187820	Inst:	VOA9.I	Dilution:	1
Run Date:	02/12/2012 20:46	Analyst:	GRB2	Purge Vol:	5 mL
Prep Date:	02/12/2012 20:46				
Data File:	021212V9\9Y723.D	Column:	DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane		54.3	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene		46.2	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene		49.6	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	41.9	50.0	83.7	(76%-127%)
Bromofluorobenzene	48.7	50.0	97.4	(80%-120%)
Toluene-d8	47.0	50.0	93.9	(80%-120%)

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SDG Number: 12-700	Matrix: WATER
Lab Sample ID: 1202597763	
Client Sample: QC for batch 1187820	Client: ARSL001
Client ID: LCS for batch 1187820	Method: SW846 8260B DOE-AL
Batch ID: 1187820	Project: QC
Run Date: 02/12/2012 12:51	SOP Ref: GL-OA-E-038
Prep Date: 02/12/2012 12:51	Dilution: 1
Data File: 021212V9\9Y706LAR.D	Purge Vol: 5 mL
	Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane		34.0	ug/L	0.300	1.00
74-87-3	Chloromethane		48.5	ug/L	0.300	1.00
75-01-4	Vinyl chloride		46.8	ug/L	0.500	1.00
74-83-9	Bromomethane		47.1	ug/L	0.300	1.00
75-00-3	Chloroethane		48.1	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane		42.3	ug/L	0.300	1.00
60-29-7	Ethyl ether		43.2	ug/L	0.300	1.00
67-64-1	Acetone		324	ug/L	3.50	10.0
75-05-8	Acetonitrile		1320	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene		49.7	ug/L	0.300	1.00
74-88-4	Iodomethane		277	ug/L	1.25	5.00
75-09-2	Methylene chloride		45.1	ug/L	3.00	10.0
75-15-0	Carbon disulfide		300	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether		47.6	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene		50.9	ug/L	0.300	1.00
108-05-4	Vinyl acetate		279	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane		50.8	ug/L	0.300	1.00
78-93-3	2-Butanone		296	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene		51.0	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane		55.1	ug/L	0.300	1.00
67-66-3	Chloroform		49.4	ug/L	0.250	1.00
74-97-5	Bromochloromethane		47.9	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane		52.7	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene		53.7	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol		5540	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride		53.0	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane		47.7	ug/L	0.250	1.00
71-43-2	Benzene		50.5	ug/L	0.300	1.00
79-01-6	Trichloroethylene		51.4	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane		51.6	ug/L	0.250	1.00
75-27-4	Bromodichloromethane		48.8	ug/L	0.250	1.00
74-95-3	Dibromomethane		48.0	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone		267	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene		51.1	ug/L	0.250	1.00
108-88-3	Toluene		47.2	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene		49.1	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane		46.6	ug/L	0.250	1.00
591-78-6	2-Hexanone		296	ug/L	1.25	5.00

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SDG Number: 12-700		Matrix:	WATER
Lab Sample ID: 1202597763			
Client Sample: QC for batch 1187820	Client: ARSL001	Project:	QC
Client ID: LCS for batch 1187820	Method: SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution:	1
Run Date: 02/12/2012 12:51	Analyst: GRB2	Purge Vol:	5 mL
Prep Date: 02/12/2012 12:51			
Data File: 021212V9\9Y706LAR.D	Column: DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane		47.9	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene		50.3	ug/L	0.300	1.00
124-48-1	Dibromochloromethane		48.3	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane		46.6	ug/L	0.250	1.00
108-90-7	Chlorobenzene		48.8	ug/L	0.250	1.00
100-41-4	Ethylbenzene		49.6	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes		101	ug/L	0.500	2.00
95-47-6	o-Xylene		49.4	ug/L	0.300	1.00
100-42-5	Styrene		49.6	ug/L	0.250	1.00
75-25-2	Bromoform		49.2	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane		47.1	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane		48.5	ug/L	0.300	1.00
108-86-1	Bromobenzene		48.9	ug/L	0.250	1.00
103-65-1	n-Propylbenzene		51.2	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene		50.3	ug/L	0.250	1.00
98-82-8	Isopropylbenzene		49.2	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene		49.7	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene		49.7	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene		49.4	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene		49.7	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene		50.0	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene		50.0	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene		48.6	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene		48.7	ug/L	0.250	1.00
104-51-8	n-Butylbenzene		49.8	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane		47.6	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene		48.6	ug/L	0.300	1.00
91-20-3	Naphthalene		43.8	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene		45.7	ug/L	0.332	1.00
107-02-8	Acrolein	U	5.00	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane	U	5.00	ug/L	1.00	5.00
107-05-1	Allyl chloride	U	5.00	ug/L	1.50	5.00
107-13-1	Acrylonitrile	U	5.00	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene	U	1.00	ug/L	0.300	1.00
107-12-0	Propionitrile	U	5.00	ug/L	1.50	5.00
126-98-7	Methacrylonitrile	U	5.00	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol	U	50.0	ug/L	12.5	50.0
80-62-6	Methyl methacrylate	U	5.00	ug/L	1.00	5.00

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SDG Number:	12-700	Matrix:	WATER
Lab Sample ID:	1202597763		
Client Sample:	QC for batch 1187820	Client:	ARSL001
Client ID:	LCS for batch 1187820	Method:	SW846 8260B DOE-AL
Batch ID:	1187820	Inst:	VOA9.I
Run Date:	02/12/2012 12:51	Analyst:	GRB2
Prep Date:	02/12/2012 12:51	Purge Vol:	5 mL
Data File:	021212V9\9Y706LAR.D	Column:	DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate	U	5.00	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane		49.8	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene		47.3	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene		46.6	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	43.8	50.0	87.6	(76%-127%)
Bromofluorobenzene	50.3	50.0	101	(80%-120%)
Toluene-d8	45.4	50.0	90.9	(80%-120%)

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SDG Number: 12-700		Matrix:	WATER
Lab Sample ID: 1202597764			
Client Sample: QC for batch 1187820	Client: ARSL001	Project:	QC
Client ID: LCS for batch 1187820	Method: SW846 8260B DOE-AL	SOP Ref:	GL-OA-E-038
Batch ID: 1187820	Inst: VOA9.I	Dilution:	1
Run Date: 02/12/2012 13:17	Analyst: GRB2	Purge Vol:	5 mL
Prep Date: 02/12/2012 13:17			
Data File: 021212V9\9Y707SHAR.D	Column: DB-624		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
75-71-8	Dichlorodifluoromethane	U	1.00	ug/L	0.300	1.00
74-87-3	Chloromethane	U	1.00	ug/L	0.300	1.00
75-01-4	Vinyl chloride	U	1.00	ug/L	0.500	1.00
74-83-9	Bromomethane	U	1.00	ug/L	0.300	1.00
75-00-3	Chloroethane	U	1.00	ug/L	0.300	1.00
75-69-4	Trichlorofluoromethane	U	1.00	ug/L	0.300	1.00
60-29-7	Ethyl ether	U	1.00	ug/L	0.300	1.00
67-64-1	Acetone	U	10.0	ug/L	3.50	10.0
75-05-8	Acetonitrile	U	25.0	ug/L	6.25	25.0
75-35-4	1,1-Dichloroethylene	U	1.00	ug/L	0.300	1.00
74-88-4	Iodomethane	U	5.00	ug/L	1.25	5.00
75-09-2	Methylene chloride	U	10.0	ug/L	3.00	10.0
75-15-0	Carbon disulfide	U	5.00	ug/L	1.25	5.00
1634-04-4	tert-Butyl methyl ether	U	1.00	ug/L	0.250	1.00
156-60-5	trans-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
108-05-4	Vinyl acetate	U	5.00	ug/L	1.50	5.00
75-34-3	1,1-Dichloroethane	U	1.00	ug/L	0.300	1.00
78-93-3	2-Butanone	U	5.00	ug/L	1.25	5.00
156-59-2	cis-1,2-Dichloroethylene	U	1.00	ug/L	0.300	1.00
594-20-7	2,2-Dichloropropane	U	1.00	ug/L	0.300	1.00
67-66-3	Chloroform	U	1.00	ug/L	0.250	1.00
74-97-5	Bromochloromethane	U	1.00	ug/L	0.300	1.00
71-55-6	1,1,1-Trichloroethane	U	1.00	ug/L	0.325	1.00
563-58-6	1,1-Dichloropropene	U	1.00	ug/L	0.250	1.00
71-36-3	n-Butyl alcohol	U	50.0	ug/L	15.0	50.0
56-23-5	Carbon tetrachloride	U	1.00	ug/L	0.300	1.00
107-06-2	1,2-Dichloroethane	U	1.00	ug/L	0.250	1.00
71-43-2	Benzene	U	1.00	ug/L	0.300	1.00
79-01-6	Trichloroethylene	U	1.00	ug/L	0.250	1.00
78-87-5	1,2-Dichloropropane	U	1.00	ug/L	0.250	1.00
75-27-4	Bromodichloromethane	U	1.00	ug/L	0.250	1.00
74-95-3	Dibromomethane	U	1.00	ug/L	0.300	1.00
108-10-1	4-Methyl-2-pentanone	U	5.00	ug/L	1.25	5.00
10061-01-5	cis-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
108-88-3	Toluene	U	1.00	ug/L	0.250	1.00
10061-02-6	trans-1,3-Dichloropropylene	U	1.00	ug/L	0.250	1.00
79-00-5	1,1,2-Trichloroethane	U	1.00	ug/L	0.250	1.00
591-78-6	2-Hexanone	U	5.00	ug/L	1.25	5.00

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SDG Number: 12-700	Matrix: WATER
Lab Sample ID: 1202597764	
Client Sample: QC for batch 1187820	Client: ARSL001
Client ID: LCS for batch 1187820	Method: SW846 8260B DOE-AL
Batch ID: 1187820	Project: QC
Run Date: 02/12/2012 13:17	SOP Ref: GL-OA-E-038
Prep Date: 02/12/2012 13:17	Dilution: 1
Data File: 021212V9\9Y707SHAR.D	Purge Vol: 5 mL
	Column: DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
142-28-9	1,3-Dichloropropane	U	1.00	ug/L	0.300	1.00
127-18-4	Tetrachloroethylene	U	1.00	ug/L	0.300	1.00
124-48-1	Dibromochloromethane	U	1.00	ug/L	0.300	1.00
106-93-4	1,2-Dibromoethane	U	1.00	ug/L	0.250	1.00
108-90-7	Chlorobenzene	U	1.00	ug/L	0.250	1.00
100-41-4	Ethylbenzene	U	1.00	ug/L	0.250	1.00
179601-23-1	m,p-Xylenes	U	2.00	ug/L	0.500	2.00
95-47-6	o-Xylene	U	1.00	ug/L	0.300	1.00
100-42-5	Styrene	U	1.00	ug/L	0.250	1.00
75-25-2	Bromoform	U	1.00	ug/L	0.250	1.00
79-34-5	1,1,2,2-Tetrachloroethane	U	1.00	ug/L	0.250	1.00
96-18-4	1,2,3-Trichloropropane	U	1.00	ug/L	0.300	1.00
108-86-1	Bromobenzene	U	1.00	ug/L	0.250	1.00
103-65-1	n-Propylbenzene	U	1.00	ug/L	0.250	1.00
95-49-8	2-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-82-8	Isopropylbenzene	U	1.00	ug/L	0.250	1.00
108-67-8	1,3,5-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
106-43-4	4-Chlorotoluene	U	1.00	ug/L	0.250	1.00
98-06-6	tert-Butylbenzene	U	1.00	ug/L	0.250	1.00
95-63-6	1,2,4-Trimethylbenzene	U	1.00	ug/L	0.250	1.00
135-98-8	sec-Butylbenzene	U	1.00	ug/L	0.250	1.00
99-87-6	4-Isopropyltoluene	U	1.00	ug/L	0.250	1.00
541-73-1	1,3-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
106-46-7	1,4-Dichlorobenzene	U	1.00	ug/L	0.250	1.00
104-51-8	n-Butylbenzene	U	1.00	ug/L	0.250	1.00
96-12-8	1,2-Dibromo-3-chloropropane	U	1.00	ug/L	0.300	1.00
87-68-3	Hexachlorobutadiene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.250	1.00
87-61-6	1,2,3-Trichlorobenzene	U	1.00	ug/L	0.332	1.00
107-02-8	Acrolein		248	ug/L	1.25	5.00
76-13-1	Trichlorotrifluoroethane		293	ug/L	1.00	5.00
107-05-1	Allyl chloride		241	ug/L	1.50	5.00
107-13-1	Acrylonitrile		254	ug/L	1.00	5.00
126-99-8	2-Chloro-1,3-butadiene		41.9	ug/L	0.300	1.00
107-12-0	Propionitrile		274	ug/L	1.50	5.00
126-98-7	Methacrylonitrile		259	ug/L	1.00	5.00
78-83-1	Isobutyl alcohol		2580	ug/L	12.5	50.0
80-62-6	Methyl methacrylate		256	ug/L	1.00	5.00

**Volatile
Certificate of Analysis
Sample Summary**

Page 3 of 3

SDG Number:	12-700	Matrix:	WATER
Lab Sample ID:	1202597764		
Client Sample:	QC for batch 1187820	Client:	ARSL001
Client ID:	LCS for batch 1187820	Method:	SW846 8260B DOE-AL
Batch ID:	1187820	Inst:	VOA9.I
Run Date:	02/12/2012 13:17	SOP Ref:	GL-OA-E-038
Prep Date:	02/12/2012 13:17	Dilution:	1
Data File:	021212V9\9Y707SHAR.D	Purge Vol:	5 mL
		Column:	DB-624

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
97-63-2	Ethyl methacrylate		247	ug/L	1.00	5.00
630-20-6	1,1,1,2-Tetrachloroethane	U	1.00	ug/L	0.300	1.00
120-82-1	1,2,4-Trichlorobenzene	U	1.00	ug/L	0.300	1.00
95-50-1	1,2-Dichlorobenzene	U	1.00	ug/L	0.250	1.00

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
1,2-Dichloroethane-d4	43.1	50.0	ug/L 86.1	(76%-127%)
Bromofluorobenzene	53.5	50.0	ug/L 107	(80%-120%)
Toluene-d8	45.8	50.0	ug/L 91.6	(80%-120%)

Semi-Volatile Analysis

Case Narrative

**Semi-Volatile Case Narrative
ARS International (ARSL)
SDG 12-700**

Method/Analysis Information

Procedure: **Analysis of Semivolatile Organic Compounds by Gas Chromatography/Mass Spectrometry**

Analytical Method: SW846 8270C

Prep Method: SW846 3510C

Analytical Batch Number: 1186837

Prep Batch Number: 1186831

Sample Analysis

The following samples were analyzed using the analytical protocol as established in SW846 8270C:

Sample ID	Client ID
295270002	CAAN-12-2031
295270004	CAAN-12-2199
295270007	CAAN-12-2201
1202595534	Method Blank (MB)
1202595535	Laboratory Control Sample (LCS)
1202595536	295270002(CAAN-12-2031) Matrix Spike (MS)
1202595537	295270002(CAAN-12-2031) Matrix Spike Duplicate (MSD)

The samples in this SDG were analyzed on an "as received" basis.

Preparation/Analytical Method Verification

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-OA-E-009 REV# 27.

Raw data reports are processed and reviewed by the analyst using the data analysis software package. False positives have been removed from the quantitation reports per standard operating procedures (SOP) section 18.2.

Calibration Information

Due to software limitations, the Form 6 may not indicate all the calibration files comprising the initial calibration. A complete list of the initial calibration data files are shown in the Calibration History report located in the Standard Data section of the data package. The various calibration mixes may not be calibrated using all of the calibration levels. In addition, not all of the mixes are calibrated using the same levels.

Diphenylamine has now superseded N-Nitroso-diphenylamine on Quantitation Reports, Initial Calibration Reports, Calibration Check Standard Reports, etc. Previous versions of EPA Methodologies referenced N-Nitroso-diphenylamine. However, as stated in EPA Methodology, "N-Nitroso-diphenylamine decomposes in the gas chromatographic inlet and cannot be separated from Diphenylamine." Studies of these two compounds at GEL, both independent of each other and together, showed that they not only co-elute, but also have similar mass spectra. N-Nitroso-diphenylamine and Diphenylamine will be reported as Diphenylamine on all reports and forms.

Initial Calibration

All initial calibration requirements have been met for this sample delivery group (SDG) in this batch. A second source initial calibration verification (ICV) was included in the standard section directly behind the initial calibration.

CCV Requirements

All associated calibration verification standard(s) (ICV or CCV) met the acceptance criteria.

Quality Control (QC) Information

Method Blank (MB) Statement

The MB analyzed with this SDG in this batch met the acceptance criteria.

Surrogate Recoveries

The MSD(1202595537(CAAN-12-2031)) displayed one biased high surrogate. Please see the QC Summary for the specific recovery value. The MB, LCS and MS satisfied batch QC acceptance criteria (with the exception of Benzidine in the MS and MSD - see Spike Recovery Statements). Therefore, it was determined that the biased high failures were limited to the MSD sample only (the surrogate and spike recoveries were in general significantly higher in the MSD compared to the MS recoveries). Since the associated parent sample was reported as non-detected for all requested target analytes, a re-extraction was considered unnecessary and the data have been reported. Please see data exception report 1048967.

Laboratory Control Sample (LCS) Recovery

The LCS(1202595535) recovered Benzidine at 17.3% (limits: 23%-131%) and Benzoic acid 28.7% (limits: 30%-142%). The failures represented less than 5% of the total requested spike analyte list. That satisfied the acceptance criteria for the client and the data were reported. Please note, both Benzidine and Benzoic acid are designated as being poor responding analytes in the analytical method. This may account for the biased low recoveries for those analytes. Please see data exception report 1048967.

QC Sample Designation

Sample 295270002 (CAAN-12-2031) was selected for analysis as the matrix spike and matrix spike duplicate.

Matrix Spike (MS) Recovery Statement

The MS(1202595536(CAAN-12-2031)) recovered Benzidine at 29.6%. The limits are 30%-120%. Since the MS(1202595536(CAAN-12-2031)) and MSD(1202595537(CAAN-12-2031)) displayed similar recoveries for Benzidine, the failures may attributed to matrix interference. Please note, Benzidine is designated as being poor responding analyte in the analytical method. This may account for the biased low recoveries for that analyte in the MS and MSD. Please see data exception report 1048967.

Matrix Spike Duplicate (MSD) Recovery Statement

The MSD(1202595537(CAAN-12-2031)) recovered Benzidine at 27.5%. The limits are 30%-120%. Since the MS(1202595536(CAAN-12-2031)) and MSD(1202595537(CAAN-12-2031)) displayed similar recoveries for Benzidine, the failures may attributed to matrix interference. Please note, Benzidine is designated as being poor responding analyte in the analytical method. This may account for the biased low recoveries for that analyte in the MS and MSD. The MSD(1202595537(CAAN-12-2031)) multiple biased high spike recoveries. Please see the QC Summary for the specific recovery values. The MB, LCS and MS satisfied batch QC acceptance criteria (with the exception of Benzidine in the MS and MSD - see Spike Recovery Statements). Therefore, it was determined that the biased high failures were limited to the MSD sample only (the surrogate and spike recoveries were in general significantly higher in the MSD compared to the MS recoveries). Since the associated parent sample was reported as non-detected for all requested target analytes, a re-extraction was considered unnecessary and the data have been reported. The MSD also recovered N-Nitrosopyrrolidine at 130% (Limits are 51-115%), Atrazine at 120% (Limits are 38-112%) and 3,3'-Dichlorobenzidine at 114% (Limits are 28-112%). The latter three analytes were not detected in the associated samples. The data is unaffected and are reported. Please see data exception report 1048967.

MS/MSD Relative Percent Difference (RPD) Statement

The MS(1202595536(CAAN-12-2031)) and MSD(1202595537(CAAN-12-2031)) displayed multiple RPD value failures. Please see the QC Summary for the specific recovery values. The RPD failures were attributed to the biased high recoveries in the MSD (see the Spike Recovery Statement).

Internal Standard (ISTD) Acceptance

The internal standard responses were within the required acceptance criteria for all samples and QC.

Technical Information:**Holding Time Specifications**

All samples in this SDG in this batch met the specified holding time. GEL assigns holding times based on the associated methodology that assigns the date and time from sample collection or sample receipt. Those holding times expressed in hours are calculated in the AlphaLIMS system. Those holding times expressed as days expire at midnight on the day of expiration.

Preparation/Analytical Method Verification

All procedures were performed as stated in the SOP. All reported compound mass spectra met the detection specifications in the method.

Sample Dilutions

The samples in this SDG in this batch did not require dilutions.

Sample Re-extraction/Re-analysis

The 1202595537(MSD) was re-analyzed. The re-analysis is reported.

Miscellaneous Information:**Data Exception (DER) Documentation**

Data exception report 1048967 was generated for this SDG.

Manual Integrations

Some initial calibration standards, continuing calibration standards, and/or samples may require manual integrations due to software limitations. Manual integrations, if any, are included with the raw data.

TIC Comment

Tentatively identified compounds (TIC) may be requested for this sample delivery group/work order. Please note that non-requested target analytes that are reported on the quantitation reports will not be present on the Form I. These detected analytes are included in the calibrated method and as a result cannot be reported on the TIC quantitation report. TIC data, if requested, are included on the Sample Data Summary (Form 1) and are also included with the sample raw data.

Additional Comments

Additional comments were not required for the SDG associated samples in this batch.

Electronic Package Comment

The following package was generated using an electronic data processing program referred to as "virtual packaging". In an effort to increase quality and efficiency, the laboratory is developing systems to eventually generate all data packages electronically. The following change from "traditional" packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. An electronic signature page inserted after the case narrative of each electronic package will indicate the analyst, reviewer, and report specialist names associated with the generation of the data and package. The data validator will always sign and date the case narrative. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

System Configuration

The Semi-Volatile-GC/MS analysis was performed on the following instrument configuration:

Instrument ID	Instrument	System Configuration	Column ID	Column Description
MSD3.I	Agilent 7890A/5975C GC/MS w/ 7683 Autosampler	HP7890A/HP5975C	DB-5MS	25m x 0.2mm, 0.33um (5% Phenylmethylpolysiloxane)

Certification Statement

Where the analytical method has been performed under NELAP certification, the analysis has met all of the requirements of the NELAC standard unless otherwise noted in the analytical case narrative.

GEL LABORATORIES LLC

2040 Savage Road Charleston SC 29407 - (843) 556-8171 - www.gel.com

Qualifier Definition Report for

ARSL001 ARS International (63641-10)

Client SDG: 12-700 GEL Work Order: 295270

The Qualifiers in this report are defined as follows:

- * A quality control analyte recovery is outside of specified acceptance criteria
- ** Analyte is a surrogate compound
- E Organics--Concentration of the target analyte exceeds the instrument calibration range
- J Value is estimated
- U Analyte was analyzed for, but not detected above the MDL, MDA, or LOD.
- DL Indicates that sample is diluted.
- RA Indicates that sample is re-analyzed without re-extraction.
- RE Indicates that sample is re-extracted.

Review/Validation

GEL requires all analytical data to be verified by a qualified data reviewer. In addition, all CLP-like deliverables receive a third level review of the fractional data package.

The following data validator verified the information presented in this data report:

Signature: 

Name: Herbert Maier

Date: 23 FEB 2012

Title: Data Validator

Sample Data Summary

**Semi-Volatile
Certificate of Analysis
Sample Summary**

Page 1 of 3

SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2031
Batch ID: 1186837
Run Date: 02/09/2012 10:28
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0907.D

Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 1000 mL
Column: DB-5ms

Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether	U	10.0	ug/L	3.00	10.0
62-75-9	N-Methyl-N-nitrosomethylamine	U	10.0	ug/L	3.00	10.0
110-86-1	Pyridine	U	10.0	ug/L	3.00	10.0
62-53-3	Aniline	U	10.0	ug/L	3.00	10.0
108-95-2	Phenol	U	10.0	ug/L	3.00	10.0
111-44-4	bis(2-Chloroethyl) ether	U	10.0	ug/L	3.00	10.0
95-57-8	2-Chlorophenol	U	10.0	ug/L	3.00	10.0
541-73-1	1,3-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
106-46-7	1,4-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
95-50-1	1,2-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
100-51-6	Benzyl alcohol	U	10.0	ug/L	3.00	10.0
95-48-7	o-Cresol	U	10.0	ug/L	3.00	10.0
65794-96-9	m,p-Cresols	U	10.0	ug/L	3.00	10.0
621-64-7	N-Nitrosodi--n-propylamine	U	10.0	ug/L	3.00	10.0
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane	U	10.0	ug/L	3.00	10.0
98-95-3	Nitrobenzene	U	10.0	ug/L	3.00	10.0
78-59-1	Isophorone	U	10.0	ug/L	3.00	10.0
88-75-5	2-Nitrophenol	U	10.0	ug/L	3.00	10.0
105-67-9	2,4-Dimethylphenol	U	10.0	ug/L	3.00	10.0
111-91-1	bis(2-Chloroethoxy)methane	U	10.0	ug/L	3.00	10.0
120-83-2	2,4-Dichlorophenol	U	10.0	ug/L	3.00	10.0
65-85-0	Benzoic acid	U	20.0	ug/L	6.00	20.0
106-47-8	4-Chloroaniline	U	10.0	ug/L	3.00	10.0
87-68-3	Hexachlorobutadiene	U	10.0	ug/L	3.00	10.0
59-50-7	Parachlorometa cresol	U	10.0	ug/L	3.00	10.0
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.300	1.00
90-12-0	1-Methylnaphthalene	U	1.00	ug/L	0.300	1.00
77-47-4	Hexachlorocyclopentadiene	U	10.0	ug/L	3.00	10.0
88-06-2	2,4,6-Trichlorophenol	U	10.0	ug/L	3.00	10.0
95-95-4	2,4,5-Trichlorophenol	U	10.0	ug/L	3.00	10.0
91-58-7	2-Chloronaphthalene	U	1.00	ug/L	0.300	1.00
88-74-4	2-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate	U	10.0	ug/L	3.00	10.0
606-20-2	2,6-Dinitrotoluene	U	10.0	ug/L	3.00	10.0

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2031
Batch ID: 1186837
Run Date: 02/09/2012 10:28
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0907.D

Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 1000 mL
Column: DB-5ms

Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene	U	10.0	ug/L	3.00	10.0
208-96-8	Acenaphthylene	U	1.00	ug/L	0.300	1.00
83-32-9	Acenaphthene	U	1.00	ug/L	0.300	1.00
51-28-5	2,4-Dinitrophenol	U	20.0	ug/L	5.00	20.0
132-64-9	Dibenzofuran	U	10.0	ug/L	3.00	10.0
58-90-2	2,3,4,6-Tetrachlorophenol	U	10.0	ug/L	3.00	10.0
84-66-2	Diethylphthalate	U	10.0	ug/L	3.00	10.0
100-02-7	4-Nitrophenol	U	10.0	ug/L	3.00	10.0
86-73-7	Fluorene	U	1.00	ug/L	0.300	1.00
7005-72-3	4-Chlorophenylphenylether	U	10.0	ug/L	3.00	10.0
100-01-6	4-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>p</i> -Nitroaniline					
534-52-1	2-Methyl-4,6-dinitrophenol	U	10.0	ug/L	3.00	10.0
122-39-4	Diphenylamine	U	10.0	ug/L	3.00	10.0
122-66-7	Azobenzene	U	10.0	ug/L	3.00	10.0
	<i>1,2-Diphenylhydrazine</i>					
101-55-3	4-Bromophenylphenylether	U	10.0	ug/L	3.00	10.0
118-74-1	Hexachlorobenzene	U	10.0	ug/L	3.00	10.0
87-86-5	Pentachlorophenol	U	10.0	ug/L	3.00	10.0
88-85-7	Dinoseb	U	10.0	ug/L	3.00	10.0
85-01-8	Phenanthrene	U	1.00	ug/L	0.300	1.00
120-12-7	Anthracene	U	1.00	ug/L	0.300	1.00
84-74-2	Di-n-butylphthalate	U	10.0	ug/L	3.00	10.0
206-44-0	Fluoranthene	U	1.00	ug/L	0.300	1.00
129-00-0	Pyrene	U	1.00	ug/L	0.300	1.00
85-68-7	Butylbenzylphthalate	U	10.0	ug/L	3.00	10.0
117-81-7	bis(2-Ethylhexyl)phthalate	U	10.0	ug/L	3.00	10.0
56-55-3	Benzo(a)anthracene	U	1.00	ug/L	0.300	1.00
218-01-9	Chrysene	U	1.00	ug/L	0.300	1.00
117-84-0	Di-n-octylphthalate	U	10.0	ug/L	3.00	10.0
205-99-2	Benzo(b)fluoranthene	U	1.00	ug/L	0.300	1.00
207-08-9	Benzo(k)fluoranthene	U	1.00	ug/L	0.300	1.00
50-32-8	Benzo(a)pyrene	U	1.00	ug/L	0.300	1.00
193-39-5	Indeno(1,2,3-cd)pyrene	U	1.00	ug/L	0.300	1.00
53-70-3	Dibenzo(a,h)anthracene	U	1.00	ug/L	0.300	1.00
191-24-2	Benzo(ghi)perylene	U	1.00	ug/L	0.300	1.00
123-91-1	1,4-Dioxane	U	10.0	ug/L	3.00	10.0
55-18-5	N-Nitrosodiethylamine	U	10.0	ug/L	3.00	10.0
930-55-2	N-Nitrosopyrrolidine	U	10.0	ug/L	3.00	10.0

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700
Lab Sample ID: 295270002

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 1000 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2031
Batch ID: 1186837
Run Date: 02/09/2012 10:28
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0907.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.0	ug/L	3.00	10.0
95-94-3	1,2,4,5-Tetrachlorobenzene	U	10.0	ug/L	3.00	10.0
608-93-5	Pentachlorobenzene	U	10.0	ug/L	3.00	10.0
1912-24-9	Atrazine	U	10.0	ug/L	3.00	10.0
92-87-5	Benzidine	U	10.0	ug/L	3.00	10.0
91-94-1	3,3'-Dichlorobenzidine	U	10.0	ug/L	3.00	10.0
120-82-1	1,2,4-Trichlorobenzene	U	10.0	ug/L	3.00	10.0

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
2,4,6-Tribromophenol	75.0	100	ug/L 75.0	(30%-128%)
2-Fluorobiphenyl	34.0	50.0	ug/L 67.9	(34%-98%)
2-Fluorophenol	41.1	100	ug/L 41.1	(21%-78%)
Nitrobenzene-d5	37.3	50.0	ug/L 74.6	(39%-117%)
Phenol-d5	26.3	100	ug/L 26.3	(14%-80%)
p-Terphenyl-d14	37.3	50.0	ug/L 74.6	(39%-129%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
No Tentatively Identified Compounds Found				ug/L		

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2199
Batch ID: 1186837
Run Date: 02/09/2012 11:43
Prep Date: 02/08/2012 17:46
Data File: S020912.B\s3b0910.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether	U	10.5	ug/L	3.16	10.5
62-75-9	N-Methyl-N-nitrosomethylamine	U	10.5	ug/L	3.16	10.5
110-86-1	Pyridine	U	10.5	ug/L	3.16	10.5
62-53-3	Aniline	U	10.5	ug/L	3.16	10.5
108-95-2	Phenol	U	10.5	ug/L	3.16	10.5
111-44-4	bis(2-Chloroethyl) ether	U	10.5	ug/L	3.16	10.5
95-57-8	2-Chlorophenol	U	10.5	ug/L	3.16	10.5
541-73-1	1,3-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
106-46-7	1,4-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
95-50-1	1,2-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
100-51-6	Benzyl alcohol	U	10.5	ug/L	3.16	10.5
95-48-7	o-Cresol	U	10.5	ug/L	3.16	10.5
65794-96-9	m,p-Cresols	U	10.5	ug/L	3.16	10.5
621-64-7	N-Nitrosodi--n-propylamine	U	10.5	ug/L	3.16	10.5
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane	U	10.5	ug/L	3.16	10.5
98-95-3	Nitrobenzene	U	10.5	ug/L	3.16	10.5
78-59-1	Isophorone	U	10.5	ug/L	3.16	10.5
88-75-5	2-Nitrophenol	U	10.5	ug/L	3.16	10.5
105-67-9	2,4-Dimethylphenol	U	10.5	ug/L	3.16	10.5
111-91-1	bis(2-Chloroethoxy)methane	U	10.5	ug/L	3.16	10.5
120-83-2	2,4-Dichlorophenol	U	10.5	ug/L	3.16	10.5
65-85-0	Benzoic acid	U	21.1	ug/L	6.32	21.1
106-47-8	4-Chloroaniline	U	10.5	ug/L	3.16	10.5
87-68-3	Hexachlorobutadiene	U	10.5	ug/L	3.16	10.5
59-50-7	Parachlorometa cresol	U	10.5	ug/L	3.16	10.5
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
91-20-3	Naphthalene	U	1.05	ug/L	0.316	1.05
90-12-0	1-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
77-47-4	Hexachlorocyclopentadiene	U	10.5	ug/L	3.16	10.5
88-06-2	2,4,6-Trichlorophenol	U	10.5	ug/L	3.16	10.5
95-95-4	2,4,5-Trichlorophenol	U	10.5	ug/L	3.16	10.5
91-58-7	2-Chloronaphthalene	U	1.05	ug/L	0.316	1.05
88-74-4	2-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate	U	10.5	ug/L	3.16	10.5
606-20-2	2,6-Dinitrotoluene	U	10.5	ug/L	3.16	10.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2199

Client: ARSL001

Project: ESHL00210

Batch ID: 1186837

Method: SW846 8270C

SOP Ref: GL-OA-E-009

Run Date: 02/09/2012 11:43

Inst: MSD3.I

Dilution: 1

Prep Date: 02/08/2012 17:46

Analyst: JLD1

Inj. Vol: 1 uL

Data File: S020912.B\s3b0910.D

Aliquot: 950 mL

Final Volume: 1 mL

Column: DB-5ms

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene	U	10.5	ug/L	3.16	10.5
208-96-8	Acenaphthylene	U	1.05	ug/L	0.316	1.05
83-32-9	Acenaphthene	U	1.05	ug/L	0.316	1.05
51-28-5	2,4-Dinitrophenol	U	21.1	ug/L	5.26	21.1
132-64-9	Dibenzofuran	U	10.5	ug/L	3.16	10.5
58-90-2	2,3,4,6-Tetrachlorophenol	U	10.5	ug/L	3.16	10.5
84-66-2	Diethylphthalate	U	10.5	ug/L	3.16	10.5
100-02-7	4-Nitrophenol	U	10.5	ug/L	3.16	10.5
86-73-7	Fluorene	U	1.05	ug/L	0.316	1.05
7005-72-3	4-Chlorophenylphenylether	U	10.5	ug/L	3.16	10.5
100-01-6	4-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>p</i> -Nitroaniline					
534-52-1	2-Methyl-4,6-dinitrophenol	U	10.5	ug/L	3.16	10.5
122-39-4	Diphenylamine	U	10.5	ug/L	3.16	10.5
122-66-7	Azobenzene	U	10.5	ug/L	3.16	10.5
	<i>1,2</i> -Diphenylhydrazine					
101-55-3	4-Bromophenylphenylether	U	10.5	ug/L	3.16	10.5
118-74-1	Hexachlorobenzene	U	10.5	ug/L	3.16	10.5
87-86-5	Pentachlorophenol	U	10.5	ug/L	3.16	10.5
88-85-7	Dinoseb	U	10.5	ug/L	3.16	10.5
85-01-8	Phenanthrene	U	1.05	ug/L	0.316	1.05
120-12-7	Anthracene	U	1.05	ug/L	0.316	1.05
84-74-2	Di-n-butylphthalate	U	10.5	ug/L	3.16	10.5
206-44-0	Fluoranthene	U	1.05	ug/L	0.316	1.05
129-00-0	Pyrene	U	1.05	ug/L	0.316	1.05
85-68-7	Butylbenzylphthalate	U	10.5	ug/L	3.16	10.5
117-81-7	bis(2-Ethylhexyl)phthalate	U	10.5	ug/L	3.16	10.5
56-55-3	Benzo(a)anthracene	U	1.05	ug/L	0.316	1.05
218-01-9	Chrysene	U	1.05	ug/L	0.316	1.05
117-84-0	Di-n-octylphthalate	U	10.5	ug/L	3.16	10.5
205-99-2	Benzo(b)fluoranthene	U	1.05	ug/L	0.316	1.05
207-08-9	Benzo(k)fluoranthene	U	1.05	ug/L	0.316	1.05
50-32-8	Benzo(a)pyrene	U	1.05	ug/L	0.316	1.05
193-39-5	Indeno(1,2,3-cd)pyrene	U	1.05	ug/L	0.316	1.05
53-70-3	Dibenzo(a,h)anthracene	U	1.05	ug/L	0.316	1.05
191-24-2	Benzo(ghi)perylene	U	1.05	ug/L	0.316	1.05
123-91-1	1,4-Dioxane	U	10.5	ug/L	3.16	10.5
55-18-5	N-Nitrosodiethylamine	U	10.5	ug/L	3.16	10.5
930-55-2	N-Nitrosopyrrolidine	U	10.5	ug/L	3.16	10.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270004

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2199
Batch ID: 1186837
Run Date: 02/09/2012 11:43
Prep Date: 02/08/2012 17:46
Data File: S020912.B\3b0910.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.5	ug/L	3.16	10.5
95-94-3	1,2,4,5-Tetrachlorobenzene	U	10.5	ug/L	3.16	10.5
608-93-5	Pentachlorobenzene	U	10.5	ug/L	3.16	10.5
1912-24-9	Atrazine	U	10.5	ug/L	3.16	10.5
92-87-5	Benzidine	U	10.5	ug/L	3.16	10.5
91-94-1	3,3'-Dichlorobenzidine	U	10.5	ug/L	3.16	10.5
120-82-1	1,2,4-Trichlorobenzene	U	10.5	ug/L	3.16	10.5

Surrogate/Tracer recovery	Result	Nominal		Recovery%	Acceptable Limits
2,4,6-Tribromophenol	87.6	105	ug/L	83.3	(30%-128%)
2-Fluorobiphenyl	40.4	52.6	ug/L	76.8	(34%-98%)
2-Fluorophenol	58.3	105	ug/L	55.4	(21%-78%)
Nitrobenzene-d5	48.1	52.6	ug/L	91.5	(39%-117%)
Phenol-d5	37.2	105	ug/L	35.4	(14%-80%)
p-Terphenyl-d14	41.7	52.6	ug/L	79.2	(39%-129%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
	No Tentatively Identified Compounds Found			ug/L		

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270007

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2201

Client: ARSL001

Project: ESHL00210

Batch ID: 1186837

Method: SW846 8270C

SOP Ref: GL-OA-E-009

Run Date: 02/09/2012 12:08

Inst: MSD3.I

Dilution: 1

Prep Date: 02/08/2012 17:46

Analyst: JLD1

Inj. Vol: 1 uL

Data File: S020912.B\s3b0911.D

Aliquot: 950 mL

Final Volume: 1 mL

Column: DB-5ms

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether	U	10.5	ug/L	3.16	10.5
62-75-9	N-Methyl-N-nitrosomethylamine	U	10.5	ug/L	3.16	10.5
110-86-1	Pyridine	U	10.5	ug/L	3.16	10.5
62-53-3	Aniline	U	10.5	ug/L	3.16	10.5
108-95-2	Phenol	U	10.5	ug/L	3.16	10.5
111-44-4	bis(2-Chloroethyl) ether	U	10.5	ug/L	3.16	10.5
95-57-8	2-Chlorophenol	U	10.5	ug/L	3.16	10.5
541-73-1	1,3-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
106-46-7	1,4-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
95-50-1	1,2-Dichlorobenzene	U	10.5	ug/L	3.16	10.5
100-51-6	Benzyl alcohol	U	10.5	ug/L	3.16	10.5
95-48-7	o-Cresol	U	10.5	ug/L	3.16	10.5
65794-96-9	m,p-Cresols	U	10.5	ug/L	3.16	10.5
621-64-7	N-Nitrosodi--n-propylamine	U	10.5	ug/L	3.16	10.5
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane	U	10.5	ug/L	3.16	10.5
98-95-3	Nitrobenzene	U	10.5	ug/L	3.16	10.5
78-59-1	Isophorone	U	10.5	ug/L	3.16	10.5
88-75-5	2-Nitrophenol	U	10.5	ug/L	3.16	10.5
105-67-9	2,4-Dimethylphenol	U	10.5	ug/L	3.16	10.5
111-91-1	bis(2-Chloroethoxy)methane	U	10.5	ug/L	3.16	10.5
120-83-2	2,4-Dichlorophenol	U	10.5	ug/L	3.16	10.5
65-85-0	Benzoic acid	U	21.1	ug/L	6.32	21.1
106-47-8	4-Chloroaniline	U	10.5	ug/L	3.16	10.5
87-68-3	Hexachlorobutadiene	U	10.5	ug/L	3.16	10.5
59-50-7	Parachlorometa cresol	U	10.5	ug/L	3.16	10.5
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
91-20-3	Naphthalene	U	1.05	ug/L	0.316	1.05
90-12-0	1-Methylnaphthalene	U	1.05	ug/L	0.316	1.05
77-47-4	Hexachlorocyclopentadiene	U	10.5	ug/L	3.16	10.5
88-06-2	2,4,6-Trichlorophenol	U	10.5	ug/L	3.16	10.5
95-95-4	2,4,5-Trichlorophenol	U	10.5	ug/L	3.16	10.5
91-58-7	2-Chloronaphthalene	U	1.05	ug/L	0.316	1.05
88-74-4	2-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate	U	10.5	ug/L	3.16	10.5
606-20-2	2,6-Dinitrotoluene	U	10.5	ug/L	3.16	10.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

Page 2 of 3

SDG Number: 12-700
Lab Sample ID: 295270007

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45

Matrix: WG

Client ID: CAAN-12-2201

Client: ARSL001

Project: ESHL00210

Batch ID: 1186837

Method: SW846 8270C

SOP Ref: GL-OA-E-009

Run Date: 02/09/2012 12:08

Inst: MSD3.I

Dilution: 1

Prep Date: 02/08/2012 17:46

Analyst: JLD1

Inj. Vol: 1 uL

Data File: S020912.B\s3b0911.D

Aliquot: 950 mL

Final Volume: 1 mL

Column: DB-5ms

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene	U	10.5	ug/L	3.16	10.5
208-96-8	Acenaphthylene	U	1.05	ug/L	0.316	1.05
83-32-9	Acenaphthene	U	1.05	ug/L	0.316	1.05
51-28-5	2,4-Dinitrophenol	U	21.1	ug/L	5.26	21.1
132-64-9	Dibenzofuran	U	10.5	ug/L	3.16	10.5
58-90-2	2,3,4,6-Tetrachlorophenol	U	10.5	ug/L	3.16	10.5
84-66-2	Diethylphthalate	U	10.5	ug/L	3.16	10.5
100-02-7	4-Nitrophenol	U	10.5	ug/L	3.16	10.5
86-73-7	Fluorene	U	1.05	ug/L	0.316	1.05
7005-72-3	4-Chlorophenylphenylether	U	10.5	ug/L	3.16	10.5
100-01-6	4-Nitroaniline	U	10.5	ug/L	3.16	10.5
	<i>p</i> -Nitroaniline					
534-52-1	2-Methyl-4,6-dinitrophenol	U	10.5	ug/L	3.16	10.5
122-39-4	Diphenylamine	U	10.5	ug/L	3.16	10.5
122-66-7	Azobenzene	U	10.5	ug/L	3.16	10.5
	<i>1,2</i> -Diphenylhydrazine					
101-55-3	4-Bromophenylphenylether	U	10.5	ug/L	3.16	10.5
118-74-1	Hexachlorobenzene	U	10.5	ug/L	3.16	10.5
87-86-5	Pentachlorophenol	U	10.5	ug/L	3.16	10.5
88-85-7	Dinoseb	U	10.5	ug/L	3.16	10.5
85-01-8	Phenanthrene	U	1.05	ug/L	0.316	1.05
120-12-7	Anthracene	U	1.05	ug/L	0.316	1.05
84-74-2	Di-n-butylphthalate	U	10.5	ug/L	3.16	10.5
206-44-0	Fluoranthene	U	1.05	ug/L	0.316	1.05
129-00-0	Pyrene	U	1.05	ug/L	0.316	1.05
85-68-7	Butylbenzylphthalate	U	10.5	ug/L	3.16	10.5
117-81-7	bis(2-Ethylhexyl)phthalate	U	10.5	ug/L	3.16	10.5
56-55-3	Benzo(a)anthracene	U	1.05	ug/L	0.316	1.05
218-01-9	Chrysene	U	1.05	ug/L	0.316	1.05
117-84-0	Di-n-octylphthalate	U	10.5	ug/L	3.16	10.5
205-99-2	Benzo(b)fluoranthene	U	1.05	ug/L	0.316	1.05
207-08-9	Benzo(k)fluoranthene	U	1.05	ug/L	0.316	1.05
50-32-8	Benzo(a)pyrene	U	1.05	ug/L	0.316	1.05
193-39-5	Indeno(1,2,3-cd)pyrene	U	1.05	ug/L	0.316	1.05
53-70-3	Dibenzo(a,h)anthracene	U	1.05	ug/L	0.316	1.05
191-24-2	Benzo(ghi)perylene	U	1.05	ug/L	0.316	1.05
123-91-1	1,4-Dioxane	U	10.5	ug/L	3.16	10.5
55-18-5	N-Nitrosodiethylamine	U	10.5	ug/L	3.16	10.5
930-55-2	N-Nitrosopyrrolidine	U	10.5	ug/L	3.16	10.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700
Lab Sample ID: 295270007

Date Collected: 02/01/2012 12:00
Date Received: 02/03/2012 08:45
Client: ARSL001
Method: SW846 8270C
Inst: MSD3.I
Analyst: JLD1
Aliquot: 950 mL
Column: DB-5ms

Matrix: WG
Project: ESHL00210
SOP Ref: GL-OA-E-009
Dilution: 1
Inj. Vol: 1 uL
Final Volume: 1 mL

Client ID: CAAN-12-2201
Batch ID: 1186837
Run Date: 02/09/2012 12:08
Prep Date: 02/08/2012 17:46
Data File: S020912.B\3b0911.D

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.5	ug/L	3.16	10.5
95-94-3	1,2,4,5-Tetrachlorobenzene	U	10.5	ug/L	3.16	10.5
608-93-5	Pentachlorobenzene	U	10.5	ug/L	3.16	10.5
1912-24-9	Atrazine	U	10.5	ug/L	3.16	10.5
92-87-5	Benzidine	U	10.5	ug/L	3.16	10.5
91-94-1	3,3'-Dichlorobenzidine	U	10.5	ug/L	3.16	10.5
120-82-1	1,2,4-Trichlorobenzene	U	10.5	ug/L	3.16	10.5

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
2,4,6-Tribromophenol	74.3	105	ug/L	70.6 (30%-128%)
2-Fluorobiphenyl	38.9	52.6	ug/L	73.9 (34%-98%)
2-Fluorophenol	50.4	105	ug/L	47.9 (21%-78%)
Nitrobenzene-d5	44.8	52.6	ug/L	85.2 (39%-117%)
Phenol-d5	31.2	105	ug/L	29.7 (14%-80%)
p-Terphenyl-d14	42.5	52.6	ug/L	80.8 (39%-129%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
No Tentatively Identified Compounds Found				ug/L		

Quality Control Summary

Semi-Volatile
Surrogate Recovery Report

Page 1 of 1

SDG Number: 12-700

Matrix Type: LIQUID

Sample ID	Client ID	2FP %REC	PHL %REC	NBZ %REC	FBP %REC	TBP %REC	TPH %REC
1202595534	MB for batch 1186831	56	35	95	83	95	84
1202595535	LCS for batch 1186831	50	31	88	79	82	80
295270002	CAAN-12-2031	41	26	75	68	75	75
1202595536	CAAN-12-2031MS	56	46	73	65	77	63
295270004	CAAN-12-2199	55	35	91	77	83	79
295270007	CAAN-12-2201	48	30	85	74	71	81
1202595537	CAAN-12-2031MSD	87 *	70	107	88	95	88

Surrogate

Acceptance Limits

2FP	= 2-Fluorophenol	(21%-78%)
PHL	= Phenol-d5	(14%-80%)
NBZ	= Nitrobenzene-d5	(39%-117%)
FBP	= 2-Fluorobiphenyl	(34%-98%)
TBP	= 2,4,6-Tribromophenol	(30%-128%)
TPH	= p-Terphenyl-d14	(39%-129%)

* Recovery outside Acceptance Limits

Column to be used to flag recovery values

D Sample Diluted

Semi-Volatile
Quality Control Summary
Spike Recovery Report

Page 1 of 4

SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1186831

Matrix: WATER

Lab Sample ID: 1202595535

Instrument: MSD3.I

Analysis Date: 02/09/2012 10:03

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
39638-32-9	LCS bis(2-Chloroisopropyl)ether	50.0	0.0	38.5	77	27-129
62-75-9	LCS N-Methyl-N-nitrosomethylam	50.0	0.0	24.9	50	30-89
110-86-1	LCS Pyridine	50.0	0.0	16.7	33	27-95
62-53-3	LCS Aniline	50.0	0.0	39.6	79	39-120
108-95-2	LCS Phenol	50.0	0.0	15.2	30	15-103
111-44-4	LCS bis(2-Chloroethyl) ether	50.0	0.0	40.2	80	39-115
95-57-8	LCS 2-Chlorophenol	50.0	0.0	38.1	76	45-105
541-73-1	LCS 1,3-Dichlorobenzene	50.0	0.0	29.3	59	31-93
106-46-7	LCS 1,4-Dichlorobenzene	50.0	0.0	29.6	59	32-93
95-50-1	LCS 1,2-Dichlorobenzene	50.0	0.0	30.3	61	33-94
100-51-6	LCS Benzyl alcohol	50.0	0.0	36.2	72	37-105
95-48-7	LCS o-Cresol	50.0	0.0	35.5	71	39-94
65794-96-9	LCS m,p-Cresols	50.0	0.0	39.0	78	37-102
621-64-7	LCS N-Nitrosodi--n-propylamine <i>N-Nitrosodipropylamine</i>	50.0	0.0	49.3	99	45-120
67-72-1	LCS Hexachloroethane	50.0	0.0	28.2	56	31-92
98-95-3	LCS Nitrobenzene	50.0	0.0	41.9	84	46-118
78-59-1	LCS Isophorone	50.0	0.0	47.6	95	48-120
88-75-5	LCS 2-Nitrophenol	50.0	0.0	40.8	82	48-110
105-67-9	LCS 2,4-Dimethylphenol	50.0	0.0	40.5	81	42-102
111-91-1	LCS bis(2-Chloroethoxy)methane	50.0	0.0	39.3	79	46-106
120-83-2	LCS 2,4-Dichlorophenol	50.0	0.0	41.6	83	49-110
65-85-0	LCS Benzoic acid	100	0.0	28.7	29 *	30-142

Semi-Volatile
Quality Control Summary
Spike Recovery Report

Page 2 of 4

SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1186831

Matrix: WATER

Lab Sample ID: 1202595535

Instrument: MSD3.I

Analysis Date: 02/09/2012 10:03

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
106-47-8	LCS 4-Chloroaniline	50.0	0.0	44.1	88	50-123
87-68-3	LCS Hexachlorobutadiene	50.0	0.0	30.1	60	29-97
59-50-7	LCS Parachlorometa cresol 4-Chloro-3-methylphenol	50.0	0.0	44.1	88	53-113
91-57-6	LCS 2-Methylnaphthalene	50.0	0.0	35.8	72	35-104
91-20-3	LCS Naphthalene	50.0	0.0	32.6	65	35-99
90-12-0	LCS 1-Methylnaphthalene	50.0	0.0	38.0	76	36-104
77-47-4	LCS Hexachlorocyclopentadiene	50.0	0.0	19.9	40	28-93
88-06-2	LCS 2,4,6-Trichlorophenol	50.0	0.0	39.9	80	48-111
95-95-4	LCS 2,4,5-Trichlorophenol	50.0	0.0	41.0	82	44-114
91-58-7	LCS 2-Chloronaphthalene	50.0	0.0	35.8	72	37-102
88-74-4	LCS 2-Nitroaniline o-Nitroaniline	50.0	0.0	43.3	87	43-125
99-09-2	LCS 3-Nitroaniline m-Nitroaniline	50.0	0.0	38.5	77	45-125
131-11-3	LCS Dimethylphthalate	50.0	0.0	43.9	88	47-127
606-20-2	LCS 2,6-Dinitrotoluene	50.0	0.0	41.6	83	52-119
121-14-2	LCS 2,4-Dinitrotoluene	50.0	0.0	39.5	79	52-122
208-96-8	LCS Acenaphthylene	50.0	0.0	37.0	74	42-108
83-32-9	LCS Acenaphthene	50.0	0.0	37.1	74	40-105
51-28-5	LCS 2,4-Dinitrophenol	50.0	0.0	31.3	63	33-126
132-64-9	LCS Dibenzofuran	50.0	0.0	39.3	79	47-111
58-90-2	LCS 2,3,4,6-Tetrachlorophenol	50.0	0.0	39.4	79	47-124
84-66-2	LCS Diethylphthalate	50.0	0.0	43.9	88	48-128
100-02-7	LCS 4-Nitrophenol	50.0	0.0	8.76	18	15-103

Semi-Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1186831

Matrix: WATER

Lab Sample ID: 1202595535

Instrument: MSD3.I

Analysis Date: 02/09/2012 10:03

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
86-73-7	LCS Fluorene	50.0	0.0	38.3	77	45-108
7005-72-3	LCS 4-Chlorophenylphenylether	50.0	0.0	37.9	76	45-107
100-01-6	LCS 4-Nitroaniline <i>p-Nitroaniline</i>	50.0	0.0	36.4	73	44-139
534-52-1	LCS 2-Methyl-4,6-dinitrophenol	50.0	0.0	37.1	74	45-122
122-39-4	LCS Diphenylamine	50.0	0.0	45.7	91	47-112
122-66-7	LCS Azobenzene <i>1,2-Diphenylhydrazine</i>	50.0	0.0	49.6	99	41-117
101-55-3	LCS 4-Bromophenylphenylether	50.0	0.0	43.3	87	44-110
118-74-1	LCS Hexachlorobenzene	50.0	0.0	45.8	92	44-114
87-86-5	LCS Pentachlorophenol	50.0	0.0	31.0	62	36-108
85-01-8	LCS Phenanthrene	50.0	0.0	41.7	83	50-111
120-12-7	LCS Anthracene	50.0	0.0	39.8	80	51-111
84-74-2	LCS Di-n-butylphthalate	50.0	0.0	46.5	93	46-130
206-44-0	LCS Fluoranthene	50.0	0.0	39.5	79	50-118
129-00-0	LCS Pyrene	50.0	0.0	40.8	82	41-121
85-68-7	LCS Butylbenzylphthalate	50.0	0.0	43.9	88	41-129
117-81-7	LCS bis(2-Ethylhexyl)phthalate	50.0	0.0	43.0	86	39-132
56-55-3	LCS Benzo(a)anthracene	50.0	0.0	41.3	83	51-111
218-01-9	LCS Chrysene	50.0	0.0	43.9	88	50-113
117-84-0	LCS Di-n-octylphthalate	50.0	0.0	45.1	90	36-134
205-99-2	LCS Benzo(b)fluoranthene	50.0	0.0	44.9	90	46-117
207-08-9	LCS Benzo(k)fluoranthene	50.0	0.0	44.1	88	46-120
50-32-8	LCS Benzo(a)pyrene	50.0	0.0	41.0	82	49-114

Semi-Volatile
Quality Control Summary
Spike Recovery Report

Page 4 of 4

SDG Number: 12-700

Sample Type: Laboratory Control Sample

Client ID: LCS for batch 1186831

Matrix: WATER

Lab Sample ID: 1202595535

Instrument: MSD3.I

Analysis Date: 02/09/2012 10:03

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
193-39-5	LCS Indeno(1,2,3-cd)pyrene	50.0	0.0	37.9	76	40-133
53-70-3	LCS Dibenzo(a,h)anthracene	50.0	0.0	44.0	88	41-133
191-24-2	LCS Benzo(ghi)perylene	50.0	0.0	38.0	76	37-133
123-91-1	LCS 1,4-Dioxane	50.0	0.0	26.5	53	39-76
930-55-2	LCS N-Nitrosopyrrolidine	50.0	0.0	50.1	100	53-108
95-94-3	LCS 1,2,4,5-Tetrachlorobenzene	50.0	0.0	37.6	75	35-97
1912-24-9	LCS Atrazine	50.0	0.0	54.4	109	47-111
92-87-5	LCS Benzidine	100	0.0	17.3	17 *	23-131
91-94-1	LCS 3,3'-Dichlorobenzidine	50.0	0.0	40.2	80	45-106
120-82-1	LCS 1,2,4-Trichlorobenzene	50.0	0.0	29.9	60	31-94

Semi-Volatile
Quality Control Summary
Spike Recovery Report

Page 1 of 8

SDG Number: 12-700

Sample Type: Matrix Spike

Client ID: CAAN-12-2031MS

Matrix: WG

Lab Sample ID: 1202595536

Instrument: MSD3.I

Analysis Date: 02/09/2012 10:53

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
39638-32-9	MS bis(2-Chloroisopropyl)ether	118	0.00 U	74.8	64	30-135
62-75-9	MS N-Methyl-N-nitrosomethylam	118	0.00 U	60.5	51	26-103
110-86-1	MS Pyridine	118	0.00 U	46.1	39	21-103
62-53-3	MS Aniline	118	0.00 U	89.6	76	29-126
108-95-2	MS Phenol	118	0.00 U	52.5	45	21-82
111-44-4	MS bis(2-Chloroethyl) ether	118	0.00 U	78.2	66	32-121
95-57-8	MS 2-Chlorophenol	118	0.00 U	74.6	63	37-111
541-73-1	MS 1,3-Dichlorobenzene	118	0.00 U	52.7	45	27-90
106-46-7	MS 1,4-Dichlorobenzene	118	0.00 U	53.4	45	28-92
95-50-1	MS 1,2-Dichlorobenzene	118	0.00 U	56.8	48	27-97
100-51-6	MS Benzyl alcohol	118	0.00 U	81.4	69	33-115
95-48-7	MS o-Cresol	118	0.00 U	77.9	66	36-102
65794-96-9	MS m,p-Cresols	118	0.00 U	91.7	78	32-117
621-64-7	MS N-Nitrosodi--n-propylamine <i>N-Nitrosodipropylamine</i>	118	0.00 U	96.6	82	39-124
67-72-1	MS Hexachloroethane	118	0.00 U	47.8	41	26-91
98-95-3	MS Nitrobenzene	118	0.00 U	82.9	70	36-131
78-59-1	MS Isophorone	118	0.00 U	93.8	80	43-120
88-75-5	MS 2-Nitrophenol	118	0.00 U	81.7	69	39-114
105-67-9	MS 2,4-Dimethylphenol	118	0.00 U	78.7	67	37-105
111-91-1	MS bis(2-Chloroethoxy)methane	118	0.00 U	78.0	66	41-108
120-83-2	MS 2,4-Dichlorophenol	118	0.00 U	83.3	71	37-116
65-85-0	MS Benzoic acid	235	0.00 U	116	49	17-96

Semi-Volatile
Quality Control Summary
Spike Recovery Report

Page 2 of 8

SDG Number: 12-700

Sample Type: Matrix Spike

Client ID: CAAN-12-2031MS

Matrix: WG

Lab Sample ID: 1202595536

Instrument: MSD3.I

Analysis Date: 02/09/2012 10:53

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
106-47-8	MS 4-Chloroaniline	118	0.00 U	93.5	80	37-125
87-68-3	MS Hexachlorobutadiene	118	0.00 U	50.5	43	22-98
59-50-7	MS Parachlorometa cresol 4-Chloro-3-methylphenol	118	0.00 U	95.9	81	46-119
91-57-6	MS 2-Methylnaphthalene	118	0.00 U	69.1	59	32-107
91-20-3	MS Naphthalene	118	0.00 U	65.1	55	29-103
90-12-0	MS 1-Methylnaphthalene	118	0.00 U	73.0	62	29-111
77-47-4	MS Hexachlorocyclopentadiene	118	0.00 U	44.8	38	19-90
88-06-2	MS 2,4,6-Trichlorophenol	118	0.00 U	84.7	72	37-116
95-95-4	MS 2,4,5-Trichlorophenol	118	0.00 U	86.9	74	37-118
91-58-7	MS 2-Chloronaphthalene	118	0.00 U	68.1	58	35-101
88-74-4	MS 2-Nitroaniline o-Nitroaniline	118	0.00 U	96.8	82	36-126
99-09-2	MS 3-Nitroaniline m-Nitroaniline	118	0.00 U	90.2	77	35-126
131-11-3	MS Dimethylphthalate	118	0.00 U	94.1	80	42-125
606-20-2	MS 2,6-Dinitrotoluene	118	0.00 U	89.5	76	48-118
121-14-2	MS 2,4-Dinitrotoluene	118	0.00 U	88.3	75	49-120
208-96-8	MS Acenaphthylene	118	0.00 U	74.2	63	35-114
83-32-9	MS Acenaphthene	118	0.00 U	72.0	61	33-111
51-28-5	MS 2,4-Dinitrophenol	118	0.00 U	88.2	75	21-131
132-64-9	MS Dibenzofuran	118	0.00 U	79.8	68	43-113
58-90-2	MS 2,3,4,6-Tetrachlorophenol	118	0.00 U	89.9	76	36-130
84-66-2	MS Diethylphthalate	118	0.00 U	94.8	81	42-128
100-02-7	MS 4-Nitrophenol	118	0.00 U	41.8	36	20-90

Semi-Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Matrix Spike

Client ID: CAAN-12-2031MS

Matrix: WG

Lab Sample ID: 1202595536

Instrument: MSD3.I

Analysis Date: 02/09/2012 10:53

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No		Parmname	Amount Added ug/L	Sample Conc. ug/L		Spike Conc. ug/L	Recovery %	Acceptance Limits
86-73-7	MS	Fluorene	118	0.00	U	76.9	65	38-115
7005-72-3	MS	4-Chlorophenylphenylether	118	0.00	U	74.9	64	41-110
100-01-6	MS	4-Nitroaniline <i>p-Nitroaniline</i>	118	0.00	U	96.0	82	34-138
534-52-1	MS	2-Methyl-4,6-dinitrophenol	118	0.00	U	89.7	76	32-124
122-39-4	MS	Diphenylamine	118	0.00	U	94.8	81	42-111
122-66-7	MS	Azobenzene <i>1,2-Diphenylhydrazine</i>	118	0.00	U	98.2	84	36-118
101-55-3	MS	4-Bromophenylphenylether	118	0.00	U	80.2	68	40-112
118-74-1	MS	Hexachlorobenzene	118	0.00	U	86.9	74	40-116
87-86-5	MS	Pentachlorophenol	118	0.00	U	74.5	63	31-114
85-01-8	MS	Phenanthrene	118	0.00	U	85.4	73	41-119
120-12-7	MS	Anthracene	118	0.00	U	83.9	71	41-118
84-74-2	MS	Di-n-butylphthalate	118	0.00	U	92.5	79	41-128
206-44-0	MS	Fluoranthene	118	0.00	U	87.0	74	40-124
129-00-0	MS	Pyrene	118	0.00	U	73.0	62	34-126
85-68-7	MS	Butylbenzylphthalate	118	0.00	U	79.8	68	37-128
117-81-7	MS	bis(2-Ethylhexyl)phthalate	118	0.00	U	79.6	68	34-132
56-55-3	MS	Benzo(a)anthracene	118	0.00	U	86.3	73	38-120
218-01-9	MS	Chrysene	118	0.00	U	91.8	78	43-116
117-84-0	MS	Di-n-octylphthalate	118	0.00	U	94.2	80	30-135
205-99-2	MS	Benzo(b)fluoranthene	118	0.00	U	87.2	74	38-123
207-08-9	MS	Benzo(k)fluoranthene	118	0.00	U	86.8	74	40-123
50-32-8	MS	Benzo(a)pyrene	118	0.00	U	86.2	73	42-117

Semi-Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Client ID: CAAN-12-2031MS

Lab Sample ID: 1202595536

Instrument: MSD3.I

Analyst: JLD1

Inj. Vol: 1 uL

Sample Type: Matrix Spike

Matrix: WG

Analysis Date: 02/09/2012 10:53

Dilution: 1

Prep Batch ID: 1186831

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits
193-39-5	MS Indeno(1,2,3-cd)pyrene	118	0.00 U	91.1	77	33-135
53-70-3	MS Dibenzo(a,h)anthracene	118	0.00 U	107	91	35-134
191-24-2	MS Benzo(ghi)perylene	118	0.00 U	88.4	75	29-134
123-91-1	MS 1,4-Dioxane	118	0.00 U	64.4	55	30-98
930-55-2	MS N-Nitrosopyrrolidine	118	0.00 U	104	88	51-115
95-94-3	MS 1,2,4,5-Tetrachlorobenzene	118	0.00 U	68.0	58	31-101
1912-24-9	MS Atrazine	118	0.00 U	104	88	38-112
92-87-5	MS Benzidine	235	0.00 U	69.5	30	30-120
91-94-1	MS 3,3'-Dichlorobenzidine	118	0.00 U	109	92	28-112
120-82-1	MS 1,2,4-Trichlorobenzene	118	0.00 U	56.5	48	25-94

Semi-Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Matrix Spike Duplicate

Client ID: CAAN-12-2031MSD

Matrix: WG

Lab Sample ID: 1202595537

Instrument: MSD3.I

Analysis Date: 02/09/2012 13:48

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
39638-32-9	MSD bis(2-Chloroisopropyl)ether	118	0.00 U	114	97	30-135	41 *	0-27
62-75-9	MSD N-Methyl-N-nitrosomethylam	118	0.00 U	101	86	26-103	50 *	0-27
110-86-1	MSD Pyridine	118	0.00 U	41.6	35	21-103	10	0-25
62-53-3	MSD Aniline	118	0.00 U	108	92	29-126	19	0-25
108-95-2	MSD Phenol	118	0.00 U	79.2	67	21-82	41 *	0-33
111-44-4	MSD bis(2-Chloroethyl) ether	118	0.00 U	119	101	32-121	41 *	0-28
95-57-8	MSD 2-Chlorophenol	118	0.00 U	115	98	37-111	43 *	0-25
541-73-1	MSD 1,3-Dichlorobenzene	118	0.00 U	76.5	65	27-90	37 *	0-28
106-46-7	MSD 1,4-Dichlorobenzene	118	0.00 U	78.9	67	28-92	39 *	0-27
95-50-1	MSD 1,2-Dichlorobenzene	118	0.00 U	81.9	70	27-97	36 *	0-28
100-51-6	MSD Benzyl alcohol	118	0.00 U	120	102	33-115	38 *	0-28
95-48-7	MSD o-Cresol	118	0.00 U	116	98	36-102	39 *	0-29
65794-96-9	MSD m,p-Cresols	118	0.00 U	138	117	32-117	40 *	0-29
621-64-7	MSD N-Nitrosodi--n-propylamine <i>N-Nitrosodipropylamine</i>	118	0.00 U	140	119	39-124	37 *	0-24
67-72-1	MSD Hexachloroethane	118	0.00 U	72.2	61	26-91	41 *	0-29
98-95-3	MSD Nitrobenzene	118	0.00 U	123	104	36-131	39 *	0-23
78-59-1	MSD Isophorone	118	0.00 U	134	114	43-120	35 *	0-23
88-75-5	MSD 2-Nitrophenol	118	0.00 U	118	101	39-114	37 *	0-32
105-67-9	MSD 2,4-Dimethylphenol	118	0.00 U	118	100	37-105	40 *	0-29
111-91-1	MSD bis(2-Chloroethoxy)methane	118	0.00 U	114	97	41-108	37 *	0-23
120-83-2	MSD 2,4-Dichlorophenol	118	0.00 U	118	100	37-116	35 *	0-25
65-85-0	MSD Benzoic acid	235	0.00 U	153	65	17-96	28 *	0-25

Semi-Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Matrix Spike Duplicate

Client ID: CAAN-12-2031MSD

Matrix: WG

Lab Sample ID: 1202595537

Instrument: MSD3.I

Analysis Date: 02/09/2012 13:48

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
106-47-8	MSD 4-Chloroaniline	118	0.00 U	129	110	37-125	32 *	0-30
87-68-3	MSD Hexachlorobutadiene	118	0.00 U	70.7	60	22-98	33 *	0-28
59-50-7	MSD Parachlorometa cresol 4-Chloro-3-methylphenol	118	0.00 U	135	115	46-119	34 *	0-27
91-57-6	MSD 2-Methylnaphthalene	118	0.00 U	91.9	78	32-107	28 *	0-26
91-20-3	MSD Naphthalene	118	0.00 U	91.3	78	29-103	34 *	0-27
90-12-0	MSD 1-Methylnaphthalene	118	0.00 U	99.6	85	29-111	31 *	0-27
77-47-4	MSD Hexachlorocyclopentadiene	118	0.00 U	53.1	45	19-90	17	0-25
88-06-2	MSD 2,4,6-Trichlorophenol	118	0.00 U	114	97	37-116	29 *	0-25
95-95-4	MSD 2,4,5-Trichlorophenol	118	0.00 U	122	104	37-118	34 *	0-25
91-58-7	MSD 2-Chloronaphthalene	118	0.00 U	88.4	75	35-101	26	0-28
88-74-4	MSD 2-Nitroaniline o-Nitroaniline	118	0.00 U	127	108	36-126	27 *	0-26
99-09-2	MSD 3-Nitroaniline m-Nitroaniline	118	0.00 U	115	98	35-126	24	0-28
131-11-3	MSD Dimethylphthalate	118	0.00 U	124	105	42-125	27 *	0-25
606-20-2	MSD 2,6-Dinitrotoluene	118	0.00 U	116	99	48-118	26 *	0-23
121-14-2	MSD 2,4-Dinitrotoluene	118	0.00 U	115	98	49-120	26 *	0-23
208-96-8	MSD Acenaphthylene	118	0.00 U	93.5	80	35-114	23	0-25
83-32-9	MSD Acenaphthene	118	0.00 U	89.2	76	33-111	21	0-29
51-28-5	MSD 2,4-Dinitrophenol	118	0.00 U	92.0	78	21-131	4	0-25
132-64-9	MSD Dibenzofuran	118	0.00 U	99.6	85	43-113	22	0-24
58-90-2	MSD 2,3,4,6-Tetrachlorophenol	118	0.00 U	113	96	36-130	22	0-25
84-66-2	MSD Diethylphthalate	118	0.00 U	124	105	42-128	27 *	0-26
100-02-7	MSD 4-Nitrophenol	118	0.00 U	41.0	35	20-90	2	0-25

Semi-Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Matrix Spike Duplicate

Client ID: CAAN-12-2031MSD

Matrix: WG

Lab Sample ID: 1202595537

Instrument: MSD3.I

Analysis Date: 02/09/2012 13:48

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
86-73-7	MSD Fluorene	118	0.00 U	97.4	83	38-115	24	0-27
7005-72-3	MSD 4-Chlorophenylphenylether	118	0.00 U	90.7	77	41-110	19	0-26
100-01-6	MSD 4-Nitroaniline <i>p</i> -Nitroaniline	118	0.00 U	113	96	34-138	17	0-25
534-52-1	MSD 2-Methyl-4,6-dinitrophenol	118	0.00 U	110	94	32-124	21	0-25
122-39-4	MSD Diphenylamine	118	0.00 U	128	109	42-111	30 *	0-27
122-66-7	MSD Azobenzene <i>1,2-Diphenylhydrazine</i>	118	0.00 U	132	112	36-118	29 *	0-26
101-55-3	MSD 4-Bromophenylphenylether	118	0.00 U	104	88	40-112	26	0-27
118-74-1	MSD Hexachlorobenzene	118	0.00 U	112	95	40-116	25	0-26
87-86-5	MSD Pentachlorophenol	118	0.00 U	85.7	73	31-114	14	0-25
85-01-8	MSD Phenanthrene	118	0.00 U	111	95	41-119	26	0-29
120-12-7	MSD Anthracene	118	0.00 U	112	95	41-118	28	0-29
84-74-2	MSD Di-n-butylphthalate	118	0.00 U	129	109	41-128	33 *	0-25
206-44-0	MSD Fluoranthene	118	0.00 U	111	95	40-124	25	0-27
129-00-0	MSD Pyrene	118	0.00 U	110	93	34-126	40 *	0-25
85-68-7	MSD Butylbenzylphthalate	118	0.00 U	120	102	37-128	40 *	0-28
117-81-7	MSD bis(2-Ethylhexyl)phthalate	118	0.00 U	118	100	34-132	39 *	0-25
56-55-3	MSD Benzo(a)anthracene	118	0.00 U	115	98	38-120	29	0-29
218-01-9	MSD Chrysene	118	0.00 U	122	104	43-116	28	0-29
117-84-0	MSD Di-n-octylphthalate	118	0.00 U	126	107	30-135	29 *	0-25
205-99-2	MSD Benzo(b)fluoranthene	118	0.00 U	116	99	38-123	29 *	0-28
207-08-9	MSD Benzo(k)fluoranthene	118	0.00 U	120	102	40-123	33 *	0-29
50-32-8	MSD Benzo(a)pyrene	118	0.00 U	115	98	42-117	29 *	0-25

Semi-Volatile
Quality Control Summary
Spike Recovery Report

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SDG Number: 12-700

Sample Type: Matrix Spike Duplicate

Client ID: CAAN-12-2031MSD

Matrix: WG

Lab Sample ID: 1202595537

Instrument: MSD3.I

Analysis Date: 02/09/2012 13:48

Dilution: 1

Analyst: JLD1

Prep Batch ID: 1186831

Inj. Vol: 1 uL

Batch ID: 1186837

CAS No	Parmname	Amount Added ug/L	Sample Conc. ug/L	Spike Conc. ug/L	Recovery %	Acceptance Limits	RPD %	Acceptance Limits
193-39-5	MSD Indeno(1,2,3-cd)pyrene	118	0.00 U	120	102	33-135	27 *	0-25
53-70-3	MSD Dibenzo(a,h)anthracene	118	0.00 U	143	121	35-134	29 *	0-25
191-24-2	MSD Benzo(ghi)perylene	118	0.00 U	122	104	29-134	32 *	0-25
123-91-1	MSD 1,4-Dioxane	118	0.00 U	107	91	30-98	50 *	0-27
930-55-2	MSD N-Nitrosopyrrolidine	118	0.00 U	153	130 *	51-115	38 *	0-26
95-94-3	MSD 1,2,4,5-Tetrachlorobenzene	118	0.00 U	86.3	73	31-101	24	0-27
1912-24-9	MSD Atrazine	118	0.00 U	142	120 *	38-112	31 *	0-24
92-87-5	MSD Benzidine	235	0.00 U	64.8	28 *	30-120	7	0-25
91-94-1	MSD 3,3'-Dichlorobenzidine	118	0.00 U	134	114 *	28-112	21	0-25
120-82-1	MSD 1,2,4-Trichlorobenzene	118	0.00 U	79.3	67	25-94	34 *	0-28

Method Blank Summary

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SDG Number:	12-700	Client:	ARSL001	Matrix:	WATER
Client ID:	MB for batch 1186831	Instrument ID:	MSD3.I	Data File:	S020912.B\s3b0905.D
Lab Sample ID:	1202595534	Prep Date:	02/08/2012 17:46	Analyzed:	02/09/12 09:38
Column:	DB-5ms				

This method blank applies to the following samples and quality control samples:

Client Sample ID	Lab Sample ID	File ID	Date Analyzed	Time Analyzed
01 LCS for batch 1186831	1202595535	S020912.B\s3b0906.D	02/09/12	1003
02 CAAN-12-2031	295270002	S020912.B\s3b0907.D	02/09/12	1028
03 CAAN-12-2031MS	1202595536	S020912.B\s3b0908.D	02/09/12	1053
04 CAAN-12-2199	295270004	S020912.B\s3b0910.D	02/09/12	1143
05 CAAN-12-2201	295270007	S020912.B\s3b0911.D	02/09/12	1208
06 CAAN-12-2031MSD	1202595537	S020912.B\s3b0915.D	02/09/12	1348

Quality Control Data

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700		Matrix: WATER
Lab Sample ID: 1202595534		
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: MB for batch 1186831	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 09:38	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 1000 mL	Final Volume: 1 mL
Data File: S020912.B\s3b0905.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether	U	10.0	ug/L	3.00	10.0
62-75-9	N-Methyl-N-nitrosomethylamine	U	10.0	ug/L	3.00	10.0
110-86-1	Pyridine	U	10.0	ug/L	3.00	10.0
62-53-3	Aniline	U	10.0	ug/L	3.00	10.0
108-95-2	Phenol	U	10.0	ug/L	3.00	10.0
111-44-4	bis(2-Chloroethyl) ether	U	10.0	ug/L	3.00	10.0
95-57-8	2-Chlorophenol	U	10.0	ug/L	3.00	10.0
541-73-1	1,3-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
106-46-7	1,4-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
95-50-1	1,2-Dichlorobenzene	U	10.0	ug/L	3.00	10.0
100-51-6	Benzyl alcohol	U	10.0	ug/L	3.00	10.0
95-48-7	o-Cresol	U	10.0	ug/L	3.00	10.0
65794-96-9	m,p-Cresols	U	10.0	ug/L	3.00	10.0
621-64-7	N-Nitrosodi--n-propylamine	U	10.0	ug/L	3.00	10.0
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane	U	10.0	ug/L	3.00	10.0
98-95-3	Nitrobenzene	U	10.0	ug/L	3.00	10.0
78-59-1	Isophorone	U	10.0	ug/L	3.00	10.0
88-75-5	2-Nitrophenol	U	10.0	ug/L	3.00	10.0
105-67-9	2,4-Dimethylphenol	U	10.0	ug/L	3.00	10.0
111-91-1	bis(2-Chloroethoxy)methane	U	10.0	ug/L	3.00	10.0
120-83-2	2,4-Dichlorophenol	U	10.0	ug/L	3.00	10.0
65-85-0	Benzoic acid	U	20.0	ug/L	6.00	20.0
106-47-8	4-Chloroaniline	U	10.0	ug/L	3.00	10.0
87-68-3	Hexachlorobutadiene	U	10.0	ug/L	3.00	10.0
59-50-7	Parachlorometa cresol	U	10.0	ug/L	3.00	10.0
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene	U	1.00	ug/L	0.300	1.00
91-20-3	Naphthalene	U	1.00	ug/L	0.300	1.00
90-12-0	1-Methylnaphthalene	U	1.00	ug/L	0.300	1.00
77-47-4	Hexachlorocyclopentadiene	U	10.0	ug/L	3.00	10.0
88-06-2	2,4,6-Trichlorophenol	U	10.0	ug/L	3.00	10.0
95-95-4	2,4,5-Trichlorophenol	U	10.0	ug/L	3.00	10.0
91-58-7	2-Chloronaphthalene	U	1.00	ug/L	0.300	1.00
88-74-4	2-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate	U	10.0	ug/L	3.00	10.0
606-20-2	2,6-Dinitrotoluene	U	10.0	ug/L	3.00	10.0

**Semi-Volatile
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Sample Summary**

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SDG Number: 12-700		Matrix: WATER
Lab Sample ID: 1202595534		
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: MB for batch 1186831	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 09:38	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 1000 mL	Final Volume: 1 mL
Data File: S020912.B\s3b0905.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene	U	10.0	ug/L	3.00	10.0
208-96-8	Acenaphthylene	U	1.00	ug/L	0.300	1.00
83-32-9	Acenaphthene	U	1.00	ug/L	0.300	1.00
51-28-5	2,4-Dinitrophenol	U	20.0	ug/L	5.00	20.0
132-64-9	Dibenzofuran	U	10.0	ug/L	3.00	10.0
58-90-2	2,3,4,6-Tetrachlorophenol	U	10.0	ug/L	3.00	10.0
84-66-2	Diethylphthalate	U	10.0	ug/L	3.00	10.0
100-02-7	4-Nitrophenol	U	10.0	ug/L	3.00	10.0
86-73-7	Fluorene	U	1.00	ug/L	0.300	1.00
7005-72-3	4-Chlorophenylphenylether	U	10.0	ug/L	3.00	10.0
100-01-6	4-Nitroaniline	U	10.0	ug/L	3.00	10.0
	<i>p</i> -Nitroaniline					
534-52-1	2-Methyl-4,6-dinitrophenol	U	10.0	ug/L	3.00	10.0
122-39-4	Diphenylamine	U	10.0	ug/L	3.00	10.0
122-66-7	Azobenzene	U	10.0	ug/L	3.00	10.0
	<i>1,2-Diphenylhydrazine</i>					
101-55-3	4-Bromophenylphenylether	U	10.0	ug/L	3.00	10.0
118-74-1	Hexachlorobenzene	U	10.0	ug/L	3.00	10.0
87-86-5	Pentachlorophenol	U	10.0	ug/L	3.00	10.0
88-85-7	Dinoseb	U	10.0	ug/L	3.00	10.0
85-01-8	Phenanthrene	U	1.00	ug/L	0.300	1.00
120-12-7	Anthracene	U	1.00	ug/L	0.300	1.00
84-74-2	Di-n-butylphthalate	U	10.0	ug/L	3.00	10.0
206-44-0	Fluoranthene	U	1.00	ug/L	0.300	1.00
129-00-0	Pyrene	U	1.00	ug/L	0.300	1.00
85-68-7	Butylbenzylphthalate	U	10.0	ug/L	3.00	10.0
117-81-7	bis(2-Ethylhexyl)phthalate	U	10.0	ug/L	3.00	10.0
56-55-3	Benzo(a)anthracene	U	1.00	ug/L	0.300	1.00
218-01-9	Chrysene	U	1.00	ug/L	0.300	1.00
117-84-0	Di-n-octylphthalate	U	10.0	ug/L	3.00	10.0
205-99-2	Benzo(b)fluoranthene	U	1.00	ug/L	0.300	1.00
207-08-9	Benzo(k)fluoranthene	U	1.00	ug/L	0.300	1.00
50-32-8	Benzo(a)pyrene	U	1.00	ug/L	0.300	1.00
193-39-5	Indeno(1,2,3-cd)pyrene	U	1.00	ug/L	0.300	1.00
53-70-3	Dibenzo(a,h)anthracene	U	1.00	ug/L	0.300	1.00
191-24-2	Benzo(ghi)perylene	U	1.00	ug/L	0.300	1.00
123-91-1	1,4-Dioxane	U	10.0	ug/L	3.00	10.0
55-18-5	N-Nitrosodiethylamine	U	10.0	ug/L	3.00	10.0
930-55-2	N-Nitrosopyrrolidine	U	10.0	ug/L	3.00	10.0

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700	Matrix: WATER
Lab Sample ID: 1202595534	
Client Sample: QC for batch 1186831	Client: ARSL001
Client ID: MB for batch 1186831	Method: SW846 8270C
Batch ID: 1186837	Inst: MSD3.I
Run Date: 02/09/2012 09:38	Analyst: JLD1
Prep Date: 02/08/2012 17:46	Aliquot: 1000 mL
Data File: S020912.B\3b0905.D	Column: DB-5ms
	Project: QC
	SOP Ref: GL-OA-E-009
	Dilution: 1
	Inj. Vol: 1 uL
	Final Volume: 1 mL

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.0	ug/L	3.00	10.0
95-94-3	1,2,4,5-Tetrachlorobenzene	U	10.0	ug/L	3.00	10.0
608-93-5	Pentachlorobenzene	U	10.0	ug/L	3.00	10.0
1912-24-9	Atrazine	U	10.0	ug/L	3.00	10.0
92-87-5	Benzidine	U	10.0	ug/L	3.00	10.0
91-94-1	3,3'-Dichlorobenzidine	U	10.0	ug/L	3.00	10.0
120-82-1	1,2,4-Trichlorobenzene	U	10.0	ug/L	3.00	10.0

Surrogate/Tracer recovery	Result	Nominal	Recovery%	Acceptable Limits
2,4,6-Tribromophenol	95.3	100	ug/L	95.3 (30%-128%)
2-Fluorobiphenyl	41.4	50.0	ug/L	82.8 (34%-98%)
2-Fluorophenol	55.8	100	ug/L	55.8 (21%-78%)
Nitrobenzene-d5	47.4	50.0	ug/L	94.8 (39%-117%)
Phenol-d5	35.3	100	ug/L	35.3 (14%-80%)
p-Terphenyl-d14	42.2	50.0	ug/L	84.3 (39%-129%)

Tentatively Identified Compound Summary

CAS No.	Tentatively Identified Compound (TIC)	RT	Estimated	Units	Fit	Qual
No Tentatively Identified Compounds Found				ug/L		

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700		Matrix: WATER
Lab Sample ID: 1202595535		
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: LCS for batch 1186831	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 10:03	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 1000 mL	Final Volume: 1 mL
Data File: S020912.B\s3b0906.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether		38.5	ug/L	3.00	10.0
62-75-9	N-Methyl-N-nitrosomethylamine		24.9	ug/L	3.00	10.0
110-86-1	Pyridine		16.7	ug/L	3.00	10.0
62-53-3	Aniline		39.6	ug/L	3.00	10.0
108-95-2	Phenol		15.2	ug/L	3.00	10.0
111-44-4	bis(2-Chloroethyl) ether		40.2	ug/L	3.00	10.0
95-57-8	2-Chlorophenol		38.1	ug/L	3.00	10.0
541-73-1	1,3-Dichlorobenzene		29.3	ug/L	3.00	10.0
106-46-7	1,4-Dichlorobenzene		29.6	ug/L	3.00	10.0
95-50-1	1,2-Dichlorobenzene		30.3	ug/L	3.00	10.0
100-51-6	Benzyl alcohol		36.2	ug/L	3.00	10.0
95-48-7	o-Cresol		35.5	ug/L	3.00	10.0
65794-96-9	m,p-Cresols		39.0	ug/L	3.00	10.0
621-64-7	N-Nitrosodi--n-propylamine		49.3	ug/L	3.00	10.0
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane		28.2	ug/L	3.00	10.0
98-95-3	Nitrobenzene		41.9	ug/L	3.00	10.0
78-59-1	Isophorone		47.6	ug/L	3.00	10.0
88-75-5	2-Nitrophenol		40.8	ug/L	3.00	10.0
105-67-9	2,4-Dimethylphenol		40.5	ug/L	3.00	10.0
111-91-1	bis(2-Chloroethoxy)methane		39.3	ug/L	3.00	10.0
120-83-2	2,4-Dichlorophenol		41.6	ug/L	3.00	10.0
65-85-0	Benzoic acid		28.7	ug/L	6.00	20.0
106-47-8	4-Chloroaniline		44.1	ug/L	3.00	10.0
87-68-3	Hexachlorobutadiene		30.1	ug/L	3.00	10.0
59-50-7	Parachlorometa cresol		44.1	ug/L	3.00	10.0
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene		35.8	ug/L	0.300	1.00
91-20-3	Naphthalene		32.6	ug/L	0.300	1.00
90-12-0	1-Methylnaphthalene		38.0	ug/L	0.300	1.00
77-47-4	Hexachlorocyclopentadiene		19.9	ug/L	3.00	10.0
88-06-2	2,4,6-Trichlorophenol		39.9	ug/L	3.00	10.0
95-95-4	2,4,5-Trichlorophenol		41.0	ug/L	3.00	10.0
91-58-7	2-Chloronaphthalene		35.8	ug/L	0.300	1.00
88-74-4	2-Nitroaniline		43.3	ug/L	3.00	10.0
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline		38.5	ug/L	3.00	10.0
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate		43.9	ug/L	3.00	10.0
606-20-2	2,6-Dinitrotoluene		41.6	ug/L	3.00	10.0

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700		Matrix:	WATER
Lab Sample ID: 1202595535			
Client Sample: QC for batch 1186831	Client: ARSL001	Project:	QC
Client ID: LCS for batch 1186831	Method: SW846 8270C	SOP Ref:	GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution:	1
Run Date: 02/09/2012 10:03	Analyst: JLD1	Inj. Vol:	1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 1000 mL	Final Volume:	1 mL
Data File: S020912.B\s3b0906.D	Column: DB-5ms		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene		39.5	ug/L	3.00	10.0
208-96-8	Acenaphthylene		37.0	ug/L	0.300	1.00
83-32-9	Acenaphthene		37.1	ug/L	0.300	1.00
51-28-5	2,4-Dinitrophenol		31.3	ug/L	5.00	20.0
132-64-9	Dibenzofuran		39.3	ug/L	3.00	10.0
58-90-2	2,3,4,6-Tetrachlorophenol		39.4	ug/L	3.00	10.0
84-66-2	Diethylphthalate		43.9	ug/L	3.00	10.0
100-02-7	4-Nitrophenol	J	8.76	ug/L	3.00	10.0
86-73-7	Fluorene		38.3	ug/L	0.300	1.00
7005-72-3	4-Chlorophenylphenylether		37.9	ug/L	3.00	10.0
100-01-6	4-Nitroaniline		36.4	ug/L	3.00	10.0
	<i>p</i> -Nitroaniline					
534-52-1	2-Methyl-4,6-dinitrophenol		37.1	ug/L	3.00	10.0
122-39-4	Diphenylamine		45.7	ug/L	3.00	10.0
122-66-7	Azobenzene		49.6	ug/L	3.00	10.0
	<i>1,2-Diphenylhydrazine</i>					
101-55-3	4-Bromophenylphenylether		43.3	ug/L	3.00	10.0
118-74-1	Hexachlorobenzene		45.8	ug/L	3.00	10.0
87-86-5	Pentachlorophenol		31.0	ug/L	3.00	10.0
88-85-7	Dinoseb	U	10.0	ug/L	3.00	10.0
85-01-8	Phenanthrene		41.7	ug/L	0.300	1.00
120-12-7	Anthracene		39.8	ug/L	0.300	1.00
84-74-2	Di-n-butylphthalate		46.5	ug/L	3.00	10.0
206-44-0	Fluoranthene		39.5	ug/L	0.300	1.00
129-00-0	Pyrene		40.8	ug/L	0.300	1.00
85-68-7	Butylbenzylphthalate		43.9	ug/L	3.00	10.0
117-81-7	bis(2-Ethylhexyl)phthalate		43.0	ug/L	3.00	10.0
56-55-3	Benzo(a)anthracene		41.3	ug/L	0.300	1.00
218-01-9	Chrysene		43.9	ug/L	0.300	1.00
117-84-0	Di-n-octylphthalate		45.1	ug/L	3.00	10.0
205-99-2	Benzo(b)fluoranthene		44.9	ug/L	0.300	1.00
207-08-9	Benzo(k)fluoranthene		44.1	ug/L	0.300	1.00
50-32-8	Benzo(a)pyrene		41.0	ug/L	0.300	1.00
193-39-5	Indeno(1,2,3-cd)pyrene		37.9	ug/L	0.300	1.00
53-70-3	Dibenzo(a,h)anthracene		44.0	ug/L	0.300	1.00
191-24-2	Benzo(ghi)perylene		38.0	ug/L	0.300	1.00
123-91-1	1,4-Dioxane		26.5	ug/L	3.00	10.0
55-18-5	N-Nitrosodiethylamine	U	10.0	ug/L	3.00	10.0
930-55-2	N-Nitrosopyrrolidine		50.1	ug/L	3.00	10.0

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700	Matrix: WATER	
Lab Sample ID: 1202595535		
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: LCS for batch 1186831	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 10:03	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 1000 mL	Final Volume: 1 mL
Data File: S020912.B\3b0906.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	10.0	ug/L	3.00	10.0
95-94-3	1,2,4,5-Tetrachlorobenzene		37.6	ug/L	3.00	10.0
608-93-5	Pentachlorobenzene	U	10.0	ug/L	3.00	10.0
1912-24-9	Atrazine		54.4	ug/L	3.00	10.0
92-87-5	Benzidine		17.3	ug/L	3.00	10.0
91-94-1	3,3'-Dichlorobenzidine		40.2	ug/L	3.00	10.0
120-82-1	1,2,4-Trichlorobenzene		29.9	ug/L	3.00	10.0

Surrogate/Tracer recovery	Result	Nominal		Recovery%	Acceptable Limits
2,4,6-Tribromophenol	82.2	100	ug/L	82.2	(30%-128%)
2-Fluorobiphenyl	39.4	50.0	ug/L	78.9	(34%-98%)
2-Fluorophenol	49.6	100	ug/L	49.6	(21%-78%)
Nitrobenzene-d5	44.1	50.0	ug/L	88.1	(39%-117%)
Phenol-d5	31.3	100	ug/L	31.3	(14%-80%)
p-Terphenyl-d14	39.8	50.0	ug/L	79.6	(39%-129%)

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 1202595536	Date Received: 02/03/2012 08:45	
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: CAAN-12-2031MS	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 10:53	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 425 mL	Final Volume: 1 mL
Data File: S020912.B\s3b0908.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether		74.8	ug/L	7.06	23.5
62-75-9	N-Methyl-N-nitrosomethylamine		60.5	ug/L	7.06	23.5
110-86-1	Pyridine		46.1	ug/L	7.06	23.5
62-53-3	Aniline		89.6	ug/L	7.06	23.5
108-95-2	Phenol		52.5	ug/L	7.06	23.5
111-44-4	bis(2-Chloroethyl) ether		78.2	ug/L	7.06	23.5
95-57-8	2-Chlorophenol		74.6	ug/L	7.06	23.5
541-73-1	1,3-Dichlorobenzene		52.7	ug/L	7.06	23.5
106-46-7	1,4-Dichlorobenzene		53.4	ug/L	7.06	23.5
95-50-1	1,2-Dichlorobenzene		56.8	ug/L	7.06	23.5
100-51-6	Benzyl alcohol		81.4	ug/L	7.06	23.5
95-48-7	o-Cresol		77.9	ug/L	7.06	23.5
65794-96-9	m,p-Cresols		91.7	ug/L	7.06	23.5
621-64-7	N-Nitrosodi--n-propylamine		96.6	ug/L	7.06	23.5
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane		47.8	ug/L	7.06	23.5
98-95-3	Nitrobenzene		82.9	ug/L	7.06	23.5
78-59-1	Isophorone		93.8	ug/L	7.06	23.5
88-75-5	2-Nitrophenol		81.7	ug/L	7.06	23.5
105-67-9	2,4-Dimethylphenol		78.7	ug/L	7.06	23.5
111-91-1	bis(2-Chloroethoxy)methane		78.0	ug/L	7.06	23.5
120-83-2	2,4-Dichlorophenol		83.3	ug/L	7.06	23.5
65-85-0	Benzoic acid		116	ug/L	14.1	47.1
106-47-8	4-Chloroaniline		93.5	ug/L	7.06	23.5
87-68-3	Hexachlorobutadiene		50.5	ug/L	7.06	23.5
59-50-7	Parachlorometa cresol		95.9	ug/L	7.06	23.5
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene		69.1	ug/L	0.706	2.35
91-20-3	Naphthalene		65.1	ug/L	0.706	2.35
90-12-0	1-Methylnaphthalene		73.0	ug/L	0.706	2.35
77-47-4	Hexachlorocyclopentadiene		44.8	ug/L	7.06	23.5
88-06-2	2,4,6-Trichlorophenol		84.7	ug/L	7.06	23.5
95-95-4	2,4,5-Trichlorophenol		86.9	ug/L	7.06	23.5
91-58-7	2-Chloronaphthalene		68.1	ug/L	0.706	2.35
88-74-4	2-Nitroaniline		96.8	ug/L	7.06	23.5
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline		90.2	ug/L	7.06	23.5
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate		94.1	ug/L	7.06	23.5
606-20-2	2,6-Dinitrotoluene		89.5	ug/L	7.06	23.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	1202595536	Date Received:	02/03/2012 08:45		
Client Sample:	QC for batch 1186831	Client:	ARSL001	Project:	QC
Client ID:	CAAN-12-2031MS	Method:	SW846 8270C	SOP Ref:	GL-OA-E-009
Batch ID:	1186837	Inst:	MSD3.I	Dilution:	1
Run Date:	02/09/2012 10:53	Analyst:	JLD1	Inj. Vol:	1 uL
Prep Date:	02/08/2012 17:46	Aliquot:	425 mL	Final Volume:	1 mL
Data File:	S020912.B\s3b0908.D	Column:	DB-5ms		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene		88.3	ug/L	7.06	23.5
208-96-8	Acenaphthylene		74.2	ug/L	0.706	2.35
83-32-9	Acenaphthene		72.0	ug/L	0.706	2.35
51-28-5	2,4-Dinitrophenol		88.2	ug/L	11.8	47.1
132-64-9	Dibenzofuran		79.8	ug/L	7.06	23.5
58-90-2	2,3,4,6-Tetrachlorophenol		89.9	ug/L	7.06	23.5
84-66-2	Diethylphthalate		94.8	ug/L	7.06	23.5
100-02-7	4-Nitrophenol		41.8	ug/L	7.06	23.5
86-73-7	Fluorene		76.9	ug/L	0.706	2.35
7005-72-3	4-Chlorophenylphenylether		74.9	ug/L	7.06	23.5
100-01-6	4-Nitroaniline		96.0	ug/L	7.06	23.5
	<i>p</i> -Nitroaniline					
534-52-1	2-Methyl-4,6-dinitrophenol		89.7	ug/L	7.06	23.5
122-39-4	Diphenylamine		94.8	ug/L	7.06	23.5
122-66-7	Azobenzene		98.2	ug/L	7.06	23.5
	<i>1,2-Diphenylhydrazine</i>					
101-55-3	4-Bromophenylphenylether		80.2	ug/L	7.06	23.5
118-74-1	Hexachlorobenzene		86.9	ug/L	7.06	23.5
87-86-5	Pentachlorophenol		74.5	ug/L	7.06	23.5
88-85-7	Dinoseb	U	23.5	ug/L	7.06	23.5
85-01-8	Phenanthrene		85.4	ug/L	0.706	2.35
120-12-7	Anthracene		83.9	ug/L	0.706	2.35
84-74-2	Di-n-butylphthalate		92.5	ug/L	7.06	23.5
206-44-0	Fluoranthene		87.0	ug/L	0.706	2.35
129-00-0	Pyrene		73.0	ug/L	0.706	2.35
85-68-7	Butylbenzylphthalate		79.8	ug/L	7.06	23.5
117-81-7	bis(2-Ethylhexyl)phthalate		79.6	ug/L	7.06	23.5
56-55-3	Benzo(a)anthracene		86.3	ug/L	0.706	2.35
218-01-9	Chrysene		91.8	ug/L	0.706	2.35
117-84-0	Di-n-octylphthalate		94.2	ug/L	7.06	23.5
205-99-2	Benzo(b)fluoranthene		87.2	ug/L	0.706	2.35
207-08-9	Benzo(k)fluoranthene		86.8	ug/L	0.706	2.35
50-32-8	Benzo(a)pyrene		86.2	ug/L	0.706	2.35
193-39-5	Indeno(1,2,3-cd)pyrene		91.1	ug/L	0.706	2.35
53-70-3	Dibenzo(a,h)anthracene		107	ug/L	0.706	2.35
191-24-2	Benzo(ghi)perylene		88.4	ug/L	0.706	2.35
123-91-1	1,4-Dioxane		64.4	ug/L	7.06	23.5
55-18-5	N-Nitrosodiethylamine	U	23.5	ug/L	7.06	23.5
930-55-2	N-Nitrosopyrrolidine		104	ug/L	7.06	23.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 1202595536	Date Received: 02/03/2012 08:45	
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: CAAN-12-2031MS	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 10:53	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 425 mL	Final Volume: 1 mL
Data File: S020912.B\3b0908.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	23.5	ug/L	7.06	23.5
95-94-3	1,2,4,5-Tetrachlorobenzene		68.0	ug/L	7.06	23.5
608-93-5	Pentachlorobenzene	U	23.5	ug/L	7.06	23.5
1912-24-9	Atrazine		104	ug/L	7.06	23.5
92-87-5	Benzidine		69.5	ug/L	7.06	23.5
91-94-1	3,3'-Dichlorobenzidine		109	ug/L	7.06	23.5
120-82-1	1,2,4-Trichlorobenzene		56.5	ug/L	7.06	23.5

Surrogate/Tracer recovery	Result	Nominal		Recovery%	Acceptable Limits
2,4,6-Tribromophenol	181	235	ug/L	76.9	(30%-128%)
2-Fluorobiphenyl	76.4	118	ug/L	64.9	(34%-98%)
2-Fluorophenol	131	235	ug/L	55.7	(21%-78%)
Nitrobenzene-d5	85.3	118	ug/L	72.5	(39%-117%)
Phenol-d5	108	235	ug/L	46.0	(14%-80%)
p-Terphenyl-d14	74.5	118	ug/L	63.4	(39%-129%)

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 1202595537	Date Received: 02/03/2012 08:45	
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: CAAN-12-2031MSD	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 13:48	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 425 mL	Final Volume: 1 mL
Data File: S020912.B\s3b0915.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
39638-32-9	bis(2-Chloroisopropyl)ether		114	ug/L	7.06	23.5
62-75-9	N-Methyl-N-nitrosomethylamine		101	ug/L	7.06	23.5
110-86-1	Pyridine		41.6	ug/L	7.06	23.5
62-53-3	Aniline		108	ug/L	7.06	23.5
108-95-2	Phenol		79.2	ug/L	7.06	23.5
111-44-4	bis(2-Chloroethyl) ether		119	ug/L	7.06	23.5
95-57-8	2-Chlorophenol		115	ug/L	7.06	23.5
541-73-1	1,3-Dichlorobenzene		76.5	ug/L	7.06	23.5
106-46-7	1,4-Dichlorobenzene		78.9	ug/L	7.06	23.5
95-50-1	1,2-Dichlorobenzene		81.9	ug/L	7.06	23.5
100-51-6	Benzyl alcohol		120	ug/L	7.06	23.5
95-48-7	o-Cresol		116	ug/L	7.06	23.5
65794-96-9	m,p-Cresols		138	ug/L	7.06	23.5
621-64-7	N-Nitrosodi--n-propylamine		140	ug/L	7.06	23.5
	<i>N-Nitrosodipropylamine</i>					
67-72-1	Hexachloroethane		72.2	ug/L	7.06	23.5
98-95-3	Nitrobenzene		123	ug/L	7.06	23.5
78-59-1	Isophorone		134	ug/L	7.06	23.5
88-75-5	2-Nitrophenol		118	ug/L	7.06	23.5
105-67-9	2,4-Dimethylphenol		118	ug/L	7.06	23.5
111-91-1	bis(2-Chloroethoxy)methane		114	ug/L	7.06	23.5
120-83-2	2,4-Dichlorophenol		118	ug/L	7.06	23.5
65-85-0	Benzoic acid		153	ug/L	14.1	47.1
106-47-8	4-Chloroaniline		129	ug/L	7.06	23.5
87-68-3	Hexachlorobutadiene		70.7	ug/L	7.06	23.5
59-50-7	Parachlorometa cresol		135	ug/L	7.06	23.5
	<i>4-Chloro-3-methylphenol</i>					
91-57-6	2-Methylnaphthalene		91.9	ug/L	0.706	2.35
91-20-3	Naphthalene		91.3	ug/L	0.706	2.35
90-12-0	1-Methylnaphthalene		99.6	ug/L	0.706	2.35
77-47-4	Hexachlorocyclopentadiene		53.1	ug/L	7.06	23.5
88-06-2	2,4,6-Trichlorophenol		114	ug/L	7.06	23.5
95-95-4	2,4,5-Trichlorophenol		122	ug/L	7.06	23.5
91-58-7	2-Chloronaphthalene		88.4	ug/L	0.706	2.35
88-74-4	2-Nitroaniline		127	ug/L	7.06	23.5
	<i>o-Nitroaniline</i>					
99-09-2	3-Nitroaniline		115	ug/L	7.06	23.5
	<i>m-Nitroaniline</i>					
131-11-3	Dimethylphthalate		124	ug/L	7.06	23.5
606-20-2	2,6-Dinitrotoluene		116	ug/L	7.06	23.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

SDG Number: 12-700	Date Collected: 02/01/2012 12:00	Matrix: WG
Lab Sample ID: 1202595537	Date Received: 02/03/2012 08:45	
Client Sample: QC for batch 1186831	Client: ARSL001	Project: QC
Client ID: CAAN-12-2031MSD	Method: SW846 8270C	SOP Ref: GL-OA-E-009
Batch ID: 1186837	Inst: MSD3.I	Dilution: 1
Run Date: 02/09/2012 13:48	Analyst: JLD1	Inj. Vol: 1 uL
Prep Date: 02/08/2012 17:46	Aliquot: 425 mL	Final Volume: 1 mL
Data File: S020912.B\s3b0915.D	Column: DB-5ms	

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
121-14-2	2,4-Dinitrotoluene		115	ug/L	7.06	23.5
208-96-8	Acenaphthylene		93.5	ug/L	0.706	2.35
83-32-9	Acenaphthene		89.2	ug/L	0.706	2.35
51-28-5	2,4-Dinitrophenol		92.0	ug/L	11.8	47.1
132-64-9	Dibenzofuran		99.6	ug/L	7.06	23.5
58-90-2	2,3,4,6-Tetrachlorophenol		113	ug/L	7.06	23.5
84-66-2	Diethylphthalate		124	ug/L	7.06	23.5
100-02-7	4-Nitrophenol		41.0	ug/L	7.06	23.5
86-73-7	Fluorene		97.4	ug/L	0.706	2.35
7005-72-3	4-Chlorophenylphenylether		90.7	ug/L	7.06	23.5
100-01-6	4-Nitroaniline		113	ug/L	7.06	23.5
	<i>p</i> -Nitroaniline					
534-52-1	2-Methyl-4,6-dinitrophenol		110	ug/L	7.06	23.5
122-39-4	Diphenylamine		128	ug/L	7.06	23.5
122-66-7	Azobenzene		132	ug/L	7.06	23.5
	<i>1,2-Diphenylhydrazine</i>					
101-55-3	4-Bromophenylphenylether		104	ug/L	7.06	23.5
118-74-1	Hexachlorobenzene		112	ug/L	7.06	23.5
87-86-5	Pentachlorophenol		85.7	ug/L	7.06	23.5
88-85-7	Dinoseb	U	23.5	ug/L	7.06	23.5
85-01-8	Phenanthrene		111	ug/L	0.706	2.35
120-12-7	Anthracene		112	ug/L	0.706	2.35
84-74-2	Di-n-butylphthalate		129	ug/L	7.06	23.5
206-44-0	Fluoranthene		111	ug/L	0.706	2.35
129-00-0	Pyrene		110	ug/L	0.706	2.35
85-68-7	Butylbenzylphthalate		120	ug/L	7.06	23.5
117-81-7	bis(2-Ethylhexyl)phthalate		118	ug/L	7.06	23.5
56-55-3	Benzo(a)anthracene		115	ug/L	0.706	2.35
218-01-9	Chrysene		122	ug/L	0.706	2.35
117-84-0	Di-n-octylphthalate		126	ug/L	7.06	23.5
205-99-2	Benzo(b)fluoranthene		116	ug/L	0.706	2.35
207-08-9	Benzo(k)fluoranthene		120	ug/L	0.706	2.35
50-32-8	Benzo(a)pyrene		115	ug/L	0.706	2.35
193-39-5	Indeno(1,2,3-cd)pyrene		120	ug/L	0.706	2.35
53-70-3	Dibenzo(a,h)anthracene	E	143	ug/L	0.706	2.35
191-24-2	Benzo(ghi)perylene		122	ug/L	0.706	2.35
123-91-1	1,4-Dioxane		107	ug/L	7.06	23.5
55-18-5	N-Nitrosodiethylamine	U	23.5	ug/L	7.06	23.5
930-55-2	N-Nitrosopyrrolidine		153	ug/L	7.06	23.5

**Semi-Volatile
Certificate of Analysis
Sample Summary**

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SDG Number:	12-700	Date Collected:	02/01/2012 12:00	Matrix:	WG
Lab Sample ID:	1202595537	Date Received:	02/03/2012 08:45		
Client Sample:	QC for batch 1186831	Client:	ARSL001	Project:	QC
Client ID:	CAAN-12-2031MSD	Method:	SW846 8270C	SOP Ref:	GL-OA-E-009
Batch ID:	1186837	Inst:	MSD3.I	Dilution:	1
Run Date:	02/09/2012 13:48	Analyst:	JLD1	Inj. Vol:	1 uL
Prep Date:	02/08/2012 17:46	Aliquot:	425 mL	Final Volume:	1 mL
Data File:	S020912.B\3b0915.D	Column:	DB-5ms		

CAS No.	Parmname	Qualifier	Result	Units	MDL/LOD	PQL/LOQ
924-16-3	N-Nitrosodi-n-butylamine	U	23.5	ug/L	7.06	23.5
95-94-3	1,2,4,5-Tetrachlorobenzene		86.3	ug/L	7.06	23.5
608-93-5	Pentachlorobenzene	U	23.5	ug/L	7.06	23.5
1912-24-9	Atrazine		142	ug/L	7.06	23.5
92-87-5	Benzidine		64.8	ug/L	7.06	23.5
91-94-1	3,3'-Dichlorobenzidine		134	ug/L	7.06	23.5
120-82-1	1,2,4-Trichlorobenzene		79.3	ug/L	7.06	23.5

Surrogate/Tracer recovery	Result	Nominal		Recovery%	Acceptable Limits
2,4,6-Tribromophenol	224	235	ug/L	95.4	(30%-128%)
2-Fluorobiphenyl	103	118	ug/L	87.8	(34%-98%)
2-Fluorophenol	204	235	ug/L	86.8	* (21%-78%)
Nitrobenzene-d5	125	118	ug/L	107	(39%-117%)
Phenol-d5	165	235	ug/L	70.1	(14%-80%)
p-Terphenyl-d14	104	118	ug/L	88.4	(39%-129%)

Miscellaneous

DATA EXCEPTION REPORT

Mo.Day Yr. 09-FEB-12	Division: Industrial	Quality Criteria: Specifications	Type: Process
Instrument Type: SEMIVOA GC/MS	Test / Method: SW846 8270C	Matrix Type: Liquid	Client Code: ESHL
Batch ID: 1186837	Sample Numbers: See Below		

Potentially affected work order(s)(SDG): 295270(12-700),295388(12-711),295392(12-718),295435(12-726)

Application Issues:

Failed Recovery for MS/PS
Failed RPD for MS/MSD, or PS/PSD
Failed Recovery for LCS/LCSD
Failed Yield for Surrogates
Failed Recovery for MSD/PSD

**Specification and Requirements
Exception Description:**

DER Disposition:

1. The LCS(1202595535) recovered Benzidine at 17.3% (limits: 23%-131%) and Benzoic acid 28.7% (limits: 30%-142%).
2. The MS(1202595536) and MSD(1202595537) recovered Benzidine at 29.6% and 27.5%, respectively. The limits are 30%-120%. The MSD also recovered N-Nitrosopyrrolidine at 130% (Limits are 51-115%), Atrazine at 120% (Limits are 38-112%) and 3,3'-Dichlorobenzidine at 114% (Limits are 28-112%).
3. The MSD(1202595537) displayed one biased high surrogate and multiple biased high spike recoveries. Please see the QC Summary for the specific recovery values.
4. The MS(1202595536) and MSD(1202595537) displayed multiple RPD value failures. Please see the QC Summary for the specific recovery values.

1. The failures represented less than 5% of the total requested spike analyte list. That satisfied the acceptance criteria for the client and the data were reported. Please note, both Benzidine and Benzoic acid are designated as being poor responding analytes in the analytical method. This may account for the biased low recoveries for those analytes.
2. Since the MS(1202595536) and MSD(1202595537) displayed similar recoveries for Benzidine, the failures may attributed to matrix interference. Please note, Benzidine is designated as being poor responding analyte in the analytical method. This may account for the biased low recoveries for that analyte in the MS and MSD. The latter three analytes were not detected in the associated samples. The data is unaffected and are reported.
3. The MB, LCS and MS satisfied batch QC acceptance criteria (with the exception of Benzidine in the MS and MSD - see item #2 above). Therefore, it was determined that the biased high failures were limited to the MSD sample only (the surrogate and spike recoveries were in general significantly higher in the MSD compared to the MS recoveries). Since the associated parent sample was reported as non-detected for all requested target analytes, a re-extraction was considered un-necessary and the data have been reported.
4. The RPD failures were attributed to the biased high recoveries in the MSD (see item #3 above).

Originator's Name:

Jennifer Dunagan Jones10-FEB-12

Data Validator/Group Leader:

Herbert Maier 22-FEB-12

Explosives by LCMSMS Analysis

Case Narrative

**LC/MS/MS Case Narrative
ARS International (ARSL)
SDG 12-700**

Method/Analysis Information

Procedure: **Definitive Low Level Analysis of Nitroaromatic Explosives Utilizing Liquid Chromatography / Mass Spectrometry / Mass Spectrometry (LC/MS/MS) by SW-846 Method 8321 Modified (8321M)**

Analytical Method: SW846 3535/8321A Modified

Prep Method: SW846 Method 3535

Analytical Batch Number: 1185693

Prep Batch Number: 1185692

Sample Analysis

The following samples were analyzed using the analytical protocol as established in SW846 3535/8321A Modified :

Sample ID	Client ID
295270003	CAAN-12-2031
295270005	CAAN-12-2199
1202592978	Method Blank (MB)
1202592979	Laboratory Control Sample (LCS)
1202592980	295270003(CAAN-12-2031) Matrix Spike (MS)
1202592981	295270003(CAAN-12-2031) Matrix Spike Duplicate (MSD)

Preparation/Analytical Method Verification

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-OA-E-056 REV# 16.

Primary Analyte Analysis

Calibration Information

Initial Calibration

All initial calibration requirements for this analysis have been met for this SDG.

Calibration Verification Standard Requirements

All associated calibration verification standard(s) (ICV or CCV) for this analysis met the acceptance criteria.

Calibration Blank Requirements

All initial or continuing calibration blanks (ICB or CCB) bracketing the analyses associated with this batch for this analysis were within acceptance criteria. Due to software limitations, the CCBs and/or the ICBs may have a concentration for target analytes in the Found column. These values should be zero.

CRI Requirements

All low level calibration verification (CRI) requirements for this analysis were met by all bracketing CRI standards.

Quality Control (QC) Information**Method Blank (MB) Statement**

The MB(s) analyzed with this SDG for this analysis met the acceptance criteria.

Surrogate Recoveries

All the surrogate recoveries were within the established acceptance criteria in this SDG in this analytical batch for this analysis.

Laboratory Control Sample (LCS) Recovery

The LCS spike recoveries were within the established acceptance limits.

QC Sample Designation

Sample 295270003 (CAAN-12-2031) was chosen for matrix spike and matrix spike duplicate analysis.

Matrix Spike (MS) Recovery Statement

The MS spike recoveries were within the established acceptance limits.

Matrix Spike Duplicate (MSD) Recovery Statement

The MSD spike recoveries were within the established acceptance limits.

MS/MSD Relative Percent Difference (RPD) Statement

The RPD(s) between the MS and MSD met the acceptance limits for this analysis.

Internal Standard (ISTD) Acceptance

The internal standard responses were within the required acceptance criteria for all samples and QC in this SDG.

Technical Information**Holding Time Specifications**

All samples in this SDG in this analytical batch met the specified holding time. GEL assigns holding times based on the associated methodology, which assigns the date and time from sample collection of sample receipt. Those holding times expressed in hours are calculated in the AlphaLIMS system. Those holding times expressed as days expire at midnight on the day of expiration.

Preparation/Analytical Method Verification

All procedures were performed as stated in the SOP.

Sample Dilutions

According to the GEL SOP for Method 8321A, all sample and QC extracts are diluted 1:1 v/v with HPLC grade water. The samples in this SDG in this analytical batch for this analysis did not require any additional dilutions.

Sample Re-extraction/Re-analysis

The SDG was re-analyzed due to a CCV failing acceptance criteria. The re-analysis passed acceptance criteria and is reported.

Secondary Analyte Analysis

Calibration Information

Initial Calibration

All initial calibration requirements for this analysis have been met for this SDG.

Calibration Verification Standard Requirements

All associated calibration verification standard(s) (ICV or CCV) for this analysis met the acceptance criteria.

Calibration Blank Requirements

All initial or continuing calibration blanks (ICB or CCB) bracketing the analyses associated with this batch for this analysis were within acceptance criteria. Due to software limitations, the CCBs and/or the ICBs may have a concentration for target analytes in the Found column. These values should be zero.

CRI Requirements

All low level calibration verification (CRI) requirements for this analysis were met by all bracketing CRI standards.

Quality Control (QC) Information

Method Blank (MB) Statement

The MB(s) analyzed with this SDG for this analysis met the acceptance criteria.

Surrogate Recoveries

All the surrogate recoveries were within the established acceptance criteria in this SDG in this analytical batch for this analysis.

Laboratory Control Sample (LCS) Recovery

The LCS spike recoveries were within the established acceptance limits.

QC Sample Designation

Sample 295270003 (CAAN-12-2031) was chosen for matrix spike and matrix spike duplicate analysis.

Matrix Spike (MS) Recovery Statement

The MS spike recoveries were within the established acceptance limits.

Matrix Spike Duplicate (MSD) Recovery Statement

The MSD spike recoveries were within the established acceptance limits.

MS/MSD Relative Percent Difference (RPD) Statement

The RPD(s) between the MS and MSD met the acceptance limits for this analysis.

Internal Standard (ISTD) Acceptance

The internal standards were not added to the secondary analyte extracts.

Technical Information

Holding Time Specifications

All samples in this SDG in this analytical batch met the specified holding time. GEL assigns holding times based on the associated methodology, which assigns the date and time from sample collection of sample receipt. Those holding times expressed in hours are calculated in the AlphaLIMS system. Those holding times expressed as days expire at midnight on the day of expiration.

Preparation/Analytical Method Verification

All procedures were performed as stated in the SOP.

Sample Dilutions

According to the GEL SOP for Method 8321A, all sample and QC extracts are diluted 1:1 v/v with HPLC grade water. The samples in this SDG in this analytical batch for this analysis did not require any additional dilutions.

Sample Re-extraction/Re-analysis

Re-extractions or re-analyses were not required in this SDG in this analytical batch for this analysis except for dilutions.

Miscellaneous Information**Data Exception (DER) Documentation**

Data exception reports (DERs) are generated to document procedural anomalies that may deviate from referenced SOP or contractual documents. A data exception report (DER) was not generated for this SDG.

Manual Integrations

Some initial calibration standards, continuing calibration standards, and/or samples required manual integrations due to software limitations..

Flagging Convention

The samples were not originally analyzed using SW-846 Method 8330.

Additional Comments

Due to software limitations, all initial calibration blanks must be designated as XIB001 in order for the forms to be correct.

Due to software limitations in the Secondary analyte analysis, false positives and analytes detected below the MDL cannot be deleted from the raw data.

Due to software limitations, file extensions such as DL, RE, etc. may not appear on the generated forms and/or raw data.

System Configuration

The laboratory utilizes a Waters LC 2795 liquid chromatography instrument for primary analyte analysis. It is coupled with either a Micromass Quattro Micro Mass Spectrometer/ Mass Spectrometer, or a Micromass Quattro Ultima Mass Spectrometer/ Mass Spectrometer. Each being designated as LCMSMS #1 and LCMSMS #2, respectively. It is fitted with an APCI (Atmospheric Pressure chemical Ionization) probe that is operated in the negative ionization mode for the primary analyte analysis. The laboratory also utilizes an Agilent 1100 liquid chromatography instrument for either primary or secondary analyte analysis. It is coupled with an Applied Biosystems 4000 Mass Spectrometer/ Mass Spectrometer, designated as either LCMSMS #3 or LCMSMS #4. It is fitted with an APCI (Atmospheric Pressure chemical Ionization) probe that is operated in the negative ionization mode for both the primary and secondary analyte analysis.

Electronic Packaging Comment

This data package was generated using an electronic data processing program referred to as virtual packaging. In an effort to increase quality and efficiency, the laboratory has developed systems to generate all data packages electronically. The following change from traditional packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. An electronic signature page inserted after the case narrative will include the data validator's signature and title. The signature page also includes the data qualifiers used in the fractional package. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

Chromatographic Columns

The detection of the primary analyte nitroaromatic and nitramines is accomplished through analysis on the following reversed phase column:

Phenomenex: Ultracarb 5u ODS (20), 250 x 4.60 mm ID.

The detection of the secondary analytes is accomplished through analysis on the following reversed phase column:

YMC: J'sphere ODS-H80, 150 x 4.6mm I.D.

Certification Statement

Where the analytical method has been performed under NELAP certification, the analysis has met all of the requirements of the NELAC standard unless otherwise noted in the analytical case narrative.

GEL LABORATORIES LLC

2040 Savage Road Charleston SC 29407 - (843) 556-8171 - www.gel.com

Qualifier Definition Report for

ARSL001 ARS International (63641-10)

Client SDG: 12-700 GEL Work Order: 295270

The Qualifiers in this report are defined as follows:

- * A quality control analyte recovery is outside of specified acceptance criteria
- ** Analyte is a surrogate compound
- U Analyte was analyzed for, but not detected above the MDL, MDA, or LOD.
- DL Indicates that sample is diluted.
- RA Indicates that sample is re-analyzed without re-extraction.
- RE Indicates that sample is re-extracted.

Review/Validation

GEL requires all analytical data to be verified by a qualified data reviewer. In addition, all CLP-like deliverables receive a third level review of the fractional data package.

The following data validator verified the information presented in this data report:

Signature: 

Name: Herbert Maier

Date: 28 FEB 2012

Title: Data Validator

Sample Data Summary

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270003

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227015a

Date Analyzed: 28-FEB-12 01:51

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
118-96-7	2,4,6-Trinitrotoluene	0.263	U	0.0842	0.263
<i>118-96-7</i>	<i>2,4,6-Trinitrotoluene</i>				
121-14-2	2,4-Dinitrotoluene	0.263	U	0.0842	0.263
<i>121-14-2</i>	<i>2,4-Dinitrotoluene</i>				
121-82-4	RDX	0.263	U	0.0842	0.263
<i>121-82-4</i>	<i>RDX</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				
2691-41-0	HMX	0.263	U	0.0842	0.263
<i>2691-41-0</i>	<i>HMX</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
606-20-2	2,6-Dinitrotoluene	0.263	U	0.0842	0.263
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
88-72-2	o-Nitrotoluene	0.263	U	0.0863	0.263
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
98-95-3	Nitrobenzene	0.263	U	0.0842	0.263
<i>98-95-3</i>	<i>Nitrobenzene</i>				
99-08-1	m-Nitrotoluene	0.263	U	0.0842	0.263
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	0.263	U	0.0842	0.263
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
99-65-0	m-Dinitrobenzene	0.263	U	0.0842	0.263
<i>99-65-0</i>	<i>m-Dinitrobenzene</i>				
479-45-8	Tetryl	0.526	U	0.0842	0.526
<i>479-45-8</i>	<i>Tetryl</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270003

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
78-11-5	PETN	0.526	U	0.105	0.526
<i>78-11-5</i>	<i>PETN</i>				
99-99-0	p-Nitrotoluene	0.526	U	0.158	0.526
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270003

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160016.wiff

Date Analyzed: 16-FEB-12 18:19

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
3058-38-6	TATB	1.05	U	0.316	1.05
<i>3058-38-6</i>	<i>TATB</i>				
618-87-1	3,5-Dinitroaniline	1.05	U	0.316	1.05
<i>618-87-1</i>	<i>3,5-Dinitroaniline</i>				
78-30-8	tris(o-cresyl) phosphate	1.05	U	0.316	1.05
<i>78-30-8</i>	<i>tris(o-cresyl) phosphate</i>				
59229-75-3	2,6-Diamino-4-nitrotoluene	2.63	U	0.526	2.63
<i>59229-75-3</i>	<i>2,6-Diamino-4-nitrotoluene</i>				
6629-29-4	2,4-Diamino-6-nitrotoluene	2.63	U	0.526	2.63
<i>6629-29-4</i>	<i>2,4-Diamino-6-nitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2199

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270005

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227018a

Date Analyzed: 28-FEB-12 03:20

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
118-96-7	2,4,6-Trinitrotoluene	0.263	U	0.0842	0.263
<i>118-96-7</i>	<i>2,4,6-Trinitrotoluene</i>				
121-14-2	2,4-Dinitrotoluene	0.263	U	0.0842	0.263
<i>121-14-2</i>	<i>2,4-Dinitrotoluene</i>				
121-82-4	RDX	0.263	U	0.0842	0.263
<i>121-82-4</i>	<i>RDX</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				
2691-41-0	HMX	0.263	U	0.0842	0.263
<i>2691-41-0</i>	<i>HMX</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	0.263	U	0.0842	0.263
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
606-20-2	2,6-Dinitrotoluene	0.263	U	0.0842	0.263
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
88-72-2	o-Nitrotoluene	0.263	U	0.0863	0.263
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
98-95-3	Nitrobenzene	0.263	U	0.0842	0.263
<i>98-95-3</i>	<i>Nitrobenzene</i>				
99-08-1	m-Nitrotoluene	0.263	U	0.0842	0.263
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	0.263	U	0.0842	0.263
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
99-65-0	m-Dinitrobenzene	0.263	U	0.0842	0.263
<i>99-65-0</i>	<i>m-Dinitrobenzene</i>				
479-45-8	Tetryl	0.526	U	0.0842	0.526
<i>479-45-8</i>	<i>Tetryl</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2199

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270005

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
78-11-5	PETN	0.526	U	0.105	0.526
<i>78-11-5</i>	<i>PETN</i>				
99-99-0	p-Nitrotoluene	0.526	U	0.158	0.526
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2199

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 295270005

Sample Amount 950 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160019.wiff

Date Analyzed: 16-FEB-12 19:09

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
3058-38-6	TATB	1.05	U	0.316	1.05
3058-38-6	TATB				
618-87-1	3,5-Dinitroaniline	1.05	U	0.316	1.05
618-87-1	3,5-Dinitroaniline				
78-30-8	tris(o-cresyl) phosphate	1.05	U	0.316	1.05
78-30-8	tris(o-cresyl) phosphate				
59229-75-3	2,6-Diamino-4-nitrotoluene	2.63	U	0.526	2.63
59229-75-3	2,6-Diamino-4-nitrotoluene				
6629-29-4	2,4-Diamino-6-nitrotoluene	2.63	U	0.526	2.63
6629-29-4	2,4-Diamino-6-nitrotoluene				

Quality Control Summary

High Explosives Surrogate Recovery Summary

Lab Name: GEL Laboratories LLCGEL Job No (SDG): 12-700Lab Code: GELHPLC Column: Phenomenex Ultracarb 5u ODS(20)

Lab Sample ID	Client Sample ID	DNT	QC Limits	Flg
295270003	CAAN-12-2031	97.9	66 - 130	
295270003	CAAN-12-2031	108	66 - 130	
295270005	CAAN-12-2199	96.4	66 - 130	
295270005	CAAN-12-2199	102	66 - 130	
1202592978	MB for batch 1185692	89.8	66 - 130	
1202592978	MB for batch 1185692	104	66 - 130	
1202592979	LCS for batch 1185692	93.1	66 - 130	
1202592979	LCS for batch 1185692	108	66 - 130	
1202592980	CAAN-12-2031(295270003MS)	95.8	66 - 130	
1202592980	CAAN-12-2031(295270003MS)	103	66 - 130	
1202592981	CAAN-12-2031(295270003MSD)	96.1	66 - 130	
1202592981	CAAN-12-2031(295270003MSD)	101	66 - 130	

DNT = 3,4-Dinitrotoluene

3B
High Explosives LCS/LCS Duplicate Summary

Lab Name: GEL Laboratories LLC

Client ID: LCS

Lab Code: GEL

GEL Job No (SDG) 12-700

Extract Batch Code: 1185692

Date Extracted: 07-FEB-12

GEL LCS ID: 1202592979

GEL LCSDUP ID: .

Analysis Date/Time: 28-FEB-12 01:21

DUP Analysis Date/Time:

Reporting Units: ug/L

QC Type: LCS/LCSD

Compound	Spike Added	LCS Conc	LCS Rec #	LCSD Conc	LCSD Rec #	RPD #	RPD	Recovery Limits
Tetryl	5	4	80					67 - 118
m-Dinitrobenzene	5	4.52	90.4					80 - 120
m-Nitrotoluene	5	4.02	80.5					67 - 111
1,3,5-Trinitrobenzene	5	4.49	89.7					69 - 120
o-Nitrotoluene	5	4.07	81.4					63 - 114
p-Nitrotoluene	5	4.36	87.2					66 - 118
2,4,6-Trinitrotoluene	5	4.73	94.6					74 - 131
2,6-Dinitrotoluene	5	4.56	91.1					80 - 120
4-Amino-2,6-dinitrotoluene	5	4.57	91.3					73 - 125
RDX	5	4.55	91					74 - 135
PETN	5	4.52	90.3					56 - 140
Nitrobenzene	5	4.26	85.2					66 - 112
HMX	5	4.22	84.4					56 - 122
2-Amino-4,6-dinitrotoluene	5	4.49	89.9					72 - 127
2,4-Dinitrotoluene	5	4.83	96.7					77 - 121

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

* Values outside of QC limits

3B
High Explosives LCS/LCS Duplicate Summary

Lab Name: GEL Laboratories LLC

Client ID: LCS

Lab Code: GEL

GEL Job No (SDG) 12-700

Extract Batch Code: 1185692

Date Extracted: 07-FEB-12

GEL LCS ID: 1202592979

GEL LCSDUP ID: .

Analysis Date/Time: 16-FEB-12 18:02

DUP Analysis Date/Time:

Reporting Units: ug/L

QC Type: LCS/LCSD

Compound	Spike Added	LCS Conc	LCS Rec #	LCSD Conc	LCSD Rec #	RPD #	RPD	Recovery Limits
2,6-Diamino-4-nitrotoluene	5	5.51	110					52 - 117
3,5-Dinitroaniline	5	5.47	109					70 - 120
TATB	10	3.73	37.3					21 - 147
tris(o-cresyl) phosphate	5	3.49	69.8					41 - 92
2,4-Diamino-6-nitrotoluene	5	4.76	95.2					51 - 100

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

* Values outside of QC limits

3
High Explosives MS/MSD Summary

Lab Name: GEL Laboratories LLC

Client ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Extract Batch Code: 1185692

Date Extracted: 07-FEB-12

GEL Spike ID: 1202592980

GEL SpikeDup ID: 1202592981

Analysis Date/Time: 28-FEB-12 02:21

MSD Analysis Date/Time: 28-FEB-12 02:50

Reporting Units: ug/L

QC Type: MS/MSD

Compound	Spike Added	Sample Conc	MS Conc	MS Rec #	MSD Conc	MSD Rec #	RPD #	RPD Limit	Rec Limits
2,4,6-Trinitrotoluene	5.31915	0	5.06	95.1	5.53	103	8.98	23	65 - 138
2,4-Dinitrotoluene	5.31915	0	5.15	96.8	5.2	96.6	.887	22	77 - 123
2,6-Dinitrotoluene	5.31915	0	4.63	87	5.08	94.5	9.33	20	78 - 117
2-Amino-4,6-dinitrotoluene	5.31915	0	4.58	86.2	4.96	92.2	7.84	24	72 - 134
4-Amino-2,6-dinitrotoluene	5.31915	0	4.8	90.3	5.06	94.1	5.18	22	70 - 134
HMX	5.31915	0	4.36	82.1	4.38	81.5	.432	31	48 - 137
Nitrobenzene	5.31915	0	4.54	85.4	4.53	84.3	.136	26	66 - 116
PETN	5.31915	0	4.8	90.2	5.14	95.6	6.85	25	50 - 148
RDX	5.31915	0	4.9	92.2	5.05	94	3.04	28	61 - 150
1,3,5-Trinitrobenzene	5.31915	0	4.59	86.4	4.69	87.3	2.16	23	51 - 120
Tetryl	5.31915	0	4.1	77.1	4.16	77.4	1.45	29	28 - 118
m-Dinitrobenzene	5.31915	0	5.06	95.2	5.2	96.6	2.61	20	70 - 130
m-Nitrotoluene	5.31915	0	4.26	80.1	4.53	84.3	6.28	26	66 - 115
o-Nitrotoluene	5.31915	0	4.13	77.7	4.56	84.8	9.73	26	66 - 118
p-Nitrotoluene	5.31915	0	4.32	81.2	4.58	85.3	5.99	24	68 - 122

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

3
High Explosives MS/MSD Summary

Lab Name: GEL Laboratories LLC

Client ID: CAAN-12-2031

Lab Code: GEL

GEL Job No (SDG) 12-700

Extract Batch Code: 1185692

Date Extracted: 07-FEB-12

GEL Spike ID: 1202592980

GEL SpikeDup ID: 1202592981

Analysis Date/Time: 16-FEB-12 18:36

MSD Analysis Date/Time: 16-FEB-12 18:52

Reporting Units: ug/L

QC Type: MS/MSD

Compound	Spike Added	Sample Conc	MS Conc	MS Rec #	MSD Conc	MSD Rec #	RPD #	RPD Limit	Rec Limits
2,4-Diamino-6-nitrotoluene	5.31915	0	5.04	94.8	5.35	99.6	6.01	26	43 - 109
2,6-Diamino-4-nitrotoluene	5.31915	0	5.61	105	5.3	98.6	5.6	26	37 - 125
3,5-Dinitroaniline	5.31915	0	5.69	107	5.62	105	1.2	20	66 - 117
TATB	10.6383	0	3.7	34.8	3.76	35	1.64	21	30 - 161
tris(o-cresyl) phosphate	5.31915	.0297	3.94	73.4	3.85	71	2.23	28	37 - 88

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

Quality Control Data

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: MB for batch 1185692

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592978

Sample Amount 1000 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227013a

Date Analyzed: 28-FEB-12 00:52

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
118-96-7	2,4,6-Trinitrotoluene	0.250	U	0.080	0.250
<i>118-96-7</i>	<i>2,4,6-Trinitrotoluene</i>				
121-14-2	2,4-Dinitrotoluene	0.250	U	0.080	0.250
<i>121-14-2</i>	<i>2,4-Dinitrotoluene</i>				
121-82-4	RDX	0.250	U	0.080	0.250
<i>121-82-4</i>	<i>RDX</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	0.250	U	0.080	0.250
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				
2691-41-0	HMX	0.250	U	0.080	0.250
<i>2691-41-0</i>	<i>HMX</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	0.250	U	0.080	0.250
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
606-20-2	2,6-Dinitrotoluene	0.250	U	0.080	0.250
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
88-72-2	o-Nitrotoluene	0.250	U	0.082	0.250
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
98-95-3	Nitrobenzene	0.250	U	0.080	0.250
<i>98-95-3</i>	<i>Nitrobenzene</i>				
99-08-1	m-Nitrotoluene	0.250	U	0.080	0.250
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	0.250	U	0.080	0.250
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
99-65-0	m-Dinitrobenzene	0.250	U	0.080	0.250
<i>99-65-0</i>	<i>m-Dinitrobenzene</i>				
479-45-8	Tetryl	0.500	U	0.080	0.500
<i>479-45-8</i>	<i>Tetryl</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: MB for batch 1185692

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592978

Sample Amount 1000 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
78-11-5	PETN	0.500	U	0.100	0.500
<i>78-11-5</i>	<i>PETN</i>				
99-99-0	p-Nitrotoluene	0.500	U	0.150	0.500
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: MB for batch 1185692

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592978

Sample Amount 1000 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160014.wiff

Date Analyzed: 16-FEB-12 17:46

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
3058-38-6	TATB	1.00	U	0.300	1.00
3058-38-6	TATB				
618-87-1	3,5-Dinitroaniline	1.00	U	0.300	1.00
618-87-1	3,5-Dinitroaniline				
78-30-8	tris(o-cresyl) phosphate	1.00	U	0.300	1.00
78-30-8	tris(o-cresyl) phosphate				
59229-75-3	2,6-Diamino-4-nitrotoluene	2.50	U	0.500	2.50
59229-75-3	2,6-Diamino-4-nitrotoluene				
6629-29-4	2,4-Diamino-6-nitrotoluene	2.50	U	0.500	2.50
6629-29-4	2,4-Diamino-6-nitrotoluene				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: LCS for batch 1185692

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592979

Sample Amount 1000 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227014a

Date Analyzed: 28-FEB-12 01:21

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
479-45-8	Tetryl	4		0.080	0.500
<i>479-45-8</i>	<i>Tetryl</i>				
99-08-1	m-Nitrotoluene	4.02		0.080	0.250
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
88-72-2	o-Nitrotoluene	4.07		0.082	0.250
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
2691-41-0	HMX	4.22		0.080	0.250
<i>2691-41-0</i>	<i>HMX</i>				
98-95-3	Nitrobenzene	4.26		0.080	0.250
<i>98-95-3</i>	<i>Nitrobenzene</i>				
99-99-0	p-Nitrotoluene	4.36		0.150	0.500
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	4.49		0.080	0.250
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	4.49		0.080	0.250
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
78-11-5	PETN	4.52		0.100	0.500
<i>78-11-5</i>	<i>PETN</i>				
99-65-0	m-Dinitrobenzene	4.52		0.080	0.250
<i>99-65-0</i>	<i>m-Dinitrobenzene</i>				
121-82-4	RDX	4.55		0.080	0.250
<i>121-82-4</i>	<i>RDX</i>				
606-20-2	2,6-Dinitrotoluene	4.56		0.080	0.250
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	4.57		0.080	0.250
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: LCS for batch 1185692

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592979

Sample Amount 1000 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
118-96-7	2,4,6-Trinitrotoluene	4.73		0.080	0.250
<i>118-96-7</i>	<i>2,4,6-Trinitrotoluene</i>				
121-14-2	2,4-Dinitrotoluene	4.83		0.080	0.250
<i>121-14-2</i>	<i>2,4-Dinitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: LCS for batch 1185692

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592979

Sample Amount 1000 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160015.wiff

Date Analyzed: 16-FEB-12 18:02

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
78-30-8	tris(o-cresyl) phosphate	3.49		0.300	1.00
78-30-8	tris(o-cresyl) phosphate				
3058-38-6	TATB	3.73		0.300	1.00
3058-38-6	TATB				
6629-29-4	2,4-Diamino-6-nitrotoluene	4.76		0.500	2.50
6629-29-4	2,4-Diamino-6-nitrotoluene				
618-87-1	3,5-Dinitroaniline	5.47		0.300	1.00
618-87-1	3,5-Dinitroaniline				
59229-75-3	2,6-Diamino-4-nitrotoluene	5.51		0.500	2.50
59229-75-3	2,6-Diamino-4-nitrotoluene				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031(295270003MS)MS

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592980

Sample Amount 940 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227016a

Date Analyzed: 28-FEB-12 02:21

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
479-45-8	Tetryl	4.1		0.0851	0.532
<i>479-45-8</i>	<i>Tetryl</i>				
88-72-2	o-Nitrotoluene	4.13		0.0872	0.266
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
99-08-1	m-Nitrotoluene	4.26		0.0851	0.266
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
99-99-0	p-Nitrotoluene	4.32		0.160	0.532
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				
2691-41-0	HMX	4.36		0.0851	0.266
<i>2691-41-0</i>	<i>HMX</i>				
98-95-3	Nitrobenzene	4.54		0.0851	0.266
<i>98-95-3</i>	<i>Nitrobenzene</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	4.58		0.0851	0.266
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	4.59		0.0851	0.266
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
606-20-2	2,6-Dinitrotoluene	4.63		0.0851	0.266
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	4.8		0.0851	0.266
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				
78-11-5	PETN	4.8		0.106	0.532
<i>78-11-5</i>	<i>PETN</i>				
121-82-4	RDX	4.9		0.0851	0.266
<i>121-82-4</i>	<i>RDX</i>				
118-96-7	2,4,6-Trinitrotoluene	5.06		0.0851	0.266
<i>118-96-7</i>	<i>2,4,6-Trinitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031(295270003MS)MS

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592980

Sample Amount 940 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
99-65-0	m-Dinitrobenzene	5.06		0.0851	0.266
99-65-0	m-Dinitrobenzene				
121-14-2	2,4-Dinitrotoluene	5.15		0.0851	0.266
121-14-2	2,4-Dinitrotoluene				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031(295270003MS)MS

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592980

Sample Amount 940 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160017.wiff

Date Analyzed: 16-FEB-12 18:36

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
3058-38-6	TATB	3.7		0.319	1.06
3058-38-6	TATB				
78-30-8	tris(o-cresyl) phosphate	3.94		0.319	1.06
78-30-8	tris(o-cresyl) phosphate				
6629-29-4	2,4-Diamino-6-nitrotoluene	5.04		0.532	2.66
6629-29-4	2,4-Diamino-6-nitrotoluene				
59229-75-3	2,6-Diamino-4-nitrotoluene	5.61		0.532	2.66
59229-75-3	2,6-Diamino-4-nitrotoluene				
618-87-1	3,5-Dinitroaniline	5.69		0.319	1.06
618-87-1	3,5-Dinitroaniline				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031(295270003MSD)MSD

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592981

Sample Amount 930 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXP0227017a

Date Analyzed: 28-FEB-12 02:50

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
479-45-8	Tetryl	4.16		0.086	0.538
<i>479-45-8</i>	<i>Tetryl</i>				
2691-41-0	HMX	4.38		0.086	0.269
<i>2691-41-0</i>	<i>HMX</i>				
98-95-3	Nitrobenzene	4.53		0.086	0.269
<i>98-95-3</i>	<i>Nitrobenzene</i>				
99-08-1	m-Nitrotoluene	4.53		0.086	0.269
<i>99-08-1</i>	<i>m-Nitrotoluene</i>				
88-72-2	o-Nitrotoluene	4.56		0.0882	0.269
<i>88-72-2</i>	<i>o-Nitrotoluene</i>				
99-99-0	p-Nitrotoluene	4.58		0.161	0.538
<i>99-99-0</i>	<i>p-Nitrotoluene</i>				
99-35-4	1,3,5-Trinitrobenzene	4.69		0.086	0.269
<i>99-35-4</i>	<i>1,3,5-Trinitrobenzene</i>				
35572-78-2	2-Amino-4,6-dinitrotoluene	4.96		0.086	0.269
<i>35572-78-2</i>	<i>2-Amino-4,6-dinitrotoluene</i>				
121-82-4	RDX	5.05		0.086	0.269
<i>121-82-4</i>	<i>RDX</i>				
19406-51-0	4-Amino-2,6-dinitrotoluene	5.06		0.086	0.269
<i>19406-51-0</i>	<i>4-Amino-2,6-dinitrotoluene</i>				
606-20-2	2,6-Dinitrotoluene	5.08		0.086	0.269
<i>606-20-2</i>	<i>2,6-Dinitrotoluene</i>				
78-11-5	PETN	5.14		0.108	0.538
<i>78-11-5</i>	<i>PETN</i>				
121-14-2	2,4-Dinitrotoluene	5.2		0.086	0.269
<i>121-14-2</i>	<i>2,4-Dinitrotoluene</i>				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031(295270003MSD)MSD

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592981

Sample Amount 930 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

Cas No.	Compound	Concentration*	Q	MDL	PQL
99-65-0	m-Dinitrobenzene	5.2		0.086	0.269
99-65-0	m-Dinitrobenzene				
118-96-7	2,4,6-Trinitrotoluene	5.53		0.086	0.269
118-96-7	2,4,6-Trinitrotoluene				

1
High Explosives Analysis Data Sheet

Lab Name: GEL Laboratories LLC

Client Sample ID: CAAN-12-2031(295270003MSD)MSD

Lab Code: GEL

GEL Job No (SDG) 12-700

Matrix: WATER

GEL Sample ID: 1202592981

Sample Amount 930 mL

Date Received: 03-FEB-12

Moisture: .

Extraction Batch ID: 1185692

Extraction Type Sol Exchange

Date Extracted: 07-FEB-12

Concentrated Extract Volume (mL) 5

Injection Volume (uL): 50

GEL data file: EXS02160018.wiff

Date Analyzed: 16-FEB-12 18:52

Dilution Factor: 2

Concentration Units: ug/L

Cas No.	Compound	Concentration*	Q	MDL	PQL
3058-38-6	TATB	3.76		0.323	1.08
<i>3058-38-6</i>	<i>TATB</i>				
78-30-8	tris(o-cresyl) phosphate	3.85		0.323	1.08
<i>78-30-8</i>	<i>tris(o-cresyl) phosphate</i>				
59229-75-3	2,6-Diamino-4-nitrotoluene	5.3		0.538	2.69
<i>59229-75-3</i>	<i>2,6-Diamino-4-nitrotoluene</i>				
6629-29-4	2,4-Diamino-6-nitrotoluene	5.35		0.538	2.69
<i>6629-29-4</i>	<i>2,4-Diamino-6-nitrotoluene</i>				
618-87-1	3,5-Dinitroaniline	5.62		0.323	1.08
<i>618-87-1</i>	<i>3,5-Dinitroaniline</i>				

Explosives Initial Calibration Blank

Lab Name: GEL Laboratories LLCGEL Job No(SDG): 12-700Lab Code: GELLab Sample ID: XIBLK01Analysis Date: 27-FEB-12 18:57GEL Data File: EXP0227001aInstrument ID: LCMSMSColumn: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
1,3,5-Trinitrobenzene	0	0
1,3-Dinitrobenzene-d4	500	610.636
2,4,6-Trinitrotoluene	0	0
2,4-Dinitrotoluene	0	0
2,6-Dinitrotoluene	0	0
2,6-Dinitrotoluene-d3	500	513.817
2-Amino-4,6-dinitrotoluene	0	0
4-Amino-2,6-dinitrotoluene	0	0
HMX	0	0
Nitrobenzene	0	0
PETN	0	0
RDX	0	0
Tetryl	0	0
m-Dinitrobenzene	0	0
m-Nitrotoluene	0	0
o-Nitrotoluene	0	0
p-Nitrotoluene	0	0

Explosives Initial Calibration Blank

Lab Name: GEL Laboratories LLCGEL Job No(SDG): 12-700Lab Code: GELLab Sample ID: XIBLK01Analysis Date: 27-FEB-12 19:26GEL Data File: EXP0227002aInstrument ID: LCMSMSColumn: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
1,3,5-Trinitrobenzene	0	0
1,3-Dinitrobenzene-d4	500	584.986
2,4,6-Trinitrotoluene	0	0
2,4-Dinitrotoluene	0	0
2,6-Dinitrotoluene	0	0
2,6-Dinitrotoluene-d3	500	603.501
2-Amino-4,6-dinitrotoluene	0	0
4-Amino-2,6-dinitrotoluene	0	0
HMX	0	0
Nitrobenzene	0	0
PETN	0	0
RDX	0	0
Tetryl	0	0
m-Dinitrobenzene	0	0
m-Nitrotoluene	0	0
o-Nitrotoluene	0	0
p-Nitrotoluene	0	0

Explosives Initial Calibration Blank

Lab Name: GEL Laboratories LLCGEL Job No(SDG): 12-700Lab Code: GELLab Sample ID: XIBLK01Analysis Date: 16-FEB-12 14:08GEL Data File: EXS02160001.wiffInstrument ID: LCMSMSColumn: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
tris(o-cresyl) phosphate	0	0
TATB	0	0
3,5-Dinitroaniline	0	0
2,4-Diamino-6-nitrotoluene	0	0
2,6-Diamino-4-nitrotoluene	0	0

Explosives Initial Calibration Blank

Lab Name: GEL Laboratories LLCGEL Job No(SDG): 12-700Lab Code: GELLab Sample ID: XIBLK01Analysis Date: 16-FEB-12 14:25GEL Data File: EXS02160002.wiffInstrument ID: LCMSMSColumn: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
tris(o-cresyl) phosphate	0	0
TATB	0	0
3,5-Dinitroaniline	0	0
2,4-Diamino-6-nitrotoluene	0	0
2,6-Diamino-4-nitrotoluene	0	0

4A
Explosives Continuing Calibration Blank

Lab Name: GEL Laboratories LLC

GEL Job No(SDG): 12-700

Lab Code: GEL

Lab Sample ID: XIBLK02

Analysis Date: 27-FEB-12 22:53

GEL Data File: EXP0227009a

Instrument ID: LCMSMS

Column: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
1,3,5-Trinitrobenzene	0	0
1,3-Dinitrobenzene-d4	500	464.531
2,4,6-Trinitrotoluene	0	0
2,4-Dinitrotoluene	0	0
2,6-Dinitrotoluene	0	0
2,6-Dinitrotoluene-d3	500	462.395
2-Amino-4,6-dinitrotoluene	0	0
4-Amino-2,6-dinitrotoluene	0	0
HMX	0	0
Nitrobenzene	0	0
PETN	0	0
RDX	0	0
Tetryl	0	0
m-Dinitrobenzene	0	0
m-Nitrotoluene	0	0
o-Nitrotoluene	0	0
p-Nitrotoluene	0	0

4A
Explosives Continuing Calibration Blank

Lab Name: GEL Laboratories LLC

GEL Job No(SDG): 12-700

Lab Code: GEL

Lab Sample ID: XIBLK03

Analysis Date: 27-FEB-12 23:53

GEL Data File: EXP0227011a

Instrument ID: LCMSMS

Column: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
1,3,5-Trinitrobenzene	0	0
1,3-Dinitrobenzene-d4	500	461.381
2,4,6-Trinitrotoluene	0	0
2,4-Dinitrotoluene	0	0
2,6-Dinitrotoluene	0	0
2,6-Dinitrotoluene-d3	500	494.767
2-Amino-4,6-dinitrotoluene	0	0
4-Amino-2,6-dinitrotoluene	0	0
HMX	0	0
Nitrobenzene	0	0
PETN	0	0
RDX	0	0
Tetryl	0	0
m-Dinitrobenzene	0	0
m-Nitrotoluene	0	0
o-Nitrotoluene	0	0
p-Nitrotoluene	0	0

4A
Explosives Continuing Calibration Blank

Lab Name: GEL Laboratories LLC

GEL Job No(SDG): 12-700

Lab Code: GEL

Lab Sample ID: XIBLK04

Analysis Date: 28-FEB-12 06:17

GEL Data File: EXP0227024a

Instrument ID: LCMSMS

Column: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
1,3,5-Trinitrobenzene	0	0
1,3-Dinitrobenzene-d4	500	433.33
2,4,6-Trinitrotoluene	0	0
2,4-Dinitrotoluene	0	0
2,6-Dinitrotoluene	0	0
2,6-Dinitrotoluene-d3	500	441.639
2-Amino-4,6-dinitrotoluene	0	0
4-Amino-2,6-dinitrotoluene	0	0
HMX	0	0
Nitrobenzene	0	0
PETN	0	0
RDX	0	0
Tetryl	0	0
m-Dinitrobenzene	0	0
m-Nitrotoluene	0	0
o-Nitrotoluene	0	0
p-Nitrotoluene	0	0

4A
Explosives Continuing Calibration Blank

Lab Name: GEL Laboratories LLC

GEL Job No(SDG): 12-700

Lab Code: GEL

Lab Sample ID: XIBLK02

Analysis Date: 16-FEB-12 16:39

GEL Data File: EXS02160010.wiff

Instrument ID: LCMSMS

Column: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
tris(o-cresyl) phosphate	0	6.76
TATB	0	0
3,5-Dinitroaniline	0	0
2,4-Diamino-6-nitrotoluene	0	0
2,6-Diamino-4-nitrotoluene	0	0

4A
Explosives Continuing Calibration Blank

Lab Name: GEL Laboratories LLC

GEL Job No(SDG): 12-700

Lab Code: GEL

Lab Sample ID: XIBLK03

Analysis Date: 16-FEB-12 17:12

GEL Data File: EXS02160012.wiff

Instrument ID: LCMSMS

Column: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
2,6-Diamino-4-nitrotoluene	0	0
3,4-Dinitrotoluene	0	0
tris(o-cresyl) phosphate	0	3.6
TATB	0	0
3,5-Dinitroaniline	0	0
2,4-Diamino-6-nitrotoluene	0	0

4A
Explosives Continuing Calibration Blank

Lab Name: GEL Laboratories LLC

GEL Job No(SDG): 12-700

Lab Code: GEL

Lab Sample ID: XIBLK04

Analysis Date: 16-FEB-12 20:49

GEL Data File: EXS02160025.wiff

Instrument ID: LCMSMS

Column: Phenomenex Ultracarb 5u ODS(20)

Compound	True	Found (ug/L)
3,4-Dinitrotoluene	0	0
tris(o-cresyl) phosphate	0	2.72
TATB	0	0
3,5-Dinitroaniline	0	0
2,4-Diamino-6-nitrotoluene	0	0
2,6-Diamino-4-nitrotoluene	0	0

General Chem Analysis

Case Narrative

**General Chemistry Narrative
ARS International (ARSL)
SDG 12-700**

Method/Analysis Information

Product: Carbon, Total Organic

Analytical Batch: 1186182

Method: SW 9060 Total Organic Carbon

Sample Analysis

The following samples were analyzed using the analytical protocol as established in SW846 9060:

Sample ID	Client ID
295270003	CAAN-12-2031
295270005	CAAN-12-2199
1202594161	Method Blank (MB)
1202594162	295075001(BuckmanPZ-12-2179) Sample Duplicate (DUP)
1202594163	295392002(CAWA-12-2018) Sample Duplicate (DUP)
1202594164	295075001(BuckmanPZ-12-2179) Post Spike (PS)
1202594165	295392002(CAWA-12-2018) Post Spike (PS)
1202594166	Laboratory Control Sample (LCS)

The samples in this SDG were analyzed on an "as received" basis.

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-GC-E-093 REV# 9.

Preparation/Analytical Method Verification

The SOP stated above has been prepared based on technical research and testing conducted by GEL Laboratories, LLC. and with guidance from the regulatory documents listed in this "Method/Analysis Information" section.

Calibration Information

The Carbon analysis was performed on a O-I Analytical Model 1010 Total Organic Carbon Analyzer.

Initial Calibration

All initial calibration requirements have been met for this SDG.

Continuing Calibration Blanks

All continuing calibration blanks (CCBs) associated with reported data from this batch were within acceptance limits.

Calibration Verification Information (CCV)

All continuing calibration verification standards (CCVs) associated with reported data from this batch were within acceptance limits.

Quality Control (QC) Information**Method Blank (MB) Statement**

The MB analyzed with this SDG met the acceptance criteria.

Laboratory Control Sample (LCS) Recovery

The LCS spike recovery met the acceptance limits.

Quality Control (QC) Designation

The following samples were selected for QC analysis: 295075001 (BuckmanPZ-12-2179) and 295392002 (CAWA-12-2018).

Matrix Spike (MS)/Post Spike (PS) Recovery Statement

The MS/PS recoveries for this sample set were within the required acceptance limits.

Duplicate Relative Percent Difference (RPD) Statement

The RPD between the sample and its duplicate met the acceptance limits.

Technical Information

GEL assigns holding times based on the date and time of sample collection. Those holding times expressed in hours are calculated in the AlphaLims system by hours. Those holding times expressed as days expire at midnight on the day of expiration.

Holding Times

All samples in this SDG met the specified holding time.

Sample Dilutions

The samples in this SDG did not require dilutions.

Sample Re-analysis

The samples in this SDG did not require re-analysis.

Miscellaneous Information**Data Exception (DER) Documentation**

A DER was not required for this SDG.

Additional Comments

Additional comments were not required for this SDG.

Electronic Packaging Comment

This data package was generated using an electronic data processing program referred to as virtual packaging. In an

effort to increase quality and efficiency, the laboratory has developed systems to generate all data packages electronically. The following change from traditional packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. The data validator will always sign and date the case narrative. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

Method/Analysis Information

Product: Ammonia Nitrogen

Analytical Batch: 1186023 **Method:** EPA 350.1 Nitrogen and Ammonia L

Prep Batch : 1186022 **Method:** EEPA 350.2 Prep

Sample Analysis

The following samples were analyzed using the analytical protocol as established in EPA 350.1:

Sample ID	Client ID
295270001	CAAN-12-2030
295270006	CAAN-12-2200
1202593853	Method Blank (MB)
1202593854	295270001(CAAN-12-2030) Sample Duplicate (DUP)
1202593855	295270001(CAAN-12-2030) Matrix Spike (MS)
1202593856	295270001(CAAN-12-2030) Matrix Spike Duplicate (MSD)
1202593857	Laboratory Control Sample (LCS)

The samples in this SDG were analyzed on an "as received" basis.

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-GC-E-106 REV# 7.

Preparation/Analytical Method Verification

The SOP stated above has been prepared based on technical research and testing conducted by GEL Laboratories, LLC. and with guidance from the regulatory documents listed in this "Method/Analysis Information" section.

Calibration Information

The Nutrient analysis was performed on a Lachat QuickChem FIA+ 8000 Series.

Continuing Calibration Blanks

All continuing calibration blanks (CCBs) associated with reported data from this batch were within acceptance limits.

Calibration Verification Information (CCV)

All continuing calibration verification standards (CCVs) associated with reported data from this batch were within acceptance limits.

Y Intercept Rule

The absolute value of the intercept is less than 3 times the MDL.

Quality Control (QC) Information

Method Blank (MB) Statement

The MB analyzed with this SDG met the acceptance criteria.

Laboratory Control Sample (LCS) Recovery

The LCS spike recovery met the acceptance limits.

Quality Control (QC) Designation

The following sample was selected for QC analysis: 295270001 (CAAN-12-2030).

Matrix Spike (MS)/Post Spike (PS) Recovery Statement

The MS/PS recovery for this sample set was within the required acceptance limits.

Matrix Spike Duplicate (MSD) Recovery Statement

The MSD recovery for this sample set was within the required acceptance limits.

MS/MSD Relative Percent Difference (RPD) Statement

The RPD between the spike and spike duplicate met the acceptance limits.

Duplicate Relative Percent Difference (RPD) Statement

The values for the sample and duplicate are less than the Practical Quantitation Limit (PQL); therefore, the RPD is not applicable. 1202593854 (CAAN-12-2030).

Technical Information

GEL assigns holding times based on the date and time of sample collection. Those holding times expressed in hours are calculated in the AlphaLims system by hours. Those holding times expressed as days expire at midnight on the day of expiration.

Holding Times

All samples in this SDG met the specified holding time.

Sample Preservation/Integrity

All the samples from this sample group met the preservation and integrity requirements of the method.

Sample Dilutions

The samples in this SDG did not require dilutions.

Sample Re-analysis

The following sample was re-analyzed due to instrument failure: 295270006 (CAAN-12-2200).

Miscellaneous Information

Data Exception (DER) Documentation

A DER was not required for this SDG.

Additional Comments

Additional comments were not required for this SDG.

Electronic Packaging Comment

This data package was generated using an electronic data processing program referred to as virtual packaging. In an effort to increase quality and efficiency, the laboratory has developed systems to generate all data packages electronically. The following change from traditional packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. The data validator will always sign and date the case narrative. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

Method/Analysis Information

Product:	Total Kjeldahl Nitrogen		
Analytical Batch:	1186025	Method:	Nitrogen and Total Kjeldahl (TKN)
Prep Batch :	1186024	Method:	EEPA 351.2 Prep

Sample Analysis

The following samples were analyzed using the analytical protocol as established in EPA 351.2:

Sample ID	Client ID
295270003	CAAN-12-2031
295270005	CAAN-12-2199
1202593858	Method Blank (MB)
1202593859	295270003(CAAN-12-2031) Sample Duplicate (DUP)
1202593860	295270003(CAAN-12-2031) Matrix Spike (MS)
1202593861	295270003(CAAN-12-2031) Matrix Spike Duplicate (MSD)
1202593862	Laboratory Control Sample (LCS)

The samples in this SDG were analyzed on an "as received" basis.

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-GC-E-104 REV# 11.

Preparation/Analytical Method Verification

The SOP stated above has been prepared based on technical research and testing conducted by GEL Laboratories, LLC. and with guidance from the regulatory documents listed in this "Method/Analysis Information" section.

Calibration Information

The Nutrient analysis was performed on a Lachat QuickChem FIA+ 8000 Series.

Continuing Calibration Blanks

All continuing calibration blanks (CCBs) associated with reported data from this batch were within acceptance limits.

Calibration Verification Information (CCV)

All continuing calibration verification standards (CCVs) associated with reported data from this batch were within acceptance limits.

Y Intercept Rule

The absolute value of the intercept is less than 3 times the MDL.

Quality Control (QC) Information

Method Blank (MB) Statement

The MB analyzed with this SDG met the acceptance criteria.

Laboratory Control Sample (LCS) Recovery

The LCS spike recovery met the acceptance limits.

Quality Control (QC) Designation

The following sample was selected for QC analysis: 295270003 (CAAN-12-2031).

Matrix Spike (MS)/Post Spike (PS) Recovery Statement

The spike recovery falls outside of the GEL acceptance limits but within the client specified limits. 1202593860 (CAAN-12-2031).

Matrix Spike Duplicate (MSD) Recovery Statement

The spike duplicate recovery falls outside of the GEL acceptance limits but within the client specified limits. 1202593861 (CAAN-12-2031).

MS/MSD Relative Percent Difference (RPD) Statement

The RPD between the spike and spike duplicate met the acceptance limits.

Duplicate Relative Percent Difference (RPD) Statement

The RPD between the sample and its duplicate met the acceptance limits.

Technical Information

GEL assigns holding times based on the date and time of sample collection. Those holding times expressed in hours are calculated in the AlphaLims system by hours. Those holding times expressed as days expire at midnight on the day of expiration.

Holding Times

All samples in this SDG met the specified holding time.

Sample Preservation/Integrity

All the samples from this sample group met the preservation and integrity requirements of the method.

Sample Dilutions

The following sample in this sample group was diluted due to matrix interference: 295270005 (CAAN-12-2199).

Sample Re-analysis

The samples in this SDG did not require re-analysis.

Miscellaneous Information

Data Exception (DER) Documentation

The following DER was generated for this SDG: 1051875 1202593860 (CAAN-12-2031) and 1202593861 (CAAN-12-2031).

Additional Comments

Additional comments were not required for this SDG.

Electronic Packaging Comment

This data package was generated using an electronic data processing program referred to as virtual packaging. In an effort to increase quality and efficiency, the laboratory has developed systems to generate all data packages electronically. The following change from traditional packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. The data validator will always sign and date the case narrative. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

Method/Analysis Information

Product:	Nitrate Nitrite by Cadmium Reduction		
Analytical Batch:	1186031	Method:	EPA 353.2 Nitrogen and Nitrate/Nitrite

Sample Analysis

The following samples were analyzed using the analytical protocol as established in EPA 353.2:

Sample ID	Client ID
295270001	CAAN-12-2030
295270006	CAAN-12-2200
1202593872	Method Blank (MB)
1202593873	295075001(BuckmanPZ-12-2179) Sample Duplicate (DUP)
1202593875	295075001(BuckmanPZ-12-2179) Post Spike (PS)
1202593877	Laboratory Control Sample (LCS)

The samples in this SDG were analyzed on an "as received" basis.

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-GC-E-128 REV# 7.

Preparation/Analytical Method Verification

The SOP stated above has been prepared based on technical research and testing conducted by GEL Laboratories, LLC. and with guidance from the regulatory documents listed in this "Method/Analysis Information" section.

Calibration Information

The Nutrient analysis was performed on a Lachat QuickChem FIA+ 8500 Series.

Continuing Calibration Blanks

All continuing calibration blanks (CCBs) associated with reported data from this batch were within acceptance limits.

Calibration Verification Information (CCV)

All continuing calibration verification standards (CCVs) associated with reported data from this batch were within acceptance limits.

Y Intercept Rule

The absolute value of the intercept is less than 3 times the MDL.

Quality Control (QC) Information

Method Blank (MB) Statement

The MB analyzed with this SDG met the acceptance criteria.

Laboratory Control Sample (LCS) Recovery

The LCS spike recovery met the acceptance limits.

Quality Control (QC) Designation

The following sample was selected for QC analysis: 295075001 (BuckmanPZ-12-2179).

Matrix Spike (MS)/Post Spike (PS) Recovery Statement

The MS/PS recovery for this sample set was within the required acceptance limits.

Duplicate Relative Percent Difference (RPD) Statement

The RPD between the sample and its duplicate met the acceptance limits.

Technical Information

GEL assigns holding times based on the date and time of sample collection. Those holding times expressed in hours are calculated in the AlphaLims system by hours. Those holding times expressed as days expire at midnight on the day of expiration.

Holding Times

All samples in this SDG met the specified holding time.

Sample Preservation/Integrity

All the samples from this sample group met the preservation and integrity requirements of the method.

Sample Dilutions

The following samples in this sample group were diluted due to high concentration: 1202593873 (BuckmanPZ-12-2179) and 1202593875 (BuckmanPZ-12-2179). The following samples in this sample group were diluted due to matrix interference: 295270001 (CAAN-12-2030) and 295270006 (CAAN-12-2200).

Sample Re-analysis

The samples in this SDG did not require re-analysis.

Miscellaneous Information

Data Exception (DER) Documentation

A DER was not required for this SDG.

Additional Comments

Additional comments were not required for this SDG.

Electronic Packaging Comment

This data package was generated using an electronic data processing program referred to as virtual packaging. In an effort to increase quality and efficiency, the laboratory has developed systems to generate all data packages electronically. The following change from traditional packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. The data validator will always sign and date the case narrative. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

Method/Analysis Information

Product:	Total Phosphorus		
Analytical Batch:	1186020	Method:	EPA 365.4 Phosphorus and Total in
Prep Batch :	1186019	Method:	EEPA 365.4 Prep

Sample Analysis

The following samples were analyzed using the analytical protocol as established in EPA 365.4:

Sample ID	Client ID
295270001	CAAN-12-2030
295270006	CAAN-12-2200
1202593841	Method Blank (MB)
1202593842	295075001(BuckmanPZ-12-2179) Sample Duplicate (DUP)
1202593844	295075001(BuckmanPZ-12-2179) Matrix Spike (MS)
1202593846	295075001(BuckmanPZ-12-2179) Matrix Spike Duplicate (MSD)
1202593848	Laboratory Control Sample (LCS)

The samples in this SDG were analyzed on an "as received" basis.

SOP Reference

Procedure for preparation, analysis and reporting of analytical data are controlled by GEL Laboratories LLC as Standard Operating Procedure (SOP). The data discussed in this narrative has been analyzed in accordance with GL-GC-E-103 REV# 8.

Preparation/Analytical Method Verification

The SOP stated above has been prepared based on technical research and testing conducted by GEL Laboratories, LLC. and with guidance from the regulatory documents listed in this "Method/Analysis Information" section.

Calibration Information

The Nutrient analysis was performed on a Lachat QuickChem FIA+ 8000 Series.

Continuing Calibration Blanks

All continuing calibration blanks (CCBs) associated with reported data from this batch were within acceptance limits.

Calibration Verification Information (CCV)

All continuing calibration verification standards (CCVs) associated with reported data from this batch were within acceptance limits.

Y Intercept Rule

The absolute value of the intercept is less than 3 times the MDL.

Quality Control (QC) Information

Method Blank (MB) Statement

The MB analyzed with this SDG met the acceptance criteria.

Laboratory Control Sample (LCS) Recovery

The LCS spike recovery met the acceptance limits.

Quality Control (QC) Designation

The following sample was selected for QC analysis: 295075001 (BuckmanPZ-12-2179).

Matrix Spike (MS)/Post Spike (PS) Recovery Statement

The MS/PS recovery for this sample set was within the required acceptance limits.

Matrix Spike Duplicate (MSD) Recovery Statement

The MSD recovery for this sample set was within the required acceptance limits.

MS/MSD Relative Percent Difference (RPD) Statement

The RPD between the spike and spike duplicate met the acceptance limits.

Duplicate Relative Percent Difference (RPD) Statement

The values for the sample and duplicate are less than the Practical Quantitation Limit (PQL); therefore, the RPD is not applicable. 1202593842 (BuckmanPZ-12-2179).

Technical Information

GEL assigns holding times based on the date and time of sample collection. Those holding times expressed in hours are calculated in the AlphaLims system by hours. Those holding times expressed as days expire at midnight on the day of expiration.

Holding Times

All samples in this SDG met the specified holding time.

Sample Preservation/Integrity

All the samples from this sample group met the preservation and integrity requirements of the method.

Sample Dilutions

The samples in this SDG did not require dilutions.

Sample Re-analysis

The following samples were re-analyzed due to instrument failure: 1202593841 (MB), 295270001 (CAAN-12-2030) and 295270006 (CAAN-12-2200). The following samples were re-analyzed due to their proximity to an overrange sample: 295270001 (CAAN-12-2030) and 295270006 (CAAN-12-2200).

Miscellaneous Information

Data Exception (DER) Documentation

A DER was not required for this SDG.

Additional Comments

Additional comments were not required for this SDG.

Electronic Packaging Comment

This data package was generated using an electronic data processing program referred to as virtual packaging. In an effort to increase quality and efficiency, the laboratory has developed systems to generate all data packages electronically. The following change from traditional packages should be noted:

Analyst/peer reviewer initials and dates are not present on the electronic data files. Presently, all initials and dates are present on the original raw data. These hard copies are temporarily stored in the laboratory. The data validator will always sign and date the case narrative. Data that are not generated electronically, such as hand written pages, will be scanned and inserted into the electronic package.

Certification Statement

Where the analytical method has been performed under NELAP certification, the analysis has met all of the requirements of the NELAC standard unless otherwise noted in the analytical case narrative.

Review Validation:

GEL requires all analytical data to be verified by a qualified data validator. In addition, all data designated for CLP or CLP-like packaging will receive a third level validation upon completion of the data package.

The following data validator verified the information presented in this case narrative:

Reviewer:  Date: 29Feb12

Sample Data Summary

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Certificate of Analysis Report for

ARSL001 ARS International (63641-10)

Client SDG: 12-700 GEL Work Order: 295270

The Qualifiers in this report are defined as follows:

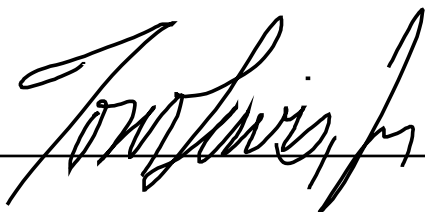
- * A quality control analyte recovery is outside of specified acceptance criteria
- ** Analyte is a surrogate compound
- E Organics--Concentration of the target analyte exceeds the instrument calibration range
- J Value is estimated
- U Analyte was analyzed for, but not detected above the MDL, MDA, or LOD.

Where the analytical method has been performed under NELAP certification, the analysis has met all of the requirements of the NELAC standard unless qualified on the Certificate of Analysis.

The designation ND, if present, appears in the result column when the analyte concentration is not detected above the detection limit.

This data report has been prepared and reviewed in accordance with GEL Laboratories LLC standard operating procedures. Please direct any questions to your Project Manager, Valerie Davis.

Reviewed by

A handwritten signature in black ink, appearing to read "Tom Davis", is written over a horizontal line.

GEL LABORATORIES LLC

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Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2030
Sample ID: 295270001
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method
Nutrient Analysis											
EPA 350.1 Nitrogen, Ammonia L "As Received"											
Nitrogen, Ammonia	U	ND	0.016	0.050	mg/L	1	KLP1	02/08/12	1542	1186023	1
EPA 353.2 Nitrogen, Nitrate/Nitrite "As Received"											
Nitrogen, Nitrate/Nitrite		0.304	0.050	0.250	mg/L	5	KLP1	02/14/12	1237	1186031	2
EPA 365.4 Phosphorus, Total in "As Received"											
Phosphorus, Total as P		0.0562	0.015	0.050	mg/L	1	KLP1	02/08/12	1417	1186020	3

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 350.2 Prep	EPA 350.1 Ammonia Nitrogen Prep	AXS5	02/07/12	1658	1186022
EPA 365.4 Prep	EPA 365.4 Phosphorus, Total in liquid PR	AXS5	02/08/12	1159	1186019

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	EPA 350.1	
2	EPA 353.2	
3	EPA 365.4	

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Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2031
Sample ID: 295270003
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method
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Carbon Analysis

SW 9060 Total Organic Carbon "As Received"

Total Organic Carbon Average	J	0.349	0.330	1.00	mg/L	1	TSM	02/07/12	1841	1186182	1
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Nutrient Analysis

Nitrogen, Total Kjeldahl (TKN) "As Received"

Nitrogen, Total Kjeldahl	U	ND	0.035	0.100	mg/L	1	KLP1	02/17/12	1038	1186025	2
--------------------------	---	----	-------	-------	------	---	------	----------	------	---------	---

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 351.2 Prep	EPA 351.2 Total Kjeldahl Nitrogen Prep	AXS5	02/14/12	1649	1186024

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	SW846 9060	
2	EPA 351.2	

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Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2199
Sample ID: 295270005
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method
Carbon Analysis											
SW 9060 Total Organic Carbon "As Received"											
Total Organic Carbon Average	J	0.359	0.330	1.00	mg/L	1	TSM	02/07/12	1915	1186182	1
Nutrient Analysis											
Nitrogen, Total Kjeldahl (TKN) "As Received"											
Nitrogen, Total Kjeldahl	U	ND	0.175	0.500	mg/L	5	KLP1	02/17/12	1049	1186025	2

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 351.2 Prep	EPA 351.2 Total Kjeldahl Nitrogen Prep	AXS5	02/14/12	1649	1186024

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	SW846 9060	
2	EPA 351.2	

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Certificate of Analysis

Report Date: February 27, 2012

Company : Los Alamos National Laboratory
Address : TA-03, SM271, Drop Pt. 02U, Rm111

Los Alamos, New Mexico 87545
Contact: Ms. Joylene Valdez
Project: LANL-WQH Water Samples

Client SDG: 12-700

Client Sample ID: CAAN-12-2200
Sample ID: 295270006
Matrix: WG
Collect Date: 01-FEB-12 12:00
Receive Date: 03-FEB-12
Collector: Client

Project: ESHL00210
Client ID: ARSL001

Parameter	Qualifier	Result	DL	RL	Units	DF	Analyst	Date	Time	Batch	Method
Nutrient Analysis											
EPA 350.1 Nitrogen, Ammonia L "As Received"											
Nitrogen, Ammonia	U	ND	0.016	0.050	mg/L	1	KLP1	02/08/12	1554	1186023	1
EPA 353.2 Nitrogen, Nitrate/Nitrite "As Received"											
Nitrogen, Nitrate/Nitrite		0.304	0.050	0.250	mg/L	5	KLP1	02/14/12	1238	1186031	2
EPA 365.4 Phosphorus, Total in "As Received"											
Phosphorus, Total as P	J	0.0356	0.015	0.050	mg/L	1	KLP1	02/08/12	1418	1186020	3

The following Prep Methods were performed:

Method	Description	Analyst	Date	Time	Prep Batch
EPA 350.2 Prep	EPA 350.1 Ammonia Nitrogen Prep	AXS5	02/07/12	1658	1186022
EPA 365.4 Prep	EPA 365.4 Phosphorus, Total in liquid PR	AXS5	02/08/12	1159	1186019

The following Analytical Methods were performed:

Method	Description	Analyst Comments
1	EPA 350.1	
2	EPA 353.2	
3	EPA 365.4	

Quality Control Summary

GEL LABORATORIES LLC

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QC Summary

Report Date: February 27, 2012

Page 1 of 3

Los Alamos National Laboratory
TA-03, SM271, Drop Pt. 02U, Rm111
Los Alamos, New Mexico

Contact: Ms. Joylene Valdez

Workorder: 295270

Parmname	NOM	Sample	Qual	QC	Units	RPD%	REC%	Range	Anlst	Date	Time
Carbon Analysis											
Batch	1186182										
QC1202594162	295075001	DUP									
Total Organic Carbon Average	J	0.654	J	0.597	mg/L	9.11	^	(+/-1.00)	TSM	02/07/12	20:40
QC1202594163	295392002	DUP									
Total Organic Carbon Average	J	0.473	J	0.484	mg/L	2.30	^	(+/-1.00)		02/07/12	23:22
QC1202594166	LCS										
Total Organic Carbon Average	10.0			9.63	mg/L			96.3 (85%-115%)		02/07/12	15:58
QC1202594161	MB										
Total Organic Carbon Average			U	ND	mg/L					02/07/12	15:49
QC1202594164	295075001	PS									
Total Organic Carbon Average	10.0	J	0.654	10.7	mg/L			100 (65%-120%)		02/07/12	21:00
QC1202594165	295392002	PS									
Total Organic Carbon Average	10.0	J	0.473	10.6	mg/L			102 (65%-120%)		02/07/12	23:42
Nutrient Analysis											
Batch	1186020										
QC1202593842	295075001	DUP									
Phosphorus, Total as P	J	0.0201	U	ND	mg/L	N/A	^		KLP1	02/08/12	13:53
QC1202593848	LCS										
Phosphorus, Total as P	1.00			1.05	mg/L			105 (84%-116%)		02/08/12	13:48
QC1202593841	MB										
Phosphorus, Total as P			U	ND	mg/L					02/08/12	13:54
QC1202593844	295075001	MS									
Phosphorus, Total as P	1.00	J	0.0201	0.943	mg/L			92.3 (54%-139%)		02/08/12	13:53
QC1202593846	295075001	MSD									
Phosphorus, Total as P	1.00	J	0.0201	0.959	mg/L	1.68		93.9 (0%-21%)		02/08/12	13:59
Batch	1186023										
QC1202593854	295270001	DUP									
Nitrogen, Ammonia	U	ND	U	ND	mg/L	N/A			KLP1	02/08/12	15:43
QC1202593857	LCS										
Nitrogen, Ammonia	1.00			1.00	mg/L			100 (90%-110%)		02/08/12	15:41
QC1202593853	MB										
Nitrogen, Ammonia			U	ND	mg/L					02/08/12	15:40
QC1202593855	295270001	MS									
Nitrogen, Ammonia	1.00	U	ND	1.01	mg/L			100 (90%-110%)		02/08/12	15:44
QC1202593856	295270001	MSD									
Nitrogen, Ammonia	1.00	U	ND	1.09	mg/L	7.62		108 (0%-15%)		02/08/12	15:48
Batch	1186025										
QC1202593859	295270003	DUP									
Nitrogen, Total Kjeldahl	U	ND	U	ND	mg/L	N/A			KLP1	02/17/12	10:39
QC1202593862	LCS										
Nitrogen, Total Kjeldahl	1.00			0.913	mg/L			91.3 (90%-110%)		02/17/12	10:37
QC1202593858	MB										
Nitrogen, Total Kjeldahl			U	ND	mg/L					02/17/12	10:36
QC1202593860	295270003	MS									

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QC Summary

Workorder: 295270

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Parmname	NOM	Sample	Qual	QC	Units	RPD%	REC%	Range	Anlst	Date	Time
Nutrient Analysis											
Batch	1186025										
Nitrogen, Total Kjeldahl	1.00	U	ND	0.889	mg/L		88.9 *	(90%-110%)		02/17/12	10:40
QC1202593861 295270003 MSD											
Nitrogen, Total Kjeldahl	1.00	U	ND	0.811	mg/L	9.18	81.1 *	(0%-20%)	KLP1	02/17/12	10:41
Batch	1186031										
QC1202593873 295075001 DUP											
Nitrogen, Nitrate/Nitrite		5.16		5.07	mg/L	1.76		(0%-20%)	KLP1	02/14/12	11:36
QC1202593877 LCS											
Nitrogen, Nitrate/Nitrite	1.00			1.02	mg/L		102	(90%-110%)		02/14/12	11:26
QC1202593872 MB											
Nitrogen, Nitrate/Nitrite			U	ND	mg/L					02/14/12	11:25
QC1202593875 295075001 PS											
Nitrogen, Nitrate/Nitrite	1.00	0.516		1.53	mg/L		101	(90%-110%)		02/14/12	11:41

Notes:

RER is calculated at the 95% confidence level (2-sigma).

The Qualifiers in this report are defined as follows:

- ** Analyte is a surrogate compound
- < Result is less than value reported
- > Result is greater than value reported
- A The TIC is a suspected aldol-condensation product
- B For General Chemistry and Organic analysis the target analyte was detected in the associated blank.
- C Analyte has been confirmed by GC/MS analysis
- D Results are reported from a diluted aliquot of the sample
- E General Chemistry--Concentration of the target analyte exceeds the instrument calibration range
- E Organics--Concentration of the target analyte exceeds the instrument calibration range
- F Estimated Value
- H Analytical holding time was exceeded
- J Value is estimated
- JNX Non Calibrated Compound
- M Matrix Related Failure
- N Organics--Presumptive evidence based on mass spectral library search to make a tentative identification of the analyte (TIC). Quantitation is based on nearest internal standard response factor
- N/A RPD or %Recovery limits do not apply.
- N1 See case narrative
- ND Analyte concentration is not detected above the detection limit
- NJ Consult Case Narrative, Data Summary package, or Project Manager concerning this qualifier
- P Organics--The concentrations between the primary and confirmation columns/detectors is >40% different. For HPLC, difference is also <70%
- Q One or more quality control criteria have not been met. Refer to the applicable narrative or DER.
- R Sample results are rejected
- U Analyte was analyzed for, but not detected above the MDL, MDA, or LOD.

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QC Summary

Workorder: 295270

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Parmname	NOM	Sample	Qual	QC	Units	RPD%	REC%	Range	Anlst	Date	Time
UJ	Compound cannot be extracted										
X	Consult Case Narrative, Data Summary package, or Project Manager concerning this qualifier										
Y	QC Samples were not spiked with this compound										
Z	Paint Filter Test--Particulates passed through the filter, however no free liquids were observed.										
^	RPD of sample and duplicate evaluated using +/-RL. Concentrations are <5X the RL. Qualifier Not Applicable for Radiochemistry.										
d	5-day BOD--The 2:1 depletion requirement was not met for this sample										
h	Preparation or preservation holding time was exceeded										

N/A indicates that spike recovery limits do not apply when sample concentration exceeds spike conc. by a factor of 4 or more.

^ The Relative Percent Difference (RPD) obtained from the sample duplicate (DUP) is evaluated against the acceptance criteria when the sample is greater than five times (5X) the contract required detection limit (RL). In cases where either the sample or duplicate value is less than 5X the RL, a control limit of +/- the RL is used to evaluate the DUP result.

* Indicates that a Quality Control parameter was not within specifications.

For PS, PSD, and SDILT results, the values listed are the measured amounts, not final concentrations.

Where the analytical method has been performed under NELAP certification, the analysis has met all of the requirements of the NELAC standard unless qualified on the QC Summary.

Miscellaneous

DATA EXCEPTION REPORT			
Mo.Day Yr. 17-FEB-12	Division: Industrial	Quality Criteria: Specifications	Type: Process
Instrument Type: LACHAT Flow Injection Analyzer	Test / Method: EPA 351.2	Matrix Type: Liquid	Client Code: ESHL
Batch ID: 1186025	Sample Numbers: See below.		
Potentially affected work order(s)(SDG): 295270(12-700),295275(12-704),295388(12-711),295392(12-718) Application Issues: Failed Recovery for MS/PS Failed Recovery for MSD/PSD			
Specification and Requirements		DER Disposition:	
Exception Description: 1. Failed Recovery for MS/MSD: QC 1202593860MS, QC 1202593861MSD		1. The spike and spike duplicate recoveries fall outside of the GEL acceptance limits but within the client specified limits.	

Originator's Name:
Kristen Parson 17-FEB-12

Data Validator/Group Leader:
Julia Hamilton 24-FEB-12